The bhsdtr package: a general purpose method of Bayesian inference for Signal Detection Theory models

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Abstract

We describe a novel method of Bayesian inference for hierarchical or non-hierarchical equal-variance normal Signal Detection Theory models with one or more criteria. The method is implemented as an open-source R package that uses the state-of-the-art platform, Stan, for sampling from posterior distributions. Our method can accommodate binary responses as well as additional ratings and an arbitrary number of nested or crossed random grouping factors associated with sensitivity or criteria effects. The SDT parameters can be regressed on additional predictors within the same model via intermediate unconstrained parameters, and the model can be extended by using automatically generated human-readable Stan code as a template. In the paper we explain how our method improves on other similar available methods, we give an overview of the package, demonstrate its ease of use by providing a real-study data analysis walk-through, and show that the model successfully recovers known parameter values when fitted to simulated data.

 $\it Keywords:$ Signal Detection Theory, rating experiments, Bayesian inference, hierarchical models

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Many tasks used in psychology studies are essentially classification tasks. For example, in a memory study participants may be required to decide if a given test item is old or new, or, in a perceptual study, an object may be either a letter or a digit. If a task requires classification, it is always possible that conclusions based on accuracy or percent correct are invalid because the ability to discriminate between stimulus classes (i.e., sensitivity) is confounded with bias, which is a tendency to classify stimuli as belonging to a particular class (Green & Swets, 1966). In principle, any effect that manifests itself in differences in classification accuracy may reflect differences in sensitivity, bias, or both. Signal Detection Theory provides a popular solution to this common problem.

However, because the SDT model is non-linear, variability in its parameters due to factors which are usually study-specific and of no direct interest to the researcher such as participants or items has to be accounted for. When they are not accounted for, e.g., by aggregating the data accros participants or stimuli, the estimates of SDT parameters are biased. As we explain later in this paper, none of the available methods of inference for hierarchical SDT models that we are aware of addresses this problem correctly in it's full generality, meaning that none of the available methods allows for fixed and random effects in both the sensitivity and the criteria parameters while restricting the parameters in accordance with SDT assumptions. Later in the paper we explain why the bhsdtr package for R (R Core Team, 2017), which we have made publicly available at https://github.com/boryspaulewicz/bhsdtr, provides a correct implementation of the general hierarchical linear regression structure defined on SDT parameters. The package repository also contains the annotated source code that was used to perform all the analyses and produce all the figures presented in this paper.

In what follows, after introducing the most common version of the SDT model, we describe its generalization, which can accommodate data from rating experiments.

Next, we explain briefly why, if a method of inference for SDT models were to be of

general use in psychology studies, it is essential that it is based on a model equipped with the hierarchical linear regression structure. The bhsdtr package meets this requirement thanks to a novel parametrization. We describe this novel parametrization and explain how reliance on some popular parametrizations leads to problems in the two other available implementations. We end the first part of this paper with a formal definition of the model as implemented in bhsdtr. The second part contains an overview of the package and a tutorial in which we demonstrate how to use our method in practice. Before we go any further, however, a note on terminology seems in order.

In the context of hierarchical modelling, factors such as participants, items, or replications are often referred to as groups. In our opinion this naming convention may be confusing; a single participant is both a group and a member of some group, while at the same time the term "group" is perhaps most strongly associated with study conditions, as in "experimental group". In this paper we use the term "sampled factor" instead because, by virtue of being a new term, it is unambiguous and seems descriptively correct: the term "sampled factor" seems to capture essential properties of such variables, i.e., a nominal scale, the fact that values are sampled from a larger population and are usually not of direct interest, as in "this is only a sample", and that conclusions of statistical analysis are meant to apply to the whole population of possible values.

Equal-variance normal SDT model with additional criteria

According to SDT, each stimulus in a classification task gives rise, by some unspecified cognitive process, to a unidimensional internal evidence value s sampled from a distribution that depends on the stimulus class. For historical reasons the two stimulus classes are often referred to as "noise" and "signal", and task performance is described in terms of hits, correct rejections, omissions, and false alarms, but this terminology is appropriate only when the model is applied to tasks that require detection, which is far from always being the case. In the most widely used version of the model, shown in Fig. 1, the two evidence distributions are normal with the same

variance, often fixed at unity to make the model identifiable. The distance d' between the means of the evidence distributions represents sensitivity. Because normal distributions are unbounded, s is always ambiguous, and so a criterion c placed on the evidence axis has to be used to reach a binary decision. A participant is assumed to decide that a stimulus belongs to the first class (e.g., an old item) if s < c, or that it belongs to the second class (e.g., a new item) if $s \ge c$. The location of the decision criterion may be interpreted in terms of bias.

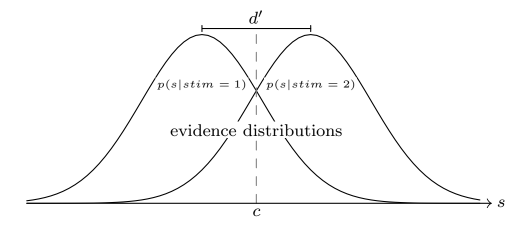


Figure 1. Equal-variance normal Signal Detection Theory model

Perhaps the simplest way of using this model is to fit it to observed response counts and use the estimated d' values in place of the percent correct scores; if the model is correct, the resulting performance measure is not contaminated by bias. The model may not be correct, which in our view is the most important reason to focus more on the generalized version shown in Fig. 2 below.

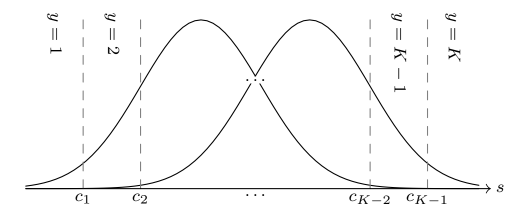


Figure 2. Equal-variance normal Signal Detection Theory model with additional criteria

This generalized model is applicable to studies in which participants are asked to rate their binary classification decisions on confidence or some other performance- or stimulus-related dimension. The ratings and the binary classification decisions can be provided together (e.g., "I am almost certain that it was a digit"), or in an arbitrary order.

