# A Pycellerator Tutorial Bruce Shapiro and Eric Mjolsness

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# A Pycellerator Tutorial

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# 12 Summary

- We present a tutorial on using Pycellerator for biomolecular simulations. Models are described in human readable (and editable) text files (UTF8 or ASCII) containing 14 collections of reactions, assignments, initial conditions, function definitions, and rate 15 constants. These models are then converted into a Python program that can optionally 16 solve the system, e.g., as a system of differential equations using ODEINT, or be run by 17 another program. The input language implements an extended version of the Cellerator 18 arrow notation, including mass action, Hill functions, S-Systems, MWC, and reactions with user-defined kinetic laws. Simple flux balance analysis is also implemented. We will demonstrate the implementation and analysis of progressively more complex models, starting from simple mass action through indexed cascades. Pycellerator can be used as a library that is integrated into other programs, run as a command line program, or in iPython notebooks. It is implemented in Python 2.7 and available under an open source GPL license.
- 26 Key words: Cellerator, SBML, Systems Biology, Python

# <sub>27</sub> 1 Introduction

Using Pycellerator typically involves the following steps steps: (a) model preparation; (b)
model instantiation; (c) model execution (simulation); and (d) simulation analysis. Model
preparation requires creation of a model in the Pycellerator model description language
(henceforth called model files). This are human-readable files that are typically
hand-written in a text-editor. A Pycellerator model consists of some combination of
reactions, assignment rules and functions, along with specifications of initial conditions,
parameter values, and constant variables. [1] In addition, the modeling language

- incorporates an extended text-based version of the full Cellerator modeling language. [2]
- 36 (However, Pycellerator is a completely separate program from Cellerator and, unlike the
- latter, does not depend on or use any features of Mathematica.) Where features are
- compatible (e.g., events and level 3 packages are not supported), models can also be read
- <sup>39</sup> from SBML files [3] and Cellerator Mathematica models.
- 40 Model instantiation means conversion of a Pycellerator model into a Python program. This
- 41 auto-generated Python program includes two parts: a main driver program that invokes
- 42 the scipy.odeint numerical solver [4], and a function that instantiates the model as a systems
- of differential equations. This function is in the format that is typically expected by odeint. The
- 44 program is saved to the file system, and modelers may choose to modify the program and/or use it
- 45 independently of the remainder of Pycellerator. This program does not depend on any
- 46 Pycellerator libraries.
- 47 Model execution and analysis involves performing simulations with the instantiated model and
- interpreting the results. Pycellerator provides functionality to execute an instantiated model and
- 49 to plot time courses of the results. In addition, the results of the numerical integration may be
- saved to numpy arrays [5], and any of the analysis functions available in Python are subsequently
- 51 available for use.
- 52 The main features of Pycellerator that differentiate it from other modeling language based
- simulators are (a) conversion to accessible, user-modifiable, executable Python model descriptions;
- and (b) the ability to incorporate standard Python expressions (such as ternary conditionals) into
- 55 assignment rules.
- With Pycellerator a modeler may perform any of the the following tasks automatically:
- 1. Generate a Python code implementation of a model.
- 2. Generate a stand-alone program that can be used to run the code produced in step 1.
- 3. Run a deterministic simulation of the model using the code generated in the step 2.

- 4. Generate Python code wrapper to perform a parametric variation (e.g., of a rate constant or initial condition over an interval) of the code implemented in step 2.
- 5. Run the parametric evaluation written in step 4.
- 6. Plot the results (e.g., state variable time courses or parametric variation) from steps 2 or 4.
- 7. Solve simple flux models for unknown fluxes.
- 8. Export results from steps 2, 4, or 7 into numpy arrays for further analysis within Python.
- These functions may be performed either in iPython notebook or from the command shell.
- 67 Auto-generated code can be modified by users and incorporated into user programs without
- 68 restriction.

# <sub>9</sub> 2 Materials

- 70 Models are composed of primarily of lists of chemical reactions and their associated rate constants
- 71 and initial conditions. An example of a simple model for an enzymatic reaction is given in listing
- 1. Models may also include equations that specify species values, mathematical functions, and to
- 73 some extent, simple Python expressions. The reactions are specified using standard arrow-like
- 74 keyboard symbols, and equations resemble standard Python expressions. In general, a model file is
- divided into six sections: reactions, parameter values, initial conditions, function definitions,
- assignment rules, and a list of constant species. Each of these sections begins with a special
- 77 keyword: \$REACTIONS, \$RATES, \$IC, \$FUNCTIONS, \$ASSIGNMENTS, \$FROZEN. The keywords
- 78 are not case sensitive. In its simplest form, a model would consist of one or more reactions, initial
- 79 conditions, and rate constants.

## $_{ t 80}$ 2.1 ${ m Arrows}$

81 The canonical arrow form is

[[reactants] arrow [products], 
$$mod[modifier]$$
,  $keyword[parameters]$ ] (1)

where reactants and products are comma-delimited sequences of one or more species names; 82 arrow is a text arrow (see table 1, column 1); keyword is a keyword that indicates how the 83 arrow is to use the list of parameters (table 1, column 2); and modifier is a one or more 84 species names that are optionally allowed with some arrow/keyword combinations. When there is only one reactant (or only one product), the square brackets around the corresponding sequence 86 (of reactants or products) is omitted. In certain cases (e.g., mass action reactions), the 87 plus-symbol ("+") is used in place of commas to delimit reactants or products, and the 88 brackets are also omitted in these situations. The entire reaction must be enclosed in square 89 brackets. Each arrow contributes terms to the system of differential equations that describe the 90 model. The following section describe how each type of arrow is understood by Pycellerator.

#### 92 2.1.1 Mass Action

The most basic reaction arrows in Pycellerator use mass action kinetics (see table 2). A numerical stoichiometry may be specified and there is no limit to either the number of reactants or products in a reaction. The standard syntax is

$$[e1*X1+e2*X2+...en*Xn -> f1*Y1 + f2*Y2 + ... + fm*Ym, k]$$
 (2)

This means that reactants X1, X2, ..., Xn are combined with stoichiometries e1, e2, ...

97 en to produce products Y1, Y2, ..., Ym with stoichiometries f1, f2, ... fm. The

98 asterisks are optional but the numerical stoichiometries must precede the symbols. For each

99 reaction in the model, a differential equation term is generated for each species (by species we

100 mean reactant or product) in that reaction. Let Z be be some species that appears with (possibly

zero) stoichiometries  $e_j$  and  $f_j$  on the left hand side and right hand side of the arrow in reaction j.

Then if species  $X_1, \ldots, X_n$  appear on the left hand side of reaction j with stoichiometries

103  $e_{j1}, \ldots, e_{jn},$ 

$$\frac{dZ}{dt} = \sum_{j \in \text{Reactions}} k_j (f_j - e_j) \prod_{a=1}^n X_a^{e_{ja}}$$
(3)

where  $k_j$  is the rate constant of reaction j. [6]

Although stoichiomety is normally integer, there is nothing preventing a modeller from using any non-integer floating point value. For example, consider the following system of biochemical reactions from the Field-Noyes (Oregonator) model, in which the original model authors use a stoichimetry of 1/2 for for the last reaction. [7]

$$Br + BrO_{3} \xrightarrow{k_{1}} HBrO_{2} + HOBr$$

$$Br + HBrO_{2} \xrightarrow{k_{2}} 2HOBr$$

$$BrO_{3} + HBrO_{2} \xrightarrow{k_{3}} 2Ce + 2HBrO_{2}$$

$$2HBrO_{2} \xrightarrow{k_{4}} BrO_{3} + HOBr$$

$$Ce \xrightarrow{k_{5}} 0.5Br$$

$$(4)$$

The Reactions section of the corresponding model file might look like this:

Each species in this system will automatically be converted to differential equations according to equation (3). If we also include BrO3 in the Frozen section of the model file (to keep its value constant), the resulting mass action equations (in Python form) are

- While it is unlikely for n nor m to be larger than two, nothing precludes modelers from
- incorporating higher order reactions in Pycellerator models.

