bcmixed: A Package for Median Inference on Longitudinal Data with the Box–Cox Transformation

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Abstract This article illustrates the use of the **bcmixed** package and focuses on the two main functions: bcmarg and bcmmrm. The bcmarg function provides inference results for a marginal model of a mixed effect model using the Box–Cox transformation. The bcmmrm function provides model median inferences based on the mixed effect models for repeated measures analysis using the Box–Cox transformation for longitudinal randomized clinical trials. Using the bcmmrm function, analysis results with high power and high interpretability for treatment effects can be obtained for longitudinal randomized clinical trials with skewed outcomes. Further, the **bcmixed** package provides summarizing and visualization tools, which would be helpful to write clinical trial reports.

Introduction

Longitudinal data are often observed in medical or biological research. One of the most popular statistical models for studying longitudinal continuous outcomes is the linear mixed effect model. Several packages are available from CRAN that allow for the implementation of linear mixed effects models (e.g., nlme (Pinheiro et al., 2021), glme (Sam Weerahandi et al., 2021), lme4 (Bates et al., 2015), CLME (Jelsema and Peddada, 2016), PLmixed (Rockwood and Jeon, 2019), MCMCglmm (Hadfield, 2010)). The linear mixed effect models assume that longitudinal outcomes follow a multivariate normal distribution. However, the distribution of the outcome is often right skewed in the medical and biological fields. Therefore, evaluating fixed effects based on the normal distribution theory may result in inefficient inferences such as power loss for some statistical tests. In addition, although a model-based mean for a certain level of the categorical exploratory variables is often estimated when applying the linear mixed effect model (e.g., the model-based mean for each treatment group of the last visit in a randomized clinical trial), the mean may be inadequate as a representative value for the skewed data.

The Box–Cox transformation (Box and Cox, 1964) is often applied to skewed longitudinal data (Lipsitz et al., 2000) to reduce the skewness of a skewed outcome and apply existing statistical models based on a normal distribution. However, it is difficult to directly interpret the model mean estimated on the scale after applying some transformations to the outcome variable.

For the sake of the interpretability of the analysis results, Maruo et al. (2015) propose a model median inference method on the original scale based on the Box–Cox transformation in the context of randomized clinical trials. Maruo et al. (2017) extend this method to the framework of mixed effects models for repeated measures (MMRM) analysis (Mallinckrodt et al., 2001) in the context of longitudinal randomized clinical trials.

The **bcmixed** package (Maruo et al., 2020) contains functions to estimate model medians for longitudinal data proposed by Maruo et al. (2017), as well as a sample data set that is used in Maruo et al. (2017). In this package, the parameter estimation is conducted partially using the gls function in the **nlme** package. This paper illustrates the usage of the package with the sample data in several contexts.

The remainder of this manuscript is organized as follows: In section Methods, we describe the methods proposed by Maruo et al. (2017). Then, we illustrate the usage of the **bcmixed** package with the example data in section Illustrations. Finally, our contributions are summarized in section Summary and discussion.

Methods

We briefly introduce the method proposed by in Maruo et al. (2017). The detailed expressions can be found in Maruo et al. (2017).

Parameter inference for the Box-Cox model

Let the outcome be a continuous and positive value. The outcomes are measured over time for each subject i = 1, ..., n, and the number of planned measurement occasions is T (occasion index: t = 1, ..., T). The outcome vector for the ith subject is denoted by $y_i = (y_{i1}, ..., y_{in_i})^T$, where n_i is the number of measurements for the ith subject. We have $n_i \le T$ because of missingness. Then, we consider applying the following model (Lipsitz et al., 2000; Box and Cox, 1964):

$$z_i = X_i \boldsymbol{\beta} + W_i \boldsymbol{b}_i + \boldsymbol{\epsilon}_i, \tag{1}$$

where z_i is the Box–Cox transformed outcome vector such that the jth element ($j = 1, ..., n_i$) is denoted by

$$z_{ij} = \begin{cases} (y_{ij}^{\lambda} - 1)/\lambda, & \lambda \neq 0, \\ \log y_{ij}, & \lambda = 0, \end{cases}$$

