

V11 – DGL-Modelle / Copasi / SBML

Aufstellen von Bilanzgleichungen

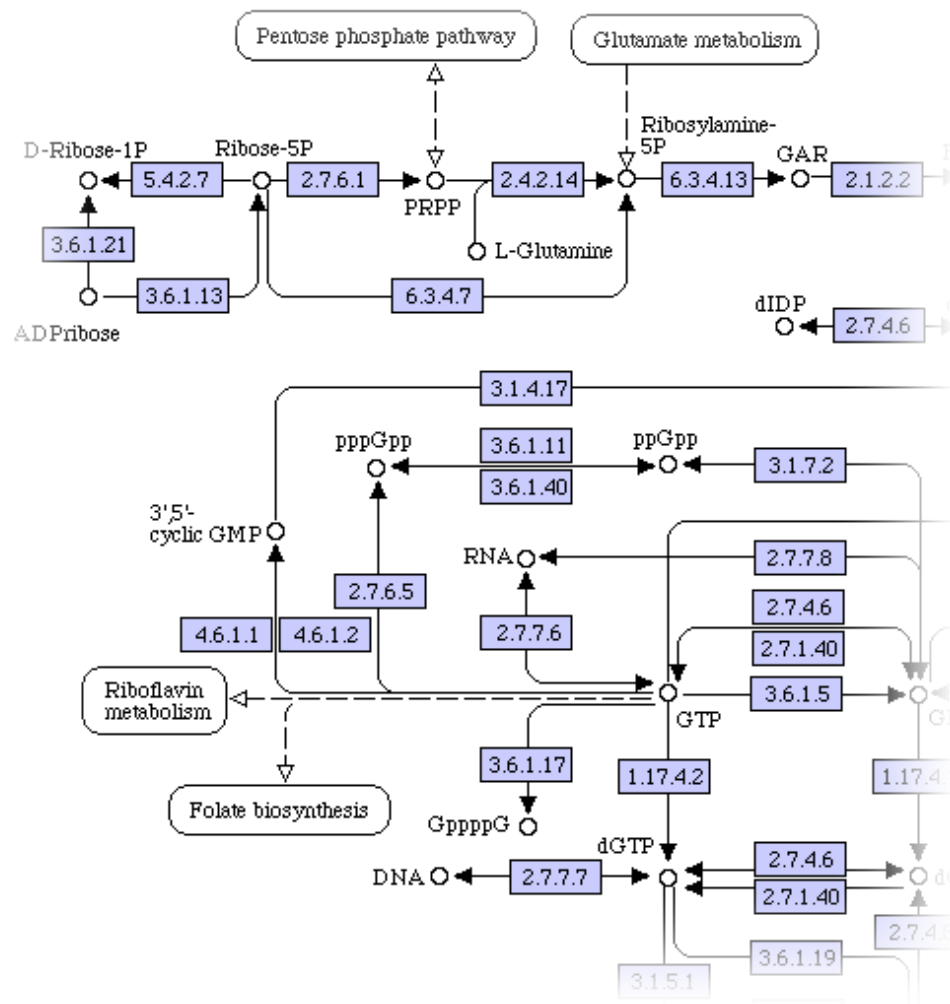
Dynamische Simulationen: Was ist das? Wozu?

Simulations-Tool: Copasi

Vereinfachte Kinetiken: MM, Inhibierung, Hill

kinetische Daten: KEGG, SABIO-RK

Wdh: über die Formel zur Formel

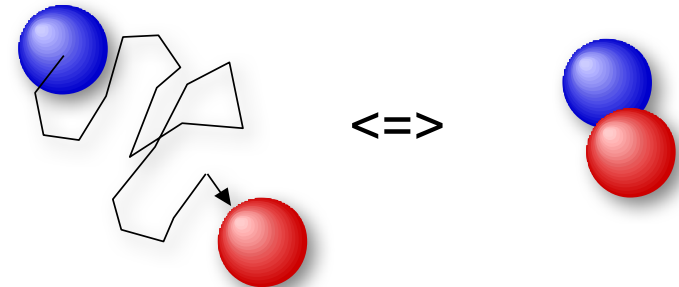
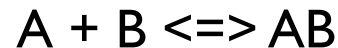


Ausschnitt aus http://www.genome.jp/dbget-bin/show_pathway?ec00230+3.6.1.9

- i) biologisches Netzwerk
- ii) Metabolite identifizieren
(incl. Konzentrationen)
- iii) Einzelreaktionen aufstellen,
Reaktionsraten?
- iv) Reaktionen in DGLs
übersetzen
- v) Anfangswerte einsetzen
und simulieren
- vi) Ergebnisse analysieren

Massenwirkungsgesetz

Einfachste chemische Reaktion



Zeitliche Änderung von [A]:

Gewinn: Dissoziation



AB zerfällt

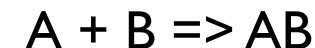
$\Rightarrow G_A$ proportional zu $[AB]$

$$G_A = k_r [AB]$$

phänomenologischer
Faktor

$$\frac{d}{dt}[A] = G_A - L_A$$

Verlust: Assoziation



A und B müssen sich finden

$\Rightarrow L_A$ abhängig von $[A]$ und $[B]$

$$L_A = k_f [A] [B]$$

$$\frac{d}{dt}[A] = k_r [AB] - k_f [A] [B]$$

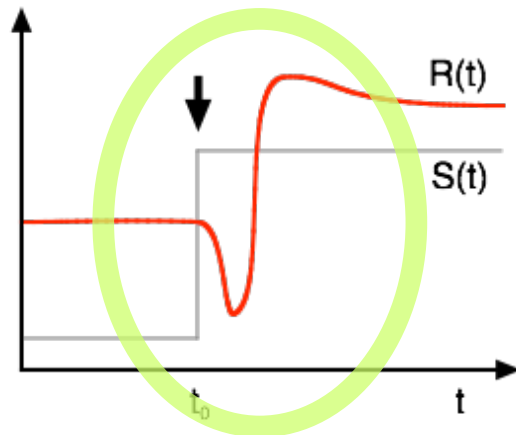
Dynamische Simulationen

Zwei Anwendungsgebiete

zeitabhängiges Verhalten

Reaktionen des Systems auf
Änderungen der äußeren Bedingungen
(Randbedingungen)

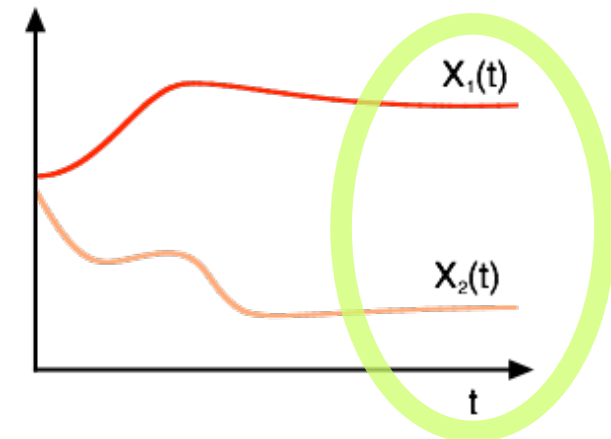
$$\frac{d\vec{X}(t)}{dt}$$



stationäre Zustände (steady state)

Suche nach Konzentrationen und
Flüssen bei konstanten
Randbedingungen

$$\frac{d\vec{X}(t)}{dt} = 0$$



Was lernt man aus dem Steady-State?

Konzentrationen = konst.

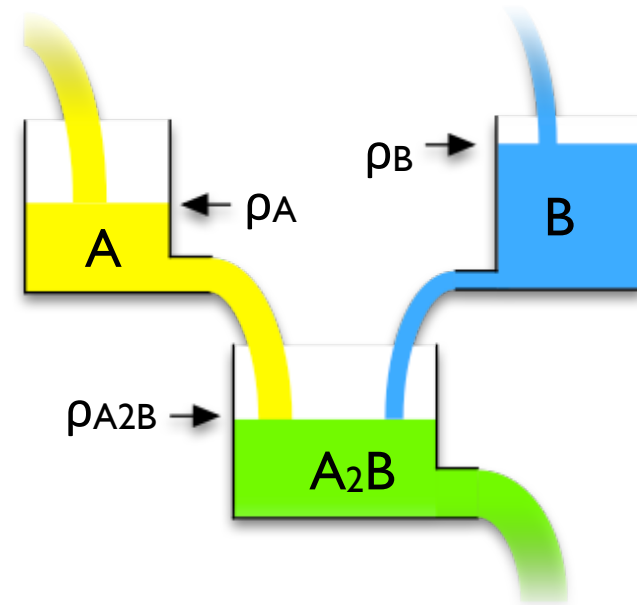
=> Zufluss = Abfluss

$$\frac{dA_2B(t)}{dt} = G_{A_2B} - L_{A_2B} = 0$$

Beispiel: $2A + B \rightleftharpoons A_2B$

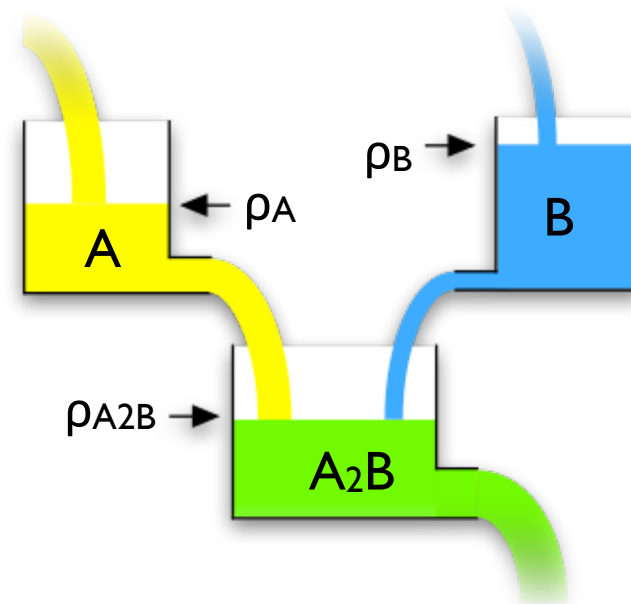
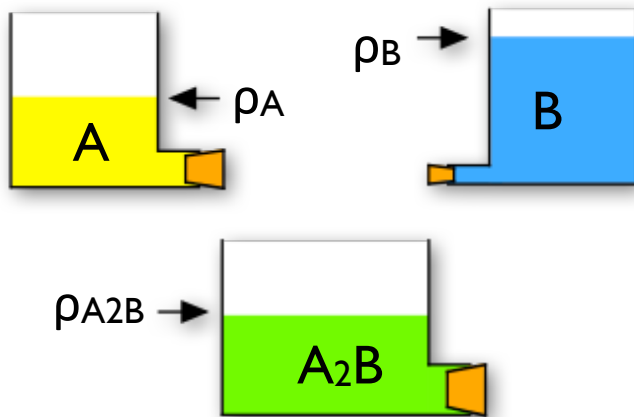
$$\frac{dA_2B(t)}{dt} = k_a A^2 B - k_d A_2B = 0$$

=> Gleichgewichts-Konzentrationen: $A_2B = \frac{k_a}{k_d} A^2 B$



- Steady state:
- + Bedingungen zwischen Konzentrationen und Raten
 - => stationäre Betriebsmodi
 - wie schnell wird der steady state erreicht?
 - absolute Mengen / effektive Volumina

Statisches vs. dynamisches Gleichgewicht



jeweils: $\frac{d\vec{X}(t)}{dt} = 0$

Infos aus zeitabhängigen Simulationen

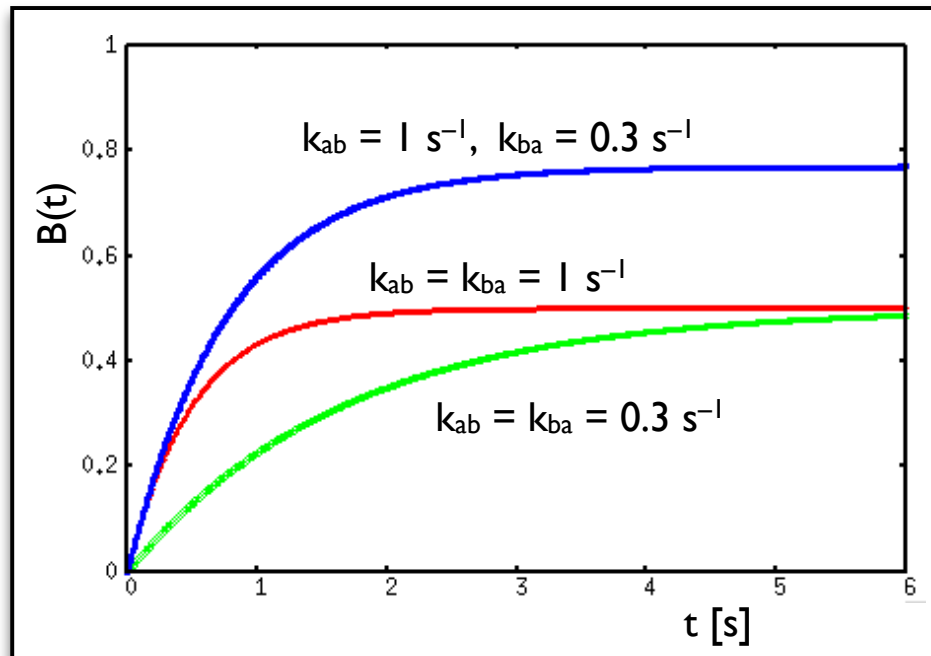
Ganz einfach: $A \rightleftharpoons B$

$$A + B = \text{const.}$$

Gleichgewicht:

$$\frac{dA(t)}{dt} = k_{ba}B - k_{ab}A$$

$$B = \frac{k_{ab}}{k_{ba}} A$$



mit Anfangsbedingungen:

$$A(t=0) = A_0$$

$$B(t=0) = 0$$

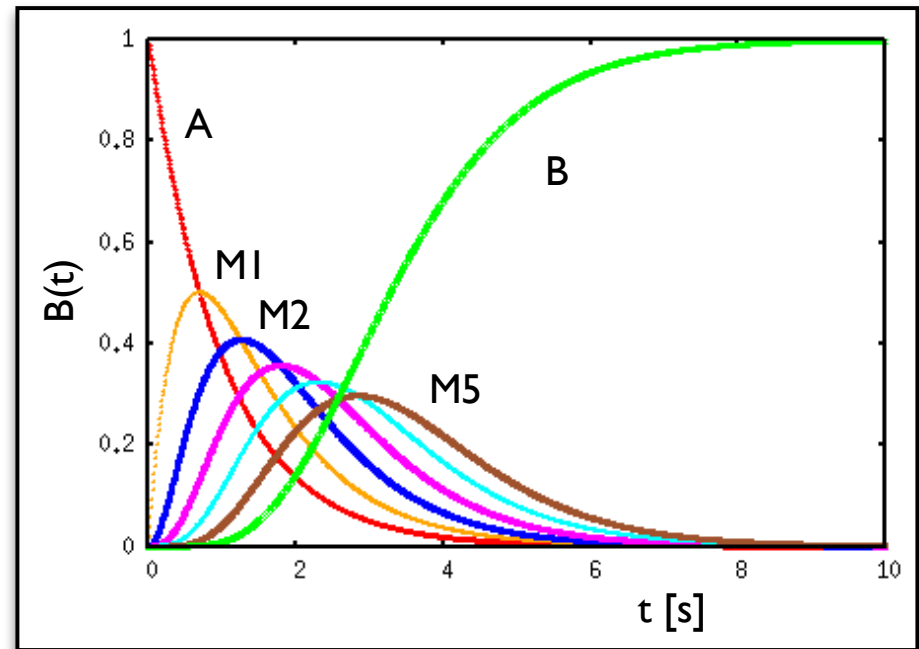
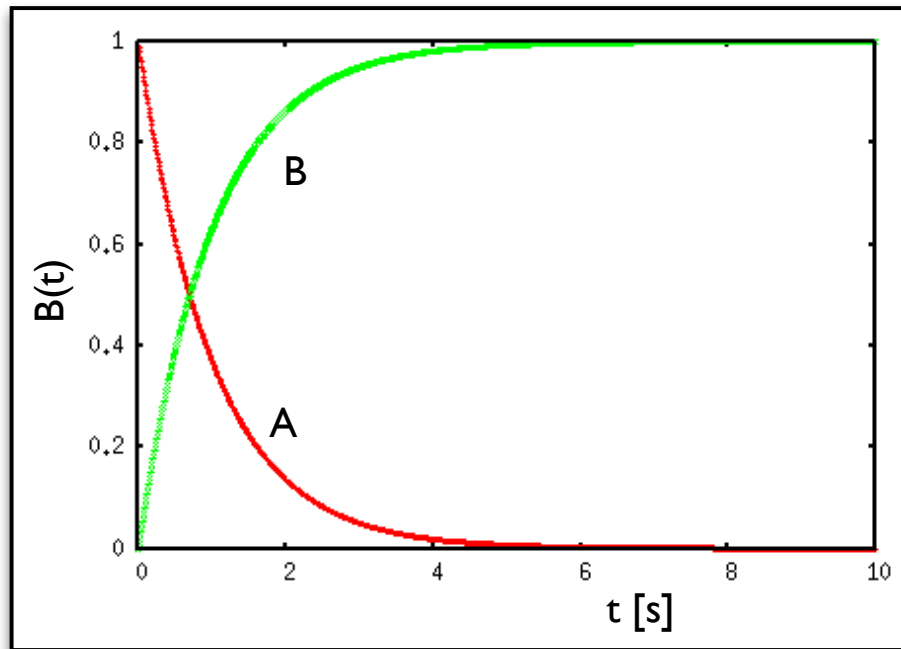
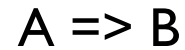
$$B(t \rightarrow \infty) = \frac{k_{ab}}{k_{ab} + k_{ba}} A_0$$

=> Gleichgewichtsverteilungen

=> wie schnell wird ein "Signal" weitergegeben?