## 115 2.1.2 Enzymatic Expansion

We define a number of catalyzed mass action reactions that are expanded into standard enzymatic reactions, e.g., simple conversion via creation of an intermediate complex. Each of the these enzymatic reactions is indicated by a single line of code in the model. Consider the following biochemical reaction:

$$S + E \underset{k_2}{\overset{k_1}{\rightleftharpoons}} SE \underset{k_4}{\overset{k_3}{\rightleftharpoons}} P + E \tag{7}$$

120 This is represented by a single arrow

$$[S=>P, mod[E], rates[k1, k2, k3, k4]]$$
 (8)

Pycellerator allows users to omit rate constants from the end of the list, defaulting their values to zero. Thus

$$[S=>P, mod[E], rates[k1,k2,k3]]$$
(9)

123 represents the reaction

$$S + E \underset{k_2}{\overset{k_1}{\longleftrightarrow}} SE \xrightarrow{k_3} P + E \tag{10}$$

Pycellerator automatically reinterprets arrow (8) as the following system of arrows. The name of
the intermediate complex is automatically generated from the names of the substrate and the
catalyst.

$$\begin{bmatrix}
S+E->S_E, k1 \\
S_E->S+E, k2 \\
S_E->P+E, k3 \\
P+E->S_E, k4 \end{bmatrix}$$
(11)

127

These arrows (as well as the other forms in table 2) are converted into differential equations as per equation (3). For example, [S<=>P, mod[F,R], rates[k1,..,k8]] represents the pair of reactions

$$S + E \underset{k_2}{\overset{k_1}{\rightleftharpoons}} SE \underset{k_4}{\overset{k_3}{\rightleftharpoons}} P + E$$

$$P + R \underset{k_6}{\overset{k_5}{\rightleftharpoons}} PR \underset{k_8}{\overset{k_7}{\rightleftharpoons}} S + R$$

$$(12)$$

Internally, this would be expanded first into two arrows of the form reaction (8) and then eight arrows of the form reaction (11). Four of these arrows would correspond to the first reaction in reactions (12) and four in the second reaction in reactions (12).

## <sup>34</sup> 2.1.3 Michaelis-Menten-Henri-Briggs-Haldane Approximation

Henri, Michaelis and Menten, and Briggs and Haldane all obtained the following formula for reaction 10, but with different assumptions.

$$\frac{dP}{dt} = \frac{vS}{K+S} = -\frac{dS}{dt} \tag{13}$$

Since they made different assumtions, the actual chemistry should be interpreted differently in
each case. Henri (in 1903) and Michaelis and Menten (in 1913) assumed fast equilibrium of the of
the catalyst/substrate reaction to form SE, which subsequently dissociates. [8, 9] Briggs and
Haldane (in 1925), on the other hand, obtained the same result by assuming that SE in
quasi-steady state. [10] The Briggs and Haldane method is usually used in elementary biology
classes. However it is interpreted, the same equation is obtained.

143 The canonical Pycellerator arrow

$$[S :-> P,MMH[K, v]]$$
 (14)

is used to produce the rate law in equation (13), which in general only has two parameters: where v is the maximum reaction rate and K is the substrate concentration at half maximum. The actual enzyme concentration E does not (normally) come into the rate law as it is absorbed into the constant v. The rate law equation (13) will be used to produce differential equation terms for the variables P and S, which will be added to other differential equation terms in the model.

Additional versions of this model implemented in Pycellerator allow K and v to be replaced by

(a) K, v, and E (this replaces v in equation (13) with vE)

$$[S :-> P, mod[E], MMH[K, v]]$$

$$(15)$$

151 (b)  $k_1$ ,  $k_2$ , and  $k_3$  (this sets  $v=k_3$  and  $K=(k_2+k_3)/k_1$  in equation (13))

$$[S : -> P, MMH[k1, k2, k3]]$$
 (16)

and (c) E,  $k_1$ ,  $k_2$ , and  $k_3$  (this sets  $v = k_3 E$  and  $K = (k_2 + k_3)/k_1$  in equation (13)):

$$[S : --> P, mod[E], MMH[k1, k2, k3]]$$
 (17)

In (a) and (c) the variable E may be any other species in the model that is controlled by its own dynamics, including other reactions or assignments, or it may be a fixed parameter. See table 3 for details.

#### $_{156}$ 2.1.4 Hill Functions

Pycellerator includes several regulatory arrows (table 4). In a regulatory arrow, only the species on the right hand side of the arrow are affected by the resulting new differential equation terms.

The species listed on the left hand side (LHS) of the arrow contribute information to the system, in the sense that they define how these terms are constructed, but the LHS species are not consumed. Regulatory arrows include Hill functions, GRN (Genetic Regulatory Network Arrows), S-Systems, and Rational functions.

Hill functions frequently arise as approximations of the cooperative binding of ligands. Because of their sigmoidal shape, Hill functions can sometimes be numerically optimized to accurately

their sigmoidal shape, Hill functions can sometimes be numerically optimized to accurately described bistable switches, where the amount, concentration, or rate of production of one species (say Y) depends on the corresponding amount or concentration of a second species (say X).[11]

The canonical form in Pycellerator is

$$[X \mid ->Y, Hill[V, n, K, a, T]]$$

$$(18)$$

which is described by the differential equations term

$$\frac{dY}{dt} = \frac{v(a+TA)^n}{K^n + (a+TX)^n} \tag{19}$$

Here v, n, K, a and T are constants that are allowed to take on any floating point or integer value. In particular, the exponent (n) is not restricted to a positive integer, and may take on negative, or even fractional values. A traditional hill function with cooperativity n and concentration and half-maximum K is obtained by setting a = 0 and T = v = 1. Multiple inducers  $X_1, X_2, \ldots, X_n$  can be combined in a single arrow,

$$[[X1, X2, ..., Xn] | ->Y, Hill[v, n, K, a, [T1, T2, ...Tn]]]$$
 (20)

174 This is described by differential equation terms

$$\frac{dY}{dt} = \frac{v(a + \sum T_j X_j)^n}{K^n + (a + \sum T_j X_j)^n}$$
(21)

A facilitated version of the Hill arrow is also available, which multiplies the corresponding differential equation terms by an optional modifier.

$$[[X1, X2, ..., Xn] | --> Y, mod[E], Hill[V, n, K, a, [T1, T2, ...Tn]]]$$
 (22)

The differential equation terms produced by equation (19) for Y and equation (21) for  $Y_1, \ldots, Y_n$  will be added to the other differential equations for those variables.

#### $_{179}$ 2.1.5 GRN Arrows

Genetic Regulatory Network (GRN) arrows are useful for modeling transcriptional networks, gene regulation, and any interactions involving bistability or switching. They can be numerically fit to 181 molecular sub-networks to describe overall input-output behavior without actually describing the 182 specific molecular mechanisms occurring within the sub-network. The GRN functions used in 183 Pycellerator are logistic functions. The slope and location of the decision/threshold boundary can be optimized to fit available data. [12] Logistic functions are commonly used in machine learning 185 to solve decision problems and as threshold functions in neural network models. The probability 186 distribution described by a logistic function can be related to a two state Boltzmann distribution 187 or softmax process. [13] 188

189 GRN arrows are summarized in Table 4. The basic GRN arrow in Pycellerator is

$$[X \mid ->Y, GRN[V, T, n, h]]$$
(23)

where v, T n, and h are constants that may be set to any floating point value. In particular, there is no restriction that exponent n be integer, and it may take on fractional or negative values. This

192 produces the differential equation term

$$\frac{dY}{dt} = \frac{v}{1 + e^{-(h+TX^n)}}\tag{24}$$

The GRN arrow does not affect the differential equation for the variables on the left side of the equation (X in this case). A standardized logistic function  $1/(1 + e^{-x})$  is obtained by setting v = T = n = 1 and h = 0. Extended forms of the GRN arrow include an optional modifier species that multiplies the rate v and the use of multiple input species.

$$[[X1, X2, ..., Xk] | ->Y, GRN[v, [T1, T2, ..., Tk], n, h]]$$

$$[[X1, X2, ..., Xk] | -->Y, mod[E], GRN[v, [T1, T2, ..., Tk], n, h]]$$

$$(25)$$

197 This changes the differential equation term to

$$\frac{dY}{dt} = \frac{vE}{1 + e^{-(h + \sum T_j X_j^n)}} \tag{26}$$

If the mod[E] is omitted in the arrow then the E is omitted from the equation.