and λ is the transformation parameter. The parameter β is the p-dimensional vector of the fixed effect, b_i is the q-dimensional vector of random effects distributed as $MVN_q(\mathbf{0}_q,D)$, ϵ_i is the n_i -dimensional vector of random errors distributed independently as $MVN_{n_i}(\mathbf{0}_{n_i},\Sigma_i)$, and X_i and W_i are $n_i \times p$ and $n_i \times q$ design matrices relating the fixed and random effects, respectively. The random variables b_i and ϵ_i are independent. From Formula (1), it is derived that $z_i|\lambda \sim MVN_{n_i}(X_i\beta,V_i)$, where $V_i = W_iDW_i^T + \Sigma_i$. In this paper, we consider situations where researchers have little interest in random effects and are interested in assessing fixed effects. In such cases, a simple formulation of the linear mixed effect model (1) can be implemented wherein the random effects are not explicitly modeled, but are rather included as part of the covariance matrix V_i . We focus on such a "marginal" mean model. The covariance parameter vector in $V = V_i$ for $n_i = T$ (i.e., subjects with no missing values) is denoted as $\alpha = (\alpha_1, \dots, \alpha_m)^T$. The dimension of α , that is m, depends on T and the specified covariance structure.

Let the model parameter vector be $\theta = (\lambda, \beta^T, \alpha^T)^T$. The maximum likelihood estimate of θ is obtained through maximizing the profile likelihood for λ (Lipsitz et al., 2000; Maruo et al., 2017). The model-based and robust variance estimators of the maximum likelihood estimator $\hat{\theta}$ are given by $V_{\theta}^{(M)} = \{-\hat{H}\}^{-1}$ and $V_{\theta}^{(R)} = \{-\hat{H}\}^{-1}\hat{f}\{-\hat{H}\}^{-1}$, respectively, where $H = \frac{\partial^2}{\partial \theta \partial \theta^T}l(\theta)$, $J = \sum_{i=1}^n \left\{\frac{\partial}{\partial \theta}l_i(\theta)\right\}\left\{\frac{\partial}{\partial \theta}l_i(\theta)\right\}^T$, $l(\theta)$ is the likelihood function for n subjects, and $l_i(\theta)$ is the likelihood function for the ith subject. The matrices \hat{H} and \hat{J} are obtained from the matrices H and J by replacing θ by $\hat{\theta}$, respectively. A robust variance estimator is an asymptotically valid estimator even when the model (1) is mis-specified (Cox, 1961).

Model median inference

We now focus on the continuous and positive outcomes of a certain disease, and consider a situation in which the efficacy of some treatments (group index: g=1,...,G) is compared based on a randomized, parallel group clinical trial, where the total number of subjects is n. The explanatory variable matrix, X_i , and the fixed effect parameter, $\boldsymbol{\beta}$ in model (1) are denoted in detail as follows. Now, X_i is given by the $n_i \times (GT+C)$ matrix that contains the intercept, G-1 group variables, T-1 occasion variables, group-by-occasion interaction variables, and C covariates, where the categorical covariates are converted into dummy variables. The fixed effect parameter vector is given by $\boldsymbol{\beta} = (\beta_I, \boldsymbol{\beta}_g^T, \boldsymbol{\beta}_t^T, \boldsymbol{\beta}_g^T, \boldsymbol{\beta}_c^T)^T$, where β_I , $\beta_g = (\beta_{g(1)}, \ldots, \beta_{g(G-1)})^T$, $\beta_t = (\beta_{t(1)}, \ldots, \beta_{t(T-1)})^T$, $\beta_{gt} = (\beta_{gt(1,1)}, \beta_{gt(1,2)}, \ldots, \beta_{gt(G-1,T-1)})^T$, and $\boldsymbol{\beta}_c = (\beta_{c(1)}, \ldots, \beta_{c(C)})^T$ are the intercept, group, occasion, group-by-occasion, and covariate parameter vectors, respectively. Although group G and occasion T is set to be at the reference level, we set $\boldsymbol{\beta}_{g(G)} = \beta_{t(T)} = \beta_{gt(G,t)} = \beta_{gt(g,T)} = 0$ for the sake of description $(g=1,\ldots,G,t=1,\ldots,T)$.

Under these settings, the model median, $\xi_{(g,t)}$, for the treatment group g at the occasion t on the original scale is given by

$$\xi_{(g,t)} = \left\{\lambda \left(\beta_I + \beta_{g(g)} + \beta_{t(t)} + \beta_{gt(g,t)} + \mathbf{x}_{\tilde{c}}^T \boldsymbol{\beta}_c\right) + 1\right\}^{1/\lambda},$$

where $x_{\bar{c}}$ is the vector of the mean of each covariate for all subjects. The model median is the inverse Box–Cox transformation of the model means on the Box–Cox transformed scale. The model median can be easily interpreted because it is the estimator of the median on the original scale.

