Killer Yeast Vs. Sensitive Yeast

Evan Cummings Intizor Aliyorov Malachi J. Cryder

MATH 445 - Statistical, Dynamical, and Computational Modeling

December 10, 2013

Proposal

The differential equation we may use for modeling the growth of yeast is the same as that used for bacterial growth in a chemostat:

$$\begin{split} \frac{dN}{dt} &= k(C)N - \frac{FN}{V}, \\ \frac{dC}{dt} &= -\alpha k(C)N - \frac{FC}{V} + \frac{FC_0}{V}, \end{split}$$

with initial conditions $C(0) = C_i$ and $N(0) = N_i$, N is the unitless optical density of yeast in the chamber, C is the unitless optical density of nutrient in the chamber, C_0 is the unitless optical density of nutrient in the reservoir, F is the in/out volume flow rate with units volume/time, V is the volume of the chamber, α is a unitless inverse of the yield constant, and k(C) is the reproduction rate for yeast in units 1/time with possible formula chosen such that $\lim_{C\to\infty} = k_{max}$, and k_{max} represents the maximum possible reproduction rate:

$$k(C) = \frac{k_{max}C}{C_n + C}.$$

where C_n is chosen such that $k(C_n) = k_{max}/2$. Because the concentration in the tank C(t) is related to the concentration in the reservoir by $C(t) \leq C_0$, C_0 may be chosen sufficiently small such that

$$k(C) = \frac{k_{max}C}{C_n + C} \approx \frac{k_{max}C}{C_n} = KC,$$

where K has units 1/time. The equations we need to solve then become

$$\frac{dN}{dt} = KCN - \frac{FN}{V},\tag{1}$$

$$\frac{dC}{dt} = -\alpha KCN - \frac{FC}{V} + \frac{FC_0}{V}. (2)$$

The quantitative measurement for fitness is a unitless measurement of optical density at steady state (N) at a given flow rate (F) in volumes/hr. In order to find the steady states, we have to

find the intersections of the null-clines at equilibrium points (\bar{N}, \bar{C}) , i.e. $dN(\bar{N}, \bar{C})/dt = 0$ and $dC(\bar{N}, \bar{C})/dt = 0$:

$$\frac{dN(\bar{N},\bar{C})}{dt} = K\bar{C}\bar{N} - \frac{F\bar{N}}{V},$$
$$= \bar{N}\left(K\bar{C} - \frac{F}{V}\right) = 0,$$

which is zero for $\bar{N}=0$ or $K\bar{C}=F/V$. Solving the other equation gives us the other steady-states:

$$\frac{dC(\bar{N},\bar{C})}{dt} = -\alpha K\bar{C}\bar{N} - \frac{F\bar{C}}{V} + \frac{FC_0}{V} = 0,$$

which is zero for $\alpha K \bar{C} \bar{N} + \frac{F\bar{C}}{V} = \frac{FC_0}{V}$.

In order to evaluate these null-clines, we need to evaluate the non-trivial cases, here for $\dot{N}=0$,

$$K\bar{C} = \frac{F}{V}$$

$$\implies \bar{C} = \frac{F}{VK}.$$
(3)

Likewise, for $\dot{C} = 0$,

$$\alpha K \bar{C} \bar{N} + \frac{F \bar{C}}{V} = \frac{F C_0}{V}$$

$$\implies \bar{N} = \frac{F C_0}{V \alpha K \bar{C}} - \frac{F}{V \alpha K} = \frac{F}{V \alpha K} \left(\frac{C_0}{\bar{C}} - 1\right). \tag{4}$$

This intersects the $\bar{N}=0$ nullcline at $\frac{F}{V\alpha K}=0$ or $\frac{C_0}{\bar{C}}=1$. However, because F is never 0, we can disregard the first equation, and we know that the only trivial steady-state is located at $\bar{N}=0$, $\bar{C}=C_0$.

