Notes YEAST_COB_Batch constraint-based approach

Try using LP in each step

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In [3]: from optlang import Model, Variable, Constraint, Objective
In [4]: # Define culture constraint-based model
        def culture(G, E):
            # LP calculation of the optimal qGr, qEr based on G and E values
            # - parameters
            q02max = 6.9e-3; kog = 2.3; koe = 1.6; YGr = 3.5; YEr = 1.32;
            alpha = 0.01; beta = 1.0
            # - transfer data from dynamic reactor model to static LP model
            qGr_opt = Variable('qGr_opt', lb=0)
            qEr_opt = Variable('qEr_opt', 1b=0)
            # - LP model constraint and objective
            mu_max = Objective(YGr*qGr_opt + YEr*qEr_opt, direction='max')
            q02lim = Constraint(kog*qGr_opt + koe*qEr_opt, ub=q02max)
            qGlim = Constraint(qGr_opt, ub=alpha*max(0,G))
            qElim = Constraint(qEr_opt, ub=beta*max(0,E))
            # - put together the LP model
            yeast_model = Model(name='Yeast bottleneck model')
            yeast_model.objective = mu_max
            yeast_model.add(q02lim)
            yeast_model.add(qGlim)
            yeast_model.add(qElim)
```

```
# - do LP optimization
    yeast_model.optimize()

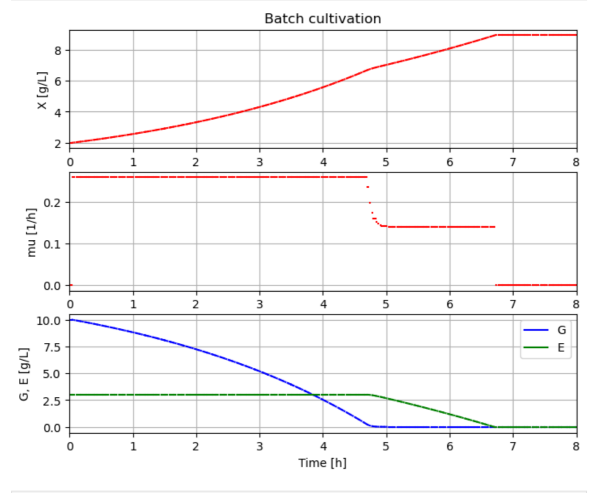
return (yeast_model.objective.value, yeast_model.variables.qGr_opt.primal, y

In [5]: # Initialization
    V_0=1.0
    init(V_0=V_0, VX_0=V_0*2.0, VG_0=V_0*10, VE_0=3.0)

In [6]: # Loop of simulations
    t_final = 8.0
    t_samp = 0.0333
    n_samp = t_final/t_samp + 1

In [7]: # Simulate n sample steps
    newplot(title='Batch cultivation', plotType='TimeSeries2')
```

```
In [7]: # Simulate n sample steps
newplot(title='Batch cultivation', plotType='TimeSeries2')
ax1.set_xlim([0, t_final]); ax2.set_xlim([0, t_final]); ax3.set_xlim([0, t_final]);
simu(t_samp, options=opts_fast)
for i in range(int(n_samp)):
    (mum_opt, qGr_opt, qEr_opt, qO2_opt) = culture(sim_res['bioreactor.c[2]'][-1
    par(mum=mum_opt, qGr=qGr_opt, qEr=qEr_opt, qO2=qO2_opt)
    simu(t_samp, 'cont', options=opts_fast)
```



```
In [8]: system_info()
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```
System information
         -OS: Windows
         -Python: 3.9.16
         -Scipy: not installed in the notebook
         -FMPy: 0.3.15
         -FMU by: JModelica.org
         -FMI: 2.0
         -Type: CS
         -Name: BPL_YEAST_COB.Batch
         -Generated: 2023-05-31T07:27:39
         -MSL: 3.2.2 build 3
         -Description: Bioprocess Library version 2.1.1
         -Interaction: FMU-explore for FMPy version 0.9.8
        !conda list optlang
In [9]:
        # packages in environment at C:\Users\janpa\miniconda3\envs\fmpylab:
        # Name
                                   Version
                                                             Build Channel
                                   1.7.0
                                                                      conda-forge
        optlang
                                                      pyhd8ed1ab 0
In [ ]:
```