Using the delta method, the variance estimator of the maximum likelihood estimator for the model median, $\hat{\xi}_{(g,t)}$, is given by $V_{\xi(g,t)}^{(\cdot)} = \Delta_{\xi(g,t)}^T V_{\theta}^{(\cdot)} \Delta_{\xi(g,t)}$, where $\Delta_{\xi(g,t)} = \frac{\partial}{\partial \theta} \xi_{(g,t)} \Big|_{\theta=\hat{\theta}}$. If we use $V_{\theta}^{(\cdot)} = V_{\theta}^{(M)}$, we obtain the model-based variance estimator, $V_{\xi(g,t)}^{(M)}$. On the other hand, the robust

variance estimator, $V_{xi(g,t)}^{(R)}$, is obtained if we use $V_{\pmb{\theta}}^{(\cdot)} = V_{\pmb{\theta}}^{(R)}$.

Thus, the model median difference between groups g_1 and g_2 at occasion t is given by $\delta_{(g_1;g_2,t)}=\xi_{(g_1,t)}-\xi_{(g_2,t)}$ and the variance estimators of the maximum likelihood estimator of the model median difference, $\hat{\delta}_{(g_1;g_2,t)}$, is given by $V_{\delta(g_1;g_2,t)}=(\Delta_{(g_1,t)}-\Delta_{(g_2,t)})^TV_{\pmb{\theta}}^{(\cdot)}(\Delta_{(g_1,t)}-\Delta_{(g_2,t)})$. The model-based and robust variance estimators are obtained in the same way as the model median estimator. Using the asymptotic normality of the maximum likelihood estimator, the $100(1-\alpha)\%$ confidence interval of $\delta_{(g_1;g_2,t)}$ is obtained as $\hat{\delta}_{(g_1;g_2,t)}\pm\Phi^{-1}(1-\alpha/2)\sqrt{V_{\delta(g_1;g_2,t)}^{(\cdot)}}$, where $\Phi^{-1}(\cdot)$ is the quantile function of the standard normal distribution. The Wald-type test for the null hypothesis, $\delta_{(g_1;g_2,t)}=0$, is performed with the test statistic $t=\hat{\delta}_{(g_1;g_2,t)}/\sqrt{V_{\delta(g_1;g_2,t)}^{(\cdot)}}$.

The performances of the above-mentioned inference procedures may be low for a small sample because these are based on the asymptotic properties of the maximum likelihood estimation. Therefore, Maruo et al. (2017) applied the following empirical small sample adjustment for the inferences of the model median differences by referring to the study in Schluchter and Elashoff (1990).

They provide an adjusted standard error (SE) for the median difference as $\sqrt{M/(M-p)}V_{\delta(g_1;g_2,t)}^{(\cdot)}$ for the compound symmetry (CS) or the first-order auto regression (AR(1)) structure and $\sqrt{n^*/(n^*-T)}\times\sqrt{V_{\delta(g_1;g_2,t)}^{(\cdot)}}$ for the unstructured (UN) structure. Further, we approximate the null distribution by the t distribution where the degrees of freedom for the CS or the AR(1) structure and the UN structure are (n-G)(T-1)-m and n^*-T , respectively.

Although Maruo et al. (2017) and Maruo et al. (2020) focus only on the three covariance structures, these three options are sufficient in the applied settings because the MMRM analysis is often applied in the following steps. The UN structure is used, and the CS or AR(1) structure with the robust variance estimation is used when the parameter estimation process is not properly converged (e.g., see Gosho et al. (2017)).

Illustrations

In this section, we describe the **bcmixed** package that provides the analysis results based on the mixed effect models with the Box–Cox transformation. The package **bcmixed** is available from the Comprehensive R Archive Network (CRAN) at https://CRAN.R-project.org/package=bcmixed. The package **bcmixed** can be installed and loaded using the following code.