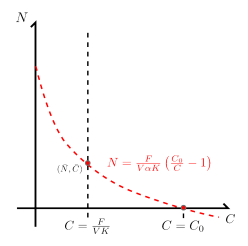


Figure 1: The $\dot{C}=0$ nullcline (dashed red) intersecting with the $\dot{N}=0$ nullcline (dashed black). The trivial and non-trivial steady-states $(0,C_0)$ and (\bar{N},\bar{C}) are shown as red dots.

By placing Eq. (3) inside Eq. (4), we can find the non-trivial steady-state, the intersection of null-clines:

$$\bar{N}(\bar{C}) = \frac{FC_0}{V\alpha K\bar{C}} - \frac{F}{V\alpha K}$$

$$= \frac{FC_0}{V\alpha K\frac{F}{VK}} - \frac{F}{V\alpha K}$$

$$= \frac{C_0}{\alpha} - \frac{F}{V\alpha K} = \frac{1}{\alpha} \left(C_0 - \frac{F}{VK} \right). \quad (5)$$

The unknown parameters in Eq (5) are α and K. We can find these parameters by fitting Eq (5) to the data by non-linear least squares fitting \bar{N}_i at F_i for i = 1, ..., n, where n is the number of observations.

After we obtain estimates of these parameters for both the killer yeast L and sensitive yeast S, (α_L , α_S and K_L , K_S respectively), we can model a "what if" scenario whereby we place both species of yeast, sensitive and killer, into one chemostat. The population of sensitive yeast S will be negatively impacted by the amount of toxin the killer yeast K can produce, so we add the term $-\beta KL$ to the differential equation describing population S. The differential equations we use to solve this three-species model is

$$\frac{dL}{dt} = K_L C L - \frac{FL}{V},\tag{6}$$

$$\frac{dS}{dt} = K_S C S - \frac{\dot{F}S}{V} - \beta S L,\tag{7}$$

$$\frac{dC}{dt} = -\alpha_L K_L C L - \alpha_S K_S C S - \frac{FC}{V} + \frac{FC_0}{V}.$$
 (8)

The data we are provided with include two sets of two separate runs, along with the concentration of nutrient in the reservoir, $C_0 = 0.02$:

1 K1 Run

Vessel One:

| Volumes/Hr | 0.028 | 0.099 | 0.142 | 0.207 | 0.269 | 0.287 | 0.352 | 0.403 |
|---------------------------------|-------|-------|-------|-------|-------|-------|-------|-------|
| Optical Density at Steady State | 0.144 | 0.151 | 0.099 | 0.069 | 0.045 | 0.02 | 0.003 | 0 |

Vessel Two:

| Volumes/Hr | 0.054 | 0.11 | 0.141 | 0.199 | 0.257 | 0.296 | 0.348 | 0.397 | 0.41 |
|---------------------------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| Optical Density at Steady State | 0.164 | 0.151 | 0.11 | 0.092 | 0.072 | 0.023 | 0.006 | 0.002 | 0.004 |

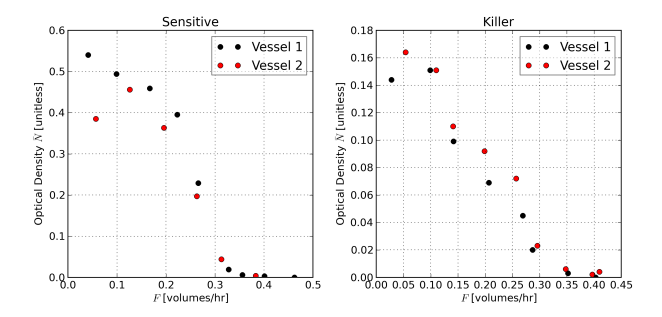
2 Sensitive Run

Vessel One:

| Volumes/Hr | 0.041 | 0.099 | 0.167 | 0.223 | 0.266 | 0.328 | 0.356 | 0.401 | 0.462 |
|---------------------------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| Optical Density at Steady State | 0.54 | 0.494 | 0.459 | 0.395 | 0.229 | 0.019 | 0.006 | 0.003 | 0 |

