SigNetSim: A web platform for building and analyzing mathematical models of molecular signaling networks.

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Introduction and objective

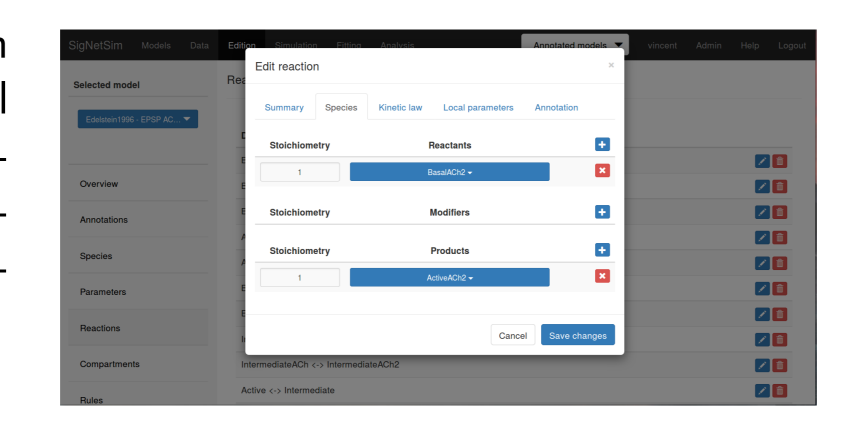
Molecular biology is experiencing a revolution, in one part thanks to new technologies to measure and perturbate biological systems in vitro, and also due to the growing importance of mathematical modeling which enables us to understand biological mechanisms in a more profound way. However, a crucial point in this transforming field is the need to provide completely new tools, which should be computationally efficient, versatile, and compatible.

To this end, we developed SigNetSim, a web platform that allows user to create, simulate, adjust and analyse biochemical reaction models. As a web platform, it is usable on multiple devices. It is designed to be installed on computation servers, with all the work executed server-side. It uses various standards to produce reproducible projects.

It is open-source, and available on GitHub and at signetsim.org

Models

Through the interface, users can easily define the mathematical model as a set of biochemical reactions, as well as other functionalites available in the SBML format.



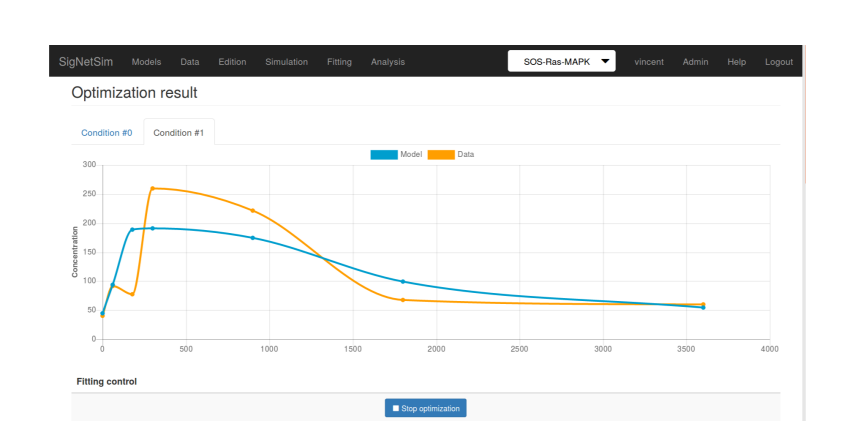
Simulations

Users can simulate their models, and obtain timeseries or steady states. It can also simulate the model according to a batch of initial conditions, and compare the results to experimental observations.



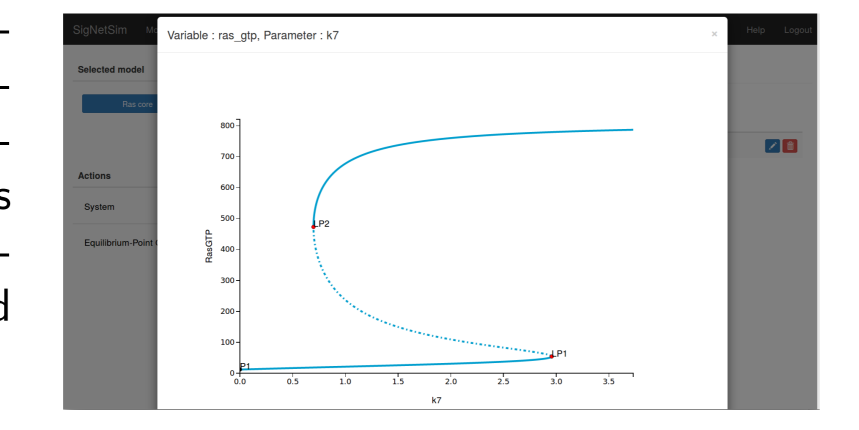
Model fitting

SigNetSim can perform curvefitting on models, using a parallelized simulated annealing. This algorithm is particularly suited for optimizing large non-linear systems.



Continuations

In order to analyse the dynamical properties of the mathematical model, SigNetSim uses continuations techniques. It allows the user to investigate the possible multiple equilibria and find the bifurcations between them.



