THEORETICAL NOTE

Factoring Out Nondecision Time in Choice Reaction Time Data: Theory and Implications

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Choice reaction time (RT) experiments are an invaluable tool in psychology and neuroscience. A common assumption is that the total choice response time is the sum of a decision and a nondecision part (time spent on perceptual and motor processes). While the decision part is typically modeled very carefully (commonly with diffusion models), a simple and ad hoc distribution (mostly uniform) is assumed for the nondecision component. Nevertheless, it has been shown that the misspecification of the nondecision time can severely distort the decision model parameter estimates. In this article, we propose an alternative approach to the estimation of choice RT models that elegantly bypasses the specification of the nondecision time distribution by means of an unconventional convolution of data and decision model distributions (hence called the D’M approach). Once the decision model parameters have been estimated, it is possible to compute a nonparametric estimate of the nondecision time distribution. The technique is tested on simulated data, and is shown to systematically remove traditional estimation bias related to misspecified nondecision time, even for a relatively small number of observations. The shape of the actual underlying nondecision time distribution can also be recovered. Next, the D’M approach is applied to a selection of existing diffusion model application articles. For all of these studies, substantial quantitative differences with the original analyses are found. For one study, these differences radically alter its final conclusions, underlining the importance of our approach. Additionally, we find that strongly right skewed nondecision time distributions are not at all uncommon.

Keywords: choice RT, diffusion model, bias, nondecision time, misspecification, D’M

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In the last decade, the use of evidence accumulation models for choice reaction time (RT) experiments has revealed fundamental insights into the process of elementary decision making (Basten, Biele, Heekeren, & Fiebach, 2010; DasGupta, Ferreira, & Miesenböck, 2014; Forstmann et al., 2008; Krajbich & Rangel, 2011; Polanía, Krajbich, Grueschow, & Ruff, 2014; Ratcliff & Dongen, 2011; Resulaj, Kiani, Wolpert, & Shadlen, 2009). A common assumption in all these studies, is that choice RT can be additively decomposed into a decision time (taken up by the process responsible for the actual choice) and a residual nondecision time. The latter is considered to be the contribution of encoding and motor response execution (Luce, 1986). In contrast to the intricate evidence accumulation or diffusion models for the decision part of choice RT (Brown & Heathcote, 2008; Ratcliff & Rouder, 1998; Usher & McClelland, 2001; Verdonck & Tuerlinckx, 2014; Wong & Wang, 2006), the models traditionally used for the nondecision time are surprisingly basic: adding a simple time constant, or, at best, adding a simple parametric distribution (mostly uniform, see Ratcliff & Tuerlinckx, 2002). However, choosing the wrong model for the nondecision time is known to be a cause for bias for the decision model’s parameter estimates, especially when data originating from a skewed nondecision time distribution are fitted under the assumption of a nonskewed nondecision time distribution (Ratcliff, 2013). Seeing a (right) skewed nondecision time distribution is at least as plausible as the current nonskewed default. Therefore, today’s golden standard for the nondecision time may be systematically delivering biased results for the decision process parameters.

In this article, we develop a novel estimation approach that bypasses the specification of the nondecision time distribution and allows us to estimate the decision model’s parameters without any nondecision time misspecification bias. Moreover, if desired, a nonparametric estimate of the residual nondecision time can be estimated. Attempts to disentangle nondecision from decision time have been proposed before (Smith, 1990), but from a different perspective: the nondecision time distribution was assumed to coincide with...
with the distribution of simple (i.e., one-choice) response times, observed in a comparable experimental setup. This distribution was then deconvolved from the distribution of total choice response times (from the original two-choice version of the experiment) to isolate the decision part. The resulting distributions could then be modeled with a decision model. The approach we propose in this article, does not use information of secondary experiments nor does it presume any particular parametric distribution (shape) for the nondecision time distribution.

In what follows we will first explain the method, then illustrate its performance based on simulated data, and finally use it to reanalyze data from three different diffusion model application articles.

The D’M Method

Assume a two-choice RT experiment with \( I \) different conditions \((i = 1, \ldots, I)\). These conditions can be stimuli of different difficulty (e.g., percentages coherently moving dots in a random dots motion task) but also other manipulations (e.g., prior expectations). During the course of the experiment, there are several trials within a condition, and each trial can result in either a correct (e.g., percentages coherently moving dots in a random dots motion task) but also other manipulations (e.g., prior expectations). During the course of the experiment, there are several trials within a condition, and each trial can result in either a correct (e.g., percentages coherently moving dots in a random dots motion task) but also other manipulations (e.g., prior expectations).