Vessel Two:

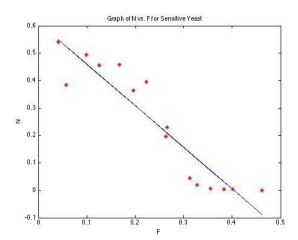
| Volumes/Hr | 0.0571 | 0.126 | 0.196 | 0.263 | 0.313 | 0.383 |
|---------------------------------|--------|-------|-------|-------|-------|-------|
| Optical Density at Steady State | 0.385 | 0.456 | 0.363 | 0.197 | 0.044 | 0.004 |



The data provided does not include the volume of the chamber, V. In order to solve Eq. (5), this quantity is needed. Here we have dimensional analysis of the problem:

$$\begin{split} \frac{dN}{dt} &= KCN - \frac{FN}{V} \\ &\equiv \left[\frac{1}{\text{time}}\right] \equiv \left[\frac{1}{\text{time}} - \frac{\text{volume}}{\text{time}} \cdot \frac{1}{\text{volume}}\right] \equiv \left[\frac{1}{\text{time}}\right], \\ \frac{dC}{dt} &= -\alpha KCN - \frac{FC}{V} + \frac{FC_0}{V} \\ &\equiv \left[\frac{1}{\text{time}}\right] \equiv \left[-\frac{1}{\text{time}} - \frac{\text{volume}}{\text{time}} \cdot \frac{1}{\text{volume}} + \frac{\text{volume}}{\text{time}} \cdot \frac{1}{\text{volume}}\right] \equiv \left[\frac{1}{\text{time}}\right]. \end{split}$$

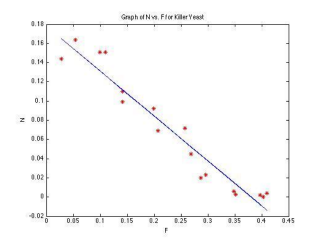
Taking V = 1, we can solve Eq. (5) using MatLab's nlinfit function. The results of this are shown below for both sensitive and killer yeast.



| | Estimates | SE | CI |
|-----------------------|-----------|--------|--------------------|
| $\overline{\alpha_S}$ | 0.0325 | 0.0024 | (0.0255, 0.0399) |
| K_S | 20.1818 | 1.0802 | (16.9281, 23.4356) |

The results on the left show that there may be a better fit to the data than Eq. (5). Notice that elimination of the last few data points corresponding to high F may be removed; this would reduce the standard error and hence provide a better estimate for α_L and K_L .

Figure 2: Sensitive yeast S steady-state best-fit line using Eq. (5).



| | Estimates | SE | CI |
|------------|-----------|--------|-------------------|
| α_L | 0.1124 | 0.0053 | (0.0969, 0.1279) |
| K_L | 19.0288 | 0.6367 | (17.1526, 20.905) |

The best-fit line on the left shows that steadystate data follows a fairly linear relationship with flow, and as such we can be confident our estimates for α_L and K_L are correct.

Figure 3: Killer yeast L steady-state best-fit line using Eq. (5).

Now that we have estimates for all values of α and K (for both killer and sensitive yeast) we can run the dynamic model (6), (7), and (8) to equilibrium. By letting β range from 0 to 0.5 (in units 1/time), and F range from 0 to 0.5 (in units volume/time), we can run the model for each β and F to determine the regions in the β , F plane where sensitive yeast S overtakes the killer yeast L and vice versa.

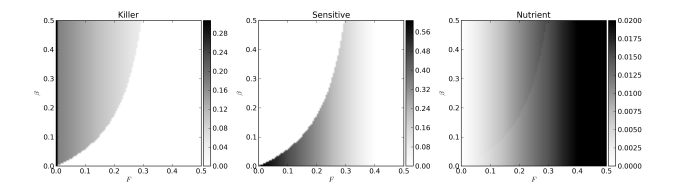


Figure 4: 250×250 steady-state optical density solution for killer yeast (left), sensitive yeast (middle) and nutrient (right) for a total run time of 40,000 hours. Equations (6), (7), and (8) were solved with the Dormand-Prince numerical integration algorithm with an absolute tolerance of 1e-6, relative tolerance of 1e-6, and timestep Δt of 500 hours. The timestep was kept high due to the $250 \times 250 = 62,500$ simulations required to complete the figure. Parallel processing was also implemented to speed up the simulation.

