StabilityChecker

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Introduction

StabilityChecker is a python package that can be utilised to identify regions of parameter space capable of producing a desired stability while handling uncertainty in biochemical rate constants and initial conditions. It can be used to design new systems of desired stability or study the stability of existing systems. StabilityChecker takes advantage of Sequential Monte Carlo methods, thus is able to harvest the power of parallelisation and speed up the algorithm. StabilityChecker is suitable for use if GPU access is available.

Overview

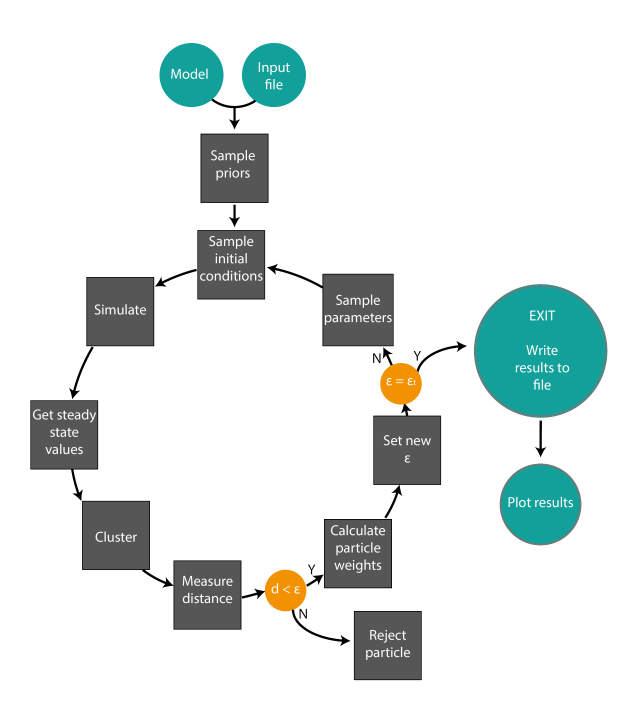


Figure 2.1: Flowchart of the algorithm used in Stability Checker. ϵ stands for the threshold for the distance from the desired value. The user provides the model and input files as inputs and the algorithm outputs the posterior distribution, the parameter space capable of exhibiting the desired stability.

Installation

StabilityChecker has been developed to work on a Linux operating system, with GPU support. The following dependencies are essential for the successful implementation of the package:

- \bullet numpy
- logging
- cuda-sim
- libsbml
- R
- \bullet ggplot2
- \$ cd StabilityChecker
- \$ python setup.py install

This will copy the module StabilityChecker into the lib/pythonXX/site-packages directory, where XX corresponds to the python version that was used. An exe=<dir> variable should also be added to the run.sh script pointing the script to the right package installation. Alternatively the path to the module can be added to the top of the run.sh file without the need for installation. The path to the cuda sim module must be added to the top of the run.sh file.

\$ export PATH=<dir>:\$PATH

Using the package

4.1 The working directory

The working directory must contain the following files. Each one is described in detail in the section following.

input_file.xml The user input file

run.sh The shell script that will initialise the scripts

 ${f plot}_{\hbox{-}}{f posterior}.{f R}$ The R code to plot the results

model file This can take two formats:

model.cu Cuda file of the model that is required to have this name

SBML model This can have any name, as long as this is provided in the input_file.xml. Cuda-sim will create the model.cu file using this SBML model.

4.2 The input file

The input.xml file contains all the necessary information that is not contained in the model itself. This file must follow the specific format showed in the 'Examples' folder. No whitespaces are to be included between the tags.

4.2.1 Required arguments

epsilons The acceptable distance from the desired value.

epsilon_t Total variance in the data

start The initial cut-off value for the distance from the desired values.

end The final cut-off value for the distance from the desired values.

epsilon_vcl Within-cluster variance

start The initial cut-off value for the distance from the desired values.

end The final cut-off value for the distance from the desired values.

eps_cl Number of clusters

start The initial cut-off value for the distance from the desired values.

end The final cut-off value for the distance from the desired values.

Desired final values The desired values the algorithm will converge to.

number_of_clusters The number of clusters in the data, describing its stability.

total_variance The total variance in the data.

cluster_variance The within-cluster variance in the data.

particles The number of accepted parameter sets required to proceed to the next iteration.

number_to_sample Number of parameter sets to sample from for iteration. This needs to be greater or equal to the particles.

initial_conditions_samples Number of initial condition sets to sample for each parameter set at each iteration.

alpha A value that describes the step size for the incremental reduction of the epsilon at each iteration. This should be set to a small value (a = 0.1) to speed up the algorithm.

times The time points for the simulation are given as a whitespace delimited list.

species_numb_to_fit Which two species numbers to be fit by the algorithm. The should be whitespace delimited.

stoch_determ The type of simulation to be used. The two options are stochastic or deterministic.

model_file Whether the model provided is in SBML or cuda format. The two options are sbml and cuda.

sbml_name The name of the SBML file that will be provided. Note that if a cuda file is provided, that must be named model.cu.

parameters The parameters included in the model.

item One item corresponds to one parameter. The number and order of the items must strictly correspond to those of the parameters in the model provided.

name The name of the parameter. Note that the parameters are read by order and not name from the model. The name provided here is used in the plotting module.

distribution From what distribution to sample values from. The two options are constant and uniform.

start The lower limit of the distribution to sample from.

end The upper limit of the distribution to sample from. Note that if the distribution is constant this value is not read.

initial_conditions The species included in the model.