Länge von Reaktionspfaden

Vergleiche:



$$A_0 = 1, \quad k = 1 \text{ s}^{-1}$$

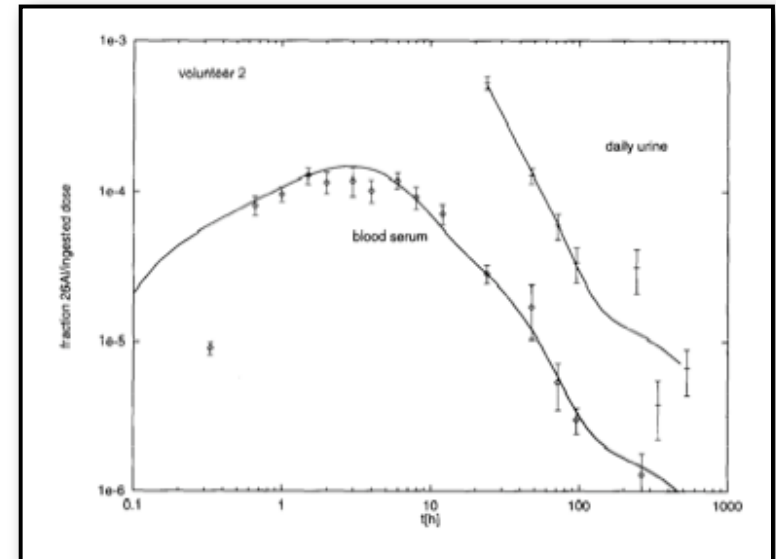
=> Zwischenprodukte verzögern die Antwort
=> Vorsicht beim Weglassen von Zwischenschritten

Puffer: Al-Metabolismus

Al ist das dritthäufigste Element (8%) und das häufigste Metall in der Erdkruste. Normalerweise ist Al harmlos, kann aber auch zu Osteopathie, Anämie oder Enzephalopathie führen.

Experimente zur Al-Aufnahme und -Ausscheidung:

- orale Gabe von 100 ng ^{26}Al ($T_{1/2} = 0.7 \text{ Myr}$)
- Blutproben nach 20 min, 40 min, ..., 46 d
- Tagesurin
- Messung der ^{26}Al -Menge



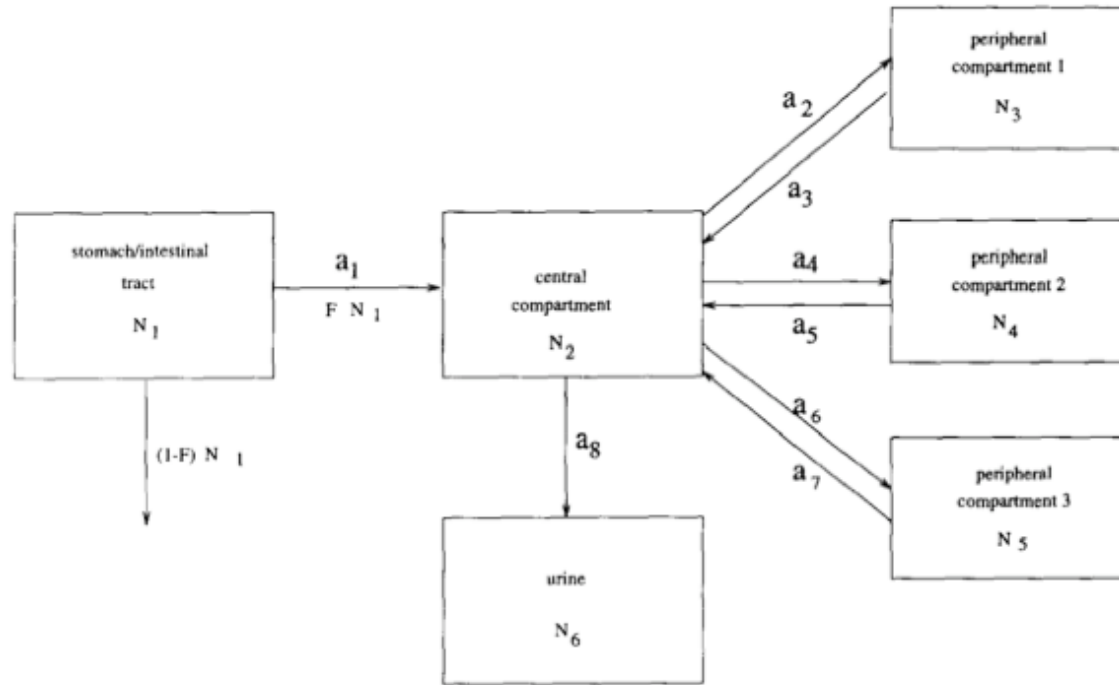
Messwerte: Blut- und Urinproben, Gewebeproben bei Ratten

=> zeitabhängige Verteilung und Speicherung in verschiedenen Geweben

=> Modellierung als Multi-Kompartiment-Modell

Modellierung des AL-Metabolismus

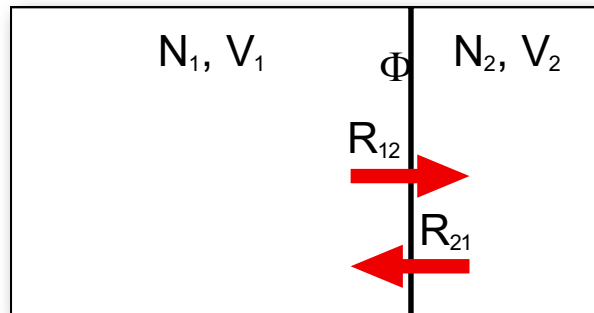
- i) AI wird aufgenommen (oral oder intravenös), kommt ins Blut
- ii) AI verteilt sich vom Blut in das umliegende Gewebe/Organe
- iii) dynamisches Gleichgewicht zwischen Blut und peripheren Gewebe-Speichern
- iv) Blut wird über Leber/Niere ausgeschieden



Hohl, ..., Nolte, Ittel, *Nucl. Inst. Meth. B* **92**
(1994) 478

Erhalte Übergangsraten zwischen (Lebensdauern) und Volumina der Kompartimente aus der Simulation durch Parameterfit
=> Hilfe für die physiologische Zuordnung

Unterschiedlich große Kompartimente



Teilchenaustausch durch Interface der Fläche Φ :

$$\frac{dN_{12}}{dt} = k_{12} \Phi \frac{N_1}{V_1} \quad \frac{dN_{21}}{dt} = k_{21} \Phi \frac{N_2}{V_2}$$

Änderungen der Anzahlen (Gesamtanzahl bleibt erhalten):

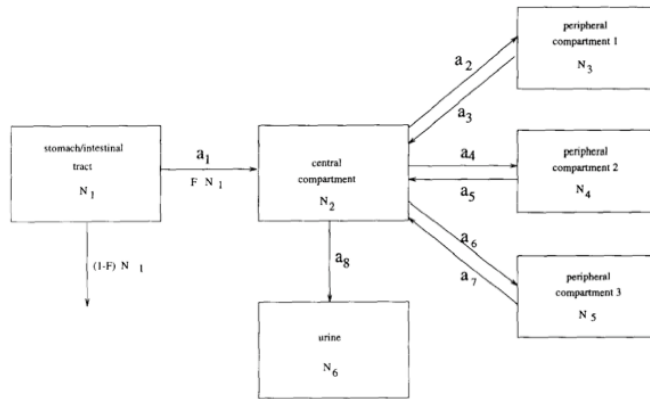
$$\frac{dN_1}{dt} = -\frac{dN_{12}}{dt} + \frac{dN_{21}}{dt} \quad \frac{dN_2}{dt} = -\frac{dN_{21}}{dt} + \frac{dN_{12}}{dt}$$

Änderungen der entsprechenden Dichten:

$$\frac{dN_1}{dt V_1} = \frac{1}{V_1} \frac{dN_1}{dt} = \frac{\tilde{k}_{21} N_2}{V_1 V_2} - \frac{\tilde{k}_{12} N_1}{V_1 V_1} \quad \frac{dN_2}{dt V_2} = \frac{V_1}{V_2} \frac{dN_1}{dt V_1}$$

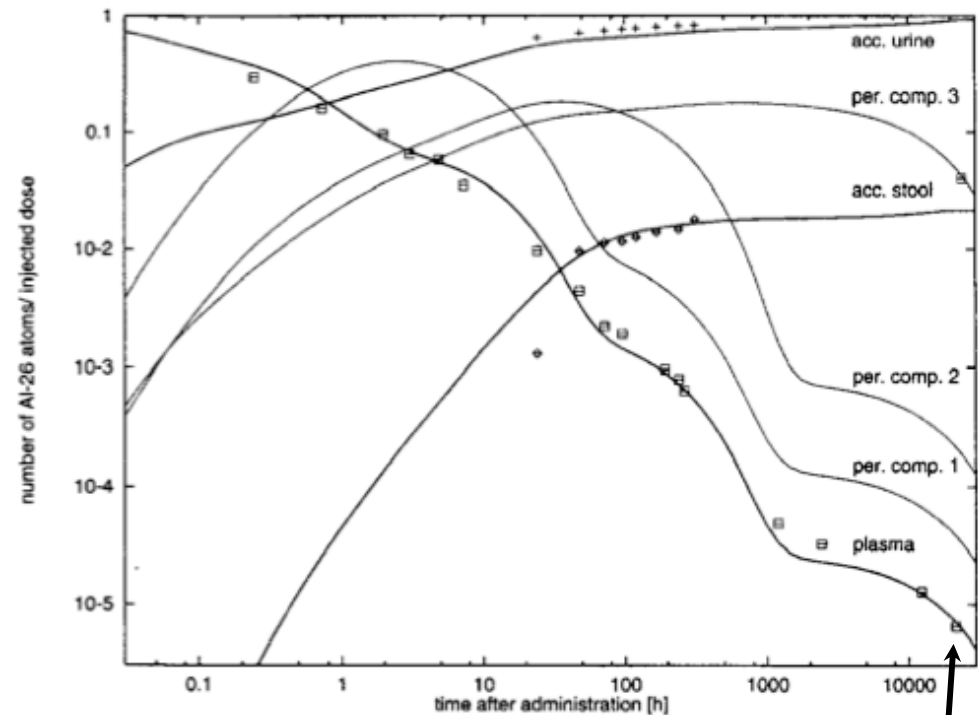
=> Simulationen mit Teilchenzahlen, Dichten "on the fly"

Ergebnisse



Drei Gewebetypen
(Kompartimente) reichen, um
die Messwerte zu beschreiben
=> schnelles, mittleres und
langsameres Kompartiment

Zeitabh. Verhalten bestimmt von
Volumen *und* Austauschraten.



2.3a

²⁶Al konnte nach mehr als zwei Jahren
immer noch im Blut nachgewiesen werden
=> Speicherung in den Knochen

Complex Pathway Simulator



Entwickelt in den Gruppen von
Pedro Mendes (Virginia Bioinf. Inst.) und Ursula Kummer (EML HD)

"COPASI is a software application for simulation
and analysis of biochemical networks."

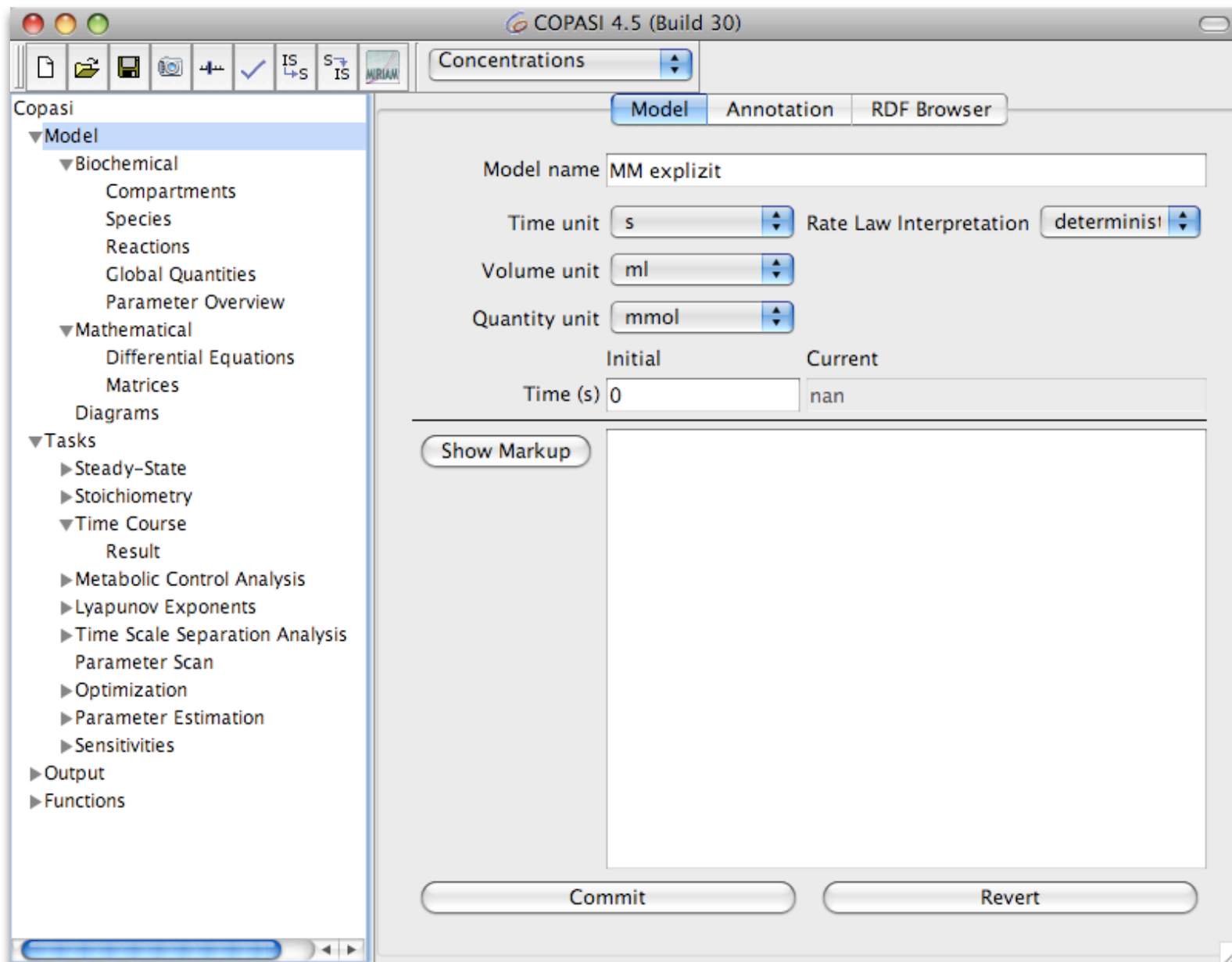
<http://www.copasi.org/>

Copasi-Features

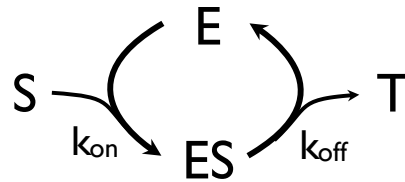
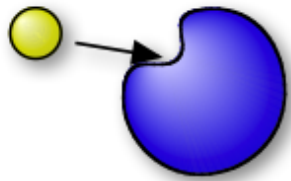
Current Features:

- Model:
 - Chemical reaction network.
 - Arbitrary kinetic functions.
 - ODEs for compartments, species, and global quantities.
 - Assignments for compartments, species, and global quantities.
 - Initial assignments for compartments, species, and global quantities.
- Analysis:
 - Stochastic and deterministic time course simulation
 - Steady state analysis (including stability).
 - Metabolic control analysis/sensitivity analysis.
 - Elementary mode analysis .
 - Mass conservation analysis.
 - Time scale separation analysis
 - Calculation of Lyapunov exponents.
 - Parameter scans.
 - Optimization of arbitrary objective functions.
 - Parameter estimation using data from time course and/or steady state experiments simultaneously.
- Graphical User Interface (CopasiUI)
 - Sliders for interactive parameter changes.
 - Plots and Histograms.
- Command Line (CopasiSE) for batch processing.
- [SBML](#) import (L1V1+2, L2V1-3) and export (L1V2, L2V1-3).
- Loading of [Gepasi](#) files.
- Export to Berkeley Madonna, XPPAUT, and C source code of the ODE system generated from the model.
- Versions for MS Windows, Linux, Mac OS X, and Solaris SPARC.