Notice in Figure 4 that as β increases the killer yeast dominates and sensitive yeast eventually dies off, while as F increases the sensitive yeast dominates and the killer yeast eventually dies out. The sensitive yeast are flushed from the container at around F = 0.4 volumes/hr.

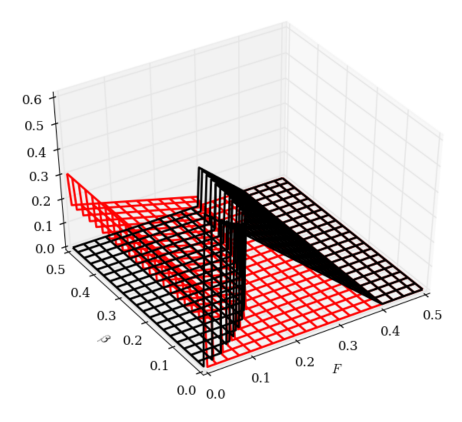


Figure 5: 3D solution depicting killer yeast (red) and sensitive yeast (black) steady-states.

Steady-State regions

In order to evaluate the steady-state solutions, we need to equate Equations (6), (7), and (8) to zero and solve:

$$\frac{d}{dt}L(\bar{L},\bar{S},\bar{C}) = K_L\bar{C}\bar{L} - \frac{F\bar{L}}{V} = 0,$$
(9)

$$\frac{d}{dt}S(\bar{L},\bar{S},\bar{C}) = K_S\bar{C}\bar{S} - \frac{F\bar{S}}{V} - \beta\bar{S}\bar{L} = 0, \tag{10}$$

$$\frac{d}{dt}C(\bar{L},\bar{S},\bar{C}) = -\alpha_L K_L \bar{C}\bar{L} - \alpha_S K_S \bar{C}\bar{S} - \frac{F\bar{C}}{V} + \frac{FC_0}{V} = 0. \tag{11}$$

From Eq. (9), we see that $\bar{L}(K_L\bar{C} - F/V) = 0$, and thus $\bar{L} = 0$ or $\bar{C} = F/(K_LV)$. For each of these steady-states, we will evaluate Equations (10) and (11) to find the qualitative behavior of the steady-states.

1
$$\bar{L} = 0$$
:

From (10),
$$\bar{S}(K_S\bar{C} - F/V - \beta\bar{L}) = 0$$
, so $\bar{S} = 0$ or $\bar{C} = \frac{F/V + \beta\bar{L}}{K_S} = \frac{F}{K_SV}$.

1. $\bar{S} = 0$:

From (11),

$$-\alpha_L K_L \bar{C}\bar{L} - \alpha_S K_S \bar{C}\bar{S} - \frac{F\bar{C}}{V} + \frac{FC_0}{V} = 0$$
$$-\frac{F\bar{C}}{V} + \frac{FC_0}{V} = 0, \quad F/V \neq 0.$$

Thus $\bar{C} = C_0$, and the first steady-state is $(0, 0, C_0)$, where both species of yeast are flushed out. Call this Steady-State I.