- **item** One item corresponds to one species. The number and order of the items must strictly correspond to those of the species in the model provided.
 - **name** The name of the species. Note that the species are read by order and not name from the model.
 - distribution From what distribution to sample values from. The two options are constant and uniform. Strictly two species must be set to uniform and the rest constant.
 - **start** The lower limit of the distribution to sample from.
 - **end** The upper limit of the distribution to sample from. Note that if the distribution is constant this value is not read.

4.2.2 Output

The outputs from running StabilityChecker are saved in the results_txt_files folder. This folder contains two files and one folder per population. The two files are the following:

- Parameter_values_final Each line contains the values of the parameters in the order set in the input_file.xml file and each line corresponds to an accepted particle in the final accepted population.
- Parameter_weights_final Contains the weights of each particle (parameter set) in the final accepted population.

Each population folder contains the following files:

- data_PopulationN.txt Each line contains the values of the parameters in the order set in the input_file.xml file and each line corresponds to an accepted particle.
- data_WeightsN.txt Contains the weights of each particle (parameter set).
- One set_resultXX per parameter set. Each line contains the steady state value of each species, in order specified in the input_file.xml file, species_numb_to_fit. Each line corresponds to one initial condition set.

The plot of the posterior is saved in the posterior.pdf file in the working directory. The plot is interpreted as follows:

- The marginal distributions of each parameter are found on the diagonal
- The pairwise joint distributions are found along the sides. The location of each pairwise joint distribution is determined by which pair of parameters are compared.

4.2.3 Running StabilityChecker

To run StabilityChecker, and once the input_file.xml file is completed, the run.sh file must be executed. The progress of the algorithm can be followed in the my_abc_scan.log file. Once Stability-Checker is finished the posterior can be seen in the posterior.pdf file.

The phase plots of the populations can be visualised by plotting the contents of the set_resultXX file for each parameter set.

Examples

5.1 The simple genetic toggle switch using ODEs

5.1.1 Background

In this example, the simple toggle switch model developed by Gardner *et al.* (2000) was used. This model consists of two mutually repressing transcription factors and is defined by the following ODEs:

$$\frac{du}{dt} = \frac{a_1}{1 + v^{\beta}} - u$$
$$\frac{dv}{dt} = \frac{a_2}{1 + v^{\gamma}} - v$$

Where a_1 and a_2 are the effective rates of synthesis of repressors 1 and 2 respectively. u is the concentration of repressor 1 and v the concentration of repressor 2. β is the cooperativity of repression of promoter 1 and γ of repressor 2. This model is capable of bistability, and StabilityChecker will be used to find the parameter space in which this behaviour is found.

5.1.2 Setting up

Model

This example does not use an SBML model but instead provides StabilityChecker with the cuda file directly. This is the model.cu file.

Input file

The input file is set up as shown in the examples folder. As this model only contains two species, u and v, these are selected for the fit as well as initial condition scan.

run file

The pythonpath must be set to point to the directory in which cuda sim was installed.

\$export PATH=<dir>:\$PATH

In addition, the exe=<dir> variable must be set to the directory in which StabilityChecker is installed.

5.1.3 Running StabilityChecker

The working directory must contain the following:

- The model.cu file
- The completed input.xml file
- The customised run.sh file
- The plot_posterior.R file

The run.sh file is then executed:

./run.sh

The progress of the algorithm can be followed in the my_abc_scan.log file.

5.1.4 Results

The posterior is found in the posterior.pdf file. It is the one shown in figure 5.1. The plots on the diagonal represent the marginal distributions for the values of each parameter that were found in the final population, thus the ones that can produce bistable behaviour. The pairwise joint distributions are found on the side plots.

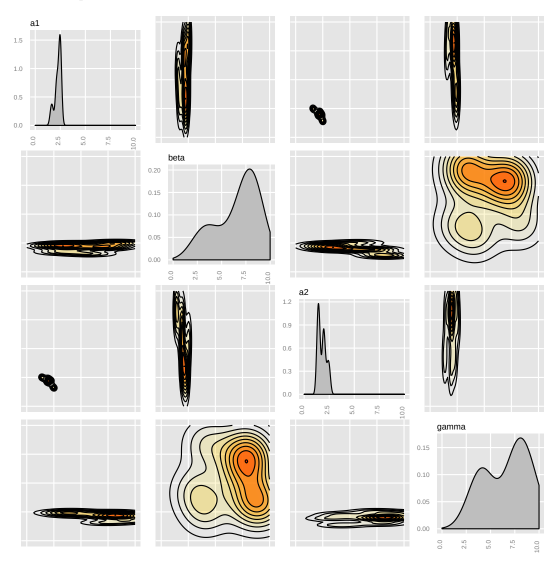


Figure 5.1: The posterior distribution of the Gardner toggle switch produced by StabilityChecker. The marginal distributions of each parameter are found on the diagonal and pairwise joint distributions along the sides.

Bibliography

Gardner, Timothy S, Cantor, Charles R, & Collins, James J. 2000. Construction of a genetic toggle switch in Escherichia coli. *Nature*.