## 199 2.1.6 Rational Functions

Rational functions produce rate laws that are described by quotients of polynomials. Each term in
the polynomial may be a product of species raised to a power. Only the species on the right hand
side of the arrow are affected by the reaction. The simple form of the rational arrow is

The corresponding contribution to the rate law is

$$\frac{dZ}{dt} = \frac{a_0^{m_0} + a_1 X_1^{m_1} + a_2 X_2^{m_2} + \dots + a_p X_p^{m_p}}{d_0^{n_0} + d_1 Y_1^{n_1} + d_2 Y_2^{n_2} + \dots + d_q Y_q^{n_q}}$$
(28)

In the more general case, each  $X_i$  or  $Y_j$  can be replaced by a product of species. For example, the arrow

206 contributes the single differential equation term (in Python)

$$A' = (A**m1*a1 + a0**m0 + a2*(B*C)**m2 + a3*(D*E*F)**m3) /$$

$$(A**n1*b1 + B**n2*b2 + b0**n0 + b3*(B*C)**n3$$

$$+ b4*(B*C*D)**n4 + b5*(B*C)**n5 + b6*(B*D)**n6)$$
(30)

An example that includes the use of rational functions is given by the implementation of plant stem cell lineage in the distribution folder (file chickarmane.model in the models folder).

## 209 2.1.7 Generalized MWC

The Monod-Wyman-Changeaux (MWC) model [14] describes allosteric enzymes with multiple
binding sites that influence one another's affinities. In addition, such an enzyme is typically
composed of multiple sub-units that may exist in different states or conformations. We follow the
"generalized" MWC model of [15], which also accounts for for multiple activator and inhibitor
factors in allosteric enzymes. The basic generalized MWC arrow is

$$[S==>P, mod[E], MWC[k, n, c, L, K]]$$
(31)

where k, n, c, L, and K are constants. This produces differential equation terms for both S and P.

$$\frac{dP}{dt} = E \frac{s(1+s)^{n-1} + Lsc(1+sc)^{n-1}}{(1+s)^n + L(1+sc)^{n-1}} = -\frac{dS}{dt}$$
(32)

where s = S/K. The generalized arrow is

$$[[S1,..] ==> P, mod[E, [A1,..], [I1,..], [[CI1,..], [CA1,..]], MWC[k, n, c, L, K]]$$

$$(33)$$

Here  $A_i$ ,  $I_i$ ,  $C_{ij}$  are optional sequences of activators, inhibitors and competitive inhibitors at the substrate and activator site, and K is a list of constants

$$[K_{S1}, K_{S2}, ..., K_{A1}, K_{A2}, ..., K_{I1}, K_{I2}, ..., K_{CI1}, ..., K_{CA1}, ..]$$
(34)

Let  $s_j = S_j/K_{Sj}$ ,  $a_j = A_j/K_{Aj}$ ,  $i_j = I_j/K_{Ij}$ ,  $\overline{s_j} = c\sum_k C_{jk}/K_{C_{jk}}$ , and  $\overline{a_j} = c\sum_k C_{jk}/K_{CA_{jk}}$ . Define the intermediate terms

$$\mathcal{A} = \prod (1 + a_j + \overline{a_j})^n \tag{35}$$

$$\mathcal{I} = \prod (1 + i_j)^n \tag{36}$$

$$S = \prod (1 + s_j + \overline{s_j})^{n-1} \tag{37}$$

$$S_c = \prod (1 + cs_j + \overline{s_j})^{n-1} \tag{38}$$

219 Then the generalized model generates terms

$$\frac{dP}{dt} = -\frac{dS_i}{dt} = E \frac{\mathcal{AS} \prod s_j + L\mathcal{IS}_c \prod (cs_j)}{\mathcal{A} \prod (1+s_j)^n + L\mathcal{IS}_c \prod (1+cs_j)}$$
(39)

220 in the system of differential equations.

#### 221 2.1.8 NHCA

The basic form for Non-hierarchical cooperative activation (NHCA)[16, 17] is

$$[[X1, X2, ...] | -->Y, mod[E], NHCA[v, [TP1, ...], [TM1, ...], [n1, ...], m, k]]$$
 (40)

where X1,.. are one or more reactants, Y is the product, E is a modifier, TP1, TP2, ...

TM1, TM2, ..., n1, n2, ..., v, m and k are numeric parameters. The corresponding rate law is

$$\frac{dY}{dt} = vE \frac{\prod_{i} (1 + T_{Pi}X_{i}^{n_{i}})^{m}}{k \prod_{i} (1 + T_{Mi}X_{i}^{n_{i}})^{m} + \prod_{i} (1 + T_{Pi}X_{i}^{n_{i}})^{m}}$$
(41)

#### 225 2.1.9 User Defined Arrows

Users can define arrows with their own rate laws. Let X1, X2, ... and Y1, Y2, ... be reactants and products, respectively, with numeric stoichiometries e1, e2, ..., and f1, f2, ... Then the basic arrow form

$$[e1*X1 + e2*X2 + ... -> f1*Y1+f2*Y2+...,using[expr]]$$
 (42)

Here using is a Pycellerator keyword and *expr* represents any evaluatable Python expression involving model species. The user arrow contributes differential equation terms

$$\frac{dZ}{dt} = (f_z - e_z) \times (expr) \tag{43}$$

for each model species Z that appears in a user reaction, where  $f_z$  and  $e_z$  are the stoichiometry of Z on the right and left hand sides of the arrow.

233 Similarly, a user-defined regulatory arrow takes the form

$$[[X1, X2, ..., Xk] | -> Y, USER[v, [T1, T2, ..., Tk], [n1, n2, ..., nk], h, f]]$$
 (44)

Here X1, X2, ..., Xk are the input variables, whose values are not affected by the arrow; Y is the output variable; v, T1, ..., Tk, n1, ..., nk, h are numeric parameters; and f is a function defined in the \$Functions section of the model file. The arrow contributes the following term to the differential equation for Y:

$$\frac{dY}{dt} = vf\left(h - \sum_{i} T_i X_i^{n_i}\right) \tag{45}$$

238 As a simple example, the following partial model file:

```
$\text{Spanning spanning span
```

This be converted to the following equations (in Python):

The lambda expansion gives a rate law term for Y of

$$\frac{dY}{dt} = -kY + \frac{1}{1 + e^{TX^n + h}} \tag{46}$$

This is similar to a generalized GRN expansion with arbitrary exponent.

#### 251 2.1.10 Cascades

A Pycellerator cascade is sequence of repeated reactions with the same arrow. For example, the enzymatic arrows

can be combined into a single arrow

$$[MAPK => MAPKp => MAPKpp, mod[KKpp], rates[a,d,k]]$$
 (48)

as shown in listing 2.

Any mass action, MMH, Hill, GRN, SSystem, or NHCA reaction can be written as a cascade to

reduce the number of arrows in the model. The reduction can be significant, especially when

cascades are combined with enzymatic expansion. A three stage MAPK cascade might be written as

```
[KKK <=> KKKp, mod[S, KKKph], rates[a1,d1,k1,0,a4,d4,k4]]

[KK <=> KKp <=> KKpp, mod[KKKp,KKph], rates[a3,d3,k3,0,a5,d5,k5]]

[MAPK <=> Kp <=> Kpp, mod[KKpp,Kph], rates[a3,d3,k3,0,a6,d6,k6]]

(49)
```

This can be further simplified with an indexed notation:

```
[K(3,0) <=> K(3,1), mod[S, KKKph], rates[a1,d1,k1,0,a4,d4,k4]]

[K(2,0) <=> K(2,1) <=> K(2,2), mod[K(3,1),KKph], rates[a3,d3,k3,0,a5,d5,k5]]

[K(1,0) <=> K(1,1) <=> K(1,2), mod[K(2,2),Kph], rates[a3,d3,k3,0,a6,d6,k6]]

(50)
```

A three stage MAPK cascade including stimulation and feedback can be written with only five arrows using <=> cascades. Using only the forward arrow cascades (=>) this can be done in eight reactions. Without any cascades, but still using enzymatic => arrows, twelve reactions are required. Each of these models would expand to the same system of 34 of simple mass action reactions, which would have to be typed in manually without using any enzymatic expansion.