Ratings are accommodated by introducing additional criteria and by modelling a combined response y, which represents both the binary classification decision and rating as a single number. The value of y increases with the strength of evidence in favor of the second stimulus class. For example, if confidence is rated on a four-point scale, then y = 1 when a subject decides that a stimulus belongs to the first class with certainty 4, and y = 5 when a subject decides that a stimulus belongs to the second class with certainty 1. More formally, if K is the number of possible combined responses, then a subject is assumed to give response k if $s \in (c_{k-1}, c_k]$, where c are the decision criteria, with c_0 and c_K fixed at $-\infty$ and $+\infty$ respectively.

There is a good reason to collect the ratings and use the generalized SDT model from Fig. 2, even when neither the ratings nor the placement of criteria are relevant to the research problem. When K=2 (no ratings), the SDT model fits perfectly, regardless of whether it is a reasonably good approximation to reality, because the data and the model have the same dimensionality. This makes the generalization to the K>2 case (i.e., decision with ratings) particularly important, as it is only when K>2 that the formal assumptions of the model (e.g., equal or unequal variance) can be tested¹, which is often done by comparing observed and predicted ROC curves.

When SDT models are used in psychology studies, researchers are usually interested not in the values of the SDT parameters themselves, but in relationships between these parameters and additional measured or manipulated variables; a good example is the dependence of d' on stimulus strength. Also, in a great majority of psychology studies in which classification tasks are used, the data have a hierarchical

¹In contrast to the formal assumptions, a psychological interpretation of the SDT parameters can be tested even when ratings are not available, e.g., by means of selective influence (Sternberg, 2001)

structure, i.e., there are repeated measures for each participant or item, and participants or items are only samples from the target population. A general-purpose method of inference for SDT models should accommodate both kinds of situations.

The importance of hierarchical regression structure

If data have a hierarchical structure, but variability due to subjects, items, or other sampled factors is not accounted for, estimates of average (fixed) effects are not guaranteed to be unbiased and conclusions are not guaranteed to generalize to the target population. The not uncommon practice of analyzing data aggregated over sampled factors represents an extreme case of ignoring hierarchical data structure. The invalidity of this approach in the context of SDT was clearly illustrated with the results of simulational studies by Morey, Pratte, and Rouder (2008); however, strictly speaking, such demonstrations are irrelevant to prooving the invalidity in question. Because SDT is a non-linear model, by definition, when estimates of its parameters are based on data aggregated over sampled factors (e.g., d' estimated for hits and false alarms averaged or summed over subjects), the expected values of these estimates (e.g., what the calculated d' actually estimates) are not in general true population averages (e.g., true d' in some condition). Such estimates are biased and inference about a target population based on them is not valid. To give a concrete example, consider two unbiased participants, one with $d'_1 = 2$ and one with $d'_2 = 4$. Their expected average accuracy is given by $(\Phi(d'_1/2) + \Phi(d'_2/2))/2 = .91$, which corresponds to d' = 2.68, whereas their true average d' is 3.

Aggregation is not the only way of ignoring a hierarchical data structure. Sometimes non-aggregated data are analyzed by using separate estimates for every participant × item × condition combination, but uncertainty due to distributions of subject or item effects is not accounted for by means of a hierarchical model structure. In such cases, conclusions – at least with respect to the uncertainties in estimates of population-average (fixed) effects are guaranteed to be valid only for the given sample, not the target population.

Another common source of problems is the separation of estimation of non-linear model parameters from regression analyses which aim to relate these parameters to measured or manipulated variables. When the SDT parameters are estimated separately for each subject and condition, and only later are these estimates regressed on predictors of interest, a number of additional issues may arise.

Firstly, the standard errors or credible intervals associated with the regression coefficients do not reflect the uncertainty in the SDT parameter estimates because the latter are treated as mere data points. The precision of parameter estimates often varies between participants, items, or conditions, but when the estimates are treated as data points, no use is made of this information. Secondly, regressing parameters on numeric predictors makes their estimates dependent on the common regression structure, and so also on each other, which can improve the quality of the estimates, just as assuming that random effects are samples from a common distribution may improve their estimates. The aforementioned problems can be dealt with by supplementing the SDT model with the hierarchical linear regression structure.

Comparison with other software implementations

Both d' and c have the virtue of being directly interpretable in terms of sensitivity and bias. However, both d' and c are constrained: d' is non-negative and, when there is more than one criterion, the elements of the c vector are order restricted ($c_{i+1} > c_i$). Hierarchical linear regression structure can only be defined on unconstrained parameters, because random effects are assumed to be normally distributed and normal distribution is unbounded and because fixed effects are represented by unconstrained parameters. We provide examples of the problems that may arise when the SDT model is not correctly reparametrized in the following summary of two hierarchical SDT implementations. One is the Gibbs sampler proposed by Morey et al. (2008) and the other is the Hierarchical Meta-d' model (HMeta-d') proposed by Fleming (2017). After describing the problems associated with those two software implementations we will briefly explain how our method compares to what the otherwise excellent brms package

has to offer.

The HMeta-d' model is a hierarchical version of the meta-d' model (Maniscalco & Lau, 2012), which in turn is a generalization of the SDT model that allows for a separate "meta-sensitivity" to account for possible discrepancies between a binary stimulus classification (referred to as a type 1 task) and the associated rating task (referred to as a type 2 or meta-cognitive task). We consider HMeta-d' here because it reduces to the SDT model with ratings when the type 1 and type 2 sensitivities are equal.