When replacing \( f_p \) and \( f_{p'} \) by \( \hat{g}_p \) and \( \hat{g}_{p'} \), the equality of Equation 1 is no longer exact. However, the discrepancy (or objective function)

\[
D_{pp'}(\theta) = d(\hat{g}_p * m_p(\theta), \hat{g}_{p'} * m_p(\theta)),
\]

where \( d(\cdot,\cdot) \) is a distance defined on the space of defective pdfs, should be small for \( \theta = \theta_0 \). Note that in Equation 2, a convolution appears between the data from condition-response pair \( p' \), represented by the smoothed estimated density \( \hat{g}_{p'} \), and the model’s density for condition-response pair \( p \), which is \( m_p(\theta) \). Therefore, we refer to our technique as the D’M method. To evaluate this crucial convolution both model and data densities are discretized using an equally spaced grid and the convolution integral is then approximated by a finite sum over this grid.

In this article we opt for a \( \chi^2 \) type of distance between the compared distributions

\[
d(a, b) = \int_0^\infty \frac{(a(t) - b(t))^2}{a(t) + b(t)} dt.
\]

For a conventional \( \chi^2 \) distance, where typically an observed distribution is compared with a model pdf, the integrand’s denominator is the pdf of the model. Because in Equation 2, both compared distributions are a convolution of a data and a model pdf, we take the sum of the two compared distributions for the integrand’s denominator in Equation 3. This distance is called the triangular discrimination (Topsoe, 2000) and can be considered as a symmetrized version of the \( \chi^2 \) distance (a “\( \chi^2 \) like distance”; Le Cam, 1986).

It is obvious that for \( \hat{g}_{p} \rightarrow K * f_p \) and \( \hat{g}_{p'} \rightarrow K * f_{p'} \), \( D_{pp'}(\theta) \) reaches its minimal value of zero, when \( \theta = \theta_0 \). Because of Equation 1, in Appendix A it is shown that if the decision model does not contain a nondecision component of its own and the total response time pdfs \( f_p \) underlying the data are not all equal across condition-response pairs, this minimum is unique, at least for the parameters pertaining to \( m_p \) and \( m_{p'} \). In this sense, \( D_{pp'}(\theta) \) can be used to estimate the parameters pertaining to \( m_p \) and \( m_{p'} \) without explicitly solving the underlying inverse convolution problem leading to an estimate of \( r(t) \). Because we are interested in the parameters pertaining to all condition-response pairs, we use the total objective function.

\[
\hat{t}(\theta) = \sum_{\theta} \left( \sum_{p} D_{pp'}(\theta) \right),
\]

where the sum runs over all possible unique combinations of two
different condition-response pairs. The estimated parameter vector \( \hat{\theta} \) is then found as: \( \hat{\theta} = \text{argmin} T(\theta) \).

If the true parameter vector \( \theta_0 \) is the absolute and unique minimum of all separate terms \( D_{ij} \) (at least for all parameters pertaining to \( m_p \) and \( m_{dp} \)), the same will hold for the sum \( T(\theta) \) (for all parameters). For \( \theta_0 \), the nondecision pdf \( r(t) \) clearly exists and can be retrieved as the deconvolution of each condition-response pair \( p \)'s total response time and decision pdfs:

\[
r = f_p \ast m_p(\theta_0)^{-1},
\]

where \(-1 \) in the exponent refers to the deconvolution operation.

However, for an arbitrary parameter vector \( \theta \), it is not sure that these deconvolutions exist. Conditions for the existence of a deconvolution are described in Appendix B and it is shown that, up to a practical degree of accuracy when dealing with noisy data, they boil down to the following condition on the variances \( \sigma^2 \):

\[
\sigma^2_{rp} \geq \sigma^2_{mp} + \sigma^2_{K},
\]

basically avoiding a negative variance for \( r(t) \). Because we use a smoothing kernel \( K \) to estimate \( f_p \), the kernel needs to be taken into account and effectively we will use the equivalent constraint:

\[
\sigma^2_{K} \geq \sigma^2_{mp} + \sigma^2_{K},
\]

where the last step follows from the additive property of variances of independent random variables. In terms of the data, the constraint is:

\[
\sigma^2_{rp} \geq \sigma^2_{mp} + \sigma^2_{K}.
\]

This condition has to be met for all condition-response pairs \( p \). Although these conditions have to be met for all condition-response pairs \( p \), the validity of each of these constraints is only as good as the estimate \( \hat{\sigma}^{2}_{K} \). For this reason, densities \( \hat{g}_p(t) \) based on only a few data points should not be included. If these constraints are imposed while minimizing \( T(\theta) \), the existence of a common \( r(t) \) for all pairs \( p \) is ensured. For the final estimation of \( r(t) \), we use the following expression:

\[
r = \hat{r} = \left( \frac{1}{2} \sum_{p=1}^{2l} m_p(\theta_0)^{-1} \right)^{-1}
\]

where both sums run over all \( 2l \) pairs \( p \). Having obtained an estimate \( \hat{\theta} \) of the true parameter values \( \theta_0 \), this results in an estimator

\[
\hat{r} = \left( \frac{1}{2} \sum_{p=1}^{2l} \hat{g}_p \right)^{-1} \left( \frac{1}{2} \sum_{p=1}^{2l} K \ast m_p(\hat{\theta}) \right)^{-1},
\]