## 266 2.2 Flux Models

A Pycellerator model may be composed either entirely of kinetic arrows as described above) or
entirely of flux arrows. The two may not be combined. Reactions that represent fluxes are a
fundamentally different type of entity than reactions used in kinetic models. This is because flux
reactions do not (necessarily) have a rate law or differential equation (of the same sort) associated
with them. What they do normally have is a total rate, given by the product of a velocity and a
stoichiometry.

273 The format of a flux arrow is

where X1, X2, ... and Y1, Y2, ... are species; e1, e2, ... and f1, f2, ... are stoichiometries;
var is an identifier used to refer to the flux variable for the reaction; low and up are numeric
lower and upper bounds for optimization; obj is a numeric objective coefficient; and fluxvalue
is an optional numeric flux value.

Users should be cautioned that the symbol "<" used in the Flux arrow is inclusive and really corresponds to the mathematical inclusive "less than or equal to" symbol, "≤." Optimization is inclusive, not exclusive. There is no less than or equal to symbol in Pycellerator, only the single "<" is used.

282 For example,

$$[ES \rightarrow E + S, Flux[0 < v < 1, 1, 0]$$
 (52)

The flux arrow means that the optimization is performed so that  $0 \le v \le 1$ , even though only the "<" is used. Attempting to include an equal sign in the expression will lead to a syntax error.

To force equality, say v = 1, in a constraint, one would use Flux[1<v<1]. The flux optimization process will solve the linear programming problem

maximize 
$$\mathbf{v^T f}$$
  
subject to  $\mathbf{Nv} = \mathbf{0}$   
and  $low_1 \leq v_1 \leq up_1$   
and  $low_2 \leq v_2 \leq up_2$   
 $\vdots$  (53)

where  $\mathbf{v}$  is the vector of fluxes  $(v_1, v_2, \dots)^T$ ,  $\mathbf{N}$  is the stoichiometry matrix, and  $\mathbf{f}$  is the vector of objective coefficients. The optimization is performed using Python's pulp package. [18]

## $_{289}$ 2.3 Functions

The \$Functions section of the model file contains a list of function definitions in standard algebraic notation. A function may have multiple arguments and these are treated as dummy parameters. When the function is instantiated the parameters are replaced with the arguments used in the function invocation. Within the function any other global parameters (such as rate constants) may be referenced. The general format is

$$f(v1, v2, \dots vn) = expr$$
 (54)

where f is the function name, as it is used elsewhere in the model; v1, ..., vn are the function dummy arguments, as they are referenced on the right hand side of the function definition; and expr is a standard arithmetic expression that is evaluable in Python. All variables and parameters must be conform to the rules for permissible Python identifiers (e.g., case sensitive, alphanumeric, must start with with a letter). When the model is converted to Python, each function is converted to a Python lambda expression.

An example with two functions is given in the minimal cascade model for a mitotic oscillator (listing 3, [19]). The function f (m, x) has two arguments,

$$f(m, x) = m * (1-x) / (K3+1-x)$$
 (55)

This function is instantiated as f (M, X) to define a concentration dependent rate constant for the reaction [Nil -> X]. The second function g (m) has one argument,

$$g(m) = (1-m) / (K1+1-m)$$
 (56)

and is instantiated as g (M) to produce a concentration dependent rate for the hill function. In
this particular model, the functions use variable names that are simular to (e.g., lower case
versions) of the variables used in their instantiations. No such restriction is actually placed on the
user, and equation (57) could just as well have been implemented as

$$f(foo, bar) = foo* (1-bar) / (K3+1-bar)$$
 (57)

## $_{ m 309}$ 2.4 Assignments

The optional \$Assignments section of the model file contains a list of species definitions as set equal to statements. The general format is

$$X = expr (58)$$

where X is the species name, as it is used elsewhere in the model, and *expr* is any Python
expression. These assignments hold at all times throughout a simulation. If X is a species that
would otherwise be define by a differential equation, then it should also be listed in the \$FROZEN
section to ensure that the differential calculation is inhibited.

## 316 2.5 Initial Conditions

Initial conditions are defined in the \$IC section. A species is not required to have an initial condition, but if an initial condition is omitted, it is assumed to be zero. The \$IC section contains a sequence of statements of the form

$$X = value$$
 (59)

where X is the species name, and value is the numeric value of the species at t = 0. If a variable is specified by an assignment rule then it should not be given an initial condition.

## 322 2.6 Parameter Values

Parameter values are defined in the \$Rates section. All constants and parameters that are
defined using an identifier in the model must be given a value in this section. The \$Rates section
contains a sequence of statements of the form

$$identifier = value$$
 (60)

where identifier is the parameter name, and value is the numeric value of the parameter.

Parameters can also be replaced with algebraic (Python) expressions in the \$ASSIGNMENT

sections. If a parameter is listed on the left-hand side of an assignment statement then its parameter value will be ignored.

## $_{30}$ 2.7 Frozen Variables

Frozen species do not contribute terms to the system of differential equations. However, if a variable is frozen but is given an assignment rule, it may still change as a function of time. This provides a convenient way to provide a time-dependent input. Frozen species are listed in the \$Frozen section of the model file. This section contains a list of the frozen species, one species per line. Only species may be frozen, not other parameters in the model.

## 336 2.8 Identifiers and Symbols

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Identifiers in the model, i.e., species representing reactants, products and modifiers; function names; and parameters (rate constants), must start with a letter and may contain any number of 338 alphanumeric characters in them. Identifiers are case sensitive, and may also contain the 339 underscore character. Users should beware that the underscore character is also used by 340 Pycellerator to join species names when auto-generating new species names (for example, see reactions (11)). 342 The special identifier Nil is used to represent the empty set and it is not converted into a differential equation term. Thus reactions such as [Nil -> X] and [X -> Nil] represent 344 cretio ex nihilo and removal from the system, respectively. 345 The special identifier t may be used in functions and assignments to define explicit time 346 dependent expressions. For example, a constant stimulation of S=1 may be turned on from

```
$\frac{349}{5}$
$$ $S$
$$ $S$
$$ $Assignments$
$$ $S = (0.0 if t<100 else (stim if t<200 else 0.0))$
```

t = 100 to t = 200 by setting S as a frozen variable and using an assignment:

- Species may be indexed using parenthesis, e.g., [X(1) -> Y(2)] or [K(2,0) => K(2,1) => 356 K(2,2)]. When the model is instantiated the index numbers are embedded into the variable name; they are not implemented as either Python arrays or lists.
- 358 If more than one statement is placed on a single line in the model file, the statements should be separated by a semicolon.

# 3 Methods

This section describes how to install the necessary software; instantiate models (generate Python functions); and run simulations using Pycellerator.

## 3.1 Requirements

## 3.1.1 Install Python

- 1. Install Python 2.7 if it is not already present (see Note 1). You will probably need to run
  the instllation in adminstrator mode (Windows) or sudo (Linux).
- On Windows, it is generally easier to install a complete scientific version. Links for several complete scientific packages are given at https://www.scipy.org/install.html.
- On Macs, Python is already installed by default, and it is not normally necessary to reinstall it.
- Linux users should be able to install Python using their package manager.
- 2. Install setuptools using pip. The program pip should automatically be installed when Python is installed.
- In a Windows command shell (it is called cmd.exe), type the following:

```
python -m pip install -U pip setuptools
375
         From the terminal program on a Mac or in Linux, type
376
                       pip install -U pip setuptools
377
         If pip fails to run in this manner, follow the instructions at
378
         https://packaging.python.org/installing to download and install get-pip.
379
         Then repeat this step.
380
      3. Install the required packages: numpy, scipy, sympy, matplotlib and pyparsing.
381
         From the command shell (any operating system),
382
                       pip install numpy scipy sympy matplotlib pyparsing
383
         Linux users may prefer to install these packages from their package repositories, but it does
384
         not matter whether you use the repository or pip
385
      4. (Optional) Install the optional packages pulp, ipython and jupyter. If pulp is not
386
         installed, then flux models cannot be solved. If ipython and jupyter are not installed,
387
         then the notebook interface will not be available. From the command shell (any operating
388
         system),
389
                       pip install pulp ipython jupyter
390
      5. (Optional) Install libsbml for Python 2.7. To be able to either read or write SBML files you
391
         must install libsbml. For most operating systems, type the following in the command shell
392
                       pip install python-libstall
393
         Before you do this, check for operating-system-specific instructions at
394
         http://sbml.org/Software/libSBML/docs/python-api/.
395
```

## 6 3.1.2 Pycellerator Installation

It is not necessary to use administrator or superuser mode to install Pycellerator.