R> install.packages("bcmixed")
R> library(bcmixed)

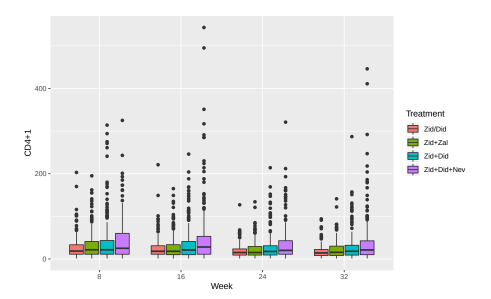


Figure 1: Longitudinal box-whisker plot of outcome variable (cd4) for each treatment group in the aidscd4 dataset.

In particular, the following main functions are demonstrated in this article:

bcmarg: Inference on the marginal model of the mixed effect model with the Box-Cox transformation.

bcmmrm: Inference on the model median for longitudinal data in randomized clinical trials.

Example data

First, we illustrate a real example for the acquired immune deficiency syndrome (AIDS) clinical trial data, which is stored as the data frame aidscd4 in the **bcmixed** package. The data are from a randomized, double-blind study of AIDS patients with advanced immune suppression (cluster of differentiation 4 (CD4) cells count of less than or equal to 50 cells/mm3) (Henry et al., 1998; Fitzmaurice et al., 2011). Patients in the AIDS clinical trial group study 193A were randomized to dual or triple combinations of human immunodeficiency virus-1 reverse transcriptase inhibitors. In particular, the patients were randomized to one of four daily regimens. The original data can be downloaded from https://content.sph.harvard.edu/fitzmaur/ala/ (Datasets->AIDS Clinical Trial Group (ACTG) Study 193A). As for the more detailed data handling process, see Maruo et al. (2017). The data frame aidscd4 contains seven variables (Table 1).

Variable	Description
id	A patient identifier; in total there are 1177 patients.
weekc	A visit variable (weeks 8, 16, 24, 32).
treatment	Allocated treatment regimens;
	1 = zidovudine alternating monthly with 400mg didanosine,
	2 = zidovudine plus 2.25mg of zalcitabine,
	3 = zidovudine plus 400mg of didanosine, and
	4 = zidovudine plus 400mg of didanosine plus 400mg of nevirapine.
age	Patients' age (years).
sex	Patients' sex (1 = male, 0 = female).
cd4.bl	A baseline value of CD4 cells count + 1.
cd4	A CD4 cells count + 1.

Table 1: Variable definition of AIDS clinical trial data

Figure 1 shows the longitudinal box-whisker plot of the values of CD4 plus 1 for each group plotted using ggplot2 package (Wickham, 2016). The distribution shapes of the measurements were heavily skewed.

bcmarg function

This function provides the inference results for the marginal model of the mixed effect model with the Box–Cox transformation described in Section Parameter inference for the Box–Cox model.

Usage: The bcmarg function is called using the following syntax.

```
bcmarg(formula, data, time = NULL, id = NULL, structure = "UN", lmdint = c(-3, 3))
```

Arguments: The bcmarg function takes arguments tabulated in Table 2.

Argument	Description
formula	A two-sided linear formula object describing the model, with the response on the left of a ~ operator and the terms, separated by + operators, on the right.
data	A data frame containing the variables used in the model.
time	A time variable name for repeated measurements. The default is NULL.
id	A subject id variable name for repeated measurements. The default is NULL.
structure	Specify the covariance structure from c("UN", "CS", "AR(1)"). The default is "UN".
lmdint	A vector containing the end points of the interval to be searched for a
	transformation parameter. The default is $c(-3,3)$.

Table 2: Auguments of bcmarg function

If time and id are not specified, an ordinary linear model with the Box–Cox transformation is applied.

Value: The bcmarg function returns an object of class "bcmarg". The objects of this class have methods for generic functions coef, logLik, vcov, fitted, print, and summary. The object includes the components for the marginal model parameter inference (Table 3).

Component	Description
lambda	A numeric value of the estimate of the transformation parameter.
beta	A vector with the estimates of the regression parameters.
alpha	A vector with the estimates of the covariance parameters.
V	A variance-covariance matrix for any subject with no missing values.
betainf	A matrix containing the inference results for beta under the assumption
	that lambda is known.
Vtheta.mod	A model-based variance-covariance matrix for MLE of the vector of all
	parameters: c(lambda,beta,alpha).
Vtheta.rob	A robust variance-covariance matrix for MLE of the vector of all parameters.
logLik	A numeric value of the maximized likelihood.
adj.prm	A vector with parameters used for the empirical small sample adjustment in
	bcmmrm: c(number of subjects, number of completed subjects, number of
	outcome observations, number of missing observations).
glsObject	An object of "gls" (or "lm" when time and id are not specified) containing
	results of gls (or lm) function on the transformed scale.

