

# DMCfun: An R package for fitting Diffusion Model of Conflict (DMC) to reaction time and error rate data

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**Abstract** Decision processes within choice reaction-time (CRT) tasks are often modelled using variations of a Diffusion Decision Model (DDM, for a review, see Ratcliff and McKoon (2008)). Ulrich, Schröter, Leuthold, and Birngruber (2015) introduced a Diffusion Model for Conflict tasks (DMC). The DMC model combines common features from within standard diffusion models with the addition of superimposed controlled and automatic activation. The DMC model is used to explain distributional reaction time (and error rate) patterns in common behavioural conflict-like tasks (e.g., Flanker task, Simon task). This paper introduces the R-package DMCfun, which implements the DMC model and provides functionality to fit the model to observed data.

**Keywords** reaction time · conflict task · Diffusion Model Conflict (DMC) · R package

## 1 Introduction

Conflict tasks (e.g., Flanker task, Simon task, Stroop task, see Eriksen & Eriksen, 1974; Lu & Proctor, 1995; MacLeod, 1991), respectively) are common experimental paradigms used to investigate decision processes under response competition (i.e., the activation of competing response alternatives). Such conflict tasks involve stimuli with multiple features that are either relevant or irrelevant to the required task. For example, a typical Flanker task consists of an array (e.g., HHHHH, HHSHH) with the central item being the “target” (task-relevant) and the surrounding items being the “flankers” (task-irrelevant). Responses are typically mapped to the left and right hand (e.g., H = left, S = right). When the target and the flankers match (mismatch), both the relevant and irrelevant dimensions indicate the same (different) response. In a typical Simon task, participants respond to a feature of a stimulus (e.g., colour, shape; relevant dimension) that is presented at different spatial locations (e.g., left vs. right side of the screen; irrelevant dimension). Again, responses are mapped to the

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left and right hands (e.g., blue = left, red = right). When a stimulus presented in a left (right) location requires a left-hand response, the relevant and irrelevant stimulus dimensions match (mismatch). In such tasks, responses are typically faster and less error-prone when the relevant and irrelevant response dimensions indicate the same response (compatible) than when they indicate different responses (incompatible). The longer reaction time and larger error rate in incompatible trials indicates that task-irrelevant information interferes with the processing of the task relevant information.

This compatibility effect pattern is consistent across all conflict-like tasks. However, when the size of the effect (incompatible - compatible) is compared across different percentiles of the RT distribution, different patterns can emerge. Such differences across the RT distribution are investigated via the use of delta functions (or delta plots) (De Jong, Liang, & Lauber, 1994; Speckman, Rouder, Morey, & Pratte, 2008). Such plots depict the compatibility effect (y-axis) at  $n$  bins (x-axis) (see Figure 1 bottom-right panel for an example of a delta plot). Such distributional analyses have indicated that the compatibility effect in some conflict tasks (e.g., Stroop task) increases with slower responses (i.e., positive-going delta slope) whereas in other conflict tasks (e.g., Simon task) smaller compatibility effects are observed with slower responses (i.e., negative-going delta slope) (see Pratte, Rouder, Morey, & Feng, 2010, for Stroop and Simon tasks).

The observation of both positive and negative-going delta slopes across different conflict tasks has strong theoretical implications for processing models of RT. For example, traditional diffusion models of RT cannot account for negative-going delta functions (Pratte et al., 2010). These models propose that task-relevant information accumulates over time until one of two decision boundaries is reached (correct vs. incorrect response boundary). Wagenmakers and Brown (2007) demonstrate that the positive relationship between the mean and standard deviation of RT excludes the possibility of observing negative-going delta functions. As summarized by Ulrich et al. (2015), traditional RT models have difficulty explaining negative-going delta functions. The DMC model is an extension of the standard diffusion model that can explain observed delta functions (both positive and negative going) across a range of conflict tasks. Since its publication, simulation and fitting procedures of the DMC model have been applied in various research context in order to provide important insights into the mechanisms at play in conflict tasks (e.g., Mittelstädt & Miller, 2020; Servant, White, Montagnini, & Burle, 2016).