We keep a list of currently [known problems](#) in COPASI.

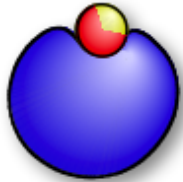


Enzyme: Michaelis-Menten-Kinetik



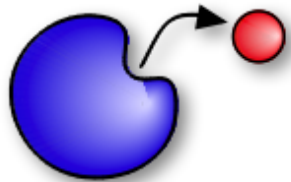
Reaktionsrate:

$$V = k_{\text{off}} ES$$



Steady state:

$$k_{\text{on}} E \cdot S = k_{\text{off}} ES$$

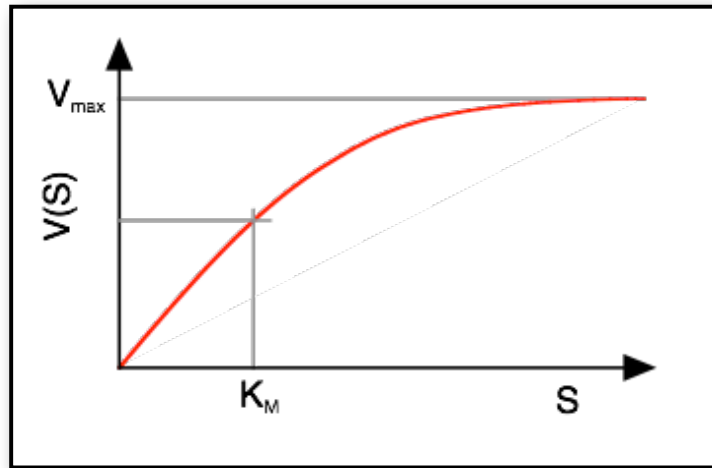


$$ES = \frac{k_{\text{on}} E \cdot S}{k_{\text{off}}} = \frac{E \cdot S}{K_M}$$

Die Gleichung

Effektiver Umsatz nach MM: $V = V_{max} \frac{S}{S + K_M}$

$$V_{max} = k_{off} E_T$$



$$K_M = \frac{k_{off}}{k_{on}}$$

- Vorteile:
- analytische Formel für den Umsatz
 - Interpretation der Kennlinie: V_{max} , K_M
 - Enzym kann ignoriert werden

Aber: weniger kinetische Informationen
 $k_{on}, k_{off}, E_T \Rightarrow V_{max}, K_M$

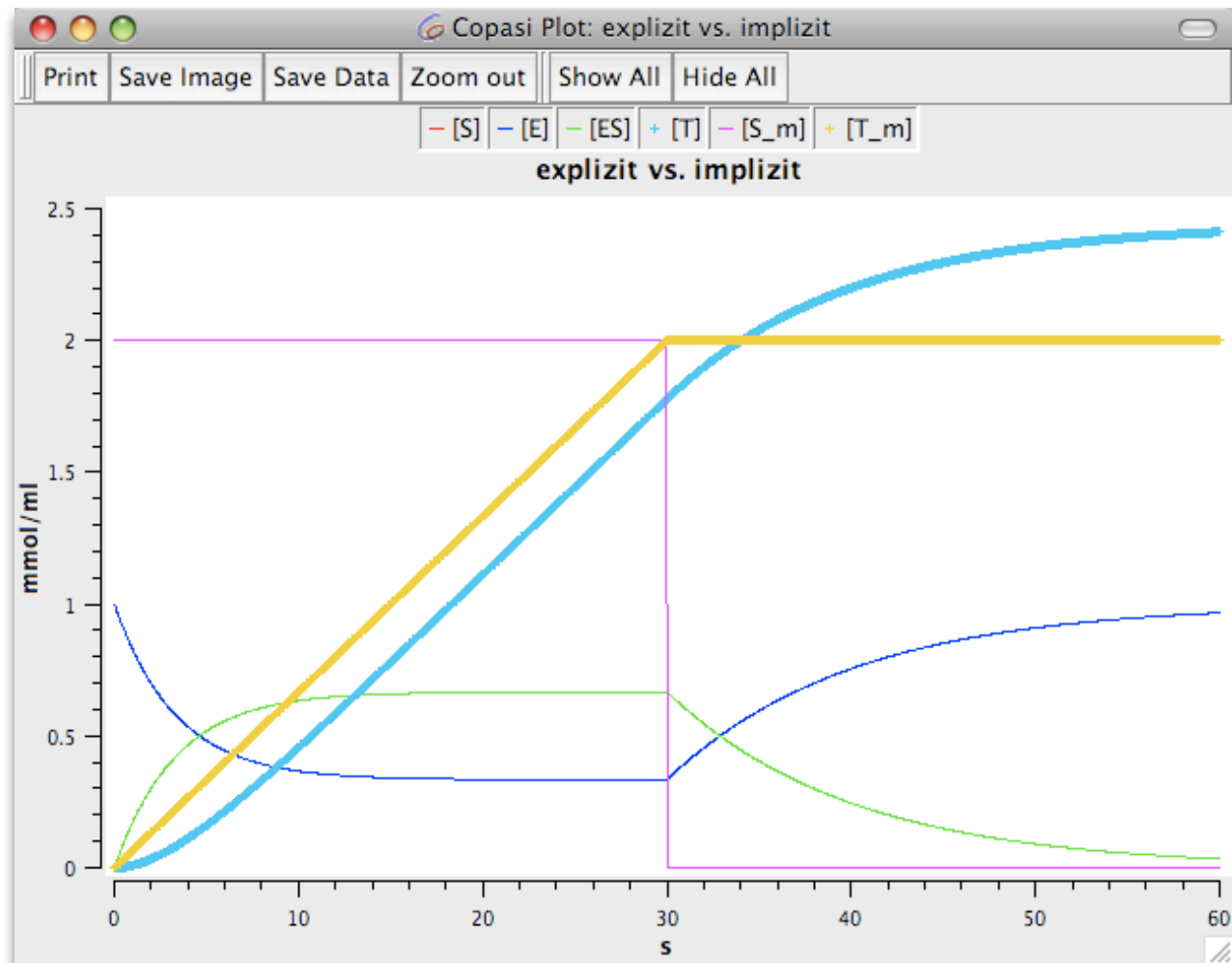
MM vs. explizite Modellierung

Wenn E verschiedene Substrate katalysiert
=> MM geht nicht

Zeitverhalten:
MM-Kinetik vs.
explizite Modellierung

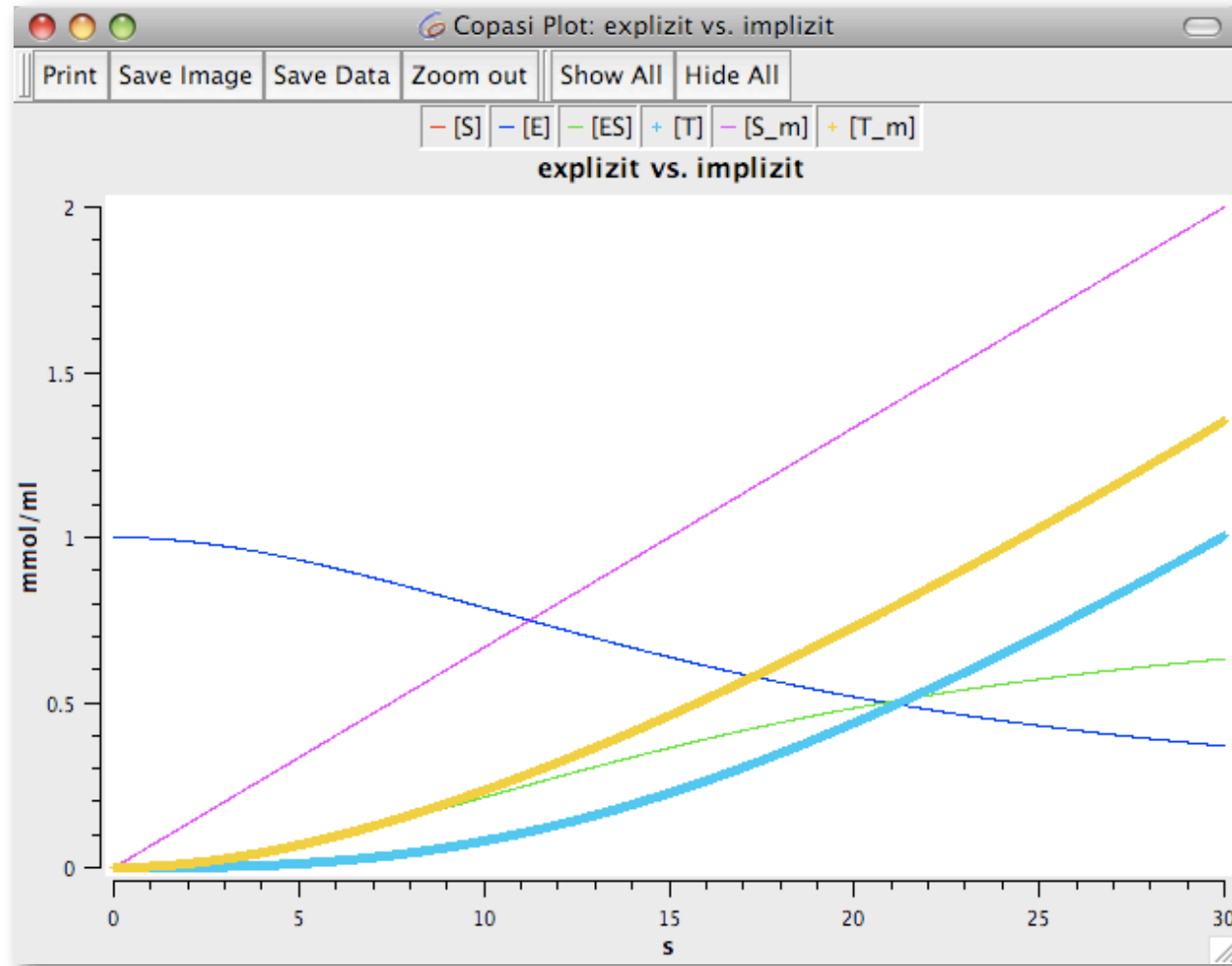
=> Einschwingen

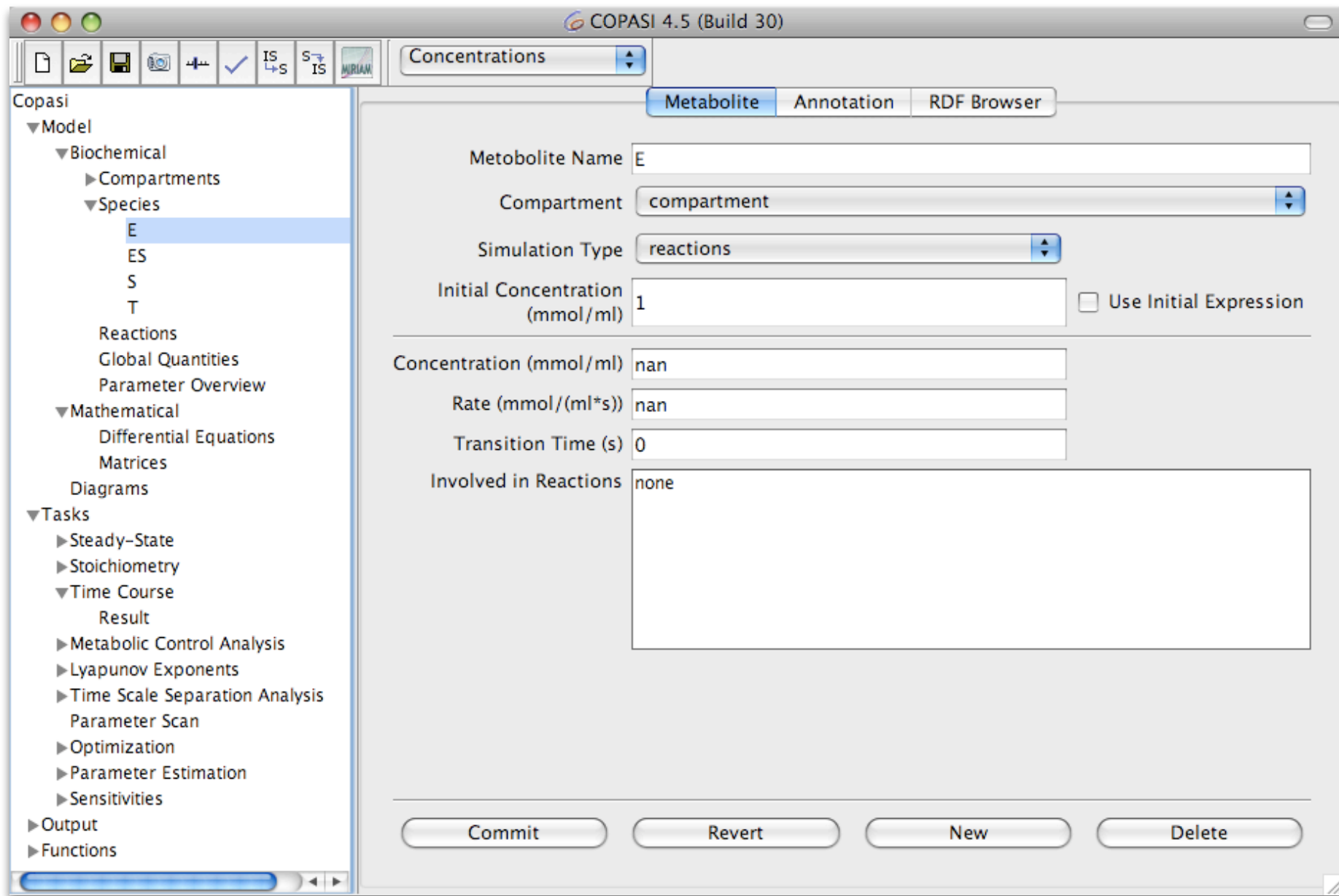
=> anderer
Gesamtumsatz



Nochmal: explizit vs. MM

linearer Anstieg von S





COPASI 4.5 (Build 30)

