BPL_TEST2_Batch_design_space - demo

In this notebook the design space for a batch cultivation process is visualized. We study it first without any measurement noise and then with measurement noise and use one separate FMU for each.

The end critersa for a batch is here when the subdstrate level has decreased below a certain predefined level Smin.

The evaluation of the batch culture is just in terms of the obtained value of cell concentration at the end in combination with how long time the culture took. The batch is accepted provided the culture fullfil the two requirements:

- X_final > X_final_min
- Time_final < time_final_max

The question is what range of process parameters Y and qSmax that can be allowed to still get accepted batches.

Here we simply use brute force and sweep through a number combinations of process parameters and evaluate by simulation the result for each parameter setting. We get rather clear-cut corners in the process parameter space that result in acceptable batches.

In the later part we introduce substrate measurement error and in this way introduce some uncertainty in the determination of end of batch. The impact of this measurement noise is that the designa space get more rounded corners.

The practical experimental approach is usually to just use a few parameter combinations and evaluate these and from that information calculate the design space. Usually linearity assumption is used. The combination of this experimental approach with brute force simulation is discussed in reference [1].

1 Batch end detection - no measurement noise

Here we load a system model without noise. Thus detection of end of batch is an event in continuous time.

Note that both disp() and describe() takes values from the last simulation

Brief information about a command by help(), eg help(simu)
Key system information is listed with the command system_info()

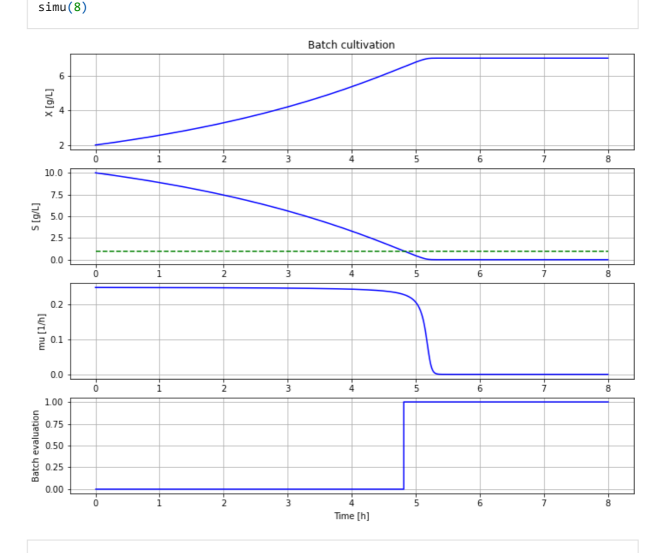
```
In [2]: # Adjust the diagrm size
    plt.rcParams['figure.figsize'] = [30/2.54, 24/2.54]
```

1.1 Batch evaluation

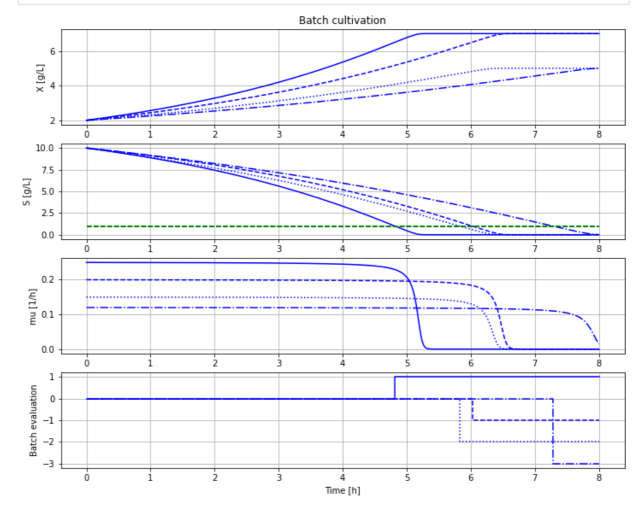
Here we see first an example of batch that has an end of batch that fulfills the criteria for acceptance. In the following diagram we see examples of impact of variation on the criteria for acceptance.