- 1. Download Pycellerator from the github repository at

  https://github.com/biomathman/pycellerator/releases (see Note 2). Look

  for the file install-pycellerator-v-X.zip (where X is some number) and download

  that file to your computer. Advanced users may be more interested in the source but you
- 2. Unzip and create working folder. Unzip the download, which will probably be in your 403 Downloads folder. Look for a folder called pycellerator in that unzipped file. Copy this 404 entire folder anywhere you want on your disk drive, such as your home folder or your 405 desktop. This is going to be your working folder for Pycellerator. It is not necessary to 406 modify your Python path so long as you run models from this folder. 407 You should see two folders inside the pycellerator folder: cellerator and models. 408 The cellerator folders contains code needed to run the program and should not be 409 modified. The models folder contains sample model files. The distribution includes 410

## 412 3.2 Model Instantiation and Simulation

don't need that to run to Pycellerator.

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## 413 3.2.1 Plot Time Course in A Notebook

documentation (see Note 3).

- Here we consider simulation and plotting of the basic model shown in listing 1. This model contains a single arrow representing reaction (??) using enzymatic expansion (arrow (8)). In the model file, only the first three of the four rate constant are specified, so the fourth rate constant defaults to zero.
- 1. Set your current working directory to the pycellerator folder that you created during installation. You can do this, e.g., by opening the folder in your desktop manager.
- 2. Using a text editor open a new text file and copy or type the contents of listing 1 into it.

  Note that if you cut and paste from an electronic version of this paper, the fonts will most

likely generate a few incompatible characters. It is best to verify that only valid text
characters (e.g., UTF-8) are in your file. Then save your file as basicmodel.model in the
current working directory.

3. Open the jupyter notebook interface. To do this, open a command shell (cmd.exe in Windows, terminal in MacOS or Linux) and type

ipython notebook

This will open the jupyter notebook interface in your default browser.

4. Create a new notebook. From the drop down menu near the top right of the jupyter window, select New > Python 2. This will open a new window labeled "untitled." From the drop down menu on the top left of the window select File > Rename. Type a name for the notebook, such as "Basic-Model" in the pop-up window and click OK. Your file will be named Basic-Model.ipynb. The file extension ipynb is required.

If you click back on the tab Home in your browser you should see a list of files. One of those files will be the file Basic-Model.ipynb that you just created. If you go to your desktop and open a folder, you will also see the file. (Note that some operating systems may suppress visibility of the file extension (the letters after the dot in the file name) when you look at your list of files this way.) You will not be able to edit or modify this file except using the jupyter interface because it is written in a special format that is called JSON. If you open it up and look at it in any other format it will probably look like nonsense to you.

5. Click on the tab for you notebook. In the first cell type in the required Python includes:

from cellerator import cellerator as c
import matplotlib.pyplot as plt
import numpy as np
%matplotlib inline

After entering code in any cell, click on enter to ensure that the code is executed.

Note that pyplot and numpy are not strictly necessary as separate imports. They are used

- inside the program, but not directly accessible by user. If you want to make modifications using pyplot or numpy features you may need to import them.
- 6. To determine the differential equations for the model and print them to the screen in

  Python form,

```
model="basicmodel.model"

c.PrintODES(model)
```

454 For this model, the output should be

```
455 E' = -E*S*a + S_E*d + S_E*k
456 S' = -E*S*a + S_E*d
457 S_E' = E*S*a - S_E*d - S_E*k
458 P' = S_E*k
```

7. Solve the model. The basic function is c.Solve:

```
t, v, s=c.Solve(model)
```

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Here model is the file name as before, and the return value is a Python tuple (t, v, s).

- t is a numpy array of times at which the solution is returned in s. it is the return value of odeint. The default setting for t=[0,1,2,..,100]. These can be changed with the keywords step and duration. Note that step only controls what is returned, and is not related to the integration step size.
- v is a list of variable names (as strings); in this case the return value would be ['E', 'S', 'S\_E', 'P']
- s is a numpy array of solution vectors, one vector per time point, as returned by odeint.
- 8. Plot the results. The basic function is c.PlotAll, which takes the three variables returned by c.Solve and returns a Pyplot axis object. As long as the line matplotlib inline was executed prior to this step (see step 5), the plot will be displayed in the next cell of your notebook.

```
ax=c.PlotAll(t,v,s)
```

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9. Tweak the run and plot parameters. To get a more precise plot, we can re-run the simulation with an output step of 0.1. Since the interesting stuff happens early (with the given values of the parameters) will also only need to run for a short time. Then we can use pyplot to add axis labels, change scales, etc.

```
t, v, s=c.Solve (model, step=.1, duration=15)
                    ax=c.PlotAll(t,v,s)
480
                    ax.set_yticks(np.arange(0,1.,.2))
481
                    ax.set_yticklabels(np.arange(0,1.01,.2),fontsize=12)
482
                    ax.set_xlim(0,15)
483
                    ax.set_xticks(np.arange(0,15.1,5))
484
                    ax.set_xticklabels(np.arange(0,15.1,5),fontsize=12)
485
                    ax.set_xlabel("Time", fontsize=14)
486
                    ax.set_ylabel("Value", fontsize=14)
                    ax.set_title("The Results of My Simulation", fontsize=16)
488
                    fig=plt.gcf()
489
                    fig.set_size_inches(6,3)
490
                    fig.tight_layout()
491
                    fig.savefig("basicmodel.pdf")
492
```

The resulting plot is shown in figure 1.

## 494 3.2.2 Run Auto-generated Code as Stand-Alone Program

model="basicmodel.model"

1. Locate the auto-generated code produced by Pycellerator. The default file name is

solver\_for\_model\_timecode.py, where model is the model name (e.g., basicmodel

in the previous section); and timecode is a time code to uniquely identify the file. The

default file name can be overridden in c.Solve with the keyword solverfile:

```
t, v, s=c.Solve (model, solverfile="foo", step=.1, duration=15)
```

- 2. Locate and examine the auto-generated code (e.g., foo.py). The code produced in this model is shown in listing 4. As the program stands right now, nothing would normally be output. The code can be modified with any standard text editor.
- 3. Modify the autogenerated code (e.g., foo.py). For example, to print a comma-separated value listing of the result to the screen, add the following code before the return statement of thesolver(), between lines 41 and 42,

```
print "t,"+",".join(variables)+","

for t,v in zip (times,sol):

print ",".join(map(str,list([t])+list(v)))
```

4. Run the program. Type the following in a command shell.

```
python foo.py
```

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#### 512 3.2.3 Run and Plot a Model From the Command Shell

1. To run the basic model with a step size of 0.1 and duration 15, plot the results, and save the results to a CSV file, type the following in the command shell on a single line

```
python pycellerator.py solve -run 15 .2 -in basicmodel.model

-plot -pyfile spam.py -out eggs.csv
```

The plot should pop up as a separate window. In some cases it might be hidden behind existing windows. The auto-generated code is written to spam.py and the results of the simulation are saved to eggs.csv. Additional options are described in the users guide.

2. Optionally modify and rerun the code. To re-run the code generated in step 1, enter

```
python spam.py
```

from the command shell. The auto-generated code for the model function is identical to the code generated in the notebook. The code generated for the driver is different, since it includes a wrapper for output. If this code is run from the command line, the plot will automatically pop up, and a new CSV file will be generated.

## 526 3.2.4 Perform Parametric Tweaks and Scans

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In listing 5 we show a toy example of the spread of disease based on the SIRS model, implementing the following system of differential equations, based on the Kermack-McKendrick model with feedback, birth, and death. [22].

$$I' = kIS - (1+d)I$$

$$S' = -kIS + bI + R(b+f) + (b-d)S$$

$$R' = I - R(d+f)$$
(61)

The populations of S (susceptible), I (infected), and R (recovered) are dimensionless; k is the ratio of infection to recovery rate (hence non-dimensional); f is some fraction of the recovered population that returns to the susceptible population; and b and d are the population birth and death rates. All newborns are assumed to be susceptible.