Concentrations

Reaction Annotation RDF Browser

Name: R1

Chemical Equation: $E + S = ES$

☒ Reversible ☐ Multi Compartment

Rate Law: Mass action (reversible) New Rate Law

Flux (mmol/s): 0

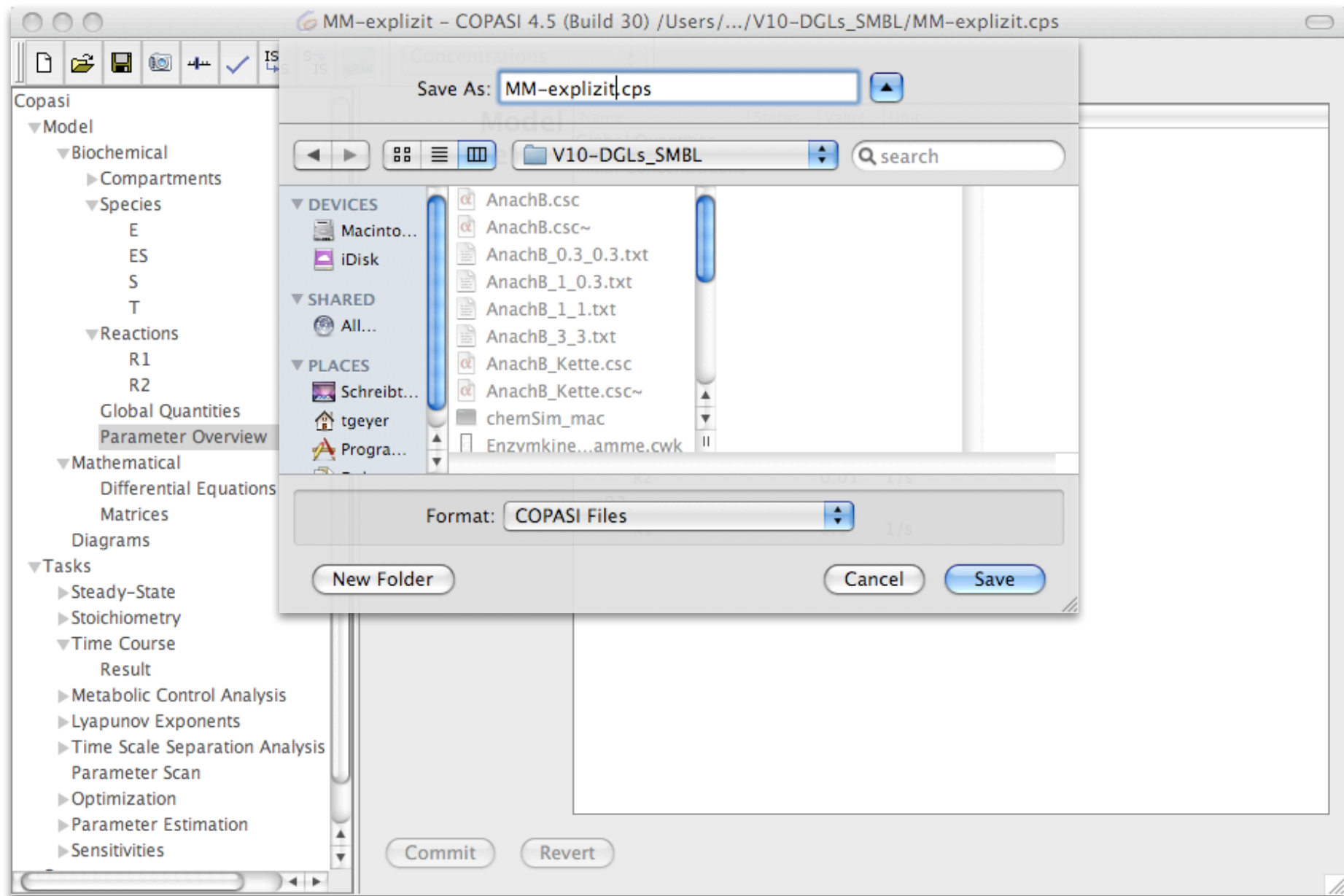
Symbol Definition

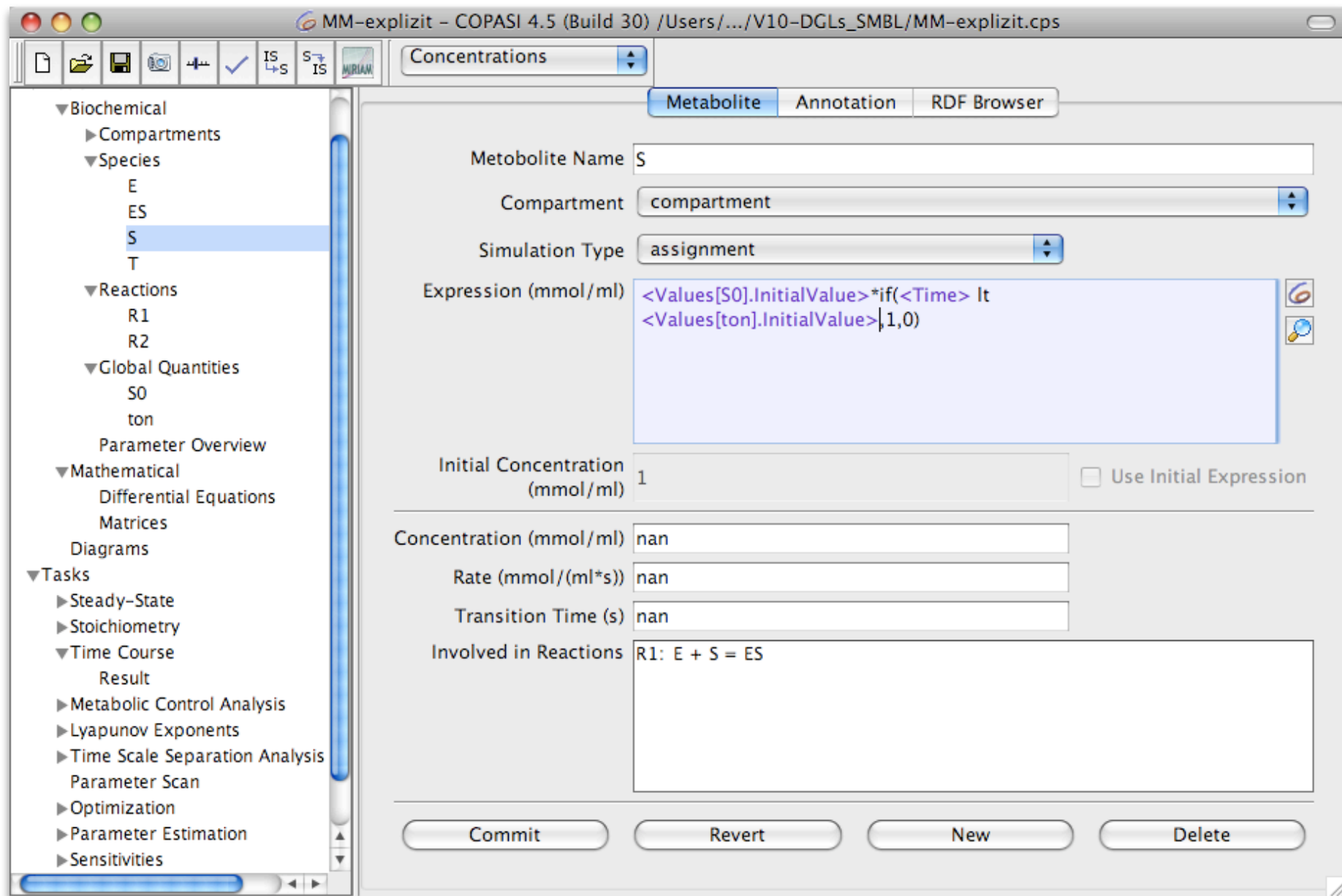
Description	Name	Value	Unit
Parameter	k1	<input type="checkbox"/> global	0.1 ml/(mmol*s)
→ Substrate	substrate		mmol/ml
	E		
	S		
Parameter	k2	<input type="checkbox"/> global	0.01 1/s
→ Product	product		mmol/ml
	ES		