where the same smoothing kernel \( K \) is used for both factors of the deconvolution. The constraints in Equation 7 ensure the existence of this deconvolution. If the deconvolution exists, then clearly for \( \hat{g}_p \rightarrow K \ast f_p \), and, therefore, \( \hat{\theta} \rightarrow \theta_0 \), \( \hat{r}(t) \rightarrow r(t) \). Note that the smoothing kernel \( K \) used in this final step can differ from the kernel used in the minimization procedure, but has to be applied in a recalculation of \( \hat{g}_p(t) \) as well (for the deconvolution, we will use a uniform distribution from 0 to 0.01 for \( K \)). In practice, we solve the deconvolution problem in Equation 9 by defining a grid (we take a grid spacing of 0.01s) and minimizing the Kullback–Leibler divergence of the model-based distribution \( r \ast \left( \sum_{p=1}^{2l} K \ast m_p(\hat{\theta}) \right) \) from the data-based distribution \( \left( \sum_{p=1}^{2l} \hat{g}_p \right) \) with respect to the nondecision time distribution probability weights assigned to the grid points. We use the same global optimizer as before to tackle this high dimensional minimization problem: for a grid spacing of 0.01s and \( \hat{r} \) clipped at 1.5s, \( \hat{r} \) consists of 150 grid points whose weights have to be estimated. Using the triangular discrimination distance instead of the Kullback-Leibler divergence results in similar estimates of \( \hat{r} \).

In the theory developed above, we have assumed that all \( 2l \) condition-response pairs share the same nondecision time distribution \( r \). Note however that the total objective function \( T(\theta) \) in Equation 4 can be easily changed to allow for multiple nondecision time distributions, each shared within a subset (with two elements or more) of condition-response pairs. This setup is the analogue of a traditional diffusion model analysis in which the nondecision time parameter \( T_r \) is allowed to vary across (some of the) conditions, while the other parameters are constrained to be equal. To implement such a situation, it suffices to limit the double sum in Equation 4 to terms produced within subsets of condition-response pairs with the same nondecision time distribution. Each nondecision time distribution can then be estimated separately by limiting the sums in Equation 9 to the subset of condition-response pairs corresponding to that nondecision time distribution.

### Performance on Simulated Data

In this section we show how well the D’M technique recovers a standard parameter set of the Ratcliff diffusion model (Ratcliff & Tuerlinckx, 2002), in conjunction with different nondecision time distributions. The quality of the estimates is systematically compared with the results obtained with a standard approach in which it is assumed that the nondecision time is uniformly distributed.

#### Data Simulation

We simulated data from a typical Ratcliff diffusion model parameter set (see Table 2) in conjunction with three different nondecision pdfs (see the right-hand side figures of the three panels in Figure 1): one right skewed, one uniform and one bimodal distribution. For all three parameter configurations, we look at data sets of 300, 1,000 and 1,000,000 RTs per stimulus. For each combination of sample size and nondecision time distribution, we simulated 100 data sets.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( a )</td>
<td>Boundary separation</td>
<td>.08</td>
</tr>
<tr>
<td>( \eta )</td>
<td>Inter-trial variability of drift rate</td>
<td>.08</td>
</tr>
<tr>
<td>( zr )</td>
<td>Relative bias</td>
<td>.5</td>
</tr>
<tr>
<td>( sz )</td>
<td>Uniform inter-trial variability of bias ( z = a \cdot zr )</td>
<td>.02</td>
</tr>
<tr>
<td>( v_i ) (i = 1, ..., 4)</td>
<td>Drift rates</td>
<td>[.4, .25, .1, 0]</td>
</tr>
</tbody>
</table>
Standard and D’M Estimation Procedures

For the standard estimation procedure we used both the standard maximum likelihood method and the quantile likelihood method.

(a) right skewed nondecision pdf

(b) uniform nondecision pdf

(c) bimodal nondecision pdf

Figure 1. In the left panels, the recovery of the Ratcliff diffusion model’s boundary separation parameter \(a\) is shown, for data simulated with either a right skewed nondecision pdf (a), a uniform nondecision pdf (b), or a bimodal combination of both (c). Estimates are shown for data sets with subsequently \(N = 10^6\), \(N = 10^3\), and \(N = 300\) observations per condition. The orange (light gray) box plots show the distributions of the estimates obtained with the traditional method, the blue (dark gray) box plots those obtained with the D’M method. (The recovery results for the other diffusion model’s parameters yield very similar conclusions and are offered as supplemental material.) In the right panels, we show for each nondecision time density, the average of the nonparametric estimates based on the D’M decision model parameter estimates as given by Equation 9. The average is represented by solid lines varying from light to dark with the increasing number of observations and compared with the original shape (dashed black line). See the online article for the color version of this figure.