 $2. \ \bar{C} = \frac{F}{K_S V}$:

From (11).

$$\begin{split} -\alpha_S K_S \left(\frac{F}{K_S V}\right) \bar{S} - \frac{F}{V} \left(\frac{F}{K_S V}\right) + \frac{FC_0}{V} &= 0 \\ -\alpha_S \left(\frac{F}{V}\right) \bar{S} - \frac{F}{V} \left(\frac{F}{K_S V}\right) + \frac{FC_0}{V} &= 0 \\ \alpha_S \left(\frac{F}{V}\right) \bar{S} &= \frac{F}{V} \left(\frac{F}{K_S V}\right) - \frac{FC_0}{V} \\ \alpha_S \bar{S} &= \frac{F}{K_S V} - C_0 \\ \bar{S} &= \frac{F}{K_S V \alpha_S} - \frac{C_0}{\alpha_S}. \end{split}$$

Therefore, the second steady-state is $\left(0, \frac{F}{K_S V \alpha_S} - \frac{C_0}{\alpha_S}, \frac{F}{K_S V}\right)$, where only the sensitive species of yeast survives. Call this Steady-State II.

2 $\bar{C} = \frac{F}{K_L V}$:

From (10),

$$\bar{S}\left(K_S\bar{C} - \frac{F}{V} - \beta\bar{L}\right) = 0$$
$$\bar{S}\left(K_S\frac{F}{K_LV} - \frac{F}{V} - \beta\bar{L}\right) = 0.$$

So $\bar{S} = 0$ or $\bar{L} = \frac{K_S F}{K_L V \beta} - \frac{F}{V \beta}$.

1. $\bar{S} = 0$:

From (11),

$$-\alpha_L K_L \bar{C} \bar{L} - \alpha_S K_S \bar{C} \bar{S} - \frac{FC}{V} + \frac{FC_0}{V} = 0$$

$$-\alpha_L \left(\frac{F}{K_L V}\right) \bar{L} - \frac{F}{V} \left(\frac{F}{K_L V}\right) + \frac{FC_0}{V} = 0$$

$$\alpha_L \left(\frac{F}{V}\right) \bar{L} = \frac{FC_0}{V} - \frac{F}{V} \left(\frac{F}{K_L V}\right)$$

$$\alpha_L \bar{L} = C_0 - \frac{F}{K_L V}$$

$$\bar{L} = \frac{C_0}{\alpha_L} - \frac{F}{K_L V \alpha_L}.$$

Thus third steady-state is $\left(\frac{C_0}{\alpha_L} - \frac{F}{K_L V \alpha_L}, 0, \frac{F}{K_L V}\right)$, where only the killer species of yeast survive. Call this Steady-State III.

2. $\bar{L} = \frac{K_S F}{K_L V \beta} - \frac{F}{V \beta}$: From (11),

$$-\alpha_L K_L \bar{C} \bar{L} - \alpha_S K_S \bar{C} \bar{S} - \frac{F \bar{C}}{V} + \frac{F C_0}{V} = 0$$
$$-\alpha_L K_L \left(\frac{F}{K_L V}\right) \left(\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta}\right) - \alpha_S K_S \left(\frac{F}{K_L V}\right) \bar{S} - \frac{F}{V} \left(\frac{F}{K_L V}\right) + \frac{F C_0}{V} = 0.$$

Solving for \bar{S} ,

$$\Rightarrow \alpha_S K_S \left(\frac{F}{K_L V} \right) \bar{S} = \frac{FC_0}{V} - \alpha_L K_L \left(\frac{F}{K_L V} \right) \left(\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} \right) - \frac{F}{V} \left(\frac{F}{K_L V} \right)$$

$$\alpha_S K_S \left(\frac{1}{K_L} \right) \bar{S} = C_0 - \alpha_L K_L \left(\frac{1}{K_L} \right) \left(\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} \right) - \frac{F}{K_L V}$$

$$\frac{\alpha_S K_S}{K_L} \bar{S} = C_0 - \alpha_L \left(\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} \right) - \frac{F}{K_L V}$$

$$\bar{S} = \left(\frac{K_L}{\alpha_S K_S} \right) C_0 - \alpha_L \left(\frac{K_L}{\alpha_S K_S} \right) \left(\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} \right) - \left(\frac{K_L}{\alpha_S K_S} \right) \left(\frac{F}{K_L V} \right)$$

$$\bar{S} = \frac{K_L C_0}{\alpha_S K_S} - \frac{\alpha_L F}{\alpha_S V \beta} + \frac{\alpha_L K_L F}{\alpha_S K_S V \beta} - \frac{F}{\alpha_S K_S V}$$

$$\bar{S} = \frac{K_L C_0}{\alpha_S K_S} + \frac{F}{V} \left(\frac{\alpha_L K_L}{\alpha_S K_S \beta} - \frac{\alpha_L}{\alpha_S \beta} - \frac{1}{\alpha_S K_S} \right).$$