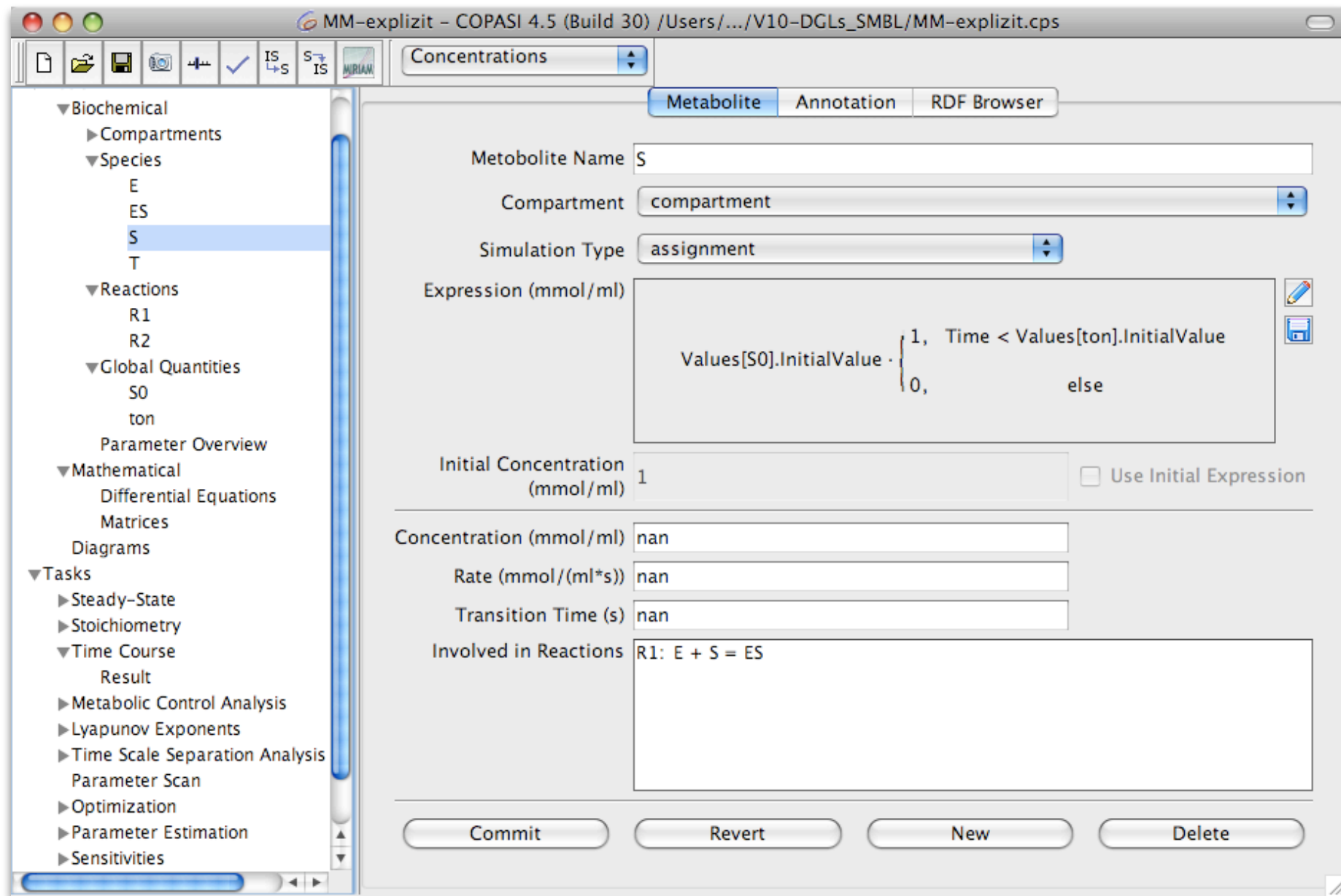
Commit Revert New Delete

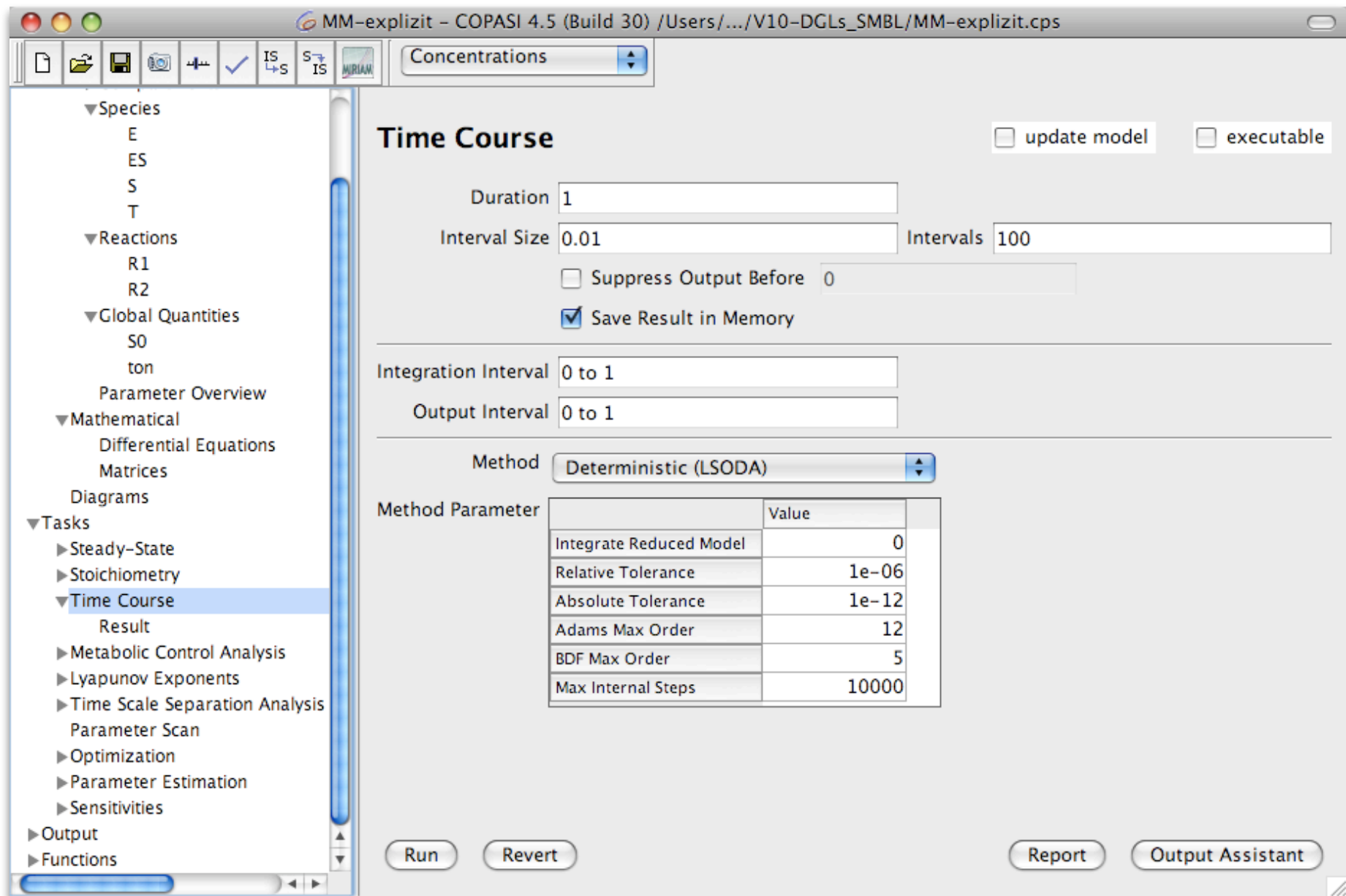
Copasi

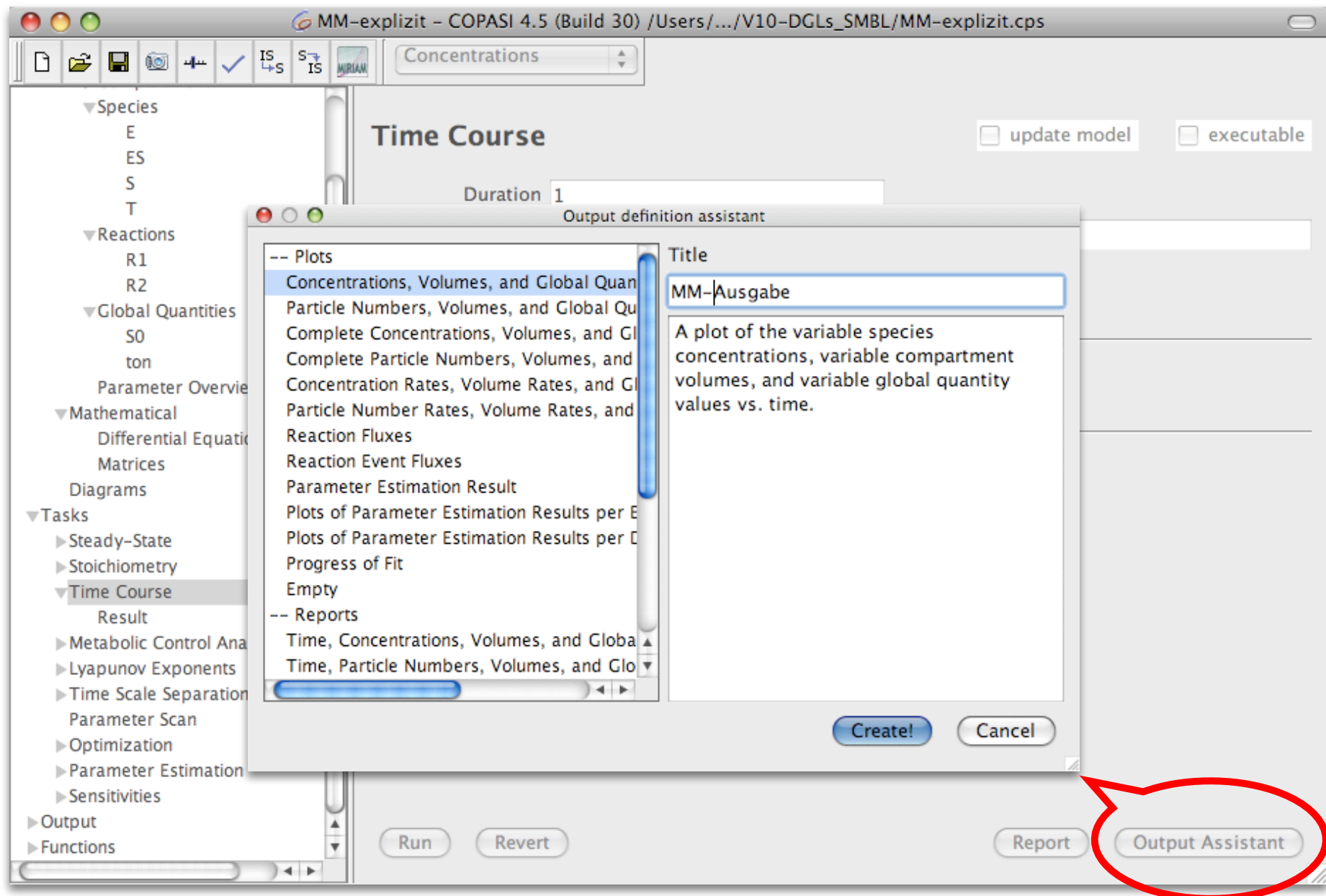
- Model
 - Biochemical
 - Compartments
 - Species
 - E
 - ES
 - Es
 - S
 - T
 - Reactions
 - R1
 - R2
 - Global Quantities
 - Parameter Overview
 - Mathematical
 - Differential Equations
 - Matrices
 - Diagrams
 - Tasks
 - Steady-State
 - Stoichiometry
 - Time Course
 - Result
 - Metabolic Control Analysis
 - Lyapunov Exponents
 - Time Scale Separation Analysis
 - Parameter Scan
 - Optimization
 - Parameter Estimation

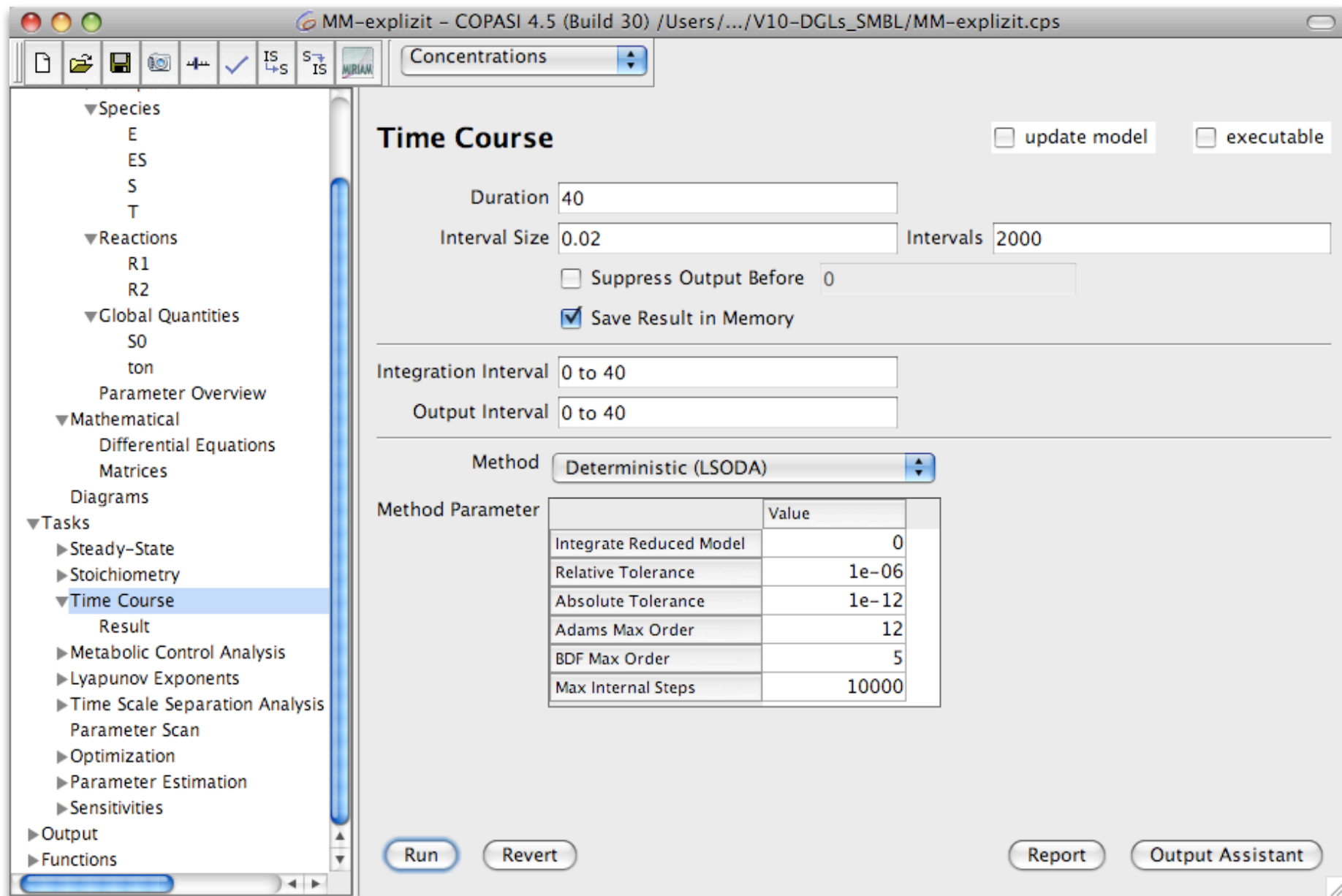


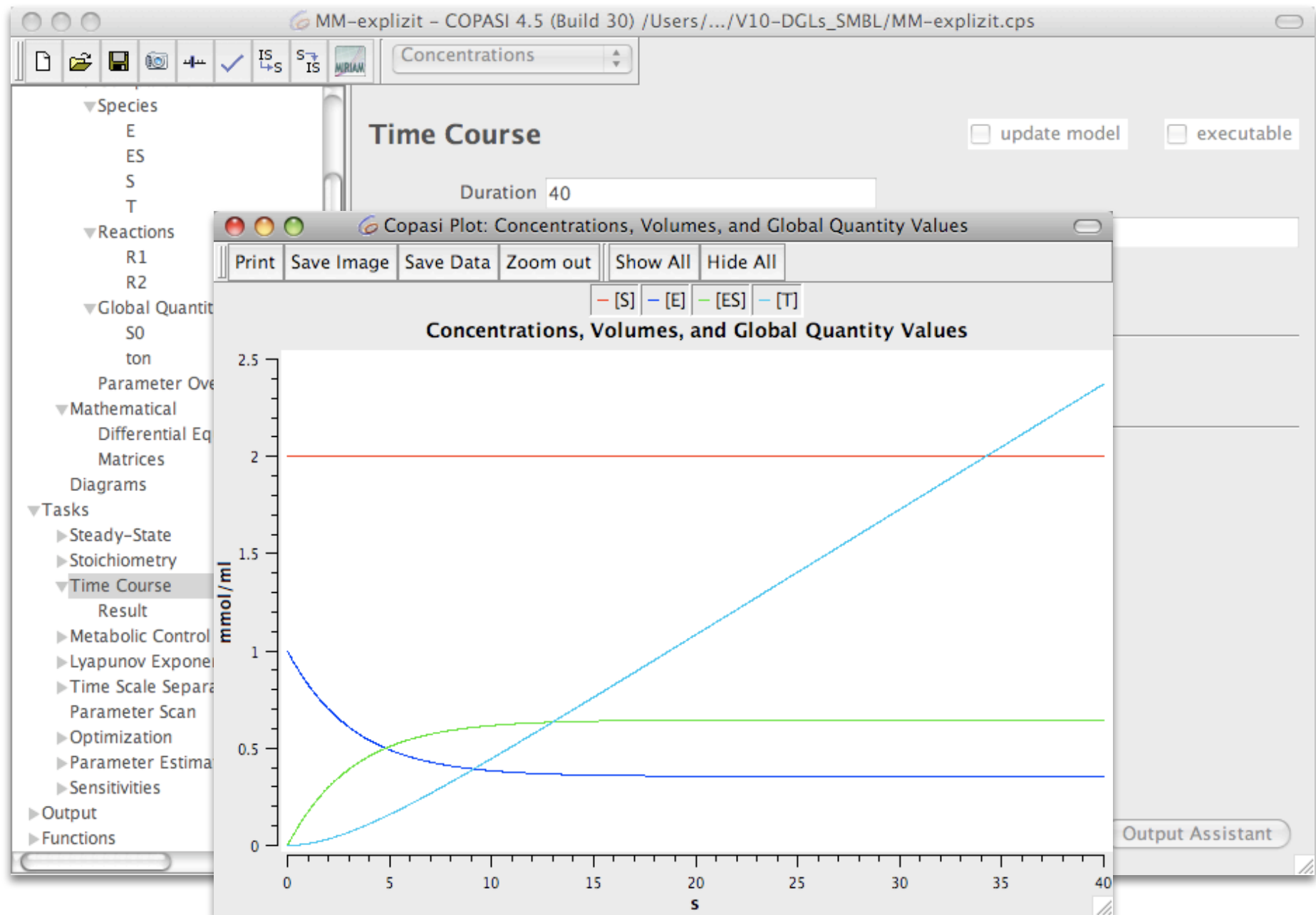


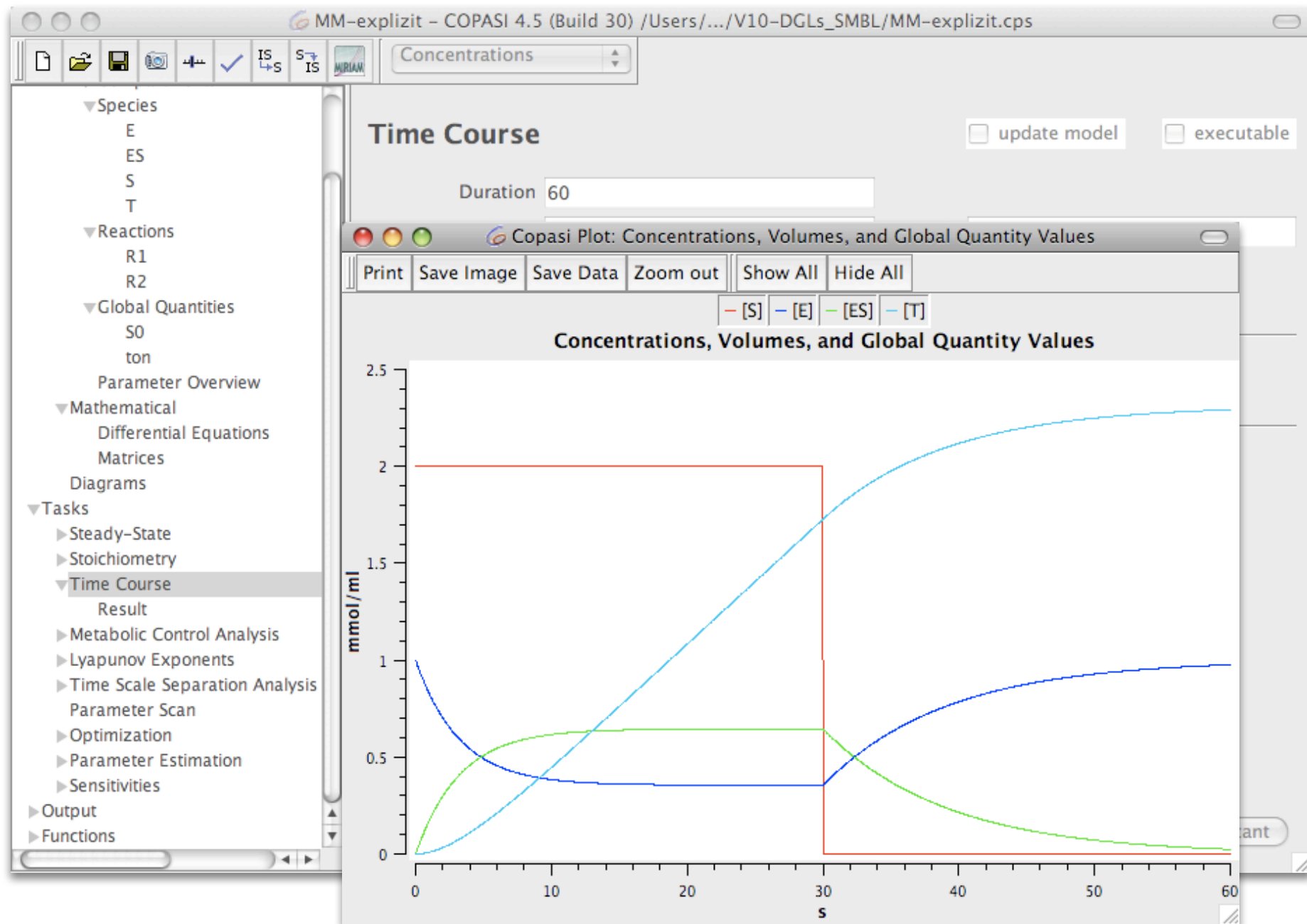


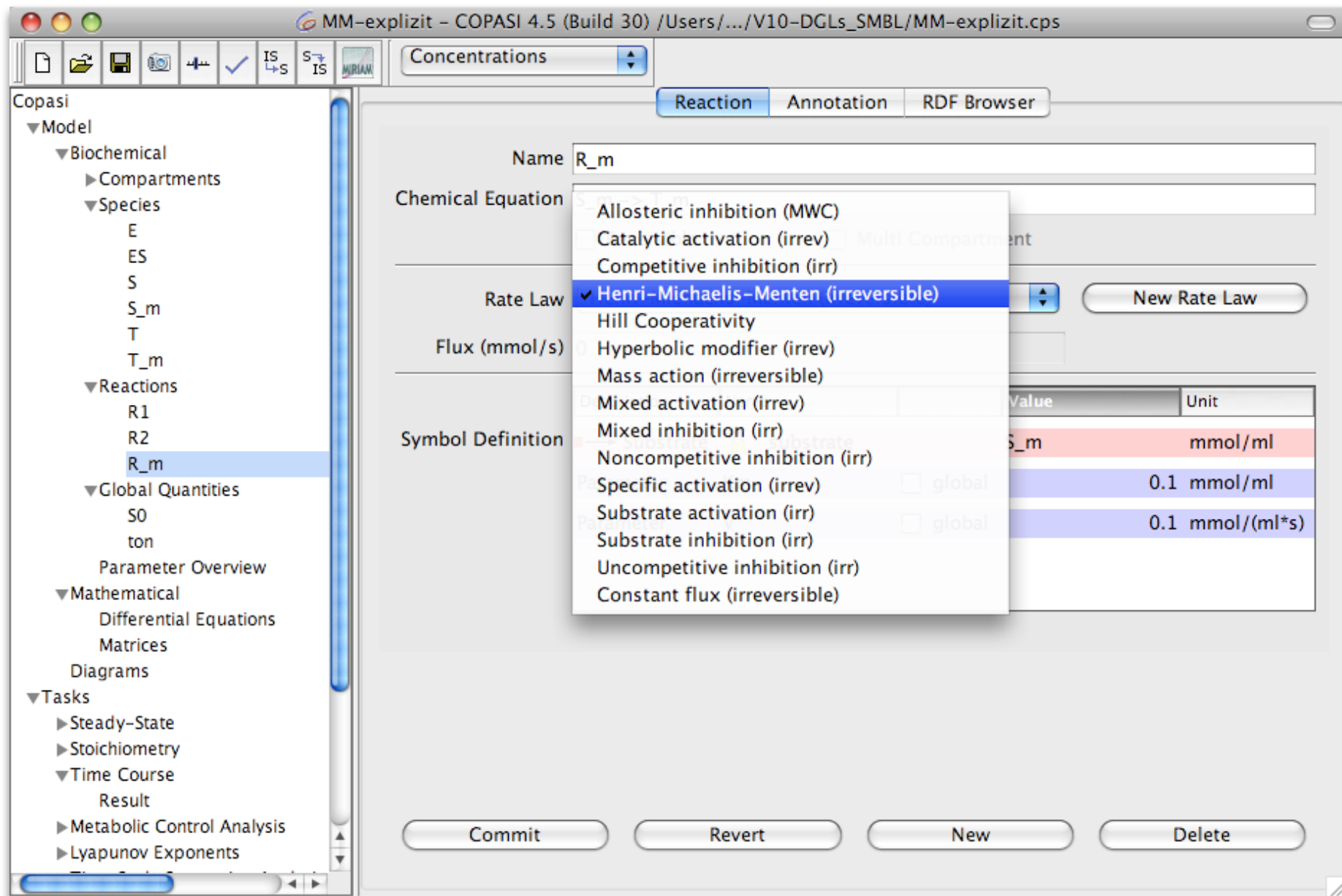








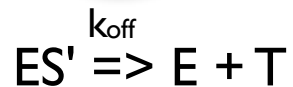
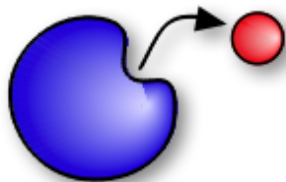
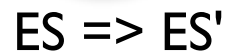
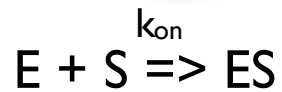
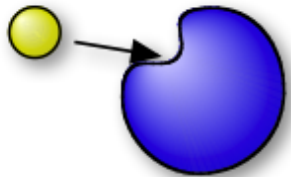