The Gibbs sampler created by Morey et al. (2008) allows for at most two sampled factors to have independent normally distributed random effects on the evidence distribution means. Unlike d', each evidence distribution mean considered in isolation is an unconstrained parameter, but the mean of the second evidence distribution is by definition greater than (d' > 0) or equal to (d' = 0) the mean of the first. The authors explicitly admit that their algorithm does not enforce this restriction because, as they say in the paper, it would greatly complicate analysis, although nothing more is said about these complications. At the same time the authors fail to mention that an immediate consequence of allowing for negative d' values is that the resulting posterior distribution loses its intended interpretation since it can have a non-zero mass when d' < 0. The outermost criteria are fixed at 0 and 1, and the ordering restriction is enforced by assuming that the likelihood is 0 whenever $c_{i+1} \leq c_i$. As the authors explain, because a sampled factor can have independent random effects on the evidence distribution means, it can have an effect on all the criteria: shifting both means by the same amount in the same direction is equivalent to keeping the sensitivity intact, while shifting the criteria relative to the evidence distributions. However, the elements of the criteria vector cannot be affected differently by the same sampled factor, which is an unrealistic restriction: participants differ in how they place the criteria just as they differ in their sensitivity.

In HMeta-d' the hierarchical structure is restricted to normally distributed random intercepts of one sampled factor. In the HMeta-d' model the d' parameter is allowed to assume negative values also, but the most problematic aspect of this

implementation is again the representation of the criteria. Each element of the criteria vector has an associated independent normal distribution, which does allow for criteria random effects, but does not enforce the necessary ordering restriction. The elements of this vector are sorted to obtain another criteria vector, and it is this sorted criteria vector that is used to model the conditional combined response distributions.

Consequently, the model does contain parameters representing the actual, order-restricted criteria, but, because sorting is not injective, the space of the actual criteria is only loosely related (i.e., not isomorphic) to the space of the unrestricted criteria vectors that are associated with the hierarchical structure. This makes posterior distributions of random criteria effects uninterpretable.

Some extensions of the SDT model can be fitted correctly using the excellent brms package, as described in Bürkner et al. (2017). The brms package is a flexible tool that shares some deep design similarities with our method. Both our package and the brms belong to a growing family of software tools that aim to provide a somewhat simplified and domain or application specific interface to one of the general purpose bayesian inference engines, in this case the stan modelling language (Carpenter et al., 2016). The brms is a highly flexible, well documented package that offers an elegant interface for fitting a large class of mixed models. Among the models that can be fitted using this package are the ordinal regression models. The hierarchical SDT model with ratings is essentially a generalization of mixed ordinal regression, since ratings are an ordinal-scale variable. However, that does not mean that a general hierarchical SDT model with ratings can be fitted using the brms package. There are three categorical distributions available at present in brms, i.e., the cumulative model, the adjacent category model and the sequential model, each with it's own set of link functions. As the author of the package explains (Bürkner et al., 2017), the only model that respects the ordering of the thresholds is the cumulative model. The thresholds in the cumulative model cannot be regressed on additional variables, consequently they cannot be associated with random effects.

Another important difference between our method and the brms package is that

brms does not utilize the fact that for ordinal regression models the data can often be significantly aggregated without loosing any information. For example, if a basic SDT model with ratings was to be fitted to an arbitrary large dataset using our method the aggregated dataset would consist of a matrix of dimension $2 \times K$. In that case each iteration of the sampling algorithm implemented in stan will consist of essentially 2 computations of the likelihood function, whereas without any aggregation step the number of such computations will be a linear function of the sample size. This is especially problematic for high-dimensional non-linear models such as hierarchical SDT with ratings, because they require relatively large datasets (i.e., many data points per participant \times condition).

Hierarchical Signal Detection Theory in an unconstrained parameter space

The general hierarchical linear regression structure can be defined on SDT parameters only if the latter are derived from unconstrained parameters. In the bhsdtr package, d' is derived from $\delta = \ln(d')$, thus random effects on d' can be modelled by assuming that δ is normally distributed. The problem of representing the criteria by unconstrained parameters is solved by mapping the R^{K-1} space of unconstrained criteria vectors to the K dimensional probability simplex space using the softmax function, and mapping the simplex space to the space of order-restricted criteria vectors by means of the inverse normal CDF:

$$c_i = \Phi^{-1}(\sum_{k=1}^i (e^{\gamma_k}) / \sum_{j=1}^K (e^{\gamma_j}))$$
 (1)

where Φ is the CDF of the standard normal distribution and $\gamma \in \mathbb{R}^K$, with γ_K fixed at 0 for identifiability. The idea is illustrated in Fig. 3 below:

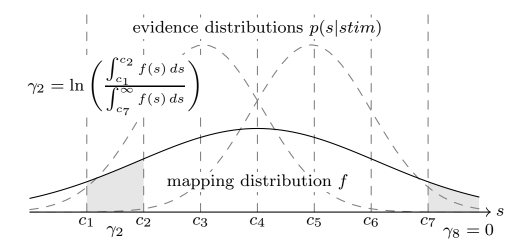


Figure 3. Mapping between the unconstrained γ vector and the criteria

Note that the normal distribution centered at the midpoint is merely a mapping device, not a third evidence distribution, and that, for the reasons that will be explained later, it is wider than the two evidence distributions. The mapping expressed by Eq. 1 is an isomorphism between the R^{K-1} space and the space of order-restricted criteria vectors. It's inverse is given by $\gamma_i = \ln \left(\int_{c_{i-1}}^{c_i} f(s) \, ds / \int_{c_{K-1}}^{\infty} f(s) \, ds \right)$, where f is the standard normal probability density function. The elements of the γ vector correspond to relative distances between pairs of adjacent criteria because their exponents represent the relative magnitudes of areas under the standard normal curve, delineated by the pairs of adjacent criteria: $e^{\gamma_i}/e^{\gamma_j} = (\Phi(c_i) - \Phi(c_{i-1}))/(\Phi(c_j) - \Phi(c_{j-1}))$. When K = 2, only γ_1 is free to vary, and its value directly represents the direction and magnitude of bias: γ_1 is 0 when the criterion is placed at the midpoint between the evidence distributions; the more negative (positive) γ_1 is, the more the criterion is shifted to the left (right) of the midpoint.

