User’s guide to Bayesian Cultural Consensus Toolbox

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Introduction

In spite of the attractive features of the Bayesian statistical framework, it has yet to gain traction among ethnographers in general. This applies for Cultural Consensus Theory (CCT) research specifically because the Bayesian General Condorcet Models (GCMs) described in the paper of Karabatsos and Batchelder (2003) have so far not been featured in any substantive ethnographic investigation. Likely one reason for that is the lack of practical exposure to the Bayesian framework. Also, the implementation of Bayesian parameter estimation routines for CCT has not yet appeared in the form of a user-friendly software package which could be easily accessible to a wide audience of ethnographers. Bayesian parameter estimation methods differ from those of the other estimation frameworks as they are based on iterative sampling schemes to obtain posterior probability distributions of the model parameters. As such, they are not implemented in common software packages or in the software packages typically used for CCT modeling like ANTHROPAC (Borgatti, 1996), and UCINET (Borgatti, Everett, & Freeman, 2002).

Here we introduce a software program called Bayesian Cultural Consensus Toolbox (BCCT) for conducting Bayesian inference for the GCM handling dichotomous data. The software is freely accessible on the first author’s website (bayesian.zitaoravecz.net) and it features an easily comprehensible user interface, so that ethnographers can simply estimate model parameters after having read the instructions in this user’s guide. The program offers a choice between several CCT models for dichotomous response data incorporating different degrees of variability in the leading parameters. We hope that this will facilitate the consideration of a greater variety of CCT models other than those handled by ANTHROPAC and UCINET, while introducing ethnographers to Bayesian methods of statistical inference. This user’s guide is intended as a supplementary material for a Field Methods paper entitled Bayesian Cultural Consensus Theory by Oravecz, Z., Vandekerckhove, J., and Batchelder, W. H., that introduces the Bayesian framework for Cultural Consensus Theory in details. However, with respect to using the BCCT software for data analysis, this documentation is self-sufficient.

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Introducing the Bayesian Cultural Consensus Toolbox through an example dataset

BCCT is a standalone application that can work solely through a graphical user interface (similar to UCINET) and does not require any programming knowledge from the user. In this section we demonstrate how straightforward it is to carry out statistical inference for several versions of the GCM. The interface allows the user to easily navigate among the different modeling choices. Moreover, BCCT offers a straightforward way to explore the posterior distribution of the parameter estimates, as well as it provides options to deal with issues of model selection and model fit. A document that downloads with the software package guides the user step by step through the installation. BCCT uses two accompanying programs that extract automatically. One of them is JAGS (Plummer, 2011) which carries out the sampling for the different GCM models. It works in the background when the BCCT software is used and is not explicitly displayed in any way for the user. Second, as the program for the interface and all supporting subprograms were written in MATLAB, a MATLAB Compiler Runtime installs when the package for the software is extracted. The user does not need a MATLAB license.1

Reading in data, specifying the model, starting the estimation

Now we demonstrate how to use BCCT for performing Bayesian statistical inference for GCM models. A flow diagram showing 10 simple steps for conducting such inference is included below in Figure 1. BCCT offers a default model, which is the GCM as it was presented in the earliest paper on CCT (Romney, Weller, & Batchelder, 1986; Batchelder & Romney, 1988), namely the model with homogeneous items and heterogenous competency and guessing bias.2 If the user successfully prepares their datafile, takes the default model and sampler settings and the sampling process works out satisfactorily, the analysis can be carried out by going through the 10 steps above. More advanced users can exploit several other options when constructing models or exploring different aspects of the results.

Hereby we present an example application to demonstrate how to carry out analysis with BCCT. We make use of a dataset that is published in Romney (1999). The dataset was collected by Weller (1984) and it has been analyzed by several other authors such as Weller (1984), Romney et al. (1986) and Batchelder and Anders (2012), however, only with models assuming item homogeneity. The original study measured knowledge about contagiousness of certain diseases in a Guatemalan population. The sample consists of 24 informants who provided dichotomous responses to 27 items about a disease being contagious or not. In the next paragraphs we present an example analysis with BCCT on this dataset, while referring to the numbering as displayed by the flowchart in Figure 1.