Therefore, the fourth and final steady-state is $\left(\frac{K_SF}{K_LV\beta} - \frac{F}{V\beta}, \frac{K_LC_0}{\alpha_SK_S} + \frac{F}{V}\left(\frac{\alpha_LK_L}{\alpha_SK_S\beta} - \frac{\alpha_L}{\alpha_S\beta} - \frac{1}{\alpha_SK_S}\right), \frac{F}{K_LV}\right)$, where both species survive. Call this Steady-State IV.

Non-Trivial Steady State Evaluation

Looking at Steady-State IV:

$$\begin{split} \bar{L} &= \frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} \\ \bar{S} &= \frac{K_L C_0}{\alpha_S K_S} + \frac{F}{V} \left(\frac{\alpha_L K_L}{\alpha_S K_S \beta} - \frac{\alpha_L}{\alpha_S \beta} - \frac{1}{\alpha_S K_S} \right) \\ \bar{C} &= \frac{F}{K_L V}. \end{split}$$

First, assume that both \bar{L} and \bar{S} are greater than zero. Then

$$\frac{K_S F}{K_L V \beta} - \frac{F}{V \beta} > 0$$

$$\frac{F}{V \beta} \left(\frac{K_S}{K_L} - 1 \right) > 0,$$

and thus because F>0 and $\beta>0,$ $\frac{K_S}{K_L}>1$ and hence $K_S>K_L.$ Next,

$$\begin{split} \frac{K_L C_0}{\alpha_S K_S} + \frac{F}{V} \left(\frac{\alpha_L K_L}{\alpha_S K_S \beta} - \frac{\alpha_L}{\alpha_S \beta} - \frac{1}{\alpha_S K_S} \right) &> 0 \\ \frac{F}{V} \left(\frac{\alpha_L K_L}{\alpha_S K_S \beta} - \frac{\alpha_L}{\alpha_S \beta} - \frac{1}{\alpha_S K_S} \right) &> -\frac{K_L C_0}{\alpha_S K_S} \\ \frac{\alpha_L K_L}{\alpha_S K_S \beta} - \frac{\alpha_L}{\alpha_S \beta} - \frac{1}{\alpha_S K_S} &> -\frac{V}{F} \cdot \frac{K_L C_0}{\alpha_S K_S} \\ \frac{\alpha_L K_L}{\beta} - \frac{\alpha_L K_S}{\beta} - 1 &> -\frac{V}{F} \cdot K_L C_0 \\ \frac{1}{\beta} (\alpha_L K_L - \alpha_L K_S) &> 1 - \frac{V}{F} \cdot K_L C_0 \\ \beta &< \frac{\alpha_L K_L - \alpha_L K_S}{1 - \frac{V}{F} \cdot K_L C_0}. \end{split}$$

A similar procedure also produces

$$\beta > \frac{\frac{F}{V}\left(\alpha_L K_S - \alpha_L K_L\right)}{C_0 K_L - \frac{F}{V}}.$$

Because $K_s > K_L$, we know that the numerator is negative. Also, because $\beta > 0$, we know that the numerator must also be negative, i.e.,