1. Tweak individual parameters using c.Solve. For example, to override the initial conditions for R and S, and the value for f in the model file, set them at run time. The options IC and RATES are Python dictionaries.

```
model="SIRS.model"

t, v, s = c.Solve(model, step=.1, duration=100,

IC={"R":.5,"S":.5},

RATES={"f":.5})

c.PlotAll(t,v,s)
```

2. Do a parametric scan. To determine the values of all the state variables at, say, t = 200, as

a function f for  $0.5 \le 5 \le 1$  in steps of 0.5, use the scan keyword.

```
variables, pscan=c.Solve(model,

scan=["f",0.05,1,0.05], duration=200)
```

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In this case c.Solve returns a 2-tuple rather than a 3-tuple. The first item is a list of the variables in the model, as strings. For this model, it will return the list ['I', 'S', 'R']. The second item is a numpy array of vectors, where each vector has the form (in this case) [f, I, S, R]. The state variables (i.e., I, S, and R) are evaluated at the very end of the simulation. Typically this would be when the simulation reaches steady state, but some knowledge of the model is required to verify this. Pyellerator does not verify steady state; it merely returns the values at the time requested.

3. Plot the parametric scan. This can be done using pyplot, or the data can be exported to a spreadsheet or plotting program. For example, to plot the fraction recovered (R) as a function of f,

```
fvals=pscan[:,0]; RVals=pscan[:,3]

plt.plot(fvals, RVals,marker="o")

plt.xlabel("f",fontsize=14)

plt.ylabel("Fraction Recovered", fontsize=14)

plt.title("SIRS model", fontsize=16)
```

The resulting parametric scan is show in figure 2.

#### 562 3.2.5 Include a Time Dependent Stimulation

The easiest way to include a time-dependent stimulation in a model is as follows.

1. Add a dummy reaction that would normally create a steady state value for your stimulation, such as

$$[Nil < -> S, rates[a0, d0]]$$
 (62)

- This tells the system to treat S as a species. If you omit this reaction, S will be considered
  an unknown variable during the simulation and the program will terminate with an error.

  Normally reaction (62) would lead to a differential equation of the form S'=a0-d0\*S.

  However, this is overridden in the following step.
- 2. Make S a frozen variable by adding a line containing S to the \$Frozen section in the model file. This tells the Pycellerator to replace the differential equation in step 1 with S'=0.
- 3. Define an assignment rule that explicitly gives the value of S as a function of time using standard Python. This tells Pycellerator to replace the differential equation for S with an algebraic expressio for S. For a square pulse, use a Python ternary operator:

```
S=(0.0 \text{ if } t<t1 \text{ else } (K \text{ if } t<t2 \text{ else } 0.0))
```

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in the \$Assignments block. The values of t1, t2, and K can be initialized in the \$Rates block. This way you can manually override the values during a simulation without editing the file. To use a more complex stimulation and make the model file more readable, use a function in the model file.

```
$Assignments

$S=(0.0 if t<t1 else (f(t) if t<t2 else 0.0))

$Functions

$f(t)=sin((t-1000.0)*pi/(3000))
```

Any function in the Python math library may be referenced.

4. Run the simulation. For a model with a large number of variables, plot the results in a grid.

```
t, v, s = c.Solve(model, step=.1, duration=5000)

c.PlotColumns(t,v,s,ncols=4,bg="white",colors=23*["black"])
```

Verify the stimulation on the plots (e.g., figure 3).

# 589 4 Notes

- 1. Pycellerator is implemented in Python 2.7. There are no plans at the present time to implement the program in Python 3.
- 2. Pycellerator can be freely downloaded from github. A public respository is located at https://github.com/biomathman/pycellerator/releases. All software is covered by a GPL version 3 license.
- 3. The complete syntax and all options are detailed in the user manual that is included with the download package.

Table 1: Arrows and keywords used in Pycellerator arrows expressions.

Arrow	Keyword	modifer	Description Typical usages
->	N/A	no	Mass action
>	N/A	yes	Mass action with modifier
<->	rates	no	Mass action
=>	rates	yes	Mass action expansion, SE complex
:=>	rates	yes	Mass action expansion, SE and PE complex
<=>	rates	yes	Mass action expansion, SE complex
->	Hill	no	Hill Function
	GRN	no	Generalized logistic rate function
	SSystem	no	S-System
	USER	no	User defined rate law
>	Hill	yes	Hill Function
	GRN	yes	Generalized logistic rate function
	NHCA	yes	Non-hierarchical cooperative activation.
	USER	yes	User defined rate law
:->	MMH	No	Michaelis-Menten-Henri-Briggs-Haldane
:>	111,111	Yes	
==>	MWC	yes	Monod-Wyman-Changeaux model.
	Rational	no	Rational Function

Table 2: Mass Action Reactions

Pycellerator Syntax	Biochemical Notation	Note
[ A + B -> C, k ]	$A + B \xrightarrow{k} C$	(a)
[ A + B <-> C, rates[k1, k2] ]	$\begin{cases} A + B \xrightarrow{k_1} C \\ C \xrightarrow{k_2} A + B \end{cases}$	(a)
[ $e1*X1 + e2*X2 + \cdots ->$ $f1*Y1 + f2*Y2 + \cdots$ , k]	$\sum_{i} e_{i} X_{i} \xrightarrow{k} \sum_{j} f_{j} Y_{j}$	(b,c)
<pre>[ e1*X1 + e2*X2 + ··· &lt;-&gt;   f1*Y1 + f2*Y2 + ···, rates[k1,k2]]</pre>	$\sum_{i} e_{i} X_{i} \stackrel{k_{1}}{\rightleftharpoons} \sum_{j} f_{j} Y_{j}$	(b,c)
[ S=>P, mod[E], rates[k1,k2,k3,k4] ]	or: $\begin{cases} S + E \stackrel{k_1}{\rightleftharpoons} SE \stackrel{k_3}{\rightleftharpoons} P + E \\ S + E \stackrel{k_1}{\rightleftharpoons} SE \\ SE \stackrel{k_2}{\rightleftharpoons} S + E \\ SE \stackrel{k_3}{\rightleftharpoons} P + E \\ P + E \stackrel{k_4}{\rightleftharpoons} PE \end{cases}$	
[S<=>P, mod[F,R], rates[k1,k2,k3,k4, k5,k6,k7,k8]]	or: $\begin{cases} S + F \stackrel{k_3}{\rightleftharpoons} SF \stackrel{k_3}{\rightleftharpoons} P + F \\ P + R \stackrel{k_5}{\rightleftharpoons} PR \stackrel{k_7}{\rightleftharpoons} S + R \\ S + F \stackrel{k_1}{\rightleftharpoons} SF & SF \stackrel{k_2}{\rightleftharpoons} S + F \\ SF \stackrel{k_3}{\rightleftharpoons} P + F & P + F \stackrel{k_4}{\rightleftharpoons} SF \\ P + R \stackrel{k_5}{\rightleftharpoons} PR & PR \stackrel{k_6}{\rightleftharpoons} P + R \\ PR \stackrel{k_7}{\rightleftharpoons} S + R & S + R \stackrel{k_8}{\rightleftharpoons} SR \end{cases}$	
[S:=>P,mod[E],rates[k1,k2,k3,k4,k5,k6]]	or: $\begin{cases} S + E \stackrel{k_1}{\rightleftharpoons} SE \stackrel{k_3}{\rightleftharpoons} PE \stackrel{k_5}{\rightleftharpoons} P + E \\ S + E \stackrel{k_1}{\Rightarrow} SE & SE \stackrel{k_2}{\Rightarrow} S + E \\ SE \stackrel{k_3}{\Rightarrow} PE & PE \stackrel{k_4}{\Rightarrow} SE \\ PE \stackrel{k_5}{\Rightarrow} P + E & P + E \stackrel{k_6}{\Rightarrow} PE \end{cases}$	

<sup>(</sup>a) May be multiple reactants and products. (b) The stoichiometries ei and fj are numeric. (c) The multiplication symbol (asterisk, "\*") between the stoichiometry and species is optional; however, the stoichiometry must come first, and be numeric. If the stoichiometry is equal to one, it may be omitted.

Table 3: Michaelis Menten type arrows in Pycellerator.