(1)(2) BCCT can read a dataset from a text file with informants represented by rows and items as columns. The data should be coded with only 0 and 1 entries. The example dataset analyzed

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1As mentioned before, BCCT can be downloaded from bayesian.zitaoravecz.net. Also, all MATLAB scripts that constitute the program are automatically downloaded as part of the BCCT package, and can be found in the folder called ‘Supplementary files’. Users who do have a MATLAB license can use these files to run BCCT in its native environment.
2The latter was fixed to 0.5 only for computational efficiency only in earlier papers, as well as in ANTHROPAC and UCINET.
Figure 1. : Flow diagram for conducting Bayesian inference for the GCM with BCCT
here downloads with the program. When BCCT opens the user can click on Browse and load the preferred dataset. Figure 2 displays a screenshot of BCCT when the example dataset in the file called contagiondata.txt has been loaded.

(3) Under the data path the program displays results from exploring the loaded data. This way the user can check whether the number of informants and items are correct. Moreover, the last box shows the ratio of the first and second eigenvalue based on the covariance matrix of the informants’ answers as described in Batchelder and Romney (1988). The same diagnostic is built in UCINET and field researchers often check the basic model axiom of a single underlying culture with the help of this number. In practice, it has been assumed that a threshold of at least 3 is needed to support the single culture assumption. However, the eigenvalue ratio can only indicate that the single culture assumption is most likely violated (if ratio is under 3), but it can never prove that this condition is met, even if it is above 3. For discussion on this problem see Hruschka, Sibley, Kalim, and Edmonds (2008). Here we offer this number only to provide an exploratory indicator of whether the dataset seems to violate the assumption. Later we show how to test the one culture assumption in the Bayesian framework as part of the posterior predictive model checking, following the guidelines of Batchelder and Anders (2012).

(4) The second panel in Figure 2 allows the user to specify which model is to be estimated. As mentioned above, by default it is set to the GCM with heterogeneous competence and guessing bias parameters, while assuming that item difficulties are neutral (homogeneous). This way a competence and a guessing bias parameter are estimated for each respondent and as many answer key variables as items. Alternative models can be set by either relaxing the neutral item difficulty constraint by clicking on the ‘heterogeneous’ option or by imposing constraints on the competence
(same for all informants) and/or the guessing bias (neutral, i.e., fixed to 0.5) parameters. For example, if we change BCCT’s default model option by fixing the guessing bias to 0.5, we fit the same GCM as UCINET and ANTHROPAC would fit but in the Bayesian statistical inference framework.

(5) The bottom panel Figure 2 contains the parameter settings for the sampler program that draws samples from the posterior distributions of the model parameters. The standard strategy is to use modern computational tools, called Markov chain Monte Carlo (MCMC) methods, to develop a sampler that can generate many random values from the posterior distribution, and use the sampled values to approximate the posterior distribution. Any MCMC sampler is designed to sample values of the parameters in proportion to how probable they are from the posterior distribution. An extensive literature exists (see e.g., Robert & Casella, 2004; Gamerman & Lopes, 2006) regarding simulation procedures to draw samples from any kind of distribution. Also, there are available software packages that automate many of the steps required to run such a sampling procedure. BCCT relies on one of these, namely JAGS. JAGS has built in heuristics to decide about the most efficient MCMC algorithms to use for the specified models and most of the time it is an amalgam of Metropolis-Hastings, Gibbs and slice sampling algorithms (Robert & Casella, 2004).

The user has to decide only four options displayed in the bottom panel, however, most often the already pre-filled values can work out satisfactorily, so the user can just leave those unmodified and click directly on the Start estimating parameters button. The reason for the options in this row is related to the standards for assuring the adequacy of the MCMC sampler which is discussed later in the Exploring the results section. In the very first box, called Adaptation, we allocate a certain number of samples to be drawn but then discarded (not used to estimate the posterior distribution), so they will not count towards our final posterior samples. The reason for this adaptation period (most often called “Burn-in”) is twofold. First of all, the first sampled values of each sample chain are highly determined by their random starting values, and therefore do not represent the posterior distribution accurately. Second, during some initial period of sampling we want the sampler program selected by JAGS to adapt its algorithms to find the most efficient one. For most CCT models the sampler needs around the first 1000 iterations to be discarded in order to avoid the effects of the initial setting and allow JAGS to select an efficient sampler.