$$\frac{V}{F} \cdot K_L C_0 < 1$$

$$K_L C_0 < \frac{F}{V}.$$

Source Code:

```
from scipy.integrate._ode    import ode
from scipy.io
                           import savemat
from pylab
                            import *
from mpl_toolkits.mplot3d
                            import Axes3D
from multiprocessing
                           import Queue, cpu_count, Process
from mpl_toolkits.axes_grid1 import make_axes_locatable
#-----
# ODE function to be integrated
def dLdt(L, S, C, params):
 INPUT:
   L - population of killer yeast.
   {\bf S} - population of sensitive yeast.
   C - population of nutrient.
   params - dLdt equation parameters.
  OUTPUT:
  dLdt - time derivative of L.
  K_L = params[0]
 F = params[1]
       = params[2]
 dLdt = K_L * C * L - F * L / V
 return array(dLdt)
def dSdt(L, S, C, params):
 INPUT:
   L - population of killer yeast.
   {\tt S} - population of sensitive yeast.
   C - population of nutrient.
   params - dSdt equation parameters.
  OUTPUT:
  dSdt - time derivative of S.
 K_S
      = params[0]
 F = params[1]
V = params[2]
 beta = params[3]
  dSdt = K_S*C*S - F*S/V - beta*S*L
 return array(dSdt)
def dCdt(L, S, C, params):
 INPUT:
   L - population of killer yeast.
   S - population of sensitive yeast.
   C - population of nutrient.
   params - dCdt equation parameters.
  OUTPUT:
  dCdt - time derivative of C.
  a_L = params[0]
  a_S = params[1]
 K_L = params[2]
 K_S = params[3]
      = params[4]
      = params[5]
 C_0 = params[6]
 dCdt = -a_L*K_L*C*L - a_S*K_S*C*S - F*C/V + F*C_0/V
 return array(dCdt)
def f(t, y, dLdt, dSdt, dCdt, L_params, S_params, C_params):
INPUT:
```

```
t - time array
           functionfunction
   dLdt
   uSdt
           - function
   dCdt.
   L_params - parameters for dLdt
   S_params - parameters for dSdt
   C_params - parameters for dCdt
 OUTPUT:
   ydot[0] = time derivative of y[0],
   ydot[1] = time derivative of y[1],
   ydot[2] = time derivative of y[2].
 L = y[0]
 S = y[1]
 C = y[2]
                                 # right hand side 1st eqn
 rhs1 = dLdt(L, S, C, L_params)
 rhs2 = dSdt(L, S, C, S_params)
rhs3 = dCdt(L, S, C, C_params)
                                 # right hand side 2nd eqn
# right hand side 3rd eqn
 return array([rhs1, rhs2, rhs3])
def model(F, beta, y0, ta, dt):
 Run model for given volume flow rate <F> and toxin coef <beta> for total
 time array <ta> in hours at timestep <dt>, also in hours. Returns the
 last solution 3-tuple for L, S, and C.
 # Additional parameters being passed to the ODE function
 a_L = 0.1124
 a_S = 0.0325
 K_L = 19.0288
 K_S = 20.1818
 V
      = 1.0
 C_0 = 0.02
 L_params = [K_L, F, V]
 S_{params} = [K_S, F, V, beta]
 C_{params} = [a_L, a_S, K_L, K_S, F, V, C_0]
 # Call function that integrates the ODE:
 r = ode(f)
 r.set_integrator('dopri5', atol=1e-6, rtol=1e-5)
 r.set_initial_value(y0, ta)
 r.set_f_params(dLdt, dSdt, dCdt, L_params, S_params, C_params)
 sol = []
 sol.append(y0)
 for t in ta[:-1]:
  r.integrate(r.t + dt)
   sol.append(r.y)
 sol = array(sol).T
 return sol[:,-1]
class solveProcess(Process):
 Process to solve the model function.
 def __init__(self, i, queue, beta_a, F_a, y0, ta, dt, p):
   Initialize the Process with ID <i>, processing queue <queue>, beta array
   parameters .
   Process.__init__(self)
   self.i = i
self.q = queue
  self.beta_a = beta_a
```

```
self.F_a = F_a