It is often a good idea to multiply all the criteria by a value greater than 1, which is equivalent to making the mapping distribution wider. This tends to even out values of γ by preventing the outermost areas under the mapping distribution curve from becoming very small relative to area delineated by adjacent pairs of non-outermost criteria. This is especially important when the criteria are widely spread, as can happen for moderate to large d' values. This feature is implemented in the **bhsdtr** package by introducing a criteria scaling factor.

Once d' and c are derived from the unconstrained δ and γ parameters, the SDT model can be supplemented with a hierarchical linear regression structure. To avoid having to deal with an even more complicated index notation², below we present only the simple case of one sampled factor.

$$oldsymbol{\delta} = oldsymbol{X}^{(\delta)}oldsymbol{eta}^{(\delta)} + oldsymbol{Z}^{(\delta)}oldsymbol{ heta}^{(\delta)} + oldsymbol{Z}^{(\delta)}oldsymbol{ heta}^{(\delta)} = e^{oldsymbol{\delta}_i} \ oldsymbol{\gamma}_{i,\cdot} = oldsymbol{X}_{i,\cdot}^{(\gamma)}oldsymbol{eta}^{(\gamma)} + oldsymbol{Z}_{i,\cdot}^{(\gamma)}oldsymbol{ heta}^{(\gamma)}oldsymbol{ heta}^{(\gamma)} \ oldsymbol{c}_{i,k} = s \ \Phi^{-1}(\sum_{l=1}^k (e^{\gamma_{i,l}})/\sum_{m=1}^K (e^{\gamma_{i,m}})) \ oldsymbol{p}(oldsymbol{y}_i = k|oldsymbol{stim}_i = 1) = \Phi(oldsymbol{c}_{i,k} + oldsymbol{d}_i'/2) - \Phi(oldsymbol{c}_{i,k-1} + oldsymbol{d}_i'/2) \ oldsymbol{p}(oldsymbol{y}_i = k|oldsymbol{stim}_i = 2) = \Phi(oldsymbol{c}_{i,k} - oldsymbol{d}_i'/2) - \Phi(oldsymbol{c}_{i,k-1} - oldsymbol{d}_i'/2) \ oldsymbol{p}(oldsymbol{y}_i = k|oldsymbol{stim}_i = 2) = \Phi(oldsymbol{c}_{i,k} - oldsymbol{d}_i'/2) - \Phi(oldsymbol{c}_{i,k-1} - oldsymbol{d}_i'/2) \ oldsymbol{e}(oldsymbol{c}_{i,k-1} - o$$

Here i=1...N is the observation number, \boldsymbol{X} is the fixed effects model matrix for the respective parameter, \boldsymbol{Z} is the random effects model matrix, $\boldsymbol{\beta}$ and $\boldsymbol{\theta}$ are the fixed and random effects, \boldsymbol{c} is an $N \times K - 1$ matrix, s is the criteria scaling factor, and \boldsymbol{y} is the combined response. Note that \boldsymbol{d}'_i is a scalar, but $\boldsymbol{\gamma}_{i,\cdot}$ is in general a vector, and so $\boldsymbol{\beta}^{(\gamma)}$ and $\boldsymbol{\theta}^{(\gamma)}$ are matrices. The j-th rows of the $\boldsymbol{\beta}^{(\gamma)}$ and $\boldsymbol{\theta}^{(\gamma)}$ matrices represent fixed and random effects on the j-th element of the $\boldsymbol{\gamma}$ vector.

Following Sorensen and Vasishth (2015) we make use of the Cholesky decomposition of the correlation matrices because it improves efficiency and admits a convenient prior on random effects correlations:

²The reader familiar with hierarchical models may be surprised by our use of superscript parenthesized Greek letters to express hierarchical relationships. We chose this convention because it allowed us to use subscripts to denote elements of vectors and matrices while minimizing the number of nested sub- or superscripts.

$$\begin{aligned} \text{vectorized}(\boldsymbol{\theta}^{(\gamma)}) &= \text{diag}(\boldsymbol{\tau}^{(\gamma)}) \boldsymbol{L}^{(\gamma)} \boldsymbol{z}^{(\gamma)} \\ \boldsymbol{\theta}^{(\delta)} &= \text{diag}(\boldsymbol{\tau}^{(\delta)}) \boldsymbol{L}^{(\delta)} \boldsymbol{z}^{(\delta)} \\ \boldsymbol{z}_i^{(\delta)} &\sim \text{Normal}(0, 1) \\ \boldsymbol{z}_i^{(\gamma)} &\sim \text{Normal}(0, 1) \end{aligned}$$

where each τ is a vector of standard deviations of random effects and each L is a Cholesky decomposition of a random effects correlation matrix, i.e., C = LL'. Thus, θ is multivariate normal with covariance matrix $\operatorname{diag}(\tau)L$.

Finally, we use weakly informative proper priors because they provide regularization and help stabilize computation. The fixed effects $\boldsymbol{\beta}^{(\delta)}$ and $\boldsymbol{\beta}^{(\gamma)}$ are given independent normal priors, the random effects standard deviations $\boldsymbol{\tau}^{(\delta)}$ and $\boldsymbol{\tau}^{(\gamma)}$ are given independent half-Cauchy priors, as recommended by Gelman (2004), and each \boldsymbol{L} is given an independent lkj prior:

$$\begin{split} \boldsymbol{\beta}_{i}^{(\delta)} &\sim \operatorname{Normal}(\boldsymbol{\mu}_{i}^{(\delta)}, \boldsymbol{\sigma}_{i}^{(\delta)}) \\ \boldsymbol{\beta}_{k,l}^{(\gamma)} &\sim \operatorname{Normal}(\boldsymbol{\mu}_{k,l}^{(\gamma)}, \boldsymbol{\sigma}_{k,l}^{(\gamma)}) \\ \boldsymbol{\tau}_{i}^{(\delta)} &\sim \operatorname{half-Cauchy}(0, \boldsymbol{\zeta}_{i}^{(\delta)}) \\ \boldsymbol{\tau}_{k,l}^{(\gamma)} &\sim \operatorname{half-Cauchy}(0, \boldsymbol{\zeta}_{k,l}^{(\gamma)}) \\ \boldsymbol{L}^{(\delta)} &\sim \operatorname{lkj}(\boldsymbol{\nu}^{(\delta)}) \\ \boldsymbol{L}^{(\gamma)} &\sim \operatorname{lkj}(\boldsymbol{\nu}^{(\gamma)}) \end{split}$$