Conditional accuracy functions (CAFs) also offer an additional tool with which to investigate compatibility effects across the RT distribution. Here, accuracy rates are calculated across  $N$  equally sized RT bins. For compatible trials, accuracy is typically high across all bins, whereas for incompatible trials, the lowest level of accuracy is observed at fastest RT bins. At the slowest RT bins, there is little difference in accuracy between compatible and incompatible conditions.

The following section introduces the DMC model Ulrich et al. (2015). It must be noted that other extensions of standard drift diffusion models with specific reference to conflict-like tasks have been proposed. For example, the dual-stage two-phase model (DSTP; Hübner, Steinhauser, and Lehle (2010)) or the shrinking spotlight model (SSP; White, Ratcliff, and Starns (2011)). A direct comparison between such models is beyond the scope of the current paper with such comparisons provided elsewhere (see Evans & Servant, 2020; White, Servant, & Logan, 2018, albeit with specific reference to the flanker task only).

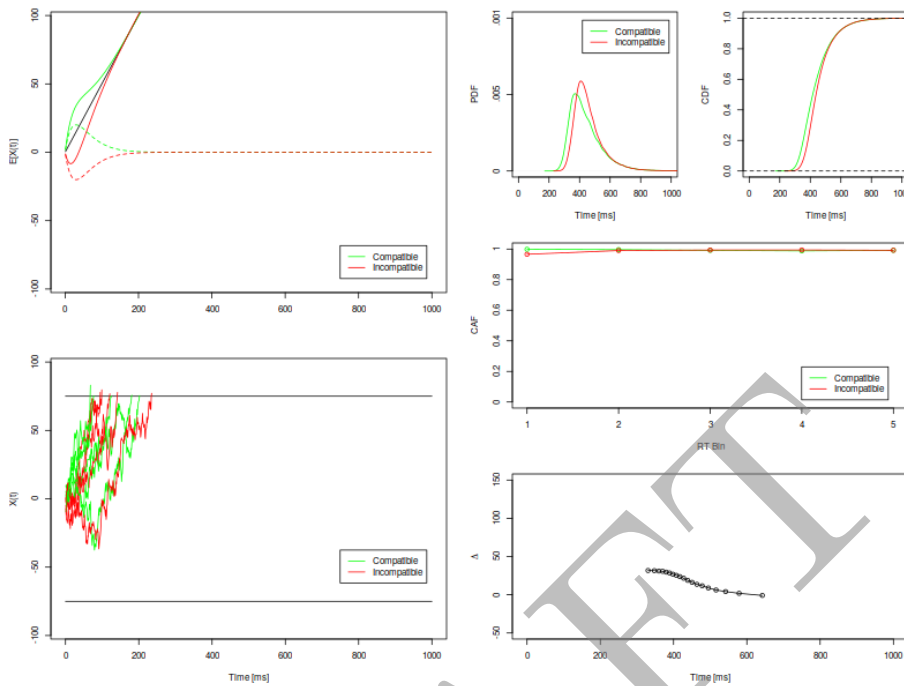
## 1.1 DMC Model Details

The DMC model of Ulrich et al. (2015) builds upon the idea that conflict tasks consist of two processes that proceed in parallel (automatic and controlled processing). Here, controlled processing concerns the task-relative information, whereas automatic processing concerns the task-irrelevant information. Whilst the architecture of such combined controlled/automatic processing within individual conflict tasks is under-specified, Ulrich et al. (2015) suggest two possible architectures (see Figure 1 Ulrich et al., 2015) under which automatic and controlled processing interact. Under both architectures, DMC assumes that a single accumulation process (combined automatic and controlled) determines the executed response. Critically, automatic processes will facilitate (impede) controlled processes in compatible (incompatible) trials. In addition, it is assumed that the activation output from automatic processes increases to a maximum before decreasing back to zero (pulse function).