In the second box, called Iterations, the number of final (useable) posterior samples for each chain is specified. It should be sufficiently large so that the sampler can explore the whole area of the conditional posterior densities of the parameters. Also, as mentioned above, this density exploration should be restarted with different initial values to further ensure the sufficient exploration of the entire parameter space, and in the box called Chains we define how many times the explorations are restarted. Our final posterior sample size is the product of these two. As a rule of thumb, 6 chains with a minimum of 1000 useable samples each should be sufficient in most cases. Finally, the very last box concerns the autocorrelation in the chains. Posterior inference should be based on chains that have little or no autocorrelation, so that we can be sure that each area of the posterior density is equally likely to be represented in our final posterior sample. The way the Thin box controls the autocorrelation is the following: it keeps only every $k^{th}$ sample and discards the others, this way decreasing correlation between the new neighboring samples. So if we put 1 in that box, that means that we do not thin and we keep every sample after the adaptation period. If we put 2 (default
setting) we discard every second sample and so on until we reach the number of useable iterations set in the Iteration box. The more we thin, the smaller the autocorrelation becomes. While for some models and some datasets probably no thinning is necessary, we will see that thinning needs to be increased in some cases. As a matter of fact, for the analysis of the contagion data, we changed the sampler setting parameters somewhat: we raised the thinning into 5, as shown in Figure 3.

(6) The screenshot in Figure 3 represents a state of the program in which the estimation has already been started by clicking the Start estimation button, the text of which is now changed into Estimating. The sampling might take several minutes, depending on the size of the dataset and the settings in the lower panel. The current example analysis for example took about one minute on a PC with a 3.20 GHz Xeon CPU and 12 GB RAM.

Figure 3. : Screenshot of BCCT after the model and the sampler options have been specified and the estimation has been started.

Exploring the results

Figure 4 displays the window which pops up after the sampling is finished, except that initially the lower panel would be empty until the user pushes the Perform checks button (which here is already changed to Checks performed).

(7) The upper panel (Check sampler) provides us with some basic information about the quality of our posterior samples. The goal of this is to make sure that the sampler is accurately tracking the correct posterior distribution on which the inferences are drawn. The first standard that needs to be met by the MCMC sampler is that the entire parameter space must be sufficiently explored. For that, sampling schemes should start multiple times, each with different starting values in different regions of the space of possible parameter values, and the generated sample chains
sequences of draws from the posterior distribution of the parameters) should all end up conveying the same information about the posterior distribution. Therefore the first box in Figure 4 reports on whether the chains appear to have converged to the same distribution. The way the program checks convergence is to calculate the $\hat{R}$ values (Gelman, Carlin, Stern, & Rubin, 2004) for each parameter, which roughly equals the ratio of the between- and within-chain variances. A generally used rule of thumb by computational statisticians is implemented in the program namely that values less than 1.1 suggest convergence. The program checks this value for each model parameter and reports convergence only if all of them are less than 1.1. While in our experience this $\hat{R}$ criterion has been accurate for indicating convergence (or lack of it) for the set of GCM models in the program, sometimes it can be reassuring to display posterior sample chains of some random parameters, and check whether the chains cover more or less the same area. Such plots can easily be generated by the first panel’s Plot chain(s) button (see later). This is an especially useful technique if we suspect that some parameters might converge slowly. To visualize, we plotted three posterior sample chains in Figure 5. The Y-axis has the sampled values of the parameter in question, while the X-axis has the iterations. As we can see, in the first 100 iterations the chains are exploring different areas. Around the 100th iteration however, they converge into the same area, around 0.5. This latter part can be considered converged and the “fury caterpillar” is the pattern the user should look for when establishing convergence of the chains based on visual assessment. In case there is no convergence we can still look at the posterior samples to try to figure out that adding more samples (and optionally also raising the number of the Adaptation period) would increase our chances of convergence. If we see that the chains are more or less in the same region, adding more iterations in the Iterations window and rerunning the model may solve the issue. However, when the model
does not fit the data, for example because the one underlying culture assumption is not met, even very long chains might not reach convergence, as reported for example in Batchelder and Anders (2012).

![Figure 5](image)

*Figure 5.* Example of posterior sample chains converging towards a region near 0.5

The second standard concerns the autocorrelation in the sequence of samples obtained from the MCMC sampler. Unlike usual random sampling from a distribution that consists of independent draws from the distribution, an MCMC sampler draws correlated values from the posterior distribution, meaning that there is autocorrelation between successive samples due to the fact the sampler satisfies the properties of a Markov chain. As it is important to make sure that all regions of the posterior density are well represented, the regions of greater density should contribute more samples than sparser regions. If there is high autocorrelation in the sampled values (lag 1 autocorrelation; i.e., the correlation of one sampled value with the next), there is a risk that some regions are oversampled, as the sampler moves too slowly in the parameter space. In Figure 6 three sample chains with high autocorrelation are displayed. We can see that these chains do not mix as well as the ones after the 100th iteration in Figure 5.

