# sybilSBML - Quick Start

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#### 1 Introduction

The package sybilSBML is an addition to SyBiL providing support for metabolic networks written in SBML (Systems Biology Markup Language), in particular those developed by Bernhard  $\emptyset$ . Palsson's lab<sup>1</sup> and those from the BiGG database<sup>2</sup>[Schellenberger et al., 2010]. The software is available from the sybilSBML homepage<sup>3</sup>.

The package sybilSBML requires one additional package rsbml available from Bioconductor<sup>4</sup>, which is itself powered by LibSBML [Bornstein et al., 2008] available from the SBML homepage<sup>5</sup> and pkg-config<sup>6</sup>.

#### 2 Installation

The following sections describe the installation process of sybilSBML on various platforms.

#### 2.1 Linux and Mac OS X

The package *sybilSBML* depends on a separate installation of *rsbml* and therefor of an installation of LibSBML (use LibSBML version 3.4.1) and pkg-config (which should be available in a Linux distribution).

If pkg-config is installed in a non-standard directory, set the two environment variables PKG\_CONFIG\_PATH and PKG\_CONFIG to appropriate values. If LibSBML is installed in a non-standard directory, set the environment variables LIBSBML3\_CFLAGS and LIBS-BML3\_LIBS, for example, if LibSBML is installed in your home directory:

```
$ export LIBSBML3_CFLAGS="-I$HOME/include"
```

\$ export LIBSBML3\_LIBS="-L\$HOME/lib -lxm12 -lz -lm -lbz2 -lsbml -lstdc++"

<sup>1</sup>http://gcrg.ucsd.edu/

<sup>2</sup>http://bigg.ucsd.edu/

<sup>3</sup>http://www.cs.uni-duesseldorf.de/AG/BI/Software/SyBiL/

<sup>4</sup>http://www.bioconductor.org/

<sup>5</sup>http://www.sbml.org/

<sup>6</sup>http://pkg-config.freedesktop.org/wiki/

For the installation of rsbml, start R and run

```
> source("http://bioconductor.org/biocLite.R")
> biocLite("rsbml")
```

Now install the source code package from a Terminal command line

```
$ R CMD INSTALL sybilSBML_X.X.X.tar.gz
```

## 2.2 Windows

There is a precompiled binary version of *rsbml* available, no additional software is required. Start R and run

```
> source("http://bioconductor.org/biocLite.R")
> biocLite("rsbml")
```

Now install the Windows binary package sybilSBML\_X.X.X.zip.

# 3 Usage

The package *sybilSBML* provides the command readSBMLmod() which reads SBML formated files and returns instances of class modelorg.

```
> library(sybil)
> library(sybilSBML)
> model <- readSBMLmod("<model>.xml")
```

## 4 Input files

The function readSBMLmod() reads metabolic network models written in SBML format (Systems Biology Markup Language). Among the models available in this de-facto standard format are in particular those developed by Bernhard Ø. Palsson's lab.

The metabolite id's are written in the format M\_<metabolite abbreviation>\_<compartment abbreviation>. The compartment abbreviation is a one letter abbreviation, e.g. c for cytosol. All metabolites outside the system boundary belong to compartment b. Those metabolites are transported into or outside the system. As long as they are mentioned, the network is closed. The function readSBMLmod() will remove them in order to produce an open network.

#### References

S. A. Becker et al. Quantitative prediction of cellular metabolism with constraint-based models: the COBRA Toolbox. *Nat Protoc*, 2(3):727–738, 2007. doi: 10.1038/nprot. 2007.99.

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- J. Schellenberger, R. Que, R. M. T. Fleming, I. Thiele, J. D. Orth, A. M. Feist, D. C. Zielinski, A. Bordbar, N. E. Lewis, S. Rahmanian, J. Kang, D. R. Hyduke, and B. Ø. Palsson. Quantitative prediction of cellular metabolism with constraint-based models: the COBRA Toolbox v2.0. Nat Protoc, 6(9):1290–1307, 2011. doi: 10.1038/nprot.2011. 308.