Specifying the prior distributions

A Bayesian model is not complete without providing fixed values of all the parameters that define prior distributions. Specifying the priors on sensitivity effects does not pose any special difficulties. The sensitivity of an unbiased classifier given percent correct (pc) is given by $2\Phi^{-1}(pc)$. When p(stim = 1) = 0.5, the greater the

bias, the lower the accuracy, meaning that an unbiased sensitivity is a lower bound on sensitivity given percent correct. Let us assume that the majority of subjects are expected to achieve percent correct within the .51 to .99 range, with negligible bias. Since $\ln(2\Phi^{-1}(.51)) = -2.99$ and $\ln(2\Phi^{-1}(.99)) = 1.54$, a reasonable weakly informative prior on δ is normal with mean (1.54 - 2.99)/2 and standard deviation (1.54 + 2.99)/2, which is the default prior on delta effects in the bhsdtr package.

Specifying the priors on criteria effects can be challenging because the criteria are order-restricted and the complexity of the mapping expressed by Eq. 1 makes it difficult to reason about priors on γ in terms of criteria effects. By default, in the **bhsdtr** package each entry in the $\sigma^{(\gamma)}$ and $\zeta^{(\gamma)}$ matrices is set to $\ln(100)$ and the criteria scaling factor is fixed at 2.

The prior on random effects standard deviations is parametrized by ζ , which represents half-width at half-maximum of the half-Cauchy distribution. In our opinion, a not-unreasonable starting point is to set ζ at the value that is greater or equal to the most likely value of the random effects standard deviation.

Finally, by default $\nu^{(\delta)} = \nu^{(\gamma)} = 1$, which implies a uniform prior on random effects correlation matrices. Because the greater the value of ν , the more emphasis is put on zero off-diagonal correlations, the researcher can force the correlations to be near-zero by choosing a large ν value.

Overview of the software implementation

The bhsdtr package implements the model in the Stan modelling language because the stan sampler uses a state-of-the-art adaptive Hamiltonian Monte Carlo algorithm which often handles high-dimensional correlated posteriors better than a Gibbs sampler.

Our package is essentially a collection of documented functions: The aggregate_responses function aggregates data as much as possible for efficiency, but without distorting the hierarchical structure. The make_stan_model function creates a model definition in the Stan language. The Stan code produced by the

make_stan_model function can be fitted as is or modified by the user if needed, e.g., to change the prior distributions or to drop the equal variance assumption. The make_stan_data function creates regression model matrices and other data structures required by the model created using the make_stan_model function. Finally, the plot_sdt_fit function can be used to visually assess the fit of the model by creating publication-ready ROC curve plots or response distribution plots with posterior predictive intervals calculated for the chosen α level.

Usage example: installing the package and testing the model on real data

To make full use of the bhsdtr package functionality, three non-standard R packages are required, namely rstan, plyr, and ggplot2. We recommend using the devtools package to install the bhsdtr package directly from the github repository. This will automatically install the package and any missing required packages:

devtools::install_git('git://github.com/boryspaulewicz/bhsdtr')
library(bhsdtr)

The essential steps of a typical data analysis process will usually involve preparing the data, creating the model code, fitting the model, assessing the fit, and possibly converting the unconstrained δ and γ parameters to d' and c.

Preparing the data. The bhsdtr package contains a dataset, gabor, from an unpublished study in which on each trial the participants had to classify a briefly presented Gabor patch as tilted to the left or to the right using the arrow keys. The participants were also asked to rate the stimuli on a 4-point Perceptual Awareness Scale (Ramsøy & Overgaard, 2004) presented at the bottom of the screen. The Gabor patch was immediately followed by a mask. The PAS ratings ranged from "no experience" to "absolutely clear image" and were provided either before (RD order condition) or after (DR order condition) the arrow keys were pressed. On each trial the Gabor patch was equally likely to be presented for 32 ms or 64 ms. Order was a between-subject variable and duration was a within-subject variable. There were 47 participants and 48 trials per condition.

In the study in question, the response was originally encoded using separate variables for accuracy and rating, so the first step was to create an appropriate response variable using the combined_response function. This function requires three variables, one encoding the stimulus class, one encoding the rating (as an integer), and one binary variable encoding the decision accuracy.

This step is required only if the ratings are available and a combined response variable is not already present in the data. In the single criterion case, the combined response variable is simply the binary classification decision. To fit a single-criterion SDT model to this dataset, the code above would have to be replaced with the following:

Next, the data has to be aggregated using the aggregate_responses function, but only to an extent that preserves all the random effects. This function requires as arguments a data frame containing all the relevant variables, the name of the stimulus class variable, the name of the combined response variable, and the vector of the names of all the variables that are to be preserved in the resulting aggregated dataset (apart from the stimulus class variable and the combined response variable), i.e., those encoding the sampled factors and those representing the independent variables used in the regression part of the model:

The main purpose of the aggregation step is to improve the efficiency of sampling from the posterior distribution. When data are aggregated in this way, the likelihood for each condition \times participant combination has to be computed only once rather than as many times as there are trials per condition per participant. Note that if there are other

sampled factors present in the data (e.g., items, replications, etc.) and the user decided to model the effects of these factors, then these factors also have to be specified at this stage to preserve the hierarchical data structure. The aggregate_responses function creates a list with three components. The data component is a data frame containing additional preserved variables, the stimulus component is the stimulus class variable, and the counts component is an $N \times K$ matrix of combined response counts, where N is the number of data points and K is the number of possible combined response values.

Creating the model code. A model is fitted using the stan function from the rstan package. This function requires a special list of data structures used by the model as well as a model specification expressed in the Stan language.