Table 3: Values of bcmarg function

In bcmarg function, lambda is estimated with the optimize function by maximizing the profile likelihood function for λ . If an error occurs in the optimize function, problems may be solved by changing the search area for λ , lmdint.

Example code: We applied a marginal model to the aidscd4 data, where the fixed effects were the treatment, visit, treatment-visit interaction, and the Box–Cox transformed baseline, where the visit was treated as a nominal variable. The covariance structure was set as unstructured (default setting). This model is frequently used for MMRM analysis. A sample code is as follows. The bct.v function returns the Box–Cox transformed vector.

The summarized analysis results on the transformed scale are obtained by applying the summary function to the "beggarg" object as follows:

```
function to the "bcmarg" object as follows.
R> summary(res1)
Box-Cox transformed mixed model analysis
 Formula: cd4 ~ as.factor(treatment) * as.factor(weekc) + cd4.bl.tr
 Time: weekc
 ID: id
 Covariance structure: "UN"
 Data: aidscd4
 Log-likelihood: -13322.96
 Estimated transformation parameter: 0.154
Coefficients on the transformed scale:
                                          Value Std.Error t-value p-value
                                         1.0849 0.1249 8.684 0.000
(Intercept)
                                                   0.1214 2.022 0.043
as.factor(treatment)2
                                         0.2454
```

```
as.factor(treatment)3
                                  0.4203
                                          0.1212 3.468
                                                        0.001
                                  0.7649
                                          0.1199 6.377
as.factor(treatment)4
                                                        0.000
                                 -0.2043
                                          0.0843 -2.424
as.factor(weekc)16
                                                        0.015
                                 -0.4498
                                          0.0899 -5.004
as.factor(weekc)24
                                                        0.000
as.factor(weekc)32
                                 -0.6750 0.0988 -6.835
                                                        0.000
cd4.bl.tr
                                  0.5782 0.0200 28.922
                                                        0.000
as.factor(treatment)3:as.factor(weekc)16 -0.0560 0.1180 -0.475 0.635
as.factor(treatment)4:as.factor(weekc)16 0.0675 0.1175 0.574 0.566
as.factor(treatment)3:as.factor(weekc)24 -0.0243
                                          0.1264 -0.193 0.847
as.factor(treatment)4:as.factor(weekc)24 0.0870 0.1263 0.689 0.491
                                          0.1414 -0.603 0.547
as.factor(treatment)2:as.factor(weekc)32 -0.0852
                                          0.1400 -0.638
as.factor(treatment)3:as.factor(weekc)32 -0.0893
                                                        0.524
as.factor(treatment)4:as.factor(weekc)32 0.1414
                                          0.1381 1.024
                                                        0.306
Covariance parameters on the transformed scale:
UN(1,1) UN(1,2) UN(1,3) UN(1,4) UN(2,2) UN(2,3) UN(2,4) UN(3,3) UN(3,4) UN(4,4)
              1.105 0.927
                          1.957
                                 1.346 1.298
                                              1.903 1.452
Correlations on the transformed scale:
         16
              24
8 1.000 0.616 0.598 0.488
16 0.616 1.000 0.697 0.655
```

The results of coefficients on the transformed scale are obtained with the gls function in the nlme package. The transformation parameter was estimated as 0.154, which suggested that the shape of the residual distribution was close to a multivariate log-normal distribution. All main effects were significant; however, treatment-by-week interaction was not significant at a level of 0.05. Note that inference results for beta under the assumption that the transformation parameter is known are provided. Although statistical tests would be asymptotically valid (e.g., see Doksum and Wong (1983)), standard errors might be underestimated (e.g., see Bickel and Doksum (1981)).

bcmmrm function

24 0.598 0.697 1.000 0.743 32 0.488 0.655 0.743 1.000

This function provides the results for the model median inferences for longitudinal randomized clinical trial data described in Section Model median inference. The parameter inference is conducted by calling the bcmarg function in the bcmmrm function.

Usage: The bcmmrm function is called using the following syntax.

```
bcmmrm(outcome, group, data, time = NULL, id = NULL, covv = NULL,
  cfactor = NULL, structure = "UN", lmdint = c(-3, 3), glabel = NULL,
  tlabel = NULL)
```

Argument: The bcmmrm function takes arguments tablated in Table 4.

