HIV-mini-project

October 18, 2019

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In [1]: # Configure Jupyter so figures appear in the notebook
      %matplotlib inline
      # Configure Jupyter to display the assigned value after an assignment
      %config InteractiveShell.ast node interactivity='last expr or assign'
      # import functions from the modsim.py module
      from modsim import *
In [2]: def make system(alpha, beta, gamma, delta, mu, pi, rho, sigma, tau):
         """Make a system object for the HIV model.
         alpha: rate of activation of latently infected cells [E/(Lt)]
         beta: rate of infection of uninfected cells per viron [1/(Vt)]
         gamma: rate of production of uninfected cells [R/t]
         delta: rate of removal of actively infected cells [1/t]
         mu: rate of cell death [1/t]
         pi: rate of production of virons by actively infected cells [V/(Et)]
         rho: proportion of cells that become latently infected upon infection [L/R]
         sigma: rate of removal of virons [1/t]
         tau: proportion of uninfected cells activated [1]
         returns: System object
         init = State(R=200, L=0, E=0, V=4 * 10**-7)
         t \ 0 = 0
         t end = 120 \# days
         dt = 1.0/20.0 \# 1.2 \text{ hour increments}
         return System(init=init, t 0=t 0, t end=t end, dt=dt, alpha=alpha, beta=beta, gamma=gamma,
                   delta=delta, mu=mu, pi=pi, rho=rho, sigma=sigma, tau=tau)
In [3]: def update func(state, t, system):
         """Update the HIV model.
         state: State with variables R, L, E, V
         t: time step
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returns: State object
        r, l, e, v = state
         # infections/activations
         infected = system.beta * r * v
        latentlyInfected = system.rho * infected
        activelyInfected = infected - latentlyInfected
        latentlyInfectedActivated = system.alpha * l
         # productions
         uninfectedProduction = system.gamma * system.tau
         vironProduction = system.pi * e
         # deaths/removals
        uninfectedDeath = system.mu * r
        latentlyInfectedDeath = system.mu*l
        activelyInfectedDeath = system.delta * e
         vironDeath = system.sigma * v
        dt = system.dt
        r += (uninfectedProduction - infected - uninfectedDeath) *dt
        1 += (latentlyInfected - latentlyInfectedActivated - latentlyInfectedDeath) *dt
        e += (activelyInfected + latentlyInfectedActivated - activelyInfectedDeath) *dt
         v += (vironProduction - vironDeath) *dt
        return State(R=r, L=l, E=e, V=v)
In [4]: def run simulation(system, update func):
         """Runs a simulation of the system.
         Adds a TimeFrame to the System: results
        system: System object
        update func: function that updates state
        init = system.init
         t 0, t end, dt = system.t 0, system.t end, system.dt
        frame = TimeFrame(columns=init.index)
        frame.row[t 0] = init
         ts = linrange(t 0, t end, dt)
        for t in ts:
           frame.row[t+dt] = update func(frame.row[t], t, system)
        return frame
In [5]: def plot results(R, L, E, V):
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system: System with parameters alpha, beta, gamma, delta, mu, pi, rho, sigma, and tau

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xNums = [0,30,60,90,120]
         fig, (pl1, pl2) = plt.subplots(1, 2, figsize=(16, 8)) # create 2 subplots, right & left
         color = 'tab:green'
         pl1.set xlabel('Days from infection')
         pl1.set ylabel('CD4 lymphocytes', color=color)
         cells, = pl1.plot(R + E + L + 800, color=color)
         pl1.tick params(axis='y', labelcolor=color)
         pl1.set vlim([0, 1200])
         plt.xticks(xNums)
         ax1 = pl1.twinx() # instantiate a second axis on pl1 that shares the same x-axis
         color = 'tab:purple'
         ax1.set ylabel('Virons V', color=color) # we already handled the x-label with ax1
         v_1 = ax1.semilogy(V, color=color)
         ax1.tick params(axis='y', labelcolor=color)
         ax1.set ylim([0.1, 10000])
         color = 'tab:red'
         pl2.set xlabel('Days from infection')
         pl2.set ylabel('R', color=color)
         r_{,} = pl2.plot(R, color=color)
         pl2.tick params(axis='y', labelcolor=color)
         pl2.set vlim([0, 250])
         plt.xticks(xNums)
         ax2 = pl2.twinx() # instantiate a second axis on pl2 that shares the same x-axis
         color = 'tab:blue'
         ax2.set ylabel('L and E', color=color) # we already handled the x-label with ax1
         l_1 = ax2.semilogy(L, color=color)
         e, = ax2.semilogy(E, color=color, linestyle='dashed')
         ax2.tick params(axis='y', labelcolor=color)
         ax2.set ylim([0.1, 100])
         pl1.legend((cells, v), ('CD4 lymphocytes', 'Cell-free virus'), fontsize=12, loc='upper right')
         pl2.legend((r, l, e), ('R', 'L', 'E'), fontsize=12, loc='upper right')
         fig.tight layout() # otherwise the right y-label is slightly clipped
In [6]: alpha = 3.6 * 10**-2 # rate of activation of latently infected cells [E/(Lt)]
      beta = 0.00027 \, \# \, \text{rate} of infection of uninfected cells per viron [1/(\text{Vt})]
      gamma = 1.36 \# rate of production of uninfected cells [R/t]
      delta = 0.33 # rate of removal of actively infected cells [1/t]
      mu = 1.36 * 10**-3 \# rate of cell death [1/t]
      pi = 100 \# rate of production of virons by actively infected cells <math>[V/(Et)]
      rho = 0.1 \# proportion of cells that become latently infected upon infection [L/R]
      sigma = 2 \# rate of removal of virons [1/t]
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"""Plot the results of the HIV model

tau = 0.2 # proportion of uninfected cells activated [1]

system = make_system(alpha, beta, gamma, delta, mu, pi, rho, sigma, tau) results = run_simulation(system, update_func) plot_results(results.R, results.L, results.E, results.V)

Out[6]:

