

# SequentialSamplingModels.jl: Simulating and Evaluating Cognitive Models of Response Times in Julia

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## ABSTRACT

Sequential sampling models (SSMs) are a widely used framework describing decision-making as a stochastic, dynamic process of evidence accumulation. SSMs' popularity across cognitive science has driven the development of various software packages that lower the barrier for simulating, estimating, and comparing existing SSMs. Here, we present a software tool, SequentialSamplingModels.jl (SSM.jl), designed to make SSM simulations more accessible to Julia users, and to integrate with the Julia ecosystem. We demonstrate the basic use of SSM.jl for simulation, plotting, and Bayesian inference.

## Keywords

cognitive models, response time, sequential sampling models, evidence accumulation, decision-making

## 1. Introduction

Sequential sampling models (SSMs) are widely used in cognitive science due to their ability to describe dissociable processes underlying a wide variety of capacities, including memory retrieval, visual perception, and decision making (Forstmann et al. (2016)). These models typically describe decision-making as a stochastic (i.e., random), dynamic evidence accumulation process which evolves until evidence for one option reaches an evidence threshold (Smith (2000); Ratcliff and McKoon; Ratcliff). By doing so, SSMs provide a generative model of joint choice-response time (RT) distributions. Figure 1 illustrates the latent evidence accumulation process for a hypothetical decision between a Margherita pizza and a pineapple pizza. Evidence accumulates stochastically until it reaches either the upper boundary, triggering the selection of the Margherita, or the lower boundary, triggering the selection of the pineapple pizza. While these models have many promising applications, their implementation and use remain a technical challenge.

The widespread interest and continuous use of SSMs across cognitive science have spurred the development of various software packages that lower the barrier for simulating, estimating, and comparing existing SSMs. These software implementations range widely, from those designed to provide maximum flexibility to end-users (Shinn et al. (2020)) to those that streamline the entire analysis of SSMs (Fengler et al.). Toolboxes have also been developed across multiple programming languages commonly used for scientific

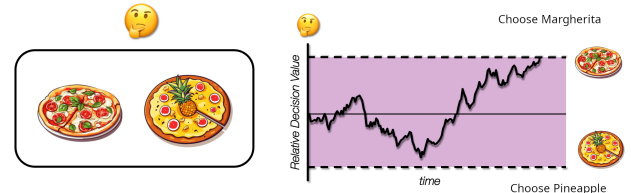


Fig. 1: An example of applying the sequential sampling modeling framework to a choice between two pizzas. The decision-maker samples evidence in favor of both options until reaching a decision boundary.

computing, including MATLAB (Vandekerckhove and Tuerlinckx (2008)), Python (Wiecki et al. (2013); Shinn et al. (2020); Fengler et al.; Murrow and Holmes (2024)), and R (Wabersich and Vandekerckhove; Stevenson et al. (2025); Ahn et al. (2017); Singmann et al.; Hartmann and Klauer (2021)).

## 2. SequentialSamplingModels.jl

While existing methods can already simulate a wide range of SSMs across multiple languages, developments in Julia have either not been established or lack maintenance. SequentialSamplingModels.jl (SSM.jl) provides the first unified interface for simulating validated SSMs, drawn from the literature to represent those most widely used models. The package offers a user-friendly method for simulating a range of SSMs of interest to researchers. It is the first attempt to integrate a package into the Julia ecosystem for simulating and evaluating a popular model of decision-making.

## 3. Availability, Development, and Documentation

SequentialSamplingModels.jl is available on the Julia package registry. The package is maintained on GitHub at <https://github.com/itsdfish/SequentialSamplingModels.jl>, and the documentation can be found at <https://itsdfish.github.io/SequentialSamplingModels.jl/dev/>. For a more comprehensive exploration of the package functionalities and further details on future developments, the user is invited to consult the package README, documentation, and issues.

## 4. Example one: simulating choices and response times

SSMs.jl is an extension of the established functionalities within the Distributions.jl package Lin et al. (2024), designed for generating multivariate choice-RT distributions. To achieve this, a new abstract

type was defined to support both continuous and discrete values. Then, a suite of constructors were designed to accommodate multivariate distributions with mixed support, which are necessary for generating discrete choices and continuous response times. Across the defined sub-types, distributions utilizing SSMs.jl will output a NamedTuple consisting of a vector of choices and RTs. With samplers designed to integrate with Distributions.jl in mind, the constructors can leverage many of the related functions for probabilistic distributions, including sampling and likelihood evaluation. In Code Block 1, using the Diffusion Decision Model (Ratcliff et al. (2016); DDM), we provide an example of these features for simulating choices and response times from a pre-defined set of parameters, as well as computing the log-likelihood of the data. The DDM consists of four parameters:

- $\nu$ : the drift rate which represents the rate of evidence accumulation towards the upper boundary
- $\alpha$ : the decision threshold which represents the amount of evidence required to make a decision
- $z$ : the starting point bias towards the upper decision boundary
- $\tau$ : non-decision time which represents the sum of encoding and motor response times.

Code 1: A code example illustrating how to create a Diffusion Decision Model object, generate simulated data, and evaluate the log likelihood of the simulated data.

```

1 using SequentialSamplingModels
2 # Create DDM distribution
3 dist = DDM(; ν = 1.00, α = 0.80, τ = 0.30, z =
0.50)
4 # Sample 10,000 simulated data from the DDM
5 sim_data = rand(dist, 10_000)
6 # compute log likelihood of simulated data
7 logpdf(dist, sim_data)
```

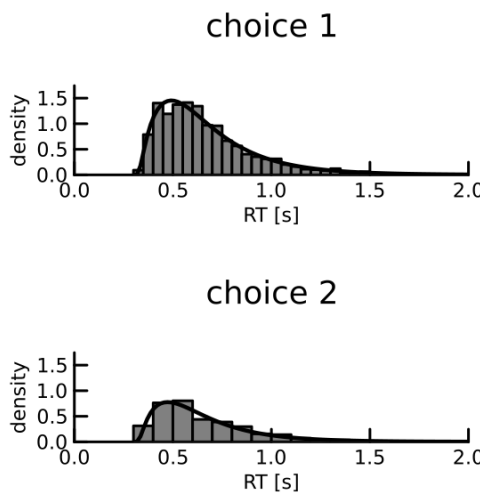


Fig. 2: RT distributions of the racing diffusion model based on Code Block 2

## 5. Example two: plotting RT-choice distributions and simulated traces

SSM.jl also enables plotting functionality for SSMs when Plots.jl (Christ et al. (2023)) is active in your Julia session. In Code Block 2, using the racing diffusion model (Tillman et al. (2020); RDM), we demonstrate how to generate two plots commonly used with SSMs. The RDM consists of the following parameters:

- $\nu$ : a vector of drift rates representing the evidence accumulation rate for each response option
- $k$ : a positive constant added to  $A$  such that  $A + k$  is the decision threshold, which represents the amount of evidence required to make a decision
- $A$ : the maximum starting point of the diffusion process
- $\tau$ : non-decision time which represents the sum of encoding and motor response times.

The first plot, shown in Figure 2, is a histogram of joint choice-RT reaction distributions paneled by choice with a probability density overlay. The second plot illustrates five simulated trajectories of the latent evidence accumulation process of the RDM, which is shown in Figure 3. In addition, users can animate the evidence accumulation trace with the function `animate` to illustrate the model's dynamics.

Code 2: SSM.jl works with Plots.jl. The first example is a histogram of joint choice-RTs (see Figure 2), and the second example is a trace of the latent evidence accumulation dynamics (see Figure 3).

```

1 using SequentialSamplingModels
2 using Plots
3 # Define parameters
4 ν = [2,1]
5 k = 0.50
6 A = 1.0
7 τ = 0.30
8 # Create RDM distribution
9 dist = RDM(;ν, k, A, τ)
10 # Plot distribution
11 histogram(dist; xlims=(0, 2))
12 plot!(dist; t_range=range(.301, 2.5, length=100))
13 # Plot model simulations traces
14 plot_model(dist, n_sim=5,
15 t_range=range(.20, 1.5, length=100), xlims=(0, 1.5))
```

## 6. Example three: Bayesian inference

In cases where closed-form likelihoods are available, SSM.jl can also facilitate parameter estimation routines available with Turing.jl, a general-purpose probabilistic programming library (Ge et al. (2018)). Below, we provide an example of writing a Turing.jl model using a SSM constructor and conducting a simple parameter estimation exercise using the linear ballistic accumulator (Brown and Heathcote (2008); LBA). The LBA has the same parameters as the RDM, but makes different assumptions about the sources of stochasticity in the evidence accumulation process. In code example 3, we use a Markov Chain Monte Carlo (MCMC) technique called the No U-Turn Sampler (Hoffman et al., 2014) to estimate the posterior distribution of the LBA parameters from simulated data.

Code 3: Use Turing.jl to conduct parameter estimation with a SequentialSamplingModels.jl constructor

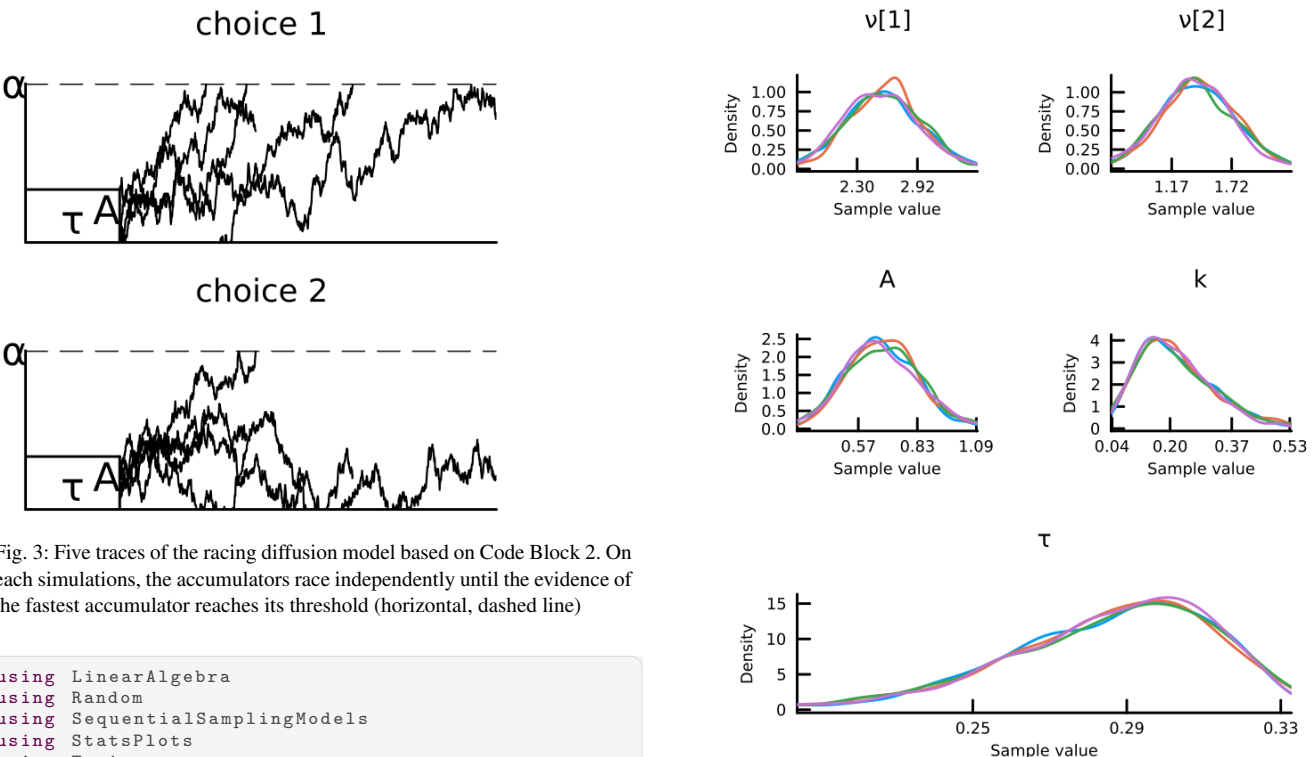


Fig. 3: Five traces of the racing diffusion model based on Code Block 2. On each simulations, the accumulators race independently until the evidence of the fastest accumulator reaches its threshold (horizontal, dashed line)

```

1  using LinearAlgebra
2  using Random
3  using SequentialSamplingModels
4  using StatsPlots
5  using Turing
6
7  Random.seed!(104)
8  # Simulate data
9  dist = LBA(ν=[3.0, 2.0], A=.8, k=.2, τ=.3)
10 data = rand(dist, 100)
11 # Define model using Turing.jl macro
12 @model function model_lba(data)
13     ν ~ MvNormal(zeros(2), I(2))
14     A ~ truncated(Normal(.8, .4), 0.0, Inf)
15     k ~ truncated(Normal(.2, .2), 0.0, Inf)
16     τ ~ Uniform(0.0, minimum(data.rt)))
17     data ~ LBA(;ν, A, k, τ)
18 end
19 # Fit model
20 chains = sample(model_lba(data), NUTS(1000, .85),
21 MCMCThreads(), 1000, 4)
22 # Examine convergence quality
23 summarystats(chains)
24 # Plot Markov Chain Monte Carlo (MCMC)
25 # trace and posterior distributions
26 plot(chains)

```

In addition, SSM.jl integrates with Pigeons.jl ((Surjanovic et al.) and ParetoSmooth.jl (<cdp49@cam.ac.uk> (2021)), via Turing.jl, allowing the user to estimate the marginal likelihood and approximate leave-one-out cross-validation for Bayesian model comparison (Vehtari et al. (2017)), respectively.

## 7. Future Applications

A major application of SSMs is to fit models to data via parameter estimation routines. However, not all SSMs have closed-form likelihoods that can be computed efficiently, and thus simulation-based inference has recently been proposed as a solution to accommodate parameter estimation for SSMs.

In the future, we aim to incorporate approximate solutions for likelihoods with no known closed form using approximate Bayesian computation (Palestro et al. (2018)) and deep learning methods

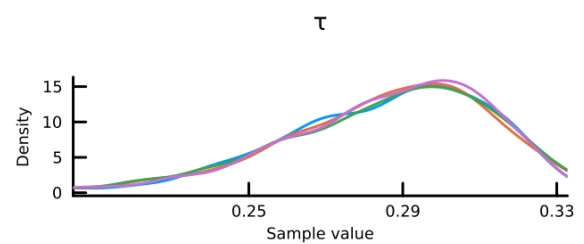


Fig. 4: Posterior distributions for the parameters of the Linear Ballistic Accumulator model based on Block 3. MCMC trace plots were removed due to space limitations.

for efficient simulation-based inference, such as likelihood approximation networks (Fengler et al. (2021)) specified in Flux.jl (Innes (2018)). This approach facilitates sampling from the posterior distribution of models without a closed-form likelihood function while maintaining a consistent syntax for end-users.

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