Argument	Description
outcome	A name of positive outcome (dependent) variable included in data.
group	A name of treatment group variable included in data.
data	A data frame that may include outcome, group, time, id, and specified covariate
	variables.
time	A name of time variable for repeated measurements included in data.
	The default is NULL.
id	A name of subject id variable for repeated measurements included in data.
	The default is NULL.
covv	A character vector with names of covariate variables included in data.
	The default is NULL.
cfactor	An integer vector including nominal variable indicators for covariate variables.
	Nominal variable: 1, continuous variable: 0. The default is NULL.

structure	Specify the covariance structure from c("UN", "CS", "AR(1)"). The default is "UN".	
conf.level	A numeric value of the confidence level for the confidence intervals.	
	The default is 0.95.	
lmdint	A vector containing end points of the interval to be searched for a transformation	
	parameter. The default is c(-3,3).	
glabel	A vector of length number of treatment groups containing the labels of group	
	variable. The default is NULL and the levels of group variable in data are used.	
tlabel	A vector of length number of repeated measures containing the labels of time	
	variable. The default is NULL and the levels of time variable in data are used.	

Table 4: Auguments of bcmmrm function

If time and id are not specified, inference results reduce to the results for the context of linear regression model provided by Maruo et al. (2015).

Value: The bcmmrm function returns an object of class "bcmmrm" representing the results of model median inference based on the Box–Cox transformed MMRM analysis. Generic functions boxplot, coef, logLik, vcov, fitted, plot, print, and summary have methods to demonstrate the results of the fit. Components tablated in Table 5 must be included in a legitimate "bcmmrm" object.

Component	Description
call	A list containing an image of the bcmmrm call that produced the object.
median.mod,	Lists including inference results for the model medians for all groups.
median.rob,	Levels of the list are time points, where the correspondence table is given
median.mod.adj,	as time.tbl\$code.mod: model-based inference, rob: robust inference,
median.rob.adj	adj: with empirical small sample adjustment.
meddif.mod,	Lists including inference results for the for the model median differences
meddif.rob,	between all pairs of groups (group1 - group0). The levels of the list are
meddif.mod.adj,	time points, where the correspondence table is given as time.tbl\$code.
meddif.rob.adj	mod: model-based inference, rob: robust inference,
	adj: with empirical small sample adjustment.
lambda	A numeric value of estimates of the transformation parameter.
R	A correlation matrix for transformed outcomes.
betainf	Inference results for beta under the assumption that lambda is known.
time.tbl	A data frame of a correspondence table for the time points.
	This object is used when applying the above generic functions.
group.tbl	A data frame of a correspondence table for treatment groups.
	This object is used when applying the above generic functions.
inf.marg	A "bcmarg" object with results of the bcmarg function called in the
	bemmrm function.
outdata	A data frame where the transformed outcome (ytr), the fitted values
	on the transformed scale (ytr.fitted), and the residuals on
	the transformed scale (res.tr) are added to input data.
conf.level	A numeric value of the specified confidence level.

Table 5: Values of bcmmrm function

lambda, R, betainf, and inf.marg are obtained from the results of the bcmarg function that is called in the bcmmrm function.

Sample code: We applied the MMRM analysis with the Box–Cox transformation described in Section bornarg function to the aidscd4 data, where the Box–Cox transformed baseline and sex were included as the covariates. The example code is as follows.

The transformed baseline and sex are continuous and categorical variables, respectively, and therefore, cfactor was set as c(0,1).