Every model has some fixed effects structure since, even when there are no predictors, the model parameters can be expressed as regressed on a vector of ones (i.e., an intercept). However, many models also have a hierarchical structure and, if that is the case, this hierarchical structure has to be specified when using the make_stan_model function. This is done by providing a list of lists of R model formulae. Each list of model formulae is composed of at least three elements and specifies the correlated random effects of one sampled factor. The group element specifies the sampled factor; the delta and gamma elements specify which effects are assumed to vary between the levels of this sampled factor. When make_stan_model is used without any arguments, it specifies a model without any random effects. Fixed effects model matrices are specified by providing a list with at least two model formulae, named delta and gamma, to the make_stan_data function that is described later in this paper. Non-default priors can be specified by adding optional elements to the random and fixed effects specification lists, as described in the make_stan_data function documentation.

In the study in question there was only one sampled factor, i.e., the participants. Because duration was a within-participant variable, in principle its effect could vary between the participants for all the SDT parameters. However, a preliminary data analysis indicated that the 32 ms difference in duration seemed to affect only the sensitivity. Thus, it was assumed that δ may depend on duration and order, but γ may

only be affected by order. Because duration was a within-subjects variable, its effect on δ was assumed to vary between the participants, but the only random effect associated with γ was the intercept.

The make_stan_data function creates fixed and random effect model matrices based on the respective model formulae using dummy contrast coding. Note that the implicit intercept was removed for the δ model matrix (the -1 term on the right hand side of the model formula). In this way, δ was estimated for every duration \times order condition. The resulting separate intercepts and slopes parametrization makes it easier to calculate arbitrary contrasts on posterior samples. A more standard parametrization was used for the γ parameter because it was initially assumed that the criteria depend only on order, which is a two-level factor, and so there was only one contrast of interest for every element of the γ vector. On the other hand, even in such cases the parametrization with separate intercepts and slopes for every condition may be more convenient if a researcher is interested in the actual criteria, as we will later explain when introducing the gamma_to_crit function. This example also illustrates how the separation of the δ and γ regression structures makes it possible to test a broad class of linear models representing the dependence of the SDT parameters on additional variables.

Fitting the model. In order to fit the model, a separate data structure used by the Stan sampler has to be created using the make_stan_data function. The obligatory arguments to this function are an aggregated data object created by the aggregate_responses function and a fixed effects specification. Importantly, if random effects are modelled, the same specification of random effects has to be provided to the

make_stan_model and make_stan_data functions.

```
sdata = make_stan_data(adata, fixed, random)
```

Finally, a vector of names of parameters of interest has to be specified when calling the stan function:

Note that since more than one sampled factor is allowed, the names of all the hierarchical parameters are indexed (e.g., delta_sd_1, delta_random_1, Corr_delta_1). The name counts_new refers to posterior predictive samples that are required by the plot_sdt_fit function. Names starting with Corr refer to random effects correlation matrices.

Assessing the model fit. As can be seen in the previous code fragment four chains of 8,000 iterations each were run simultaneously; the first half of the posterior samples, which served as a warm-up period for tuning the parameters of the sampling algorithm, was discarded. Part of the resulting Stan output is presented in Table 1 below.

Table 1 $Model \ fit \ summary \ statistics$

	mean	SE_{mean}	SD	2.5%	97.5%	No. eff. samples	\hat{R}
delta_fixed[1]	-0.10	0.00	0.15	-0.41	0.18	4626	1.00
$delta_fixed[2]$	1.12	0.00	0.09	0.95	1.29	4764	1.00
$delta_fixed[3]$	-0.39	0.00	0.20	-0.80	-0.03	5427	1.00
$delta_fixed[4]$	1.28	0.00	0.11	1.06	1.50	5464	1.00
$\operatorname{gamma_fixed}[1,\!1]$	-0.14	0.00	0.06	-0.27	-0.02	2925	1.00
${\rm gamma_fixed}[1,\!2]$	-0.22	0.00	0.10	-0.42	-0.01	8199	1.00
${\rm gamma_fixed}[2,\!1]$	-0.70	0.00	0.18	-1.05	-0.35	3060	1.00
${\rm gamma_fixed}[2,\!2]$	0.49	0.00	0.29	-0.06	1.04	3452	1.00
${\tt gamma_fixed[3,1]}$	-0.54	0.00	0.22	-0.96	-0.11	3103	1.00
${\rm gamma_fixed}[3,\!2]$	0.83	0.01	0.35	0.14	1.51	3086	1.00
${\rm gamma_fixed[4,1]}$	0.28	0.00	0.25	-0.20	0.76	3090	1.00
${\rm gamma_fixed}[4,\!2]$	0.42	0.01	0.40	-0.37	1.22	3291	1.00
${\rm gamma_fixed}[5,\!1]$	-0.21	0.01	0.30	-0.80	0.37	3440	1.00
${\rm gamma_fixed}[5,\!2]$	0.83	0.01	0.48	-0.10	1.78	3637	1.00
${\rm gamma_fixed[6,1]}$	-0.77	0.00	0.24	-1.23	-0.31	3378	1.00
${\rm gamma_fixed}[6,\!2]$	0.80	0.01	0.38	0.05	1.56	3316	1.00
${\rm gamma_fixed[7,1]}$	-0.32	0.00	0.16	-0.64	0.01	3346	1.00
${\rm gamma_fixed}[7,\!2]$	0.42	0.00	0.28	-0.11	0.98	3333	1.00
${\rm delta_sd_1[1]}$	0.66	0.00	0.11	0.48	0.90	6373	1.00
$delta_sd_1[2]$	0.44	0.00	0.05	0.35	0.56	5463	1.00
${\rm gamma_sd_1[1]}$	0.16	0.00	0.08	0.02	0.31	2071	1.00
$gamma_sd_1[2]$	0.82	0.00	0.11	0.62	1.06	5852	1.00
${\rm gamma_sd_1[3]}$	1.08	0.00	0.12	0.86	1.33	4293	1.00
$gamma_sd_1[4]$	1.27	0.00	0.14	1.03	1.57	3723	1.00
$gamma_sd_1[5]$	1.55	0.00	0.17	1.24	1.91	4297	1.00

Given the complexity of the model the chains exhibited good mixing and seemed to have converged; there were enough effective posterior samples for the fixed effect parameters to estimate 95% credible intervals well and none of the Gelman-Rubin \hat{R} statistics crossed the conventional 1.01 threshold, suggesting negligible sensitivity to the initial values.