The formal model specification for DMC is provided in Ulrich et al. (2015) and thus, will only be briefly described here. The time-course of automatic activation is modelled using a Gamma density function with a shape parameter (aaShape, with  $\alpha > 1$ ) and a scale parameter (tau,  $\tau$ ) multiplied by a constant (amp, +ve for compatible, -ve for incompatible trials), with the drift rate ( $\mu$ ,  $\mu$ ) being the first derivative of the density function. This gives DMC 7 parameters that can vary: amp, aaShape, tau,  $\mu$ , bnds, resMean, and resSD. Like standard diffusion models (see Ratcliff, 2013), DMC also allows increased trial-to-trial variability by allowing the starting point to vary. This starting point is sampled from a general beta distribution (spShape  $\alpha > 0$ ) centered around zero within the boundary range. DMC assumes that total RT is a combination of decision (D) and residual non-decision (R) processes and assumes that compatibility effects influence D processes only. The predictions of DMC are independent of the distributional properties of R. However, to provide more realistic RTs, the non-decisional component (R) was sampled from a normal distribution with a given mean and standard deviation. Within the DMC model, activation accumulates over  $t$  ( $t > 0$ ), with  $t$  being the time since stimulus onset.  $\mathbf{X}_c(t)$  represents the accumulation of noisy information from controlled processes, whereas  $\mathbf{X}_a(t)$  is the accumulation of noisy information from automatic processes. For controlled activation, drift rate ( $\mu_c$ ) is assumed to be time-independent, whereas for automatic activation, drift rate ( $\mu_a$ ) depends on  $t$ . The critical assumption is that total accumulation of information is the sum of the controlled and automatic processes ( $\mathbf{X}(t) = \mathbf{X}_c(t) + \mathbf{X}_a(t)$ ) since it is assumed that automatic activation spills over to the controlled process. Responses are generated whenever total activation exceeds some boundary ( $b$ ).

Figure 1 (top-left panel) illustrates the model dynamics for automatic and controlled activation averaged over 100,000 compatible and incompatible trials. The solid black line represents controlled activation. The green and red dotted lines represent automatic activation in compatible and incompatible trials, respectively. Critically, the automatic activation induced by the task-irrelevant dimension is brief and time-dependent. The solid green and red lines represent the summed automatic and controlled activation. For compatible trials, automatic and controlled activation both accumulate in the same direction as the correct response, whereas for incompatible trials, automatic and controlled activation produce opposite activation levels. This results in delayed boundary level activation for incompatible compared to compatible trials.

= Ulrich et al. (2015) collected data from both a flanker task and a simon task and fitted this data to DMC. Here, the delta (RT) and CAF (accuracy) values from the observed and predicted data were calculated. The model was fitted by minimising the root-mean-square



**Fig. 1** DMC simulation. This simulation involved 100,000 trials for each compatibility condition with the following default simulation parameters:  $\text{amp} = 20$ ,  $\text{tau} = 30$ ,  $\mu = 0.5$ ,  $\text{bnds} = 75$ ,  $\text{resMean} = 300$ ,  $\text{resSD} = 30$ ,  $\text{aaShape} = 2$ . The upper-left panel shows the mean activation functions with the black line representing controlled activation. The dotted green and red lines represent the automatic activation for compatible and incompatible trials, respectively. The solid green and red lines represent the sum of controlled and compatible and incompatible automatic activation, respectively. The bottom-left panels shows 5 individual compatible and incompatible trials. A response is generated when activation exceeds the boundary level (horizontal straight lines). The right upper two panels show the probability distribution function (PDF) and the cumulative distribution function (CDF) for both compatible and incompatible trials. The middle right panel show the conditional accuracy function (CAF) for compatible and incompatible trials. The bottom right panel show the delta function for the compatibility effect (incompatible - compatible).

error (RMSE) weighted difference between the predicted and observed delta and CAF values using the SIMPLEX minimization routine of Nelder and Mead (1965).

## 2 R Package: DMCfun

The R-package DMCfun provides functionality allowing both the simulation of the DMC model and fitting the model to observed data. The simulation code is written in C++ utilising the R-package Rcpp (Eddelbuettel et al., 2011). The use of C++ gives a significant performance boost over base R whilst the Rcpp interface allows convenient use from within the standard R environment and also direct access to plotting facilities within R. The fitting procedure uses the R-package optimr Nash (2016) with method = "Nelder-Mead". As in Ulrich et al. (2015) the model fit was evaluated by calculating the RMSE weighted difference between the predicted and observed delta/CAF values. The following sections will provide a

tutorial-type overview of using the package, both for running the basic simulation and also fitting the observed data from Ulrich et al. (2015).