Pycellerator Syntax	Rate Law
[S:->P,MMH[K, v]]	$\frac{dP}{dt} = -\frac{dS}{dt} = \frac{vS}{K+S}$
[S:->P, mod[E], MMH[K, v]]	$\frac{dP}{dt} = -\frac{dS}{dt} = \frac{vSE}{K+S}$
$[S:->P,MMH[k_1,k_2,k_3]]$	$\frac{dP}{dt} = -\frac{dS}{dt} = \frac{k_3 S}{(k_2 + k_3)/k_1 + S}$
[S:->P, mod[E], MMH[k <sub>1</sub> , k <sub>2</sub> , k <sub>3</sub> ]]	$\frac{dP}{dt} = -\frac{dS}{dt} = \frac{k_3 SE}{(k_2 + k_3)/k_1 + S}$

Table 4: Regulatory arrows in Pycellerator. Regulatory arrows only affect the variables on the right-hand side of the arrow symbol; they do not contribute differentiatial equation terms to variables on the left hand side.

Type	Pycellerator arrow	Differential equation term
	[X ->Y, Hill[v,n,K,a,T]]	$Y' = \frac{v(a+TX)^n}{K^n + (a+TX)^n}$
Hill	[[X1,X2,,Xn] ->Y, Hill[v,n,K,a,[T1,T2,,Tn]]]	$Y' = \frac{v(a + \sum T_j X_j)^n}{K^n + (a + \sum T_j X_j)^n}$
	[[X1,X2,,XN] >Y, mod[E],Hill[v,n,K,a, [T1,T2,,Tn]]	$Y' = \frac{vE(a + \sum T_j X_j)^n}{K^n + (a + \sum T_j X_j)^n}$
GRN	[X ->Y,GRN[v,T,n,h]]	$Y' = \frac{v}{1 + e^{-(h + TX^n)}};$
	[[X1,X2,,Xn] ->Y, GRN[v,[T,],n,h]]	$Y' = \frac{v}{1 + e^{-(h + \sum T_j X_j^n)}}$
	[[X1,X2,,Xn] >Y, mod[E],GRN[v,[T,],n,h]]	$Y' = \frac{vE}{1 + e^{-(h + \sum T_j X_j^n)}}$
S-System	[[X1,,Xn] ->Y,SSystem[tau, a,b,[g1,,gn],[h1,,hn]]	$Y' = \frac{1}{\tau} \left( a \prod_{i} X_i^{g_i} - b \prod_{i} X_i^{h_i} \right)$
Rational	<pre>[[[X1, X2,], [Y1, Y2,]] ==&gt;Z,</pre>	$Z' = \frac{a_0^{m_0} + \sum_i a_i X_i^{m_i}}{d_0^{m_0} + \sum_i d_i Y_i^{n_i}}$
	<pre>[[[X11*X12*, X21*X22*, ],     [Y11*Y12*, Y21*Y22* ]]] ==&gt;Z,rational[[a0,a1,a2,],     [d0,d1,d2,],[m0,m1,m2,],     [n0,n1,n2,]]]</pre>	$Z' = \frac{a_0^{m_0} + \sum_i a_i (X_{i1} X_{i2} \cdots)^{m_i}}{d_0^{m_0} + \sum_i d_i (Y_{i1} Y_{i2} \cdots)^{n_i}}$

Listing 1: A basic model describing the reaction  $S+E \overset{a}{\underset{d}{\rightleftarrows}} SE \overset{k}{\xrightarrow{}} P+E$  using three elementary reactions.

```
597
                                         $REACTIONS
598
                                          [S+E \rightarrow SE, a]
599
                                          [SE \rightarrow S+E, d]
600
                                          [SE \rightarrow P+E, k]
601
                                         $IC
602
                                          S = 1
603
                                          E = 1
604
                                          P = 0
605
                                         $Rates
606
                                          a = 1
607
                                          d = 1
608
609
610
```

Listing 2: Model of MAPK oscillation demonstrating the use of cascades and an external stimulation. Stimulation is provided by species S.[20, 21].

```
611
   $Reactions
612
    [Nil<->S, rates[a0, d0]]
613
    [KKK <=> KKKp, mod[S, KKKph], rates[a1,d1,k1,0,a4,d4,k4]]
614
    [KK <=> KKp <=> KKpp, mod[KKKp, KKph], rates[a3,d3,k3,0,a5,d5,k5]]
615
    [MAPK <=> Kp <=> Kpp, mod[KKpp,Kph], rates[a3,d3,k3,0,a6,d6,k6]]
616
    [KKK_S + Kpp <-> KKK_S_Kpp, rates[a7, d7]]
617
618
     KKK = 100;
                   KKKp = 0
619
     KK = 300;
                   KKp = 0;
                                KKpp = 0
620
             300; Kp = 0;
     MAPK =
                                Kpp = 0
621
                   KKph = 1; KKKph = 10
     Kph = 1;
622
   $Frozen
623
    S
624
   $Assignments
625
    S=0. if t<1000 else (1.0 if t<4000 else 0.0)
626
   $Rates
627
    a0 = 1; d0 = 1
628
    a1 = 1; d1 = 7.5; k1 = 2.5
629
    a3 = 1; d3 = 10;
                        k3 = 0.025
630
    a4 = 1; d4 = 1;
                        k4 = 1
631
    a5 = 1; d5 = 1;
                        k5 = 1
632
    a6 = 1; d6 = 1;
                        k6 = 1
633
    a7 = 1; d7 = 1
634
635
```

Listing 3: Goldbeter's Minimal cascade model for a mitotic oscillator.[19] This file is included in the distribution as sample model Gold1; an alternative version called Goldbeter does not use functions.

```
636
                $REACTIONS
637
                  [C <-> Nil, rates[kd, vi]]
638
                  [C \mid --> Nil, mod[X], Hill[vd, 1, Kd, 0,1]]
639
                  [M \mid --> Nil, mod[Nil], Hill[v2, 1, K2, 0, 1]]
640
                  [X \mid --> Nil, mod[Nil], Hill[v4, 1, K4, 0, 1]]
641
                 [Nil \rightarrow X, "vm3*f(M,X)"]
642
                 [C \mid -> M, Hill["vm1*g(M)", 1, Kc, 0, 1]]
643
                $IC
644
                 C = 0.1
645
                 M = 0.2
646
                 X = 0.3
647
                $FUNCTIONS
648
                 f(m, x) = m * (1-x) / (K3+1-x)
                 g(m) = (1-m)/(K1+1-m)
650
                $RATES
651
                 vd = 0.1;
                                vi = 0.023; v2 = 0.167;
652
                                             kd = 0.00333; K1 = 0.1
                 vm1 = 0.5;
                                vm3 = 0.2;
653
                 K2 = 0.1;
                                K3 = 0.1;
                                              K4 = 0.1;
                                                              Kc = 0.3
654
                 Kd = 0.02
655
656
```

Listing 4: Auto-generated Python code for the model shown in listing 1.

```
657
       import numpy as np
658
    2
       from scipy.integrate import odeint
659
    3
660
    4
       from math import *
661
       def ode_function_rhs(y,t):
662
    6
663
    7
         # this odeint(..) compatible function was
664
         # automatically generated by Cellerator 2016-07-31 12:27:17
665
    9
         # 2.7.6 (default, Jun 22 2015, 17:58:13) [GCC 4.8.2]
666
   10
         # linux2
667
   11
668
   12
669
   13
         # Model:
670
   14
   15
         \# [S \Rightarrow P, mod[E], rates[a,d,k]]
672
   16
         673
   17
         # rate constants
674
         a = 1.0
   18
675
   19
         d = 1.0
676
   20
         k = 1.0
677
   21
       # pick up values from previous iteration
   22
        E = \max(0, y[0])
679
   23
         S = \max(0, y[1])
680
   24
         S_E = max(0, y[2])
681
   25
         P = \max(0, y[3])
682
   26
       # calculate derivatives of all variables
683
   27
         yp=[0 for i in range(4)]
684
   28
         yp[0] = -E*S*a + S_E*d + S_E*k
685
   29
         yp[1] = -E*S*a + S_E*d
686
   30
         yp[2] = E*S*a - S_E*d - S_E*k
687
         yp[3] = S_E * k
   31
688
   32
         return yp
689
   33
690
   34
       def thesolver():
691
   35
           filename ="/home/mathman/Desktop/pycellerator/basicmodel.model"
692
   36
           variables=['E', 'S', 'S_E', 'P']
693
   37
           runtime = 15
694
   38
           stepsize = 0.1
695
   39
           times = np.arange(0, runtime+stepsize, stepsize)
696
   40
           y0 = [1.0, 1.0, 0.0, 0.0]
697
           sol = odeint(ode_function_rhs, y0, times, mxstep=50000)
   41
698
   42
           return sol
699
   43
700
   44
       if __name__=="__main__":
701
   45
           thesolver()
703
```