Figs. 4 and 5 below contain normal quantile-quantile plots of random effects. The plots indicate that the distributions of random δ and γ effects can be approximated by normal distributions and that – at least in this particular example – these parameters seem to be good candidates for representing variability in the sensitivity and criteria parameters due to the sampled factors.

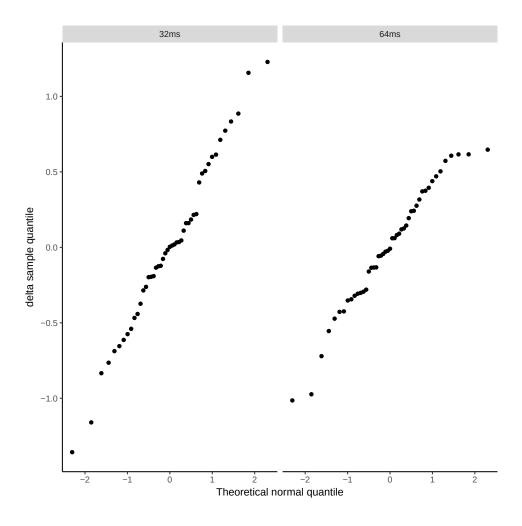


Figure 4. Normal quantile-quantile plots of δ random effects

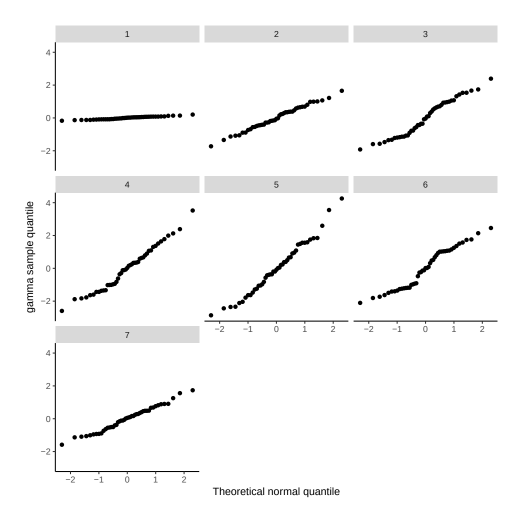


Figure 5. Normal quantile-quantile plots of γ random effects

Once enough good quality posterior samples are obtained for the parameters of interest, the inference process can be carried out by calculating credible intervals or HPD intervals for any function of the parameters or Bayes factors for each parameter separately. However, even when the stan output summary does not indicate sampler convergence issues, before drawing any further conclusions the researcher should first check if the model fits the data. The plot_sdt_fit function can be used for this purpose:

This function requires at least three arguments: a stanfit object, an aggregated data list produced by the aggregate_responses function that was used to produce the stanfit object, and a vector of names of variables that will determine how the data will

be partitioned before plotting. We recommend assessing the fit at the individual level by including the participant identification number in the list of conditioning variables, but we did not do it here because the resulting plot would take up too much space.

As can be seen in Fig. 6, which shows the ROC curves produced by the code above, the model seemed to fit the data well in all but one condition (decision after rating, 64 ms duration), in which three out of seven relevant³ points were outside the two-dimensional 95% posterior predictive regions.

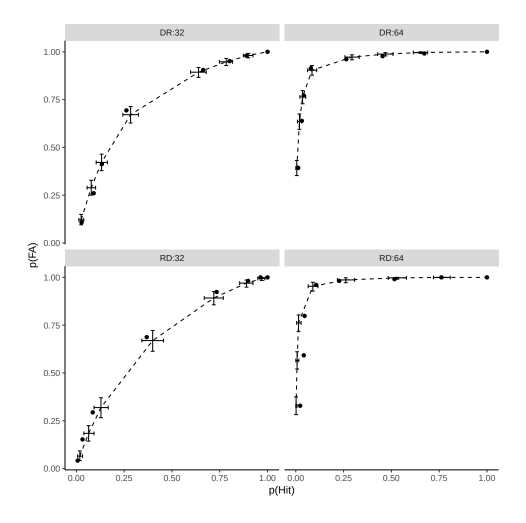


Figure 6. ROC curve fit

Another way to assess model fit visually is by inspecting the conditional response distributions (p(y|stim)), such as those shown in Fig. 7, which was also created using the plot_sdt_fit function.

³The point in the upper-right corner of a ROC curve is always in the (1,1) position

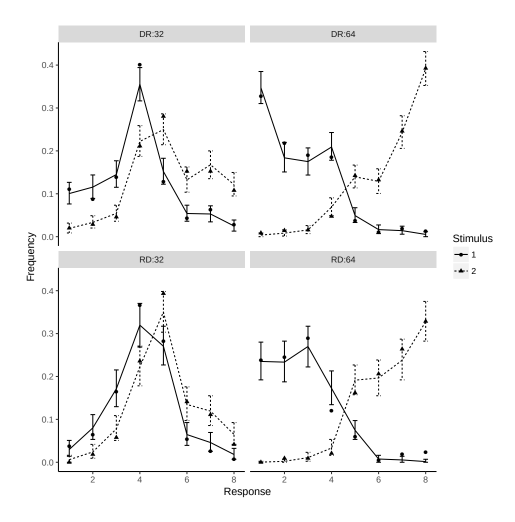


Figure 7. Response distribution fit

Both plots can be informative about the reasons why a model does not fit the data. In this particular case, the plot seems to suggest that it may be a good idea to inspect the fit at the individual level and see if there are some participants with unusual p(y|stim=1) distributions in the RD × 64 ms condition. On the other hand, it is also possible that the lack of fit is mainly a consequence of the assumption that duration had zero effect on γ , or that more substantial modifications are necessary, such as dropping the equal variance assumption.