## 2.1 Getting Started

The package is currently hosted on GitHub (<https://github.com/igmmgi/DMCfun>). It is anticipated that the package will also be submitted to the R CRAN repository. The package can be installed directly from GitHub using the devtools package (via `install_github`, see Code Example 1).

R Code Example 1: installation

```
> # install.packages("devtools")
> devtools::install_github("igmmgi/DMCfun")
> library(DMCfun)
> help(package = "DMCfun")
```

## 2.2 DMC Simulation

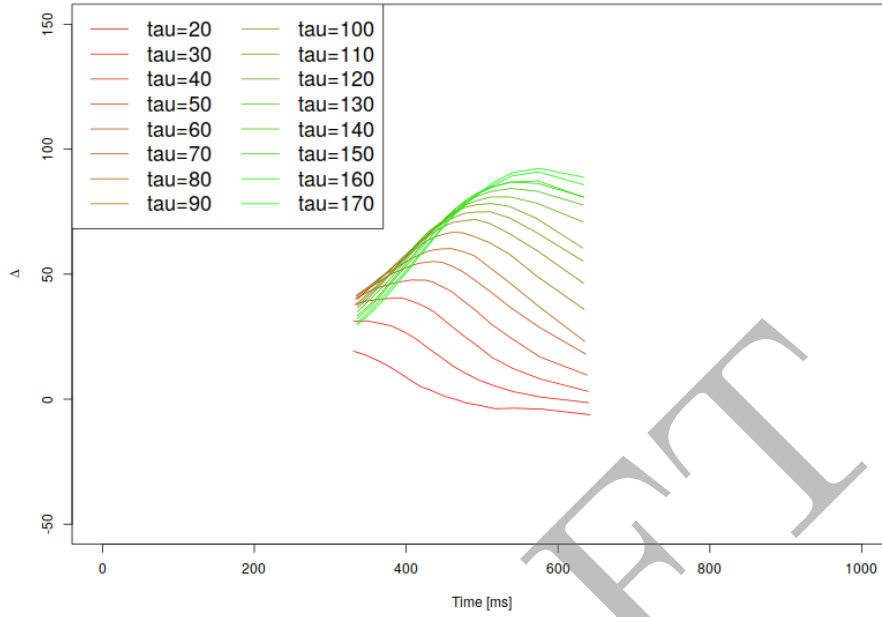
The simulation code is within the functions `dmcSim` (see Code Example 2). The function `dmcSim()` allows the user to specify parameter values using named arguments (`?dmcSim`). A single simulation run with 100,000 trials in each compatibility condition takes approximately < 150 ms (standard desktop PC). The function `dmcSims` allows the user to specify a list of input parameters in order to run multiple simulations. For example, Ulrich et al. (2015) observed that the parameter tau ( $\tau$ ) largely determined the slope to the delta function. In Code Example 3, the DMC simulation is run using tau starting parameters from 20 to 170 in steps of 10 (see Figure 2).

R Code Example 2

```
> dmc <- dmcSim(fullData = TRUE) # Fig 3 from Ulrich et al. (2015)
> dmc <- dmcSim(fullData = TRUE, tau = 150) # Fig 4 from Ulrich et al. (2015)
> plot(dmc)
> plot(dmc, figType = "caf")
> plot(dmc, figType = "delta")
>
> dmc <- dmcSim()
> plot(dmc) # Fig 3 from Ulrich et al.
```

## 2.3 DMC Fit: Real data

The package includes the raw data from Ulrich et al. (2015) (see Code Example 4). This data can be fitted to DMC via the function `dmcFitAgg` (see Code Example 5). This will fit DMC to the data averaged across all participants, whereas the function `dmcFitVPs` will fit DMC to individual participants. A summary of the fit is provided. By default, `dmcFit*` fits the following parameters: `amp`, `tau`, `mu`, `bnds`, `resMean`, `resSD`, `aaShape`, `spShape`. The function `dmcFit*`