(Heathcote & Brown, 2004; Heathcote, Brown, & Mewhort, 2002). As the results are very comparable, we only show results of the latter. As is customary, a uniform distribution for the nondecision time distribution was used and the location and spread of this uniform distribution was estimated alongside the pure decision model parameters.

In the D’M procedure no nondecision pdf has to (nor should) be included. To evaluate the convolution integral, we choose a grid ranging from 0 to 5 s (well beyond any observed RT) with equally spaced nodes at every 0.01 s.

For both methods (standard and D’M), the model density \(m(\theta)\) needs to be calculated. To do this, we use code from the fast-dm project (Voss & Voss, 2007). Also in both methods, an objective function has to be minimized (the negative quantile likelihood for the standard method and Equation 4 for D’M). To find this minimum, we use a global optimizer (i.e., differential evolution; Storn & Price, 1997).

Results

Figure 1 shows the estimates for the boundary separation \(a\) (results for the other parameters are offered as supplemental material), in conjunction with three different shapes of nondecision time densities. The traditional approach results in a systematic estimation bias, except for the uniform nondecision pdf, in which case the nondecision model is perfectly specified for the data. This systematic bias is resolved with the D’M method, which gives better estimates (or comparable in the unlikely case that the actual nondecision time density is a uniform distribution), even for a number of data points as low as 300 per condition. In addition, the average recovered nondecision time densities match remarkably well with the true nondecision time densities.

As an additional test, we have repeated this analysis for different values of boundary separation \(a\). Increasing values of \(a\) correspond to a larger proportion of the total RT (and its variance) being accounted for by the decision model (keeping everything else constant). The results are presented in Figure 2. For the right skewed nondecision time distribution, the D’M method always outperforms (or matches) the traditional method, regardless of the value of \(a\) or the number of observations. For the uniform and bimodal nondecision time distributions, D’M still performs better or equal for normal and high values of \(a\), but can have higher biases for small values of \(a\), if there is a limited number of observations. This is what one would expect: As the proportion of total response time variance generated by the nondecision time process gets larger (the nondecision time distribution proportions of total variance at \(a = 0.04\) are 0.91, 0.47 and 0.87, for right-skewed, uniform and bimodal, respectively), the decision part becomes increasingly obfuscated by the nondecision part. As the D’M method only uses a parametric model for the decision part, more observations are needed to correctly disentangle the two components. Thus, if (a) the uniform distribution is a good enough proxy for the actual nondecision time distribution, (b) the actual nondecision time distribution contributes enough to the total response time, and (c) there is not a lot of data, the traditional method can outperform the D’M method. It has to be noted, however, that unless we have some prior information about the real shape of the nondecision time distribution, the first condition can never be checked in a real world problem. If one wants
to avoid results that depend on any particular presupposed shape of the nondecision time distribution, the D*M approach is by definition the better choice. Figure 3 shows the nonparametric estimates of the nondecision time distributions following the D*M estimates shown in Figure 2. Only for the smallest boundary separation $a = 0.04$ and a limited number of observations, the nonparametric estimates of the nondecision time distribution is somewhat biased, but even then the main features of the distributions are recovered.

For the estimation of the decision model parameters, the repeated calculation of the complete decision model pdfs on a reasonably detailed grid is by far the most computationally

Figure 2. Estimation biases of boundary separation $a$, for varying $a$. The same nondecision time distributions were used as in Figure 1: a right skewed nondecision pdf (a), a uniform nondecision pdf (b), or a bimodal combination of both (c). Orange (light gray) lines correspond to traditional estimates, blue (dark gray) lines to D*M estimates. The estimates are based on data sets with subsequently $N = 10^2$ (solid lines), $N = 10^3$ (dashed lines), and $N = 300$ (dotted-dashed lines) observations per condition. If the number of trials is high (solid lines), the D*M method always outperforms (or matches) the traditional method, no matter what the value is of $a$. For less data (dashed lines, dotted-dashed lines), D*M still performs better for normal and high values of $a$. For a small amount of data and small values of $a$, however, and if the nondecision time distribution is more in line with the traditional uniform assumption like it is in (b) and (c), the D*M method shows a higher bias than the traditional method. Similar patterns are found for the other parameters. See the online article for the color version of this figure.