Nevertheless, if sufficiently many samples are obtained from a MCMC sampler, a binned frequency distribution of values closely approximates the desired posterior distribution. This leads to a decision about ‘thinning’ the sample by retaining only every kth sample, which the user has to make in the bottom of the previous panel, as shown in Figure 3. Once the sampling is done, the second textbox of the top line in Figure 4 tells us about the autocorrelation. The program is set to the somewhat liberal level of 0.3 to decide about reporting a problem with the autocorrelations.
However, the program can return information on the autocorrelations in the samples of any of the parameters (see next paragraph), hence users more concerned about autocorrelation can always check these values and rerun the analysis with an increased number of thinning to reduce autocorrelation. That strategy is also advised for cases when this textbox reports autocorrelations higher than 0.3. As mentioned above, we can decrease autocorrelation by increasing thinning (see box called Thin in Figure 3, lower right corner). If the autocorrelation levels are only moderately higher than 0.3, increasing Thin to 5 should solve the issue. Larger autocorrelation values might require setting thinning to 50. Alternatively, instead of increasing thinning, one can add more chains.

(8) The middle panel of Figure 4 contains options to explore the results. First of all, all estimated parameters, indexed by persons or items when applicable, are loaded into a textbox. The user can make multiple selections of parameters of interest and then inspect the statistics of their posterior density. Then by clicking on the ‘Display posterior statistics’ under this selection, a new window will pop out showing some summary statistics of the posterior samples. An example of this can be seen in Figure 7.

Each line corresponds to the statistics of one of the model parameters named in the first column. The second and third columns show the posterior mean and median, which can be thought of as point estimates of the parameters. The fourth column displays the uncertainty in the parameter estimates in terms of posterior standard deviations. The fifth and sixth columns concern the spread of the posterior density by specifying its 2.5<sup>th</sup> and 97.5<sup>th</sup> percentiles, or in other words
the 95% Bayesian credible interval. Then the next column shows the exact $\hat{R}$ values on which the convergence check (see above) is based. Finally, the last column reports the level of autocorrelation. The results displayed in this table can be written out into a standard text file by clicking the button below the table. Looking at the first line for a competence parameter in Figure 7, we can see that a possible point estimate for the competence parameter of informant 001 is 0.93. Also, its 95% BCI lies above 0.7, so it is very likely that this informant’s competence level is above 0.7. For deriving how probably that is exactly, we go back to the main window, select our parameter of interest and push Plot cumulative distribution button. It then displays its cumulative posterior density curve, as it is shown in Figure 8. Since this plot is interactive, by pointing at the particular parameter value on the x-axis, the plot shows us its place on the curve and displays numerically how likely it is that the parameter value is lower than that value. For $D_1 < 0.7$ it shows 0.0065, so it is rather unlikely that this informant has low competence level.

The parameter selection in Figure 7 is only an example, the user can set all of the parameters or any subset of them. However, at the end of the displayed table, there is always some summary about informant and/or item specific parameters. More specifically, based on the point estimates (posterior means) of the competence (or ability), guessing bias and item difficulty (if applicable) parameters, summary statistics such as mean, median, standard deviation, and 2.5th and 97.5th percentiles are calculated. This way the user can have an idea about the average level of these parameters in the sample (mean, median), as well as about how much variability there is among
informants and items (standard deviation, percentiles). These quantities would be the usual sorts of statistics about the competence level of a group of informants that one would publish in a research paper. For example, we see that the mean competence level is 0.74 and the standard deviation is 0.11. In the case of the guessing bias, the mean is 0.55, which is close to the neutral guessing value of 0.50 assumed in earlier analysis of the contagion data by ANTHROPAC. However, the standard deviation of the guessing bias is 0.20, which suggests that there is a lot of heterogeneity in guessing bias in the group of informants. Certainly, one can look not only at the point estimates but the BCIs as well, and can calculate for example how many informants are most likely to have competency above 0.7, or how many of them will most likely always answer “False” etc. Moreover, although it is not displayed in the current example, ethnographers might also find it interesting to compare item difficulty levels for the different questions of the questionnaire, when the GCM with item difficulty is estimated. In a nutshell, BCCT provides a complex output with a potential to investigate various research questions.