**Fig. 2** DMC Simulation delta functions with varying tau parameter only.

uses the R-package `optimr` Nash (2016) with method = "Nelder-Mead" and allow the specification of a number of options including which parameters to fit and their starting values. Ulrich et al. (2015) observed that model fit was highly influenced by the tau parameter. The function `dmcFit*` first searches a grid space ( $N = 10$ ) within this single parameter to find the "best" starting value of tau before the simplex minimisation routine is implemented. An initial search of a parameter space has been shown to improve parameter recovery (Hübner & Pelzer, 2020). Such an initial search grid and subsequent fitting procedure takes < 1 minute. It is also possible to search the initial grid space across multiple parameters with the search space being determined by the input arguments `minVals`, `maxVals`, and `fitInitialGridN`. It should be noted that increasing the size of the initial grid space will increase computational time prior to the simplex minimisation routine dramatically. For example, searching 6 parameters within a grid space of 6, results in 46,656 initial parameter sets. Although these are independent and are run in parallel, such an initial grid space takes approximately 20 minutes (tested on 12 core desktop PC). As a result, the default setting is to perform the initial search in the tau space only. Such a procedure produced consistently good fits with the data of Ulrich et al. (2015) ( $RMSE < 15$ ), although this does not exclude the possibility that other datasets would benefit from an extended search grid (or alternative starting values) regarding the initial values for the simplex routine.

## R Code Example 3

```
> dmc <- dmcSims(list(tau = seq(20,170,10)))
> plot(dmc[[1]]) # full plot first combination
> plot(dmc)      # plot delta functions for each combination
```

Users can fit DMC to their own data using the function `dmcObservedData` (see Code Example 6). This function takes an R dataframe (or a \*.txt file with tab separated values, one header line) (see Table 1) and calculates the required variables. Four data columns are required: one for participant number, one for compatibility, one for reaction time, and one for error rate. The specific column names and column codings can be specified using function arguments.

## R Code Example 4

```
flankerDataRow      # raw flanker data
flankerData         # summarised flanker data
flankerData$summary # aggregated flanker data
plot(flankerData)
simonDataRow        # raw simon data
simonData           # summarised simon data
simonData$summary   # aggregated simon data
plot(simonData)
```

## R Code Example 5

```
> fit<-dmcFitAgg(flankerData) # flanker data from Ulrich et al. (2015)
> plot(fit,flankerData)
> summary(fit)

# output values from flanker data
# A tibble: 1 x 9
  amp   tau   mu  bnds resMean resSD aaShape spShape rmse
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1  17.7  106.  0.599  55.7   326.   28.8   2.02   3.04   7.50

> fit<-dmcFitAgg(simonData) # simon data from Ulrich et al. (2015)
> plot(fit,simonData)
> summary(fit)

# output values from simon data
# A tibble: 1 x 9
  amp   tau   mu  bnds resMean resSD aaShape spShape rmse
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1  17.9  27.9  0.595  58.1   315.   31.5   2.32   3.20  11.6
```