Listing 5: Simple SIRS disease model described in equations (61).

```
704
    $REACTIONS
705
      [S + I -> I + I, k]
706
      [R \rightarrow Nil, d]; [I \rightarrow Nil, d]; [S \rightarrow Nil, d]
707
      [R \rightarrow R+S, b]; [I \rightarrow I + S, b]; [S \rightarrow S+S, b]
708
      [I \rightarrow R, 1]
709
     [R \rightarrow S, f]
710
    $IC
711
     S = 0.99999999
712
     I = 1.0E-7
713
     R = 0
714
    $RATES
715
     k = 5.0
716
     d = 0.0005
717
     b = 0.0005
     f = 0.05
719
720
```

## 721 References

- 722 [1] Shapiro, BE, Mjolsess E (2016)Pycellerator: an arrow-based reaction-like modelling language
- for biological simulations. Bioinformatics.32(4):629-31. doi: 10.1093/bioinformatics/btv596.
- <sup>724</sup> [2] Shapiro, BE, Levchenko, A, Meyerowitz, EM, Wold, BJ, Mjolsness, ED (2003) Cellerator:
- extending a computer algebra system to include biochemical arrows for signal transduction
- simulations. Bioinformatics 19:677-678, doi: 10.1093/bioinformatics/btg042.
- [3] Hucka, M, Finney, A, Sauro, HM, et al. (2003) The systems biology markup language (SBML):
- a medium for representation and exchange of biochemical network models. Bioinformatics,
- 729 19:513-523. doi: 10.1093/bioinformatics/btg015
- 730 [4] Jones E, Oliphant E, Peterson P, et al. (2001) SciPy: Open Source Scientific Tools for Python.
- http://www.scipy.org/ (Online; accessed 2016-08-01).
- 732 [5] van der Walt, S, Colbert, SC, Varoquaux g. (2011) The NumPy Array: A Structure for
- 733 Efficient Numerical Computation Computing in Science & Engineering. 13:22-30.
- doi:10.1109/MCSE.2011.37
- 735 [6] Waage, P, Guldherg, CM (1864) Forhandlinger i Videnskabs Selskabet i Christiania, 37.
- 736 [7] Field, RJ, Noyes, RM (1974) Oscillations in chemical systems. IV. Limit cycle behavior in a
- model of a real chemical reaction. J. Chem. Phys. 60: 1877-1884. doi:10.1064/1.1681288
- 738 [8] Henri, V (1903) Lois Générales de l'action des Distases. Hermann, Paris.
- 739 [9] Michaelis, L, and Menten, M L (1913) Die Kinetik der Invertinwirkung, Biochemische
- 740 Zeitschrift 49:333-369.
- <sup>741</sup> [10] Briggs, GE, Haldane, JBS (1925) A Note on the Kinetics of Enzyme Action. Biochemical
- Journal 19(2):338-339. doi: 10.1042/bj0190338
- [11] Hill, AV (1910) The possible effects of the aggregation of the molecules of haemoglobin on its
- dissociation curve. Proc. Phsyiol. Soc. 40 (suppl): 4-7. doi: 10.1113/jphysiol.1910.sp001386

- <sup>745</sup> [12] Mjolsness, E, Sharp, DH, Reinitz, J (1991) A Connectionist Model of Development. J. theor.
- 746 Biol. 152: 429-453. doi:10.1016/S0022-5193(05)80391-1
- 747 [13] Murphy, KP (2012) Machine Learning, A Probabilistic Perspective. MIT Press, Cambridge.
- 748 [14] Monod, J, Wyman, J, Changeux, JP (1965) On the Nature Of Allosteric Transitions: A
- 749 Plausible Model. J. Mol. Biol. 12:88-118. doi:10.1016/S0022-2836(65)80285-6
- 750 [15] Najdi, T, Yang, C-R, Shapiro, BE, Hatfield, W, Mjolsness, E(2006) Application of a
- 751 Generalized MWC Model for the Mathematical Simulation of Metabolic Pathways Regulated by
- Allosteric Enzymes. J Bioinform Comput Biol 4(2):335-55. doi:10.1142/S0219720006001862
- <sup>753</sup> [16] Mjolsness, E. (2000) Trainable gene regulation networks with applications to Drosophila
- pattern formation. In: Bower, JM, Bolouri, H (ed) Computational Models of Genetic and
- <sup>755</sup> Biochemical Networks, MIT Press, Cambridge.
- 756 [17] Shapiro BE, Mjolsness ED (2001) Developmental Simulations with Cellerator. Paper
- presented at Second International Conference on Systems Biology, Pasadena, 4-7 Nov. 2001
- 758 [18] Mitchell, S, O'Sulivan, M, Dunning, I (2011) PuLP: A Linear Programming Toolkit for
- Python. http://www.optimization-online.org/DB\_FILE/2011/09/3178.pdf.
- 760 Accessed 31 July 2016.
- 761 [19] Goldbeter, A (991) A minimal cascade model for the mitotic oscillator involving cyclin and
- cdc2 kinase. Proc. Natl. Acad. Sci. USA 88:9107-1101 (1991). doi 10.1073/pnas.88.20.9107
- 763 [20] Huang, CY, Ferrell, JE Jr (1996) Ultrasensitivity in the mitogen-activated protein kinase
- cascade. Proc Natl Acad Sci U S A. 93(19):10078-83.
- 765 [21] Kholodenko BN (2000) Negative feedback and ultrasensitivity can bring about oscillations in
- the mitogen-activated protein kinase cascades. Eur J Biochem. 267(6):1583-8 doi
- 767 10.1046/j.1432-1327.2000.01197.x
- <sup>768</sup> [22] Kermack, WO, McKendrick, AG (1927) A Contribution to the Mathematical Theory of
- 769 Epidemics. Proc. Royal Soc. A. 115(772):700-721. doi 10.1098/rspa.1927.0118

Figure 1: Time course of simulation of basic model shown in listing 1.

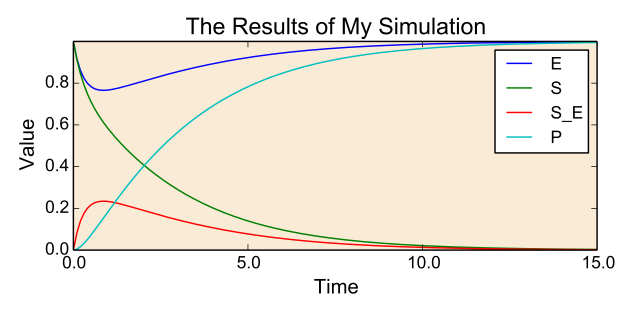


Figure 2: Parametric scan of the SIRS model (listing 5).

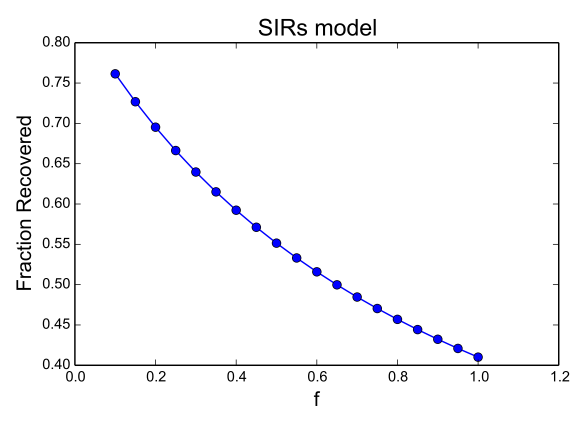


Figure 3: Oscillations in MAPK cascade with feedback and square wave stimulation. The model is shown in listing 2.