Converting unconstrained δ and γ parameters to sensitivities and criteria. Posterior δ and γ samples have to be transformed in order to work with the d' and c parameters. This is straightforward only when fixed effects represent average parameter values in separate conditions, not when they represent differences between conditions or regression slopes. In our example, because separate intercepts and slopes

parametrization was used for the δ fixed effects model matrix, all four delta_fixed parameters represent condition averages and can easily be transformed to sensitivities by applying the exponential function. It is important to remember that because this is a non-linear transformation, the δ to d' conversion step should be done first before applying any other transformations to the posterior samples; for example, the exponential of a point and interval estimate of δ is not equal to the point and interval estimate calculated after first transforming the δ posterior samples to the d' samples.

In this case the first column of the gamma_fixed parameter matrix (the intercept) corresponds to the values of the γ vector in the DR condition, but the second column corresponds to the effect of order on γ . For this reason, in this particular case the posterior criteria samples can be obtained using the gamma_to_crit function only for the first column of the gamma_fixed matrix. That's because the second column represents the difference in γ between conditions and a nonlinear transformation from γ to c will not give a correct difference in c.

Testing the model on simulated data

We simulated the data from a hypothetical exact replication of the previously described experiment using the point estimates from the previous fit as known realistic parameter values. Mixing performance was similar to the real data case. All the model parameters were correctly recovered in a sense that the true values were outside the 95% credible intervals no more than 5% of the time.

As we have repeatedly stressed the models that lack the necessary hierarchical structure may easily show reliable effects where none exist or fail to detect true differences. To illustrate this problem, an SDT model that differed from the true model only in that it did not have any hierarchical structure was fitted to the same simulated dataset. Tring to obtain general theoretical results on the typical extent of bias in models that ignore the hierarchical structure would be a serious undertaking, but even this simple simulational example shows how misleading can the results of such analyses be. Since the non-hierarchical model was much simpler and the data consisted of only

eight vectors of response counts, the mixing of the chains was excellent.

The 95% credible intervals calculated for the fixed effects based on each model are compared in Fig. 8 below. The estimates were centered on the true values to simplify the presentation. As can be seen, the true model correctly recovered the known parameter values, but the estimates based on the simplified, non-hierarchical model were severely biased; the credible intervals were not only about three times shorter on average, but also failed to contain most of the true values.

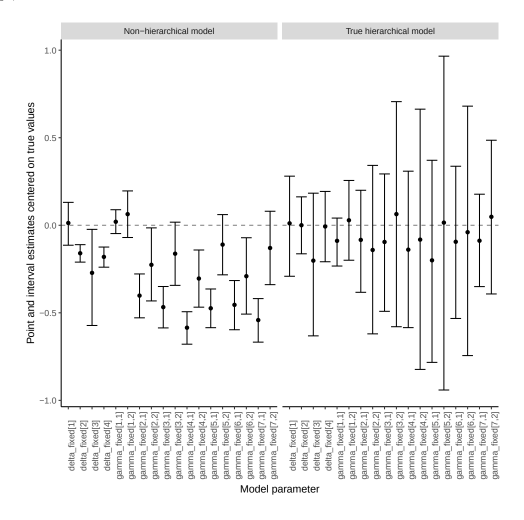


Figure 8. Comparison of the point and interval posterior estimates based on the true hierarchical and the simplified non-hierarchical models

To avoid bias, the hierarchical structure present in the data can be reduced or ignored only if the fit of the full hierarchical model indicates that the random effect variances or correlations are likely to be equal to zero. Importantly, this rule is valid regardless of the apparent fit of the model, as illustrated by the contents of Fig. 9

below; even though the simplified model provided strongly biased results the observed ROC curves seemed to fit the model's predictions quite well, giving a false impression of model validity.

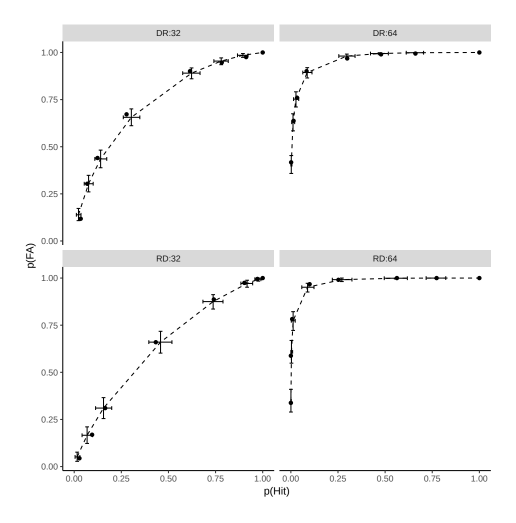


Figure 9. ROC curve fit for the non-hierarchical model

Concluding remarks

The importance of SDT to psychology stems from the fact that given weak assumptions about an underlying decision process, it promises to deconfound sensitivity from bias in arbitrary binary classification tasks - a problem almost as common in psychology studies as the usage of classification tasks. To the best of our knowledge, at present the bhsdtr package provides the only method of Bayesian inference for SDT models with or without ratings that can be recommended as a default choice in typical applications. Our parametrization forces the sensitivity to be non-negative and the

criteria to be order-restricted, while the isomorphisms between the d' and c parameters and the unconstrained δ and γ parameters make it possible to supplement the SDT model with the general hierarchical linear regression structure. There is no limit to the number of sampled factors except for the one imposed by available computational resources; correlations of random effects of the same sampled factor are accounted for, all the SDT parameters can be modelled by linear regression within the same model, and all the effects on all the SDT parameters estimable within the levels of the sampled factors can have associated random effects. If the need arises to relax a built-in restriction, experienced users can extend the model in arbitrary ways by using automatically generated human-readable Stan code as a template.

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