Vereinfachte Kinetiken

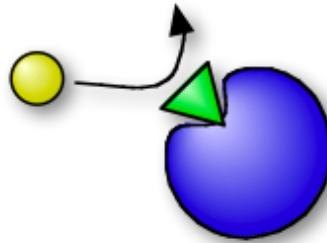
Enzymreaktion:

Michaelis-Menten



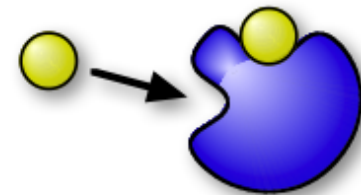
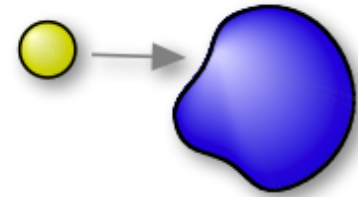
kompetitive Inhibition:

Inhibitor vs. Substrat



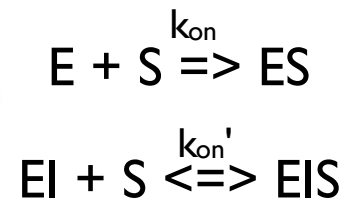
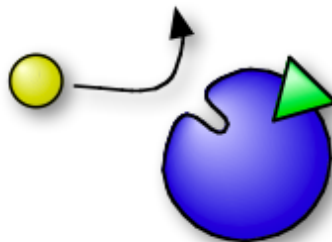
Kooperative Bindung:

Hill-Kinetik

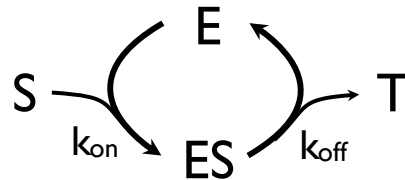
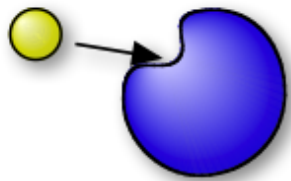


nicht-kompetitive Inhibition:

Inhibitor verändert Enzym



Enzyme: Michaelis-Menten-Kinetik



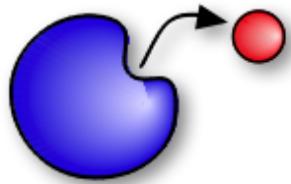
Reaktionsrate:

$$V = k_{\text{off}} ES$$



Steady state: $k_{\text{on}} E \cdot S = k_{\text{off}} ES$

$$ES = \frac{k_{\text{on}} E \cdot S}{k_{\text{off}}} = \frac{E \cdot S}{K_M}$$



Gesamtmenge an Enzym ist konstant:

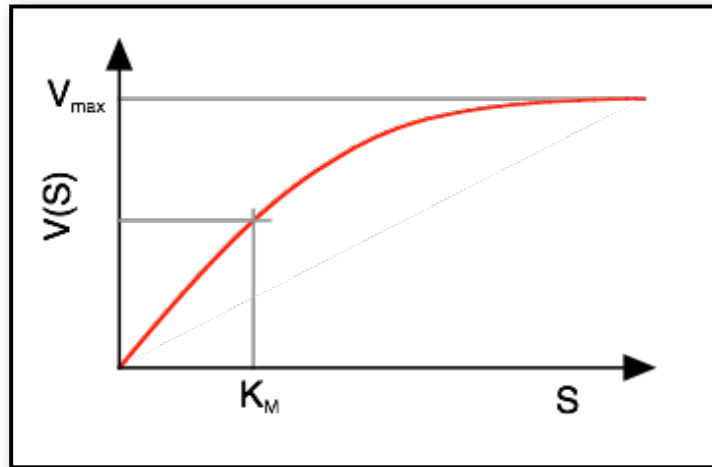
$$E_T = E + ES \quad \Rightarrow \quad ES = E_T \frac{S}{S + K_M}$$

Umsatz: $V = V_{\text{max}} \frac{S}{S + K_M}$

Die Gleichung

Effektiver Umsatz nach MM: $V = V_{max} \frac{S}{S + K_M}$

$$V_{max} = k_{off} E_T$$



$$K_M = \frac{k_{off}}{k_{on}}$$

- Vorteile:
- analytische Formel für den Umsatz
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Aber: weniger kinetische Informationen
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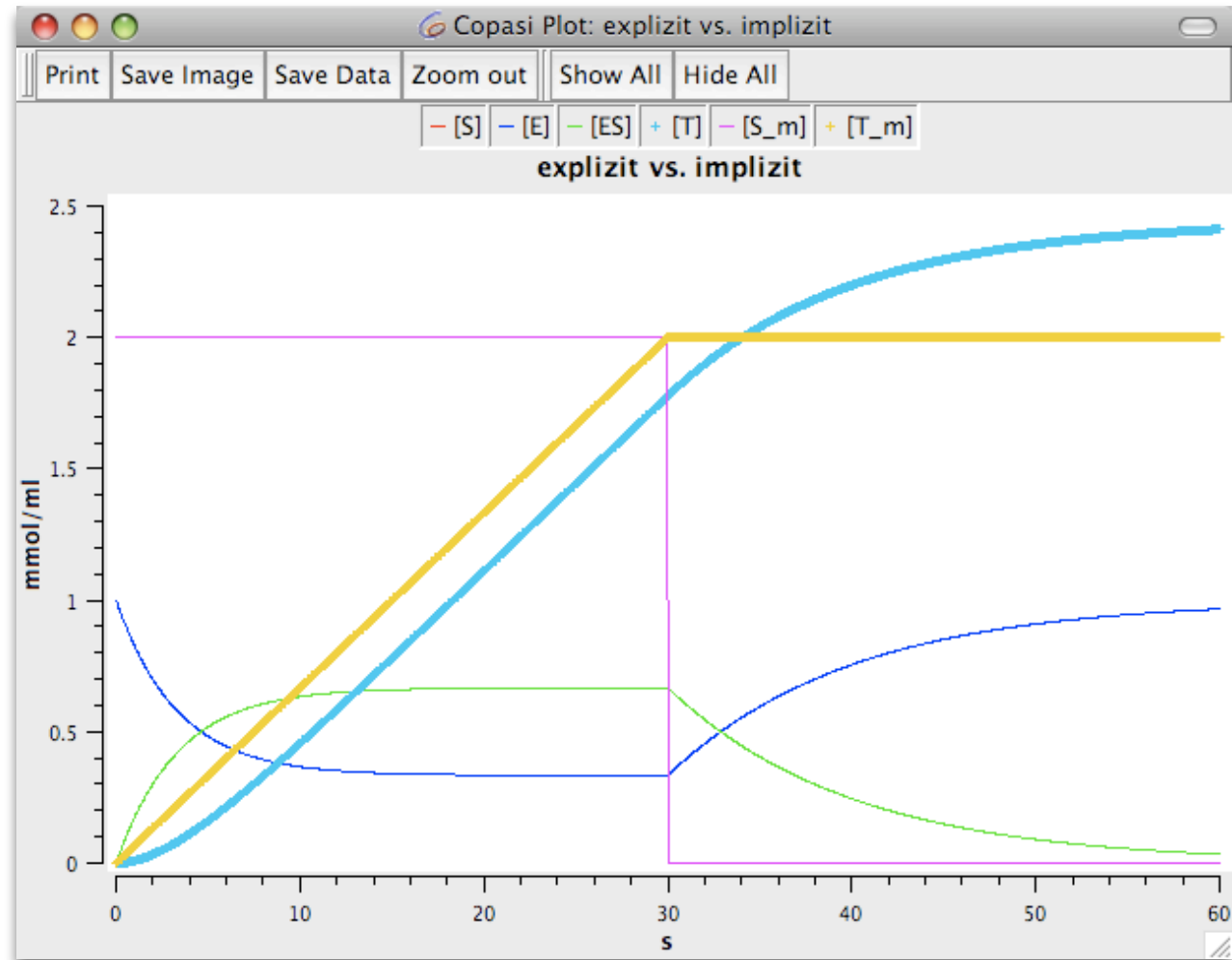
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Wenn E verschiedene Substrate katalysiert
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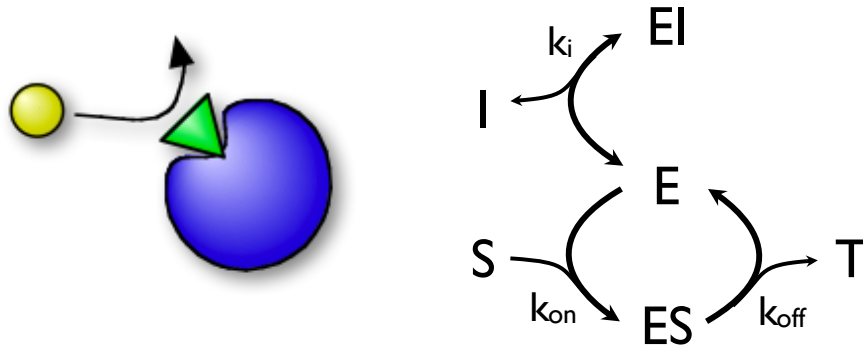
Zeitverhalten:
MM-Kinetik vs.
explizite Modellierung

=> Einschwingen

=> anderer
Gesamtumsatz



Kompetitive Hemmung



Zwei Pfade:

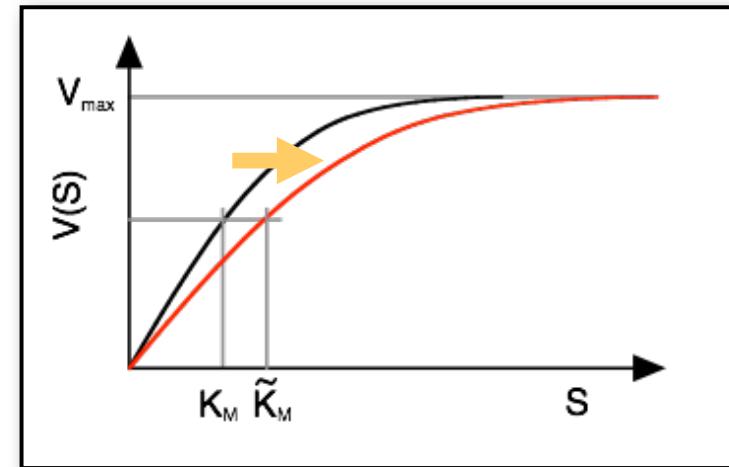


=> I verdrängt S

$S \ll I$: weniger freies E verfügbar
=> weniger ES
=> V reduziert

$$\tilde{K}_M = K_M (1 + I/K_I)$$

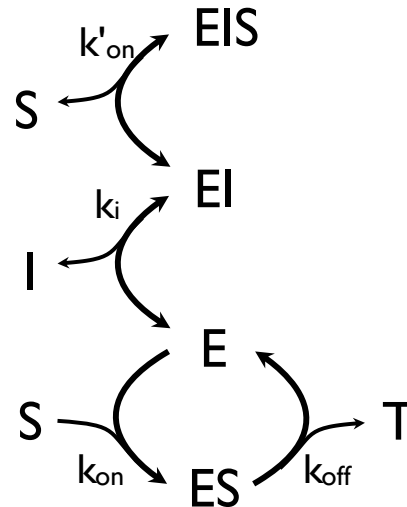
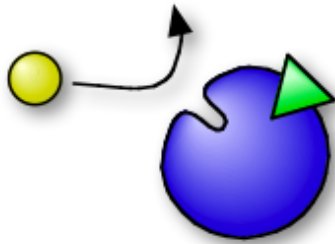
$S \gg I$: S verdrängt I
=> Inhibition unterdrückt
=> V_{max} unverändert



$$V = V_{max} \frac{S}{S + K_M (1 + I/K_I)}$$

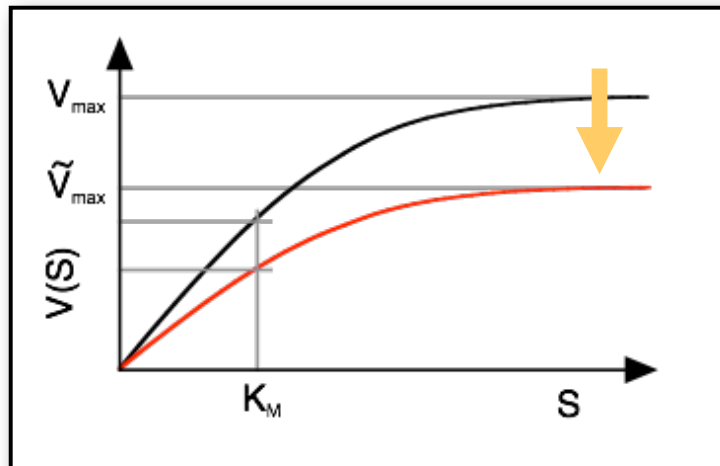
Nichtkompetitive Inhibition

Inhibitor blockiert Enzym



=> I reduziert effektives E_T

$$\Rightarrow \tilde{V}_{\max} = \frac{V_{\max}}{1 + I/K_I}$$



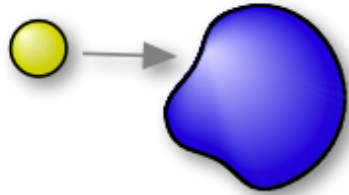
Analytische Formeln

=> Wirkungsweise von I aus steady state

Anzahl Parameter:

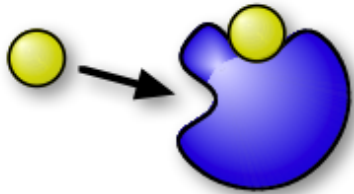
- explizit: $k_{\text{on}}, k_{\text{off}}, E_T, k_{i,\text{on}}, k_{i,\text{off}}, k'_{\text{on}}, k'_{\text{off}}$
- effektiv: V_{\max}, K_M, K_i

Kooperativität: Hill-Kinetik



Archibald Hill (1913): "Bindung des ersten Metaboliten vereinfacht Bindung des/der nächsten."