Figure 3. Recovery of the respective nondecision time distributions based on data sets generated with varying boundary separation $a$, with $N = 300$ (left) and $N = 1,000$ (right) observations per condition. Red (gray) lines indicate the nondecision time distributions obtained with the lowest value of boundary separation $a = 0.04$; the rest of the nondecision time distributions (with $a$ somewhere between 0.06 and 0.12) are in black. See the online article for the color version of this figure.
expensive part for both the traditional and the D*M method. In our implementation, a single evaluation of the D*M or traditional objective function takes up a comparable amount of time. We opted for a global minimization routine to reduce potential problems concerning local minima, but this resulted in optimization times that are probably a lot slower than strictly necessary. Running single-threaded on an i7 core clocked at 3.60 GHz, both a single D*M estimate and traditional estimate take around 5 min to complete. The estimation of the nondecision time (once D*M parameter estimates have been obtained) takes about 30 min for a grid with 0.01 s spacing but only about 15 s for a grid with 0.05 s spacing. To be confident we reached convergence, we repeated every estimation multiple times (with a different population of starting values). The D*M estimates seemed somewhat more robust than the traditional estimates: for the D*M objective function almost every repetition resulted in the same global minimum; for the traditional objective function, ending up in a local minimum was a bit more frequent. By repeating the minimization procedure multiple times (5 times for D*M and 10 times for the traditional method), we repeatedly found a lowest minimum and were convinced of convergence. An alternative for the global optimizer may be a local optimizer in combination with a rational starting point (e.g., using EZ diffusion, Wagenmakers, Maas, & Grasman, 2007). Such a routine may yield equally good results and will be much faster. However, because of the novelty of the D*M method, we were more concerned with accuracy (i.e., avoiding local optima) than with speed.

Existing Diffusion Model Analyses Revisited

To illustrate how the D*M parameter estimation method can lead to fundamentally different findings than the traditional method, we reanalyze the data of three choice RT studies in which a diffusion model is used. These articles can be seen as typical examples of today’s common practice of using diffusion models parameters to explain differences between (groups of) participants and/or experimental conditions. We only present the main results; more detailed results are offered as supplemental material. The first two panels show the estimates of a selection of decision model parameters, obtained with either the traditional (a) or D*M method (b). In each of the two panels, the darker box plots show the estimates for the elevated prior likelihood condition, the lighter plots for the larger potential pay-off condition. If for the group of participants Δσ or Δν is significantly different from 0 (two-sided sign test), this is indicated with a green (gray; p < .001) or hatched green (hatched gray; p < .01) marking of the label. Panel (c) shows the nondecision time densities inferred from the D*M estimates in Panel (b). The densities for the elevated likelihood condition are shown in the upper half of the plot and those for the larger potential pay-off condition are shown, mirrored, in the lower half. The solid lines show the mean nondecision pdfs across participants, the lighter areas display the double SE interval. Panel (d) is a quantile-quantile plot of the data in Panel (c), and is better suited to look at the differences between the nondecision pdfs from the two conditions. The gray area represents a 95% confidence interval of the mean quantile-quantile values (black crosses). See the online article for the color version of this figure.

Application 1: A Diffusion Model Analysis of Choice Bias (Mulder, Wagenmakers, Ratcliff, Boekel, & Forstmann, 2012)

Mulder et al. (2012) investigate, using a moving-dot perceptual decision making experiment, how different types of pretrial information biases people toward one of two choice alternatives. One type of pretrial information concerns the elevated prior likelihood of the occurrence of one of two choices, the other type involves a larger potential pay-off for one of the two choices. The diffusion model is used to study which decision process aspects are affected by each form of pretrial information (elevated prior likelihood or larger pay-off, each requiring a separate diffusion model analysis). One possibility is that the bias is caused by a shift Δν in the diffusion model’s drift rate parameter, the other possibility is that the bias is caused by a shift Δσ in the starting position parameter. The results are shown in Figure 4.

The authors concluded that it is mainly the starting position parameter that is responsible for the bias introduced by both elevated prior likelihood and larger potential pay-off conditions. Upon reanalysis we come to the same qualitative conclusion, for both the traditional and D*M procedure. Quantitatively, however, there are clear differences between the D*M and the traditional estimates. From Figure 4, it can be seen that the boundary separation parameter a is systematically lower for the D*M estimates and the resulting between-person variance (as indicated by the box plots) is considerably smaller than that of the traditional estimates. Clearly, the assumption of a uniform nondecision time density is not valid; instead, the nondecision time density estimates indicate a strong right skew. However, in this case, this misspecification does not yield a different qualitative conclusion.
Application 2: A Diffusion Model Analysis of Posterror Slowing (Dutilh et al., 2011)