            = y0
= ta
   self.y0
   self.ta
              = dt
   self.dt
   self.m
              = len(beta_a)
   self.n
              = len(F_a)
   self.p
               = p
 def run(self):
   solve the differential equations for all beta_a and F_a.
   p = self.p
   m = self.m
   n = self.n
   SS_L = zeros((m,n))
   SS_S = zeros((m,n))
   SS_C = zeros((m,n))
   for i, beta in enumerate(self.beta_a):
     for j, F in enumerate(self.F_a):
       print 'Process %i solving: beta=%f, F=%f' % (self.i, beta, F)
       sol = model(F=F, beta=beta, y0=self.y0, ta=self.ta, dt=self.dt)
       SS_L[i,j] = sol[0]
       SS_S[i,j] = sol[1]
       SS_C[i,j] = sol[2]
   self.q.put(array([SS_L, SS_S, SS_C])) # add the result to the queue.
def plot_sol(ax, f, extent, tit, cmap='Greys'):
  plot the 2D solution <f> to axes <ax>.
        = ax.imshow(f[::-1,:], extent=extent, cmap=cmap)
 divider = make_axes_locatable(ax)
        = divider.append_axes("right", size="5%", pad=0.05)
 ax.set_title(tit)
 ax.set_ylabel(r'$\beta$')
 ax.set_xlabel(r'$F$')
 colorbar(im, cax=cax)
# parameters :
m = 100
          # number of beta discretizations.
n = 100
            # number of F discretizations.
p = 3
            # number of parameters
t0 = 0.0
            # initial time
tf = 40000  # final time
dt = 500
            # time step
# Initial conditions
y0 = [0.3, 0.3, 0.001]
# range of beta, flow, and time to model :
betaMin = 0.0
betaMax = 0.5
Fmin = 0.0
     = 0.5
Fmax
beta_a = linspace(betaMin, betaMax, m)
       = linspace(Fmin,
                           Fmax,
F_a
      = arange(t0, tf+dt, dt)
ta
# multiprocessing data structures :
solvers = []
queue = []
numCpus = cpu_count()
Fs = array_split(F_a, numCpus)
```

```
# create a solver for each processor and begin solving each :
for i in range(numCpus):
  q = Queue()
  queue.append(q)
  solver = solveProcess(i, q, beta_a, Fs[i], y0, ta, dt, p)
  solvers.append(solver)
  solver.start()
# wait until solver (started above) finishes :
for s in solvers:
 s.join()
# retrive the results :
sols = []
for q in queue:
 while q.empty() == False:
    sols.append(q.get())
# put the results from the individual cores back together :
for i, s in enumerate(sols):
  if i == 0:
    L_sol = s[0]
    S_sol = s[1]
    C_{sol} = s[2]
  else:
    L_sol = hstack((L_sol, s[0]))
    S_sol = hstack((S_sol, s[1]))
    C_{sol} = hstack((C_{sol}, s[2]))
Flow, Beta = meshgrid(F_a, beta_a)
data = {'Beta' : Beta,
        'Flow' : Flow,
        'L_sol' : L_sol,
        'S_sol' : S_sol,
        'C_sol' : C_sol}
savemat('.../../killer_yeast/data/results.mat', data)
# plot the results :
fig = plt.figure()
ax = fig.add_subplot(111, projection='3d')
{\tt ax.plot\_wireframe(Flow,\ Beta,\ L\_sol,\ color='r',\ lw=2.0,\ rstride=5,\ cstride=5)}
ax.plot_wireframe(Flow, Beta, S_sol, color='k', lw=2.0, rstride=5, cstride=5)
ax.set_ylabel(r'$\beta$')
ax.set_xlabel(r'$F$')
show()
fig = plt.figure(figsize=(15,5))
ax1 = fig.add_subplot(131)
ax2 = fig.add_subplot(132)
ax3 = fig.add_subplot(133)
extent = [Fmin, Fmax, betaMin, betaMax]
plot_sol(ax1, L_sol, extent, r'Killer',
                                            cmap='Greys')
plot_sol(ax2, S_sol, extent, r'Sensitive', cmap='Greys')
plot_sol(ax3, C_sol, extent, r'Nutrient', cmap='Greys')
tight_layout()
savefig('.../../killer_yeast/doc/images/sols.png', dpi=300)
show()
```