The print function only provides information about model detail, the estimated transformation parameter, the maximized log-likelihood, and the model median estimate for each time point and group as follows.

```
R> print(res2)
Model median estimation based on MMRM with Box-Cox transformation
 Outcome: cd4
 Group: treatment
 Time: weekc
 ID: id
 Covariate(s): c("cd4.bl.tr", "sex")
 Covariance structure: "UN"
 Data: aidscd4
 Estimated transformation parameter: 0.154
 Log-likelihood: -13322.36
Model median estimates (row: group, col: time):
  treatment | weekc 8 16 24 32
           Zid/Did 18.9 16.5 14.1 12.1
2
           Zid+Zal 22.0 17.9 14.6 13.5
3
           Zid+Did 24.5 20.9 18.2 15.1
4
       Zid+Did+Nev 30.1 27.8 24.2 21.8
   The summary function provides more detailed analysis results as follows.
R> summary(res2)
Model median inference based on MMRM with the Box-Cox transformation
Data and variable information:
 Outcome: cd4
 Group: treatment
 Time: weekc
 ID: id
 Covariate(s): c("cd4.bl.tr", "sex")
 Data: aidscd4
Analysis information:
 Covariance structure: "UN"
 Robust inference: TRUE
 Empirical small sample adjustment: TRUE
 Confidence level: 0.95
Analysis results:
 Estimated transformation parameter: 0.154
Model median inferences for weekc = 8
    treatment median SE lower.CL upper.CL
1
     Zid/Did 18.9 0.862
                             17.2 20.6
     Zid+Zal 22.0 1.124
                              19.8
                                       24.2
2
     Zid+Did 24.5 1.465
                              21.6
                                       27.4
4 Zid+Did+Nev 30.1 1.597
                              27.0
                                       33.3
(...Omitted for weeks 16 and 24...)
Model median inferences for weekc = 32
    treatment median
                     SE lower.CL upper.CL
1
     Zid/Did 12.1 0.662
                           10.8
                                       13.4
                              11.9
                                       15.1
     Zid+Zal
               13.5 0.813
3
     Zid+Did
               15.1 1.019
                              13.1
                                       17.1
4 Zid+Did+Nev 21.8 1.376
                              19.1
                                       24.5
```

```
Inferences of model median difference between groups
 ( treatment 1 - treatment 0 ) for weekc = 8
 treatment 1 treatment 0 delta SE lower.CL upper.CL t.value p.value
    Zid+Zal Zid/Did 3.12 1.40 0.363 5.87
                                                  2.22 0.027
1
    Zid+Did Zid/Did 5.64 1.69 2.325 8.96
                                                   3.34 0.001
3 Zid+Did+Nev Zid/Did 11.25 1.80 7.711 14.80
                                                  6.24 0.000
    Zid+Did Zid+Zal 2.53 1.83 -1.059 6.12 1.39 0.167
5 Zid+Did+Nev Zid+Zal 8.14 1.93 4.349 11.93 4.22 0.000
6 Zid+Did+Nev Zid+Did 5.61 2.16 1.372 9.85 2.60 0.010
(...Omitted for weeks 16 and 24...)
Inferences of model median difference between groups
 ( treatment 1 - treatment 0 ) for weekc = 32
 treatment 1 treatment 0 delta SE lower.CL upper.CL t.value p.value
     Zid+Zal Zid/Did 1.35 1.04 -0.692 3.40 1.30 0.194
     Zid+Did
               Zid/Did 3.00 1.20
                                  0.633
                                           5.37
                                                   2.49
                                                         0.013
                                  6.705 12.69
    Zid+Did
+Did+Nev
+Did+Nev
3 Zid+Did+Nev
               Zid/Did 9.70 1.52
                                                   6.37
                                                         0.000
               Zid+Zal 1.65 1.30 -0.907
                                           4.20
                                                   1.27
                                                         0.206
                                 5.206
               Zid+Zal 8.34 1.60
5 Zid+Did+Nev
                                          11.48
                                                   5.23
                                                         0.000
                                  3.338 10.06
6 Zid+Did+Nev
               Zid+Did 6.70 1.71
                                                   3.92
                                                         0.000
```

Significant differences were detected for the following pairs Zid+Did vs. Zid/Did, Zid+Did+Nev vs. Zid/Did, Zid+Did+Nev vs. Zid/Did, Zid+Did+Nev vs. Zid+Did at week 32.

The summary function provides the results using the robust variance and the small sample adjustment in the default settings. If users want to summarize results using the model variance and not using the small sample adjustment, specify summary(bcmmrmObject,robust = F,ssadjust = F). Further details of the summary function for the "bcmmrm" object can be obtained with ?summary.bcmmrm.