The Plot density curve button opens a plot of the smoothed posterior distribution of the selected parameter. Finally the Plot chain(s) button generates plots showing all the sample chains for each of selected parameters.

As mentioned earlier, BCCT is a standalone software that does not require a MATLAB licence to run. However, if the user does have a MATLAB licence, BCCT can be run from MATLAB by using the supplementary files that can be downloaded from the same website as the standalone program. This way the user can investigate even more characteristics of the posterior sample chains.

3 For example, the user can explore whether the different person-specific parameters seem to be correlated. For that this script would be run: plotmatrix([reshape(samples.theta(:,:,1),[],1) reshape(samples.g(:,:,1),[],1) ], xlabel('person 1 ability'),ylabel('person 1 guessing'))
Checking model fit

The Perform check button of the lower panel (already pushed in Figure 4) carries out several goodness-of-fit tests. As the tests may take a few minutes, this option is not executed automatically, but only at the user’s request. The lower panel of Figure 4 shows the results of these checks for the example dataset.

Bayesian \textit{p}-value. The posterior \textit{p}-value reported in that box represents a global fit statistic of the current CCT model to the data. Calculation of this value is based on Bernoulli deviance functions (Karabatsos & Batchelder, 2003) of the raw data and replicated datasets based on the posterior distributions of the model parameters (set by a maximum 1000 samples in BCCT). As this statistic summarizes how likely it is that the current data, or data that are more extreme, would occur under the assumption that the model is correct, low \textit{p}-values (under 0.05) indicate a poor fit.

Posterior predictive check of the one underlying culture assumption. The usual rule of thumb used by ethnographers (Weller, 2007) for supporting the single culture assumption is that this ratio of the first and the second eigenvalues is above 3. This rule is actually problematic, as the ratio is very dependent on the overall level of competence on the group of informants. The posterior predictive check option for the single culture assumption in the BCCT software improves upon the classical eigenvalue test based on the recent work of Batchelder and Anders (2012). With BCCT we can investigate the posterior predictive distribution of the ratio of the first to second eigenvalues of several simulated datasets based on the posterior samples of model parameters. Then we can check whether the data eigenvalue ratio is within the 95\% posterior predictive distribution of the model based eigenvalue ratios. If not, BCCT returns an error message saying: ‘PPC failed’, otherwise it returns ‘PPC passed’. By pushing the button under the statistic we can see the smoothed posterior predictive distribution of these eigenvalues with the data’s eigenvalue ratio represented by a straight line, and this plot enables the user to exactly where the data eigenvalue ratio is with respect to the distribution of simulated values.

Model selection with Deviance Information Criterion. An experienced user might analyze the same dataset with several different GCM models which all pass the single culture posterior predictive assumption as well as the Bayesian \textit{p}-value indicates a reasonable model fit. Then the researcher is faced with the question of which model should they base their final conclusions. While the Bayesian \textit{p}-value and the eigen-ratio test provide goodness-of-fit tests in the absolute sense (the model either fits or does not), with the Deviance Information Criterion (DIC; Spiegelhalter, Best, Carlin, & van der Linde, 2002) one can evaluate the relative goodness of fit of different models. DIC takes into account two important features of the model: the complexity (based on the number of parameters) and the fit (measured by a deviance statistic). Naturally, more complex models fit better, therefore DIC penalizes based on the effective number of parameters in the model to arrive to a fair measure of fit. The model with the lowest DIC is supposed to be the best fitting one. For further discussion of DIC in the context of CCT modeling please consult Karabatsos and Batchelder (2003).
Model selection in general. Besides DIC, one can also consider more substantive arguments when it comes to model selection. Let us consider first the case of whether or not one should introduce heterogeneity in the competence parameter. Cultural Consensus Theory was generally based on the assumption that different people have different level of cultural knowledge. Therefore we recommend always keeping this parameter heterogenous. In theory however, such a model constraint might be useful when dealing with a group of cultural experts, or informants that are otherwise very homogeneous in their competence. Also it was this homogenous competence constraint that allowed Batchelder and Romney (1988, Table 5) to construct conservative power tables of the minimum number of informants needed to achieve pre-specified levels of confidence in the accuracy of the estimated answer key.