A well documented observation in choice RT experiments is that the RT increases on a trial immediately after an incorrect choice. This phenomenon is called posterror slowing (PES). Dutilh et al. (2011) have attempted to isolate the diffusion model parameters that can account for the differences between postcorrect and posterror trials of a lexical-decision task. Participants have to classify a string of letters as a word or nonword. Besides the obvious word versus nonword manipulation, there are six different word frequency types. In their diffusion model analyses, the authors estimate a separate nondecision parameter $T_{cr}$ for every stimulus condition. Analogously, the D*M method is set-up to allow for a separate nondecision time distribution for each stimulus condition (grouping choice RT distributions that share a nondecision time distribution) as explained in the last paragraph of the theory section. The results are shown in Figure 5. Based on their analysis, which involves the estimation of two separate diffusion models (one for the posterror and one for the postcorrect condition), the authors concluded that posterror slowing is very much associated with an increase in response caution. A similar analysis with the D*M method, however, shows no association between PES and response caution, but reveals other associations, namely with $\eta_w$ and most word drift rates $v_i$ ($i = 2, \ldots , 6$). More specifically, after an error, the drift rates for words become smaller (i.e., closer to zero) and the trial-to-trial variability of word drift rates also decreases. The reason for the difference between the original and the D*M analysis lies with the specification of the nondecision pdf. The authors allow a different mean nondecision time for all stimuli, but assume an equal, uniform width. As can be seen in Figure 5, Panel d, the D*M method suggests both mean and variance of the nondecision time increase after an error trial when judging a word stimulus (shown in black in Panel d in Figure 5), but the nondecision time does not change after an error trial when judging a nonword stimulus (shown by the red [gray] dashed line in Panel d in Figure 5). These effects could not be accommodated by the author’s particular specification of the nondecision time distribution, so other parameters had to compensate, with different results and conclusions as a consequence. Based on our analysis, we have to conclude that in the context of this particular diffusion model, there is a fundamental difference in the processing of word and nonword stimuli after an error trial.

As for all our reanalyses, we rigorously implemented the model assumptions of the original article. However, it is tempting to wonder if just allowing the width of the uniform nondecision time distribution to vary across stimuli in the traditional approach, would be sufficient to get traditional results comparable to the D*M analysis. To answer this question, we performed the extra analysis and a similar picture as Figure 5, Panel a was obtained (see Supplemental Material), showing that the absence of skew in the nondecision time distribution specification is an essential component of the misspecification. As a possibly viable traditional alternative to our method, one could now suggest to have mean, skew and width as parameters of some new nondecision specification, but this would mean estimating 35 parameters (21 nondecision parameters on top of the 14 decision model parameters), while D*M can handle the problem with only the 14 decision model parameters. Furthermore, there is no guarantee that this would be sufficient to get traditional results comparable to the D*M version of Panel (a). Panel (c) shows the participant averaged nondecision time densities inferred from the D*M estimates, separately for all six different word types and nonwords ($v_{w1}, \ldots , v_6$ and $v_{NW}$). Statistically significant effects for the differences (two-sided sign test) are indicated with a green/red (light gray/dark gray) marking (opaque for $p < 0.001$, hatched for $p < 0.01$) of the label: Green (light gray) means a positive effect or a larger value posterior compared to postcorrect, red (dark gray) a negative effect or a smaller value posterior compared to postcorrect. Panel (b) is the D*M version of Panel (a). Panel (c) shows the participant averaged nondecision time densities inferred from the D*M estimates, separately for all six different word types (shades of gray) and nonwords (red [gray] and dashed). The nondecision time densities of the posterror condition are shown in the upper half, those of the postcorrect condition are shown, mirrored, in the lower half. Panel (d) is a quantile-quantile plot of the data in Panel (c), and is better suited to look at the differences between the nondecision time densities from the two conditions. (Because of the many nondecision time pdfs, no confidence intervals are shown in Panels (c) and (d).) See the online article for the color version of this figure.
alternative parametric nondecision time distribution will (always) suffice.

**Application 3: A Diffusion Model Analysis of Task Switching Costs (Schmitz & Voss, 2012)**

Schmitz and Voss (2012) investigate which diffusion model parameters can best explain task-switching costs. We limit our analyses to the first experiment in the article, which compares task-switching and task-repeating trials from a classical alternating runs paradigm to each other and to pure task trials (no task-switching within an experimental block). In these analyses, three separate diffusion models are estimated (with both methods): one for the pure task trials, one for the task-repeating trails and one for the task-switching trials. Partial results are shown in Figure 6. The main qualitative findings of the original article pertaining to this experiment, namely a positive change in boundary separation \(a\) (or caution) and a negative change in drift rate between task-switching and pure task trials, were confirmed by our reanalysis, for both the traditional and the D'M method. Participants are more cautious in the task switch condition and at the same time, they process the information less well. Quantitatively, the parameter estimates again differ considerably from the original study. It can also be seen that the nondecision time distribution in the task switching condition has a much more outspoken skew to the right (compared with the pure task nondecision time pdf).

**Discussion**

Traditional parameter estimates of decision models to choice RT data have been shown to be vulnerable to the misspecification of the extra nondecision component (Ratcliff, 2013). In this article, we have proposed a solution to this problem. By means of a handy convolution between data and decision model distributions, hence called the D'M method, we were able to factor out the nondecision time distribution from the estimation procedure. Through theory and a simulation study, the method was shown to remediate the traditional bias related to the misspecification of the nondecision time distribution, even for a limited number of data points. Additionally, the actually simulated nondecision time distributions could systematically be recovered. To illustrate the method's relevance, we applied it to three existing diffusion model application articles. For all studies, we found substantial differences with the traditional parameter estimates; in one case, using the D'M method radically altered the conclusions of the original article, clearly demonstrating the necessity of the method.