The inference results for the median differences at week 32 (fourth visit) can also be called as follows although the levels of the group variable in the data frame are used without formatting.

```
R> res2$meddif.rob.adj[[4]]
```

```
group1 group0 delta SE lower.CL upper.CL t.value p.value
1 2 1 1.354438 1.041338 -0.6922404 3.401117 1.300672 1.940595e-01
2 3 1 3.000942 1.204919 0.6327547 5.369129 2.490575 1.312570e-02
3 4 1 9.697631 1.522831 6.7046091 12.690653 6.368159 4.869820e-10
4 3 2 1.646503 1.299231 -0.9070469 4.200054 1.267291 2.057293e-01
5 4 2 8.343192 1.596236 5.2058987 11.480486 5.226792 2.682862e-07
6 4 3 6.696689 1.709027 3.3377114 10.055667 3.918421 1.034880e-04
```

The "bcmmrm" object can be plotted with the plot function as follows (Figure 2).

```
R> plot(res2, xlab = "Week", ylab = "CD4+1")
```

The plot function provides a longitudinal plot in the default settings. However, a plot at a specified time point can be drawn with the following code (Figure 3):

```
R> plot(res2, timepoint = 32, xlab = "Treatment", ylab = "CD4+1", col = 1:4)
```

Further, the plot function provides the results using the robust variance and the small sample adjustment in the default setting. Many other options such as main and legend can be used in the plot function. Further details can be obtained with <code>?plot.bcmmrm</code>.

A diagnosis of a model fitting can be conducted with the boxplot function, which provides a box-whisker plot of the Box–Cox transformed residuals for each group. A sample code is provided as follows (Figure 4):

```
R> boxplot(res2)
```

The shape of the transformed residual for each group is not skewed and the median and mean are close to each other, which suggests that the median would not be biased at week 32. The boxplot function provides the results at the last time points in the default settings. A box-whisker plot at another time point can be obtained by specifying boxplot(bcmmrmObject,timepoint = xx).

Summary and discussion

We demonstrated the use of the **bcmixed** package. The bcmarg function provides the analysis results for a marginal model of a mixed effect model with the Box–Cox transformation. The results for the statistical test of the bcmarg function are meaningful. However, it is difficult to interpret coefficients (β) on the transformed scale.

The bcmmrm function provides the model median inferences based on the MMRM with the Box–Cox transformation for longitudinal randomized clinical trials. Using this function, treatment effects can be interpreted as median differences between treatment groups at specified time points. Maruo et al.

Longitudinal plot for model median of each group 40 Zid/Did Zid+Zal 35 Zid+Did Zid+Did+Nev 30 25 20 15 9 8 16 24 32

Figure 2: Longitudinal plot of model median for each group, created by applying plot function to bcmmrm object.

Week

Zid/Did Zid+Zal Zid+Did Zid+Did+Nev Treatment

Figure 3: Plot of model median for each group at week 32, created by applying plot function to bcmmrm object with timepoint option.

Plot for model median of each group

(2017) also show that this method controls the type I error of the statistical test for the model median difference, and it has moderate or high performance for power compared with the existing methods (ordinary MMRM, MMRM for the log-transformed outcome, etc.) from their simulation studies (the simulation program is provided in https://github.com/kzkzmr/Maruo_2017_StatMed_Simulation with penalized power results proposed by Cavus et al. (2019)). Thus, bcmmrm function analysis results with high power and high interpretability for longitudinal randomized clinical trials with skewed outcomes. Further, the bcmmrm function provides summarizing and visualization tools, which would be helpful to write clinical trial reports.

Although the **bcmixed** package can be used for data other than that of randomized clinical trials, the performances of these methods have not been evaluated well yet. Therefore, users should use them carefully. Although a model fitting can be diagnosed with the boxplot function, more model diagnosis tools may be implemented in the future. Users might apply the **MissMech** package(Jamshidian et al., 2014, 2015), which diagnoses multivariate normality and heteroscedasticity, to the transformed residuals stored in "bcmarg" object. Note that the **MissMech** package assumes missing mechanisms are missing completely at random (MCAR), and statistical tests for model fittings may lead to significant results for a medium-to-large sample even when the models fit the data adequately.

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Box-whisker plot for transformed residuals at weekc = 32

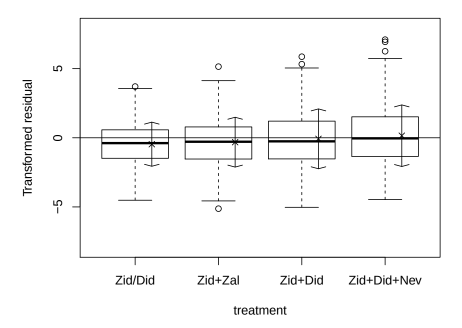


Figure 4: Box-whisker plot of transformed residuals for each group at week 32, created by applying boxplot function to bcmmrm object.

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