In CCT applications the guessing bias parameter has been fixed to a neutral level of $g = 0.50$ for all respondents in order to be able to fit the data in ANTHROPAC or UCINET. While this may be a good approximation, in signal detection studies (Macmillan & Creelman, 2005) the guessing bias found in data fits is not at a neutral level. For example, one can imagine situations where sizable differences in guessing bias are to be expected (e.g., individual differences in how a person responds when they do not know something or aspects of the wording of the particular items), in which case the assumption of homogeneity in this parameter might be untenable.

Finally, researchers might have constructed a survey with items of varying difficulty, in which case it is appropriate to allow for item heterogeneity in the model.

Saving results

(10) Finally results on the model parameters can be saved from the Summary of the requested results window as shown in Figure 7 by clicking the Write results to file button. This will result in an excel file with its first sheet the posterior statistics on the selected parameters and with its second sheet the group results.

Testing BCCT

Here we provide a simulation study to demonstrate the parameter recovery accuracy of BCCT. A hundred datasets – 25 informants and 50 items each – were generated based on the all heterogeneous model. ‘True’ parameter values were drawn from a uniform distribution between 0.1 and 0.99 for the ability, guessing bias and item difficulty parameters. The answer key was sampled from a Bernoulli distribution with probability 0.5. In the BCCT window the competence, guessing bias and item difficulty were set to “heterogeneous”. We checked convergence and autocorrelation in each analysis and they met our standards defined above.

To evaluate the parameter recovery in the simulation study, two measures were calculated based on the posterior samples of each run. First, we checked whether the 95% posterior BCIs of the estimates of abilities, guessing biases and item difficulties contained the simulated ‘true’ parameter values. We calculated the proportion of times when it was outside of this region over persons (for ability and guessing bias) and items (item difficulty) for each run. The first row of Table 1 shows these percentages averaged over the 100 runs. Second, we calculated average absolute deviation (AAD) between simulated and estimated values. That is to say that we took a posterior mean.

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4Right now such power tables can be constructed with BCCT in a timely manner only if BCCT is run from script.
Table 1: Results of the simulation study with BCCT.

<table>
<thead>
<tr>
<th></th>
<th>Ability</th>
<th>Guessing bias</th>
<th>Item difficulty</th>
</tr>
</thead>
<tbody>
<tr>
<td>‘True’ value outside 95% BCI</td>
<td>7.0 %</td>
<td>4.2 %</td>
<td>7.7 %</td>
</tr>
<tr>
<td>Average Absolute Difference</td>
<td>0.0993 (0.0191)</td>
<td>0.0886 (0.0151)</td>
<td>0.1223 (0.0159)</td>
</tr>
</tbody>
</table>

estimate for each parameter in each run, we calculated the absolute value of the deviation between this estimate and the simulated value, and we averaged it over persons or items. The second row of Table 1 displays the mean and the standard deviation of this value over the 100 runs.

Another measure of parameter recovery takes into account the uncertainty around the point estimates as well. As mentioned before, around each point estimate we can calculate Bayesian credible interval. Then we can set up different criteria of how large of that interval should contain the ‘true’ (simulated) parameter. A frequently applied approach is that the ‘true’ values should lie outside of the 95% credible interval about 5% of the time. The third row of Table 1 shows that how many percent of the estimates (averaged over respondents/items) lie outside of this 95% credible interval. As we can see, these values are quite close 5% for all three parameter classes.

With respect to the answer keys, on average over the 100 runs only 1.30 (std = 0.09) items were incorrectly recovered. These incorrectly recovered items were all rather difficult: their average item difficulty level was 0.84 (std = 0.09). As the average of the simulated ability parameters were only slightly above 0.5, these results are coherent with our expectations based on the simulation settings.

Conclusions

The BCCT software has been recently developed and although it has been thoroughly tested it might still run into unexpected errors. We encourage all users to send e-mails to the first author about problems as they arise, so that these issues can be mended. In turn updates for BCCT will be available for download.

We intend the current software package to be a starting point towards developing a large variety of user-friendly BCCT toolboxes for CCT models for questionnaires requiring other types of responses, e.g. Likert, ranking, matching, and continuous response scales. Based on feedback from users more complex BCCT packages will be written to enable researchers to investigate even more complex CCT modeling strategies, including hierarchical models and covariate analysis.

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References