In two of the three applications we reanalyzed, nonparametric estimates of the nondecision time distributions revealed a clear right skew (Application 1 and 2). In our simulation study we have shown that ignoring such a right skew may cause severe biases in the diffusion model parameter estimates. Therefore, in the absence of a good model for nondecision time, we strongly advise the use of the D'M method for estimating choice RT models. Additionally, the nonparametric estimates of the resulting nondecision time distributions give a detailed picture of what nondecision time can look like (assuming the decision model itself is well-specified that is), which could in turn be used as a guide or template for constructing more intricate models for nondecision time.

**References**


Luce, R. D. (1986). *Response times: Their role in inferring elementary mental organization*. New York, NY: Oxford University Press.


Appendix A

Unique Global Optimum

In this appendix, we show under which conditions the objective function $D_{pp}(\theta)$ of Equation 2, for $g_p \to K * f_p$ and $g_{p'} \to K * f_{p'}$, has a unique global optimum at the true parameter vector $\theta_0$. We will start from the reasonable assumption that the decision model does not contain a nondecision component of its own at the true parameter vector $\theta_0$, or, more specifically, that for the true parameter vector $\theta_0$, there exists no distribution $u(t)$ and parameter vector $\theta_0 \neq \theta_0$ such that for every $p$:

$$m_p(\theta_0) = m_p(\theta_0) * u. \quad (10)$$

Imagine for a moment that such a $u(t)$ and $\theta_0 \neq \theta_0$ do exist. The fundamental Equation 1 may then be transformed as (for every $p, p'$):

$$f_p * m_p(\theta_0) = f_{p'} * m_{p'}(\theta_0)$$
$$f_p * m_p(\theta_0) * u = f_{p'} * m_{p'}(\theta_0) * u$$
$$f_p * m_p(\theta_0) = f_{p'} * m_{p'}(\theta_0)$$
$$m_p(\theta_0) * r * m_{p'}(\theta_0) = m_{p'}(\theta_0) * r * m_p(\theta_0)$$
$$m_p(\theta_0) * m_{p'}(\theta_0) = m_{p'}(\theta_0) * m_p(\theta_0),$$

where we make use of the fact convolution is a commutative operator and that all factors have a Laplace transform (see below in Section A.2 for more information) such that this identity can be safely deconvolved from both sides. Thus, the final identity is a direct consequence of the presence of a nondecision component inside the decision model at $\theta_0$. Note that another way this final identity can be reached is when for all $p, p'$: $m_p(\theta_0) = m_{p'}(\theta_0)$, and, as a consequence of Equation 10, $m_p(\theta_0) = m_{p'}(\theta_0)$. This would mean that all total response time pdfs $f_p$ would be equal across condition-response pairs. Now we can move to the actual proposition.

Proposition

If the decision model itself does not contain a common nondecision component that can be factored out, or more specifically, if there does not exist a parameter $\theta_0 \neq \theta_0$ such that for every $p, p'$:

$$m_p(\theta_0) * m_{p'}(\theta_0) = m_p(\theta_0) * m_{p'}(\theta_0), \quad (11)$$

the minimum of

$$D_{pp}(\theta) = d(\hat{g}_p * m_p(\theta), \hat{g}_{p'} * m_{p'}(\theta)),$$

for $g_p \to K * f_p$ and $g_{p'} \to K * f_{p'}$ at $\theta = \theta_0$ is unique.

Proof

Assume there does exist another global minimum at $\theta_0 \neq \theta_0$ for which

$$D_{pp}(\theta) = d(\hat{g}_p * m_p(\theta), \hat{g}_{p'} * m_{p'}(\theta)) = 0$$

for $g_p \to K * f_p$ and $g_{p'} \to K * f_{p'}$.

Then, for every $p, p'$

$$K * f_p * m_p(\theta_0) = K * f_{p'} * m_{p'}(\theta_0)$$
$$K * r * m_p(\theta_0) * m_{p'}(\theta_0) = K * r * m_{p'}(\theta_0) * m_p(\theta_0)$$

Because every factor of this equation has a Laplace transform (see below in Appendix B for more information), we can safely deconvolve $K$ and $r$ from both sides:

$$m_p(\theta_0) * m_{p'}(\theta_0) = m_{p'}(\theta_0) * m_p(\theta_0) \quad (12)$$

This is in contradiction with the initial assumption of Equation 11, which states that there is no such parameter $\theta_0 \neq \theta_0$. This proves the proposition.

(Appendices continue)
To formulate conditions for the existence of the deconvolution in Equation 5, we make use of cumulant-generating functions. The cumulant-generating function \( [G(d)](s) \) of a probability density function \( d(t) \) is defined as

\[
[G(d)](s) = \log \left( \int_0^\infty e^{st}d(t)dt \right).
\]

Thus, the cumulant-generating function is the logarithm of the moment generating function. Because the range of integration is the positive half of the real line, we can write the cumulant-generating function also as the logarithm of the Laplace transform (Kreyszig, 2010). It follows that for a probability density function \( d(t) \) that is piecewise continuous and of exponential order, that is, \( d(t) \leq Me^{-at} \) with \( M \) and \( r \) real positive numbers, the cumulant generating function \( [G(d)](s) \) exists, at least for \( s < k \). When we do not need the argument \( s \), we will also write the cumulant-generating function in short as \( G(d) \).