Wurde formuliert um die kooperative Bindung von Sauerstoff an Hämoglobin zu erklären ($n = 2.8 \dots 3.0$)



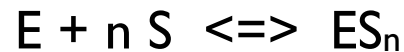
Zum Vergleich: $E + S \rightleftharpoons ES$

$$K = \frac{E \cdot S}{ES}$$

$$Y = \frac{ES}{E + ES} = \frac{S}{S + K}$$

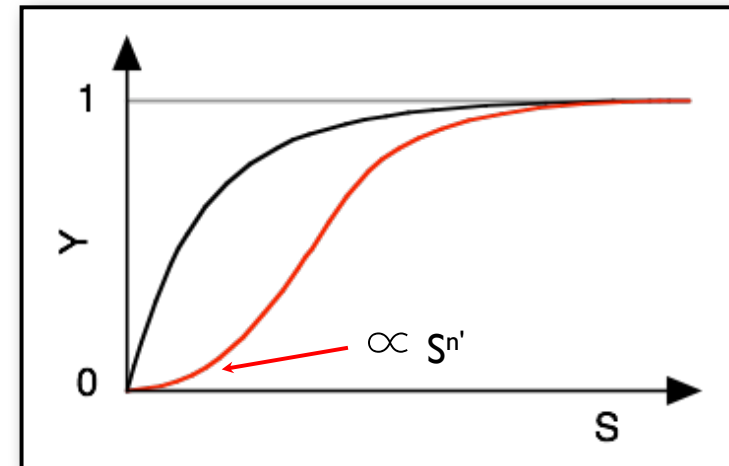
Anteil an besetzten Bindungstaschen

Mehrere Substrat-Moleküle gleichzeitig:



$$Y = \frac{S^{n'}}{S^{n'} + K^{n'}}$$

Hill-Koeffizient: $1 \leq n' \leq n$



Wann effektive Kinetiken?

Pro:

- weniger Aufwand
(Modell, Simulation, Parameter)
- analytische Lösungen für
einfache Systeme
- korrekter Steady state

Contra:

- weniger Parameter
=> weniger kinetische Informationen
- falsches dynamisches Verhalten

"Effektive Kinetiken brauchbar für langsame Signale"

"langsam" = Relaxationszeiten aller Zwischenschritte deutlich
kürzer als Änderungen des Signals

Woher bekommt man die Daten?

- Experten fragen
- Originalartikel lesen
- lesen lassen:
 - => Student, HiWi
 - => Datenbanken

Pfade: KEGG

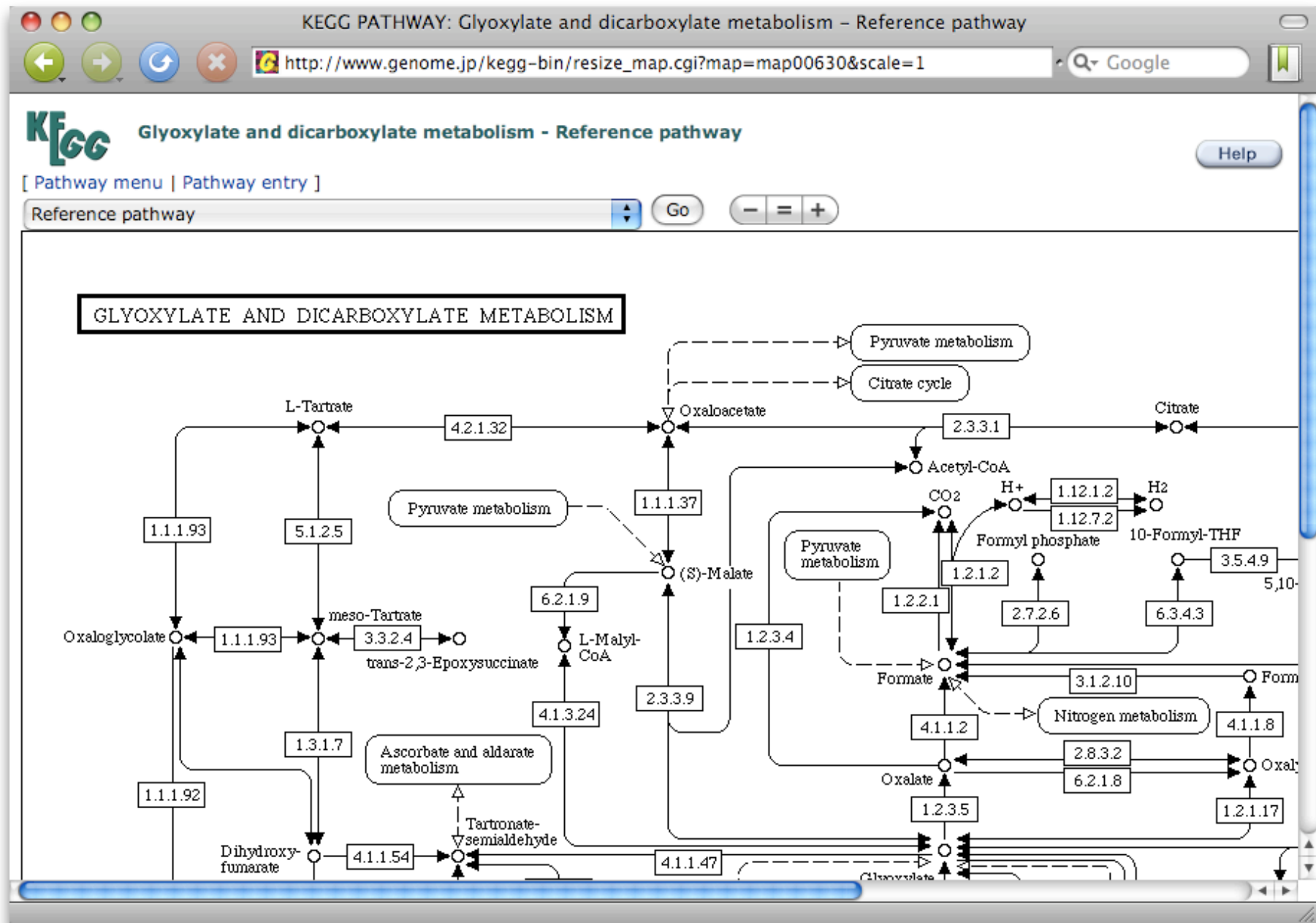
<http://www.genome.jp/kegg/>



kinetische Daten: SABIO-RK

http://sabio.villa-bosch.de/welcome_new.jsp?

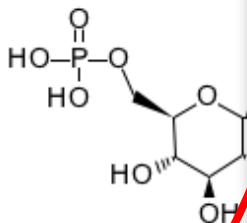




Inside KEGG

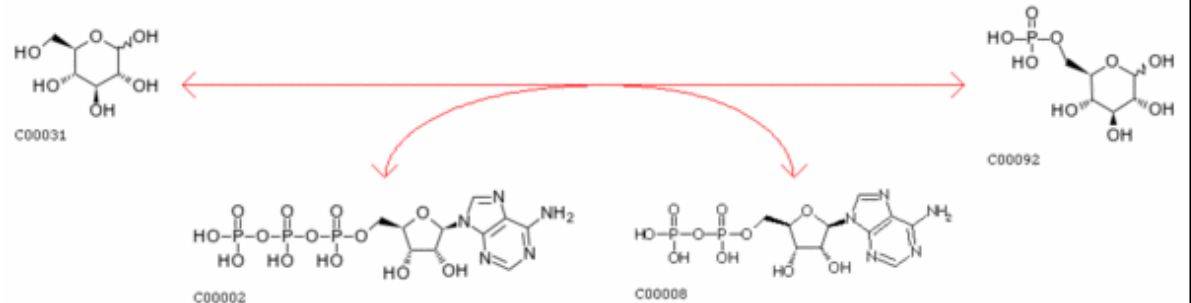
KEGG COMPOUND: C00092

Help

Entry	C00092
Name	D-Glucose 6-phosphat Glucose 6-phosphat Robison ester
Formula	C6H13O9P
Mass	260.0297
Structure	 <p>C00092</p> <p>Mol file KEGG file</p>
Reaction	R00299 R00303 R007 R00838 R00839 R008 R05804 R06043 R060 R08125 R08404 R086
Pathway	PATH: ko00500 Stan PATH: ko00521 Stre PATH: ko00562 Inos PATH: map01062 Bic PATH: ko02020 Two PATH: ko02060 Phosphotransferase system (PTS)
Enzyme	1.1.1.49 1.1.1.200 2.4.1.1 (E) 2.4.1.15 2.4.1.36 2.4.1.216 2.7.1.1 2.7.1.2 2.7.1.61 2.7.1.63 2.7.1.142 2.7.1.147 3.1.3.9 3.1.3.58 3.2.1.86 3.2.1.93 3.2.1.122 3.5.-.- 5.3.1.9 5.4.2.2 5.4.2.5 5.5.1.4
Other DBs	CAS: 56-73-5

KEGG REACTION: R00299


Help

Entry	R00299	Reaction
Name	ATP:D-glucose 6-phosphotransferase	
Definition	ATP + D-Glucose <=> ADP + D-Glucose 6-phosphate	
Equation	C00002 + C00031 <=> C00008 + C00092	
		
RPair	RP: RP00003 C00002_C00008 main RP: RP00060 C00031_C00092 main RP: RP08556 C00002_C00092 trans	
Pathway	PATH: rn00521 Streptomycin biosynthesis	
Enzyme	2.7.1.1 2.7.1.2	
LinkDB	All DBs	

Reaktionsraten?


Raten: SABIO-RK

What is SABIO-RK? [TOP](#)

The SABIO-RK (System for the Analysis of Biochemical Pathways - Reaction Kinetics) is a web-based application based on the SABIO relational database that contains information about biochemical reactions, their kinetic equations with their parameters, and the experimental conditions under which these parameters were measured. It aims to support modellers in the setting-up of models of biochemical networks, but it is also useful for experimentalists or researchers with interest in biochemical reactions and their kinetics. Information about reactions and their kinetics can be exported in SBML ([Systems Biology Mark-Up Language](#) ) format.

This project is sponsored by the [Klaus Tschira Foundation](#)  and partially by the German BMBF (Bundesministerium für Bildung und Forschung).

Data Provenance [TOP](#)

There are two main sources for the data contained in SABIO. Most of the reactions, their association with pathways and their enzymatic classification, is extracted from the [KEGG \(KEGG: Kyoto Encyclopedia of Genes and Genomes\)](#)  database. The other main source of data are publications. The reaction kinetics data are obtained by manual extraction from literature sources and curated.

Suche in SABIO-RK

Reaction Search

Specify Search Criteria:

with **Reactant(s)**

✖

 D-Glucose 6-phosphate

in **Pathway(s)**

having **Enzyme(s)**

✖

 2.7.1.1:Hexokinase

in **Publication**

related to **Protein** (UniProtID)

in **Organism(s)**

✖

 Homo sapiens

Submit Search

Reset Form

Search Results

Total number of reactions found for specified search criteria: 2

[Click here to view your search criteria](#)

Modify Search

Number of results per page: 10

Display

Show only reactions having kinetic data matching the search criteria ☒

Send Selected Reactions to SBML File

Reactions	Select Reaction(s) (De)Select All	Kinetic Data for this reaction (Click to View)	Enzyme EC#	Kinetic data for enzymes (Click to View)
ITP + D-Glucose <-> D-Glucose 6-phosphate + IDP	<input type="checkbox"/>	<div>view</div>	2.7.1.1	<div>view</div>
D-Glucose + ATP <-> D-Glucose 6-phosphate + ADP	<input type="checkbox"/>	<div>view</div>	2.7.1.1 2.7.1.2	<div>view</div> <div>view</div>

Pages: 1

Previous

Next

Entry Nr. 2362

⊕

⊖

Select

Organism:	Homo sapiens		
Tissue:	erythrocyte		
EC Class: 2.7.1.1	wildtype		

Substrates

name	location	comment
ATP	-	-
D-Glucose	-	-

Products

name	location	comment
ADP	-	-
D-Glucose 6-phosphate	-	-

Modifiers

name	location	effect	comment	protein complex
Mg2+	-	Modifier-Cofactor	-	-
Hexokinase(Enzyme)	-	Modifier-Catalyst	-	-
2,3-Diphosphoglycerate	-	Modifier-Inhibitor	-	-

Enzyme (protein data)

	UniProt-ID	name	mol. weight (kDa)	deviation (kDa)
subunit	-	-	-	-
complex	-	-	-	-

Kinetic Law

type	formula
Uncompetitive inhibition	unknown

Parameters

name	species	type	start value	end value	deviation	unit	comment
B	ATP	concentration	1	-	-	mM	-
C	Mg2+	concentration	0.25	3	-	mM	-
I	2,3-Diphosphoglycerate	concentration	0	5	-	mM	-
Km_Mg	Mg2+	Km	0.0023	-	-	M	-
Km_Glu	D-Glucose	Km	0.000093	-	-	M	-
A	D-Glucose	concentration	0.3	1	-	mM	-

Experimental conditions

	start value	end value	unit
pH	8	-	-
temperature	23	24	°C

buffer: 50 mM Tris chloride, 1 mM NADP+, 0.1 mg glucose 6-phosphate dehydrogenase

Zusammenfassung

Dynamische Simulationen:

- zeitliches Verhalten
- steady state = stationäre Lösung des DGL-Systems
- Puffergrößen und Reaktionsraten

Copasi:

- Simulation und Analyse chemischer Reaktionen

Vereinfachte Kinetiken:

- hilft im steady state, problematisch bei zeitabh. Prozessen
- Bsp: kinetische Isolierung von Signalpfaden

Simulationsparameter?

- KEGG – Pfade
- SABIO-RK: hand-kurierte Reaktionsparameter

Systems Biology Markup Language



XML-Dialekt für Speicherung und Austausch
biochemischer Modelle

=> Archivierung

=> Transfer von Modellen in andere Softwaretools

Acknowledgements

We are deeply indebted to the many funding agencies and organizations that have supported SBML over the years.

Since 2003, primary support for the continued development of SBML, as well as supporting software and activities, has come from the **National Institute of General Medical Sciences (NIH/NIGMS)** [\[link\]](#) under grants R01 GM070923 and R01 GM077671. Additional support is provided by the **California Institute of Technology** [\[link\]](#) (USA) and **Keio University** [\[link\]](#) (Japan).

The development of SBML from its inception through 2003 was principally funded by the **Japan Science and Technology Agency** [\[link\]](#) under the **ERATO Kitano Symbiotic Systems Project** [\[link\]](#).

Additional support has in the past been provided by the following organizations and agencies: the **Systems Biology Institute** [\[link\]](#) (Japan), the **University of Hertfordshire** [\[link\]](#) (UK), the **Molecular Sciences Institute** [\[link\]](#) (USA), the **National Human Genome Research Institute** [\[link\]](#) (USA), the **International Joint Research Program of NEDO** [\[link\]](#) (Japan), the **ERATO-SORST** [\[link\]](#) Program of the Japan Science and Technology Agency (Japan), the **Ministry of Agriculture** [\[link\]](#) (Japan), the **Ministry of Education, Culture, Sports, Science and Technology** [\[link\]](#) (Japan), the **BBSRC e-Science Initiative** [\[link\]](#) (UK), the **DARPA IPTO Bio-Computation Program** [\[link\]](#) (USA), the Army Research Office's **Institute for Collaborative Biotechnologies** [\[link\]](#) (USA), and the **Air Force Office of Scientific Research** [\[link\]](#) (USA).