Acknowledgements

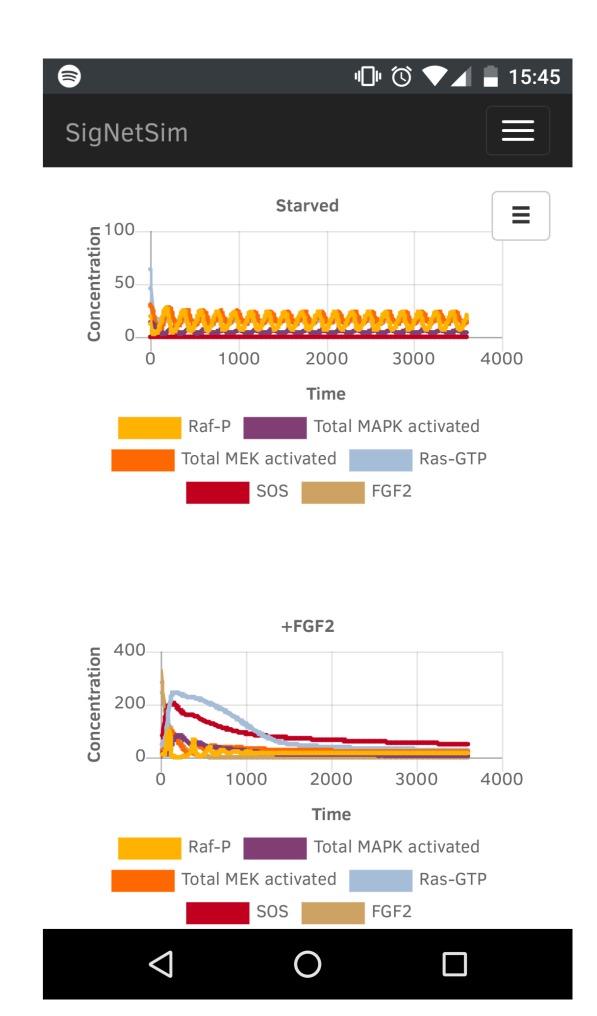




Multi-Device

SigNetSim is a web interface written mostly in python with the Django framework, which uses Bootstrap as a front-end framework. It is designed to function in various devices, from smartphone to desktop computer. The computer-intensive computations are performed server-side.

An important part of the code is written in JavaScript, which makes the interface fast and dynamic. SigNetSim uses a collection of JavaScript libraries to make the interface more responsive. It can draw reaction diagrams using Cytoscape.js, simulations as interactive plots using Chart.js, or render mathematical formulas using Mathjax.js.



Standards

SigNetSim is compatible with most of the SBML format. It also uses the SBML comp package to represent hierarchical model composition.



It is partially compatible with SED-ML format, which stores simulation settings and allows users to easily reproduce simulations from literature.



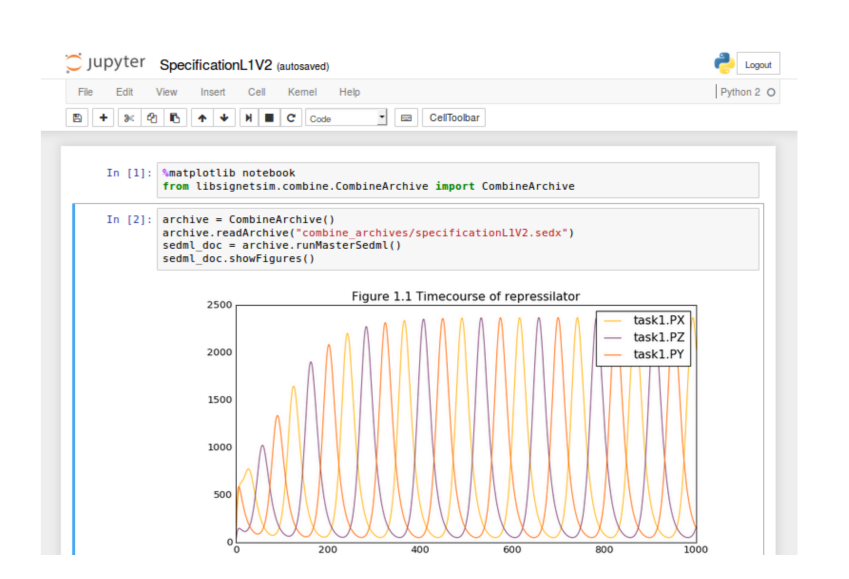
Moreover, SigNetSim can use COMBINE archive format, which stores both the SBML model and the SED-ML simulation file into one file. It also can also store experimental data from its database into the NuML format. Loading an existing combine archive will automatically load a new project



existing combine archive will automatically load a new project, with models and simulations.

libSigNetSim

SigNetSim is modular, and its core library is usable directly from jupyter notebooks. Users can build models with a few lines of code. One advantage of this use case is to be able to analyse the model directly with the computer algebra system Sympy.



Open source

SigNetSim, and its core library libSigNetSim, are available on GitHub. They are distributed under AGPLv3 and GPLv3 licences, respectively. We use continuous integration to perform tests on every update, making the software more robust. Installation scripts are provided for debian-based linux distributions, as well as Docker images.

Future work

- SBGN viewer/editor
- Stochastic simulations
- Database interaction, annotations

https://signetsim.org