The cumulant-generating function \( G(d) \) is determined by the coefficients \( \kappa_n^d (n > 0) \) of its MacLaurin expansion, and these are called the cumulants:

\[
[G(d)](s) = \sum_{n=1}^\infty \frac{\kappa_n^d}{n!} s^n.
\]

Note that \( \kappa_2^d \) is the variance of \( d(t) \), denoted as \( \sigma_d^2 \) in the main text.

The cumulant-generating function is a convenient tool when dealing with convolutions because it treats them additively. Consider another pdf \( e(t) \), with cumulant generating function \( G(e) \), then

\[
G(d \ast e) = G(d) + G(e).
\]

Obviously, the additivity of the cumulant-generating function is transferred to the cumulant coefficients:

\[
\kappa_n^{d \ast e} = \kappa_n^d + \kappa_n^e,
\]

that holds for all \( n (n > 0) \).

In our approach, we assume that for both the model \( m_j(r; \theta_0) \) and the nondecision time distribution \( r(t) \), the corresponding cumulant-generating function exists. Most choice RT models are diffusion models and have an exponential decay (Noble, Ricciardi, & Sacerdote, 1985), thereby satisfying the exponential order condition. In these cases, \( m_j(r; \theta_0) \) always has a corresponding cumulant-generating function \( [G(m)](s; \theta_0) \) or \( G(m; \theta_0) \) short. For the unknown nondecision time distribution \( r(t) \) we will simply assume the existence of a cumulant generating function \( G(r) \) implying the existence of a cumulant-generating function \( G(f_r) \) as well.

Because the cumulants, and therefore, also the cumulant-generating function, can be expressed in terms of moments, sufficient conditions for a valid cumulant-generating function can be seen as sufficient conditions for a set of moments to lead to a valid pdf. The latter conditions, for a nonnegative random variable, are a well known set of inequalities (Cressie & Holland, 1983; Karlin & Studden, 1966). More specifically, given a set of \( J + 1 \) numbers \( \mu = (\mu_0, \mu_1, \ldots, \mu_J) \), if

\[
\Delta_{2k} = \det \begin{bmatrix} \mu_0 & \mu_1 & \cdots & \mu_k \\ \mu_1 & \mu_2 & \cdots & \mu_{k+1} \\ \vdots & \vdots & \ddots & \vdots \\ \mu_k & \mu_{k+1} & \cdots & \mu_{2k} \end{bmatrix} \geq 0
\]

(for \( 2k \leq J \)) and

\[
\Delta_{2k+1} = \det \begin{bmatrix} \mu_1 & \mu_2 & \cdots & \mu_{k+1} \\ \mu_2 & \mu_3 & \cdots & \mu_{k+2} \\ \vdots & \vdots & \ddots & \vdots \\ \mu_{k+1} & \mu_{k+2} & \cdots & \mu_{2k+1} \end{bmatrix} \geq 0
\]

(for \( 2k + 1 \leq J \)), there exists a density function \( d(t) \), so that \( \int_0^1 d(t)dt = \mu_j \) for \( j = 0, \ldots, J \).

For the first three inequalities (i.e., \( J = 2 \)) this means:

\[
\Delta_0 = \mu_0 \geq 0 \\
\Delta_1 = \mu_1 \geq 0 \\
\Delta_2 = \mu_2 - \mu_1 \mu_0 = \mu_2 - \mu_1^2 = \kappa_2 \geq 0.
\]

The first two inequalities are trivial and always fulfilled in this context. The third inequality reduces to a constraint on the second-order cumulants that are being subtracted:

\[
\kappa_2^d = \kappa_2^f - \kappa_2^p \geq 0,
\]

or

\[
\kappa_2^f \geq \kappa_2^p.
\]

In terms of variances \( \sigma_2^2 \) (the terminology and notation used in the main text), this inequality becomes:

\[
\sigma_f^2 \geq \sigma_p^2.
\]

More intricate constraints apply to higher moments and cumulants. For our purposes, however, the higher order criteria are ignored during the minimization process, as their estimates become increasingly volatile for a finite number of data points.

\[\text{Appendix B}\]

Conditions for the Existence of a Deconvolution

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\[\text{1} \] The importance of the exponential order condition can be shown as follows:

\[ G(d)(s) = \log(\int_0^1 e^{st}d(t)dt) \leq \log(\int_0^1 e^{Me^{-at}}dt) = \log(Me^{-at}) - \log(k - s), \text{which shows that the integral exists for all } s < k. \]