[von http://sbml.org/Acknowledgments](http://sbml.org/Acknowledgments)

SBML <= XML

XML = eXtensible Markup Language

- hierarchische Baumstruktur:
=> Schachtelung von `<Object> ... </Object>` oder `<Objekt [Parameter...]/>`
- genau ein Wurzelobjekt: `<sbml...>`

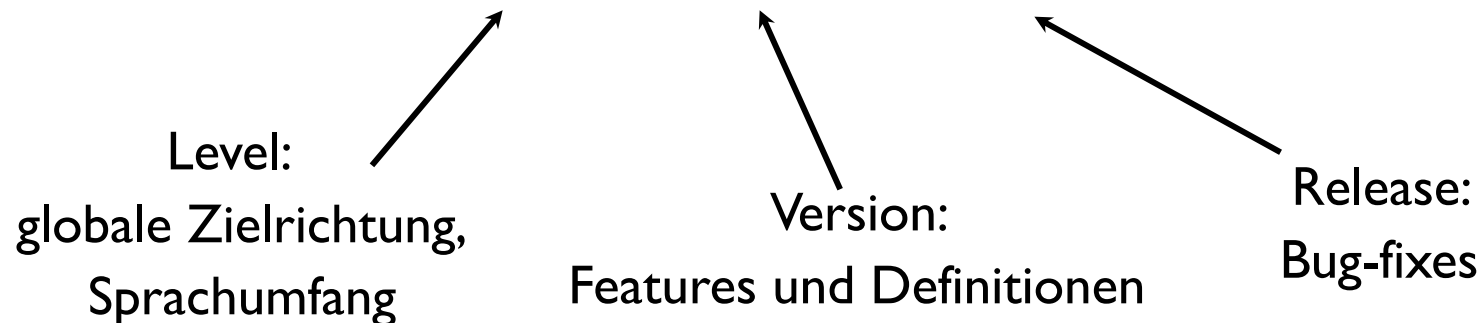
Aktuelle Dialekte: [siehe http://sbml.org/Documents/Specifications](http://sbml.org/Documents/Specifications)

SBML Level 1, Version 2

<http://www.sbml.org/specifications/sbml-level-1/version-2/sbml-level-1-v2.pdf>

SBML Level 2, Version 4, Release 1

<http://precedings.nature.com/documents/2715/version/1>



Was ist enthalten?

beginning of model definition

list of function definitions (optional)

list of unit definitions (optional)

list of compartment types (optional)

list of species types (optional)

list of compartments (optional)

list of species (optional)

list of parameters (optional)

list of initial assignments (optional)

list of rules (optional)

list of constraints (optional)

list of reactions (optional)

list of events (optional)

end of model definition

Ein Beispiel



```
<?xml version="1.0" encoding="UTF-8"?>
<sbml level="2" version="3" xmlns="http://www.sbml.org/sbml/level2/version3">
  <model name="EnzymaticReaction">
    <listOfUnitDefinitions>
      <unitDefinition id="per_second">
        <listOfUnits>
          <unit kind="second" exponent="-1"/>
        </listOfUnits>
      </unitDefinition>
      <unitDefinition id="litre_per_mole_per_second">
        <listOfUnits>
          <unit kind="mole" exponent="-1"/>
          <unit kind="litre" exponent="1"/>
          <unit kind="second" exponent="-1"/>
        </listOfUnits>
      </unitDefinition>
    </listOfUnitDefinitions>
    <listOfCompartments>
      <compartment id="cytosol" size="1e-14"/>
    </listOfCompartments>
    <listOfSpecies>
      <species compartment="cytosol" id="ES" initialAmount="0" name="ES"/>
      <species compartment="cytosol" id="P" initialAmount="0" name="P"/>
      <species compartment="cytosol" id="S" initialAmount="1e-20" name="S"/>
      <species compartment="cytosol" id="E" initialAmount="5e-21" name="E"/>
    </listOfSpecies>
    <listOfReactions>
      <reaction id="veq">
        <listOfReactants>
          <speciesReference species="E"/>
          <speciesReference species="S"/>
        </listOfReactants>
        <listOfProducts>
          <speciesReference species="ES"/>
        </listOfProducts>
        <kineticLaw>
          <math xmlns="http://www.w3.org/1998/Math/MathML">
            <apply>
              <times/>

```

```

              <ci>cytosol</ci>
            </apply>
            <minus/>
            <apply>
              <times/>
              <ci>kon</ci>
              <ci>E</ci>
              <ci>S</ci>
            </apply>
          </math>
        </kineticLaw>
      </reaction>
      <reaction id="vcat" reversible="false">
        <listOfReactants>
          <speciesReference species="ES"/>
        </listOfReactants>
        <listOfProducts>
          <speciesReference species="E"/>
          <speciesReference species="P"/>
        </listOfProducts>
        <kineticLaw>
          <math xmlns="http://www.w3.org/1998/Math/MathML">
            <apply>
              <times/>
              <ci>cytosol</ci>
              <ci>kcat</ci>
              <ci>ES</ci>
            </apply>
          </math>
          <listOfParameters>
            <parameter id="kon" value="1000000" units="litre_per_mole_per_second"/>
            <parameter id="koff" value="0.2" units="per_second"/>
          </listOfParameters>
        </kineticLaw>
      </reaction>
    </listOfReactions>
  </model>
</sbml>

```

Nochmal:



```
<?xml version="1.0" encoding="UTF-8"?>
<sbml level="2" version="3" xmlns="http://www.sbml.org/sbml/level2/version3">
  <model name="EnzymaticReaction">
    <listOfUnitDefinitions>
      :
    </listOfUnitDefinitions>
    <listOfCompartments>
      <compartment id="cytosol" size="1e-14"/>
    </listOfCompartments>
    <listOfSpecies>
      <species compartment="cytosol" id="ES" initialAmount="0" name="ES"/>
      <species compartment="cytosol" id="P" initialAmount="0" name="P"/>
      <species compartment="cytosol" id="S" initialAmount="1e-20" name="S"/>
      <species compartment="cytosol" id="E" initialAmount="5e-21" name="E"/>
    </listOfSpecies>
    <listOfReactions>
      :
    </listOfReactions>
  </model>
</sbml>
```

Details: Einheiten

```
<listOfUnitDefinitions>  
  <unitDefinition id="per_second">  
    <listOfUnits>  
      <unit kind="second" exponent="-1"/>  
    </listOfUnits>  
  </unitDefinition>
```

per_seconds := s⁻¹

```
<unitDefinition id="litre_per_mole_per_second">  
  <listOfUnits>  
    <unit kind="mole" exponent="-1"/>  
    <unit kind="litre" exponent="1"/>  
    <unit kind="second" exponent="-1"/>  
  </listOfUnits>  
</unitDefinition>  
</listOfUnitDefinitions>
```

$\frac{\text{litre}}{\text{mol s}}$

SBML Software Guide/SBML Software Matrix – SBML.org

http://sbml.org/SBML_Software_Guide/SBML_Software_Matrix

SBML Software Matrix

This matrix provides an at-a-glance summary of software known to us to provide some degree of support for reading, writing, or otherwise working with SBML. The columns' meanings are explained below. For a list of longer descriptions grouped into themes, please see our [SBML Software Summary](#) page.

	Capabilities					Frameworks							API	Dep.	Platforms	SBML		Availabil.		
	Creation	Simulation	Analysis	Database	Utility	ODE	DAE	PDE	Stochastic	Events	Logical	Other				Import	Export	Open source	Academic use	Commercial use
Cellware	•	•				•									L,W,M	•		•	F	\$
CL-SBML					•							•	LISP	LISP	L	•		•	F	F
CLEML												•			L,W	•	•		F	F
COBRA			•		•	•						•		MATLAB	L,W,M	•	•	•	F	F
ConsensusPathDB					•										B	•	•	•	F	F
COPASI	•	•	•		•	•			•				C++, Java, Python		L,W,M	•	•	•	F	\$
Cyto-Sim		•			•				•						L,W,M				F	F
Cytoscape	•				•	•							Java		L,W,M	•		•	F	F
DBSolve		•	•		•	•										•	•		F	F
Dizzy		•				•			•						L,W,M	•	•	•	F	F
E-CELL	•	•				•			•						L,W		•	•	F	F
ecellJ					•											•			F	F
EPE	•			•		•							Java		L,W, M	•			F	F
ESS		•							•					BSP				•	F	F

Import nach Copasi

The screenshot shows the COPASI 4.5 (Build 30) interface. The left sidebar displays the model hierarchy: Model > Biochemical > Compartments (cytosol) > Species (E, ES, P, S) > Reactions (vcat, **veq**). The main window is titled 'enzymatic - COPASI 4.5 (Build 30) /Users/.../V11/enzymatic.cps' and has tabs for 'Concentrations', 'Reaction' (selected), 'Annotation', and 'RDF Browser'.

In the 'Reaction' tab, the following details are visible:

- Name:** veq
- Chemical Equation:** $E + S = ES$
- Reversible:** ☒ (Multi Compartment: ☐)
- Rate Law:** Mass action (reversible) (New Rate Law button)
- Flux (mol/s):** 0

The 'Symbol Definition' table is shown below:

Description	Name	Value	Unit
Parameter	k1	<input type="checkbox"/> global	1e+06 l/(mol*s)
→ Substrate	substra		mol/l
	E		
	S		
Parameter	k2	<input type="checkbox"/> global	0.2 1/s

Buttons at the bottom include 'Commit', 'Revert', 'New', 'Delete', 'Clear', 'Delete/Undelete', and 'New'.

Details: eine Reaktion

```

<listOfReactions> :
  <reaction id="vcat" reversible="false">
    <listOfReactants>
      <speciesReference species="ES"/>
    </listOfReactants>
    <listOfProducts>
      <speciesReference species="E"/>
      <speciesReference species="P"/>
    </listOfProducts>
    <kineticLaw>
      <math
xmlns="http://www.w3.org/1998/Math/MathML">
        <apply>
          <times/>
          <ci>cytosol</ci>
          <ci>kcat</ci>
          <ci>ES</ci>
        </apply>
      </math>
      <listOfParameters>
        <parameter id="kcat" value="0.1"
units="per_second"/>
      </listOfParameters>
    </kineticLaw>
  </reaction>
</listOfReactions>

```



$$\Rightarrow \frac{dN}{dt} = V_{\text{cytosol}} k_{\text{cat}} [ES]$$

lokaler Parameter!

SBML lesbar machen



convert

SBML file:

Report options

MIRIAM annotations: ☒ Check SBML consistency: ☒ Include predefined unit declarations: ☒

Layout options

Convert to: Set name in equations: ☐ Landscape: ☐

Font size: Reaction participants in one table: ☐ Set identifiers in typewriter font: ☒

Paper size: Create a title page: ☐

<http://webservices.cs.uni-tuebingen.de/>

Dräger A, Planatscher H, Wouamba DM, Schröder A, Hucka M, Endler L, Golebiewski M, Müller W, and Zell A: "SBML2LaTeX: Conversion of SBML files into human-readable reports", Bioinformatics 2009

Drei Minuten später:

convert

Please download your result here:
[07ff0064-6af4-4eb5-bea1-906da1fbcd86-request.pdf](#)

Submit another file

SBML Model Report

Model name: “EnzymaticReaction”



June 30, 2009

1 General Overview

This is a document in SBML Level 2 Version 3 format. Table 1 gives an overview of the quantities of all components of this model.

Table 1: The SBML components in this model.
All components are described in more detail in the following sections.

Element	Quantity	Element	Quantity
compartment types	0	compartments	1
species types	0	species	4
events	0	constraints	0
reactions	2	function definitions	0
global parameters	0	unit definitions	2
rules	0	initial assignments	0

2 Unit Definitions

This is an overview of seven unit definitions. The units `substance`, `volume`, `area`, `length`, and `time` are predefined by SBML and not mentioned in the model.

2.1 Unit `per_second`

Definition s^{-1}

2.2 Unit `litre_per_mole_per_second`

Definition $\text{mol}^{-1} \cdot \text{l} \cdot s^{-1}$

2.3 Unit `substance`

Notes Mole is the predefined SBML unit for `substance`.

Definition `mol`

2.4 Unit `volume`

Notes Litre is the predefined SBML unit for `volume`.

Definition `l`

2.5 Unit `area`

Notes Square metre is the predefined SBML unit for `area` since SBML Level 2 Version 1.

Definition m^2

2.6 Unit `length`

Notes Metre is the predefined SBML unit for `length` since SBML Level 2 Version 1.

Definition `m`

2.7 Unit `time`

Notes Second is the predefined SBML unit for `time`.

Definition `s`

3 Compartment

This model contains one compartment.

Table 2: Properties of all compartments.

Id	Name	SBO	Spatial Dimensions	Size	Unit	Constant	Outside
cytosol			3	10^{-14}	l	<input checked="" type="checkbox"/>	

3.1 Compartment cytosol

This is a three-dimensional compartment with a constant size of 10^{-14} litre.

4 Species

This model contains four species. Section 6 provides further details and the derived rates of change of each species.

Table 3: Properties of each species.

Id	Name	Compartment	Derived Unit	Constant	Boundary Condition
ES	ES	cytosol	$\text{mol} \cdot \text{l}^{-1}$	<input type="checkbox"/>	<input type="checkbox"/>
P	P	cytosol	$\text{mol} \cdot \text{l}^{-1}$	<input type="checkbox"/>	<input type="checkbox"/>
S	S	cytosol	$\text{mol} \cdot \text{l}^{-1}$	<input type="checkbox"/>	<input type="checkbox"/>
E	E	cytosol	$\text{mol} \cdot \text{l}^{-1}$	<input type="checkbox"/>	<input type="checkbox"/>

5 Reactions

This model contains two reactions. All reactions are listed in the following table and are subsequently described in detail. If a reaction is affected by one or more modifiers, the identifiers of the modifier species are written above the reaction arrow.

Table 4: Overview of all reactions

Nº	Id	Name	Reaction Equation	SBO
1	veq		$E + S \rightleftharpoons ES$	
2	vcat		$ES \longrightarrow E + P$	

5.1 Reaction veq

This is a reversible reaction of two reactants forming one product.

Reaction equation



Reactants

Table 5: Properties of each reactant.

Id	Name	SBO
E	E	
S	S	

Product

Table 6: Properties of each product.

Id	Name	SBO
ES	ES	

Kinetic Law

Derived unit $s^{-1} \cdot \text{mol}$

$$v_1 = \text{vol}(\text{cytosol}) \cdot (\text{kon} \cdot [E] \cdot [S] - \text{koff} \cdot [ES]) \quad (2)$$

Table 7: Properties of each parameter.

Id	Name	SBO	Value	Unit	Constant
kon			1000000.0	$\text{mol}^{-1} \cdot \text{l} \cdot \text{s}^{-1}$	<input checked="" type="checkbox"/>
koff			0.2	s^{-1}	<input checked="" type="checkbox"/>

6 Derived Rate Equations

When interpreted as an ordinary differential equation framework, this model implies the following set of equations for the rates of change of each species.

6.1 Species ES

Name ES

Initial amount 0 mol

This species takes part in two reactions (as a reactant in *vcat* and as a product in *veq*).

$$\frac{d}{dt}ES = v_1 - v_2$$

(5)

6.2 Species P

Name P

Initial amount 0 mol

This species takes part in one reaction (as a product in *vcat*).

$$\frac{d}{dt}P = v_2$$

(6)

6.3 Species S

Name S

Initial amount 10^{-20} mol

This species takes part in one reaction (as a reactant in *veq*).

$$\frac{d}{dt}S = -v_1$$

(7)

6.4 Species E

Name E

Initial amount $5 \cdot 10^{-21}$ mol

This species takes part in two reactions (as a reactant in *veq* and as a product in *vcat*).

$$\frac{d}{dt}E = v_2 - v_1$$

(8)

es gibt bereits sehr viele Modelle

The screenshot shows the BioModels Database website in a web browser. The browser's address bar displays <http://www.ebi.ac.uk/biomodels-main/>. The website header includes the EMBL-EBI logo, an "EB-eye Search" bar with a dropdown menu set to "All Databases" and a search input field, and a "Go" button. Navigation links for "Databases", "Tools", "EBI Groups", "Training", "Industry", "About Us", and "Help" are present, along with a "Site Index" link. A secondary navigation bar contains links for "BioModels Home", "Browse models", "Submit", "Sign in", "Support", and "About BioModels".

The main content area is titled "BioModels Database - A Database of Annotated Published Models". Below this title, a paragraph describes the database as a resource for storing, searching, and retrieving published mathematical models of biological interests, which are annotated and linked to relevant data resources like publications, compound databases, and pathways.

On the left side, there is a search section with a text input field, "Search", "Go to the model", and "Advanced search" buttons. Below this, the "Browse models" section lists three categories: "Curated models (216)", "Browse models using GO", and "Non-curated models (196)". Further down are links for "Simulate in JWS Online" and "Submit a model". At the bottom left, a mirror site is mentioned: "Mirror at California Institute of Technology <http://biomodels.caltech.edu>".

On the right side, the "Model of the month" section for May 2009 features a text description of sucrose accumulation in sugar cane and a small diagram of a metabolic pathway. The diagram shows a central node labeled "Suc" with an incoming arrow from the left and two outgoing arrows to the right, labeled "11" and "8". Below the text is a "Read more..." link. The "News" section at the bottom right reports on the "Fourteenth release" dated 16th June 2009, with a link to "Download All Models Under SBML Format".

Klausur-relevanter Vorlesungsstoff

Vorlesung	Folien
1	14-22, 27, 35
2	3-43
3	3-22, 25, 34-46
4	13
5	1-34, 39, 41
6	1-11, 15-36
7	5-6, 9-12, 16-18
8	9-39
9	7-10, 16-20, 30-35
10	1, 4,5, 7-9, 12-19
11	3-8, 16-18, 31-33, 38
12	8-10, 13
13	4, 22, 24-28, 30-31

Am Ende von VI3 können zur Klausurvorbereitung Fragen zur gesamten Vorlesung gestellt werden.