## R Code Example 6

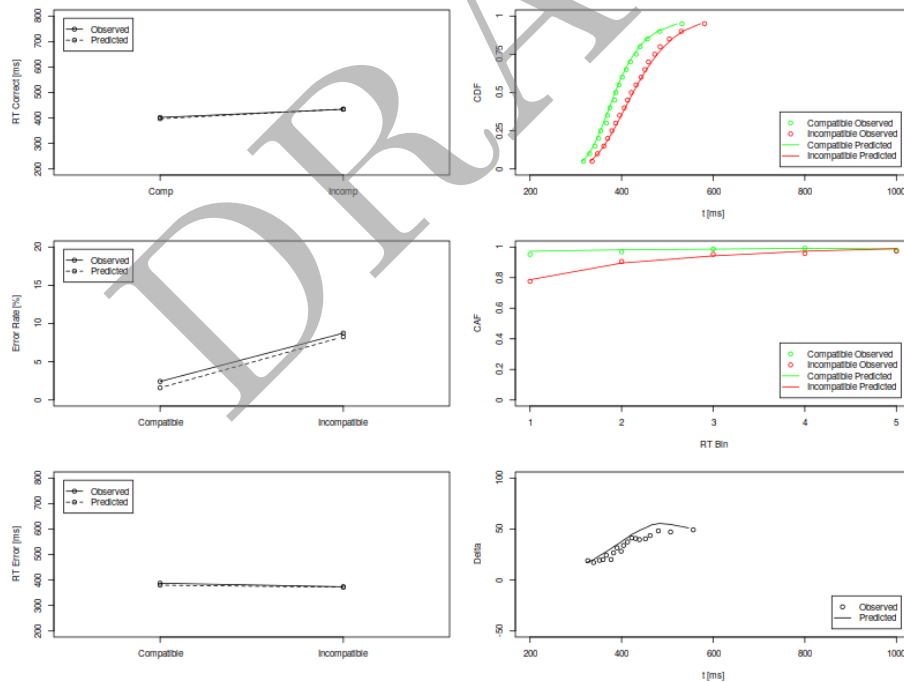
```

> ?dmcObservedData
> dat0b <- dmcObservedData(datframe,
                           columns = c("SNo", "comp", "rt", "err"),
                           compCoding = c(1, 2),
                           errorCoding = c(0, 1))
> plot(dat0b)
> dmcFit <- dmcFitAgg(dat0b)
> plot(dmcFit, dat0b)

```

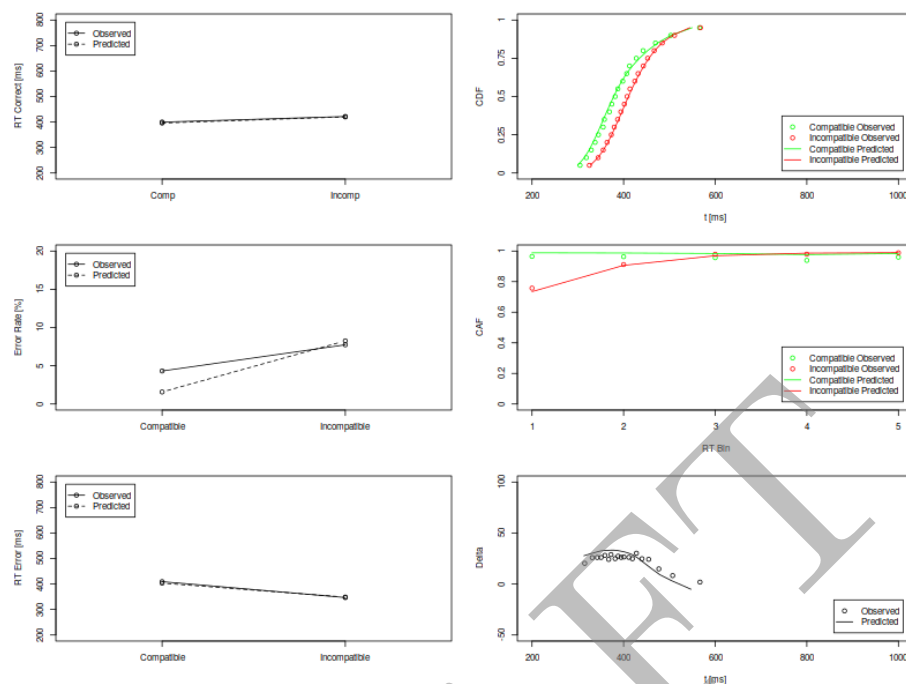
Table 1: Raw Data file example

VP	Comp	RT	Error
1	comp	601.657	0
1	comp	451.429	0
1	comp	418.097	0
1	incomp	451.436	0
1	comp	401.312	0
1	incomp	434.749	0



**Fig. 3** DMC Fit from the Flanker data presented in Ulrich, Schröter, Leuthold, and Birngruber (2015). Refer to Code Example 5.





**Fig. 4** DMC Fit from the Simon data presented in Ulrich, Schröter, Leuthold, and Birngruber (2015). Refer to Code Example 5.

### 3 Conclusions

To summarize, the R-package DMCfun provides users with an easy-to-use interface for both simulating the DMC model with various input parameters and fitting this model to observed behavioural data from within conflict-like tasks.

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### Conflict of interest

The authors declare that they have no conflict of interest.

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