The variable batch_evaluation goes from 0 to either 1 or a negative value when end of batch is detected. A positive value 1 means that the acceptance criteria is fullfilled and a negative value -1, -2 or -3 is obtained if one or more criteria for acceptance is broken.

```
In [3]: # Nominal parameters
    par(S_min=1.0, time_final_max=6.0, X_final_min=5.0)
    init(VX_0=2, VS_0=10)
    par(Y=0.5, qSmax=0.5, Ks=0.1)
In [4]: # Simulation of nominal parameters that gives a batch that meed the end criteria
    newplot(plotType='TimeSeries_2')
    rim(0)
```



```
In [5]: # Exammple of process parameter changes and how they meet the end criteria
newplot(plotType='TimeSeries_2')
par(Y=0.50, qSmax=0.50); simu(8) # - pass (solid line)
par(Y=0.50, qSmax=0.40); simu(8) # - fail criteria time_final < 6.0 (dashed line)
par(Y=0.30, qSmax=0.50); simu(8) # - fail criteria X_final > 5.0 (dotted line)
par(Y=0.30, qSmax=0.40); simu(8) # - fail both criteria (dash dotted line)
```



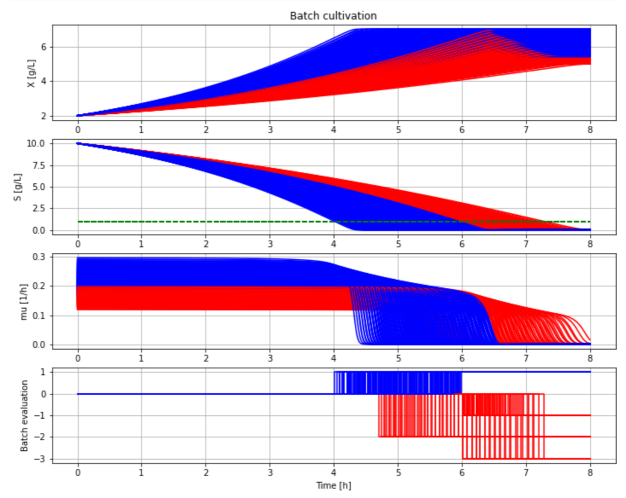
We see that the accepted batch (solid line) finish first. The batches that fail take longer time and two of them has also lower cell concentration at the end.

1.2 Batch evaluation under process variation - parameter sweep

Now let us systematically sweep through a number of combinations of process parameters Y and qSmax and evaluate the batches and visualise their result.

```
In [6]: # Define sweep ranges and storage of final data
    nY = 20
    nqSmax = 20
    Y_range = np.linspace(0.3,0.5,nY)
    qSmax_range = np.linspace(0.4,0.6,nqSmax)
    data = np.zeros([nY,nqSmax,5])
In [7]: # Run parameter sweep - takes a few minutes
    newplot(plotType='TimeSeries_2_diagrams')
    init(VX_0=2, VS_0=10)
    for j in range(nY):
```

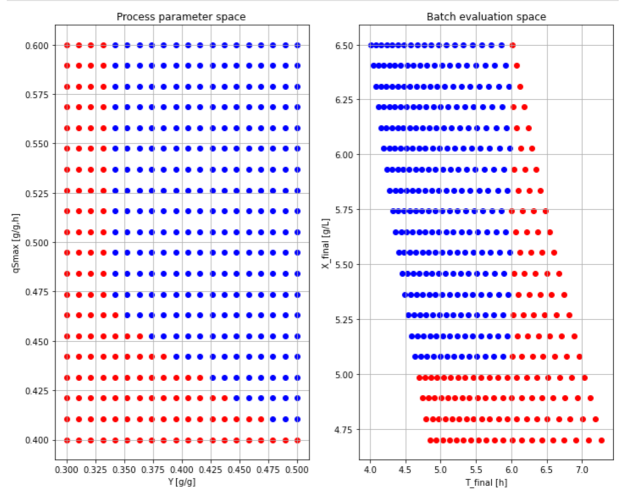
```
for k in range(nqSmax):
        par(Y=Y_range[j])
        par(qSmax=qSmax_range[k])
        simu(8)
        # Store final results
        data[j,k,0] = Y_range[j]
        data[j,k,1] = qSmax_range[k]
        data[j,k,2] = sim_res['monitor.time_final'][-1]
        data[j,k,3] = sim_res['monitor.X_final'][-1]
        data[j,k,4] = sim_res['monitor.batch_evaluation'][-1]
        # Plot simulation results
        if sim_res['monitor.batch_evaluation'][-1] > 0:
            ax1.plot(sim_res['time'], sim_res['bioreactor.c[1]'],'b-')
            ax2.plot(sim_res['time'], sim_res['bioreactor.c[2]'],'b-')
            ax2.plot([0, simulationTime], [model.get('monitor.S_min'), model.get('mo
            ax3.plot(sim_res['time'], sim_res['bioreactor.culture.q[1]'],'b-')
            ax4.step(sim_res['time'],sim_res['monitor.batch_evaluation'],where='post
            ax1.plot(sim_res['time'], sim_res['bioreactor.c[1]'],'r-')
            ax2.plot(sim_res['time'], sim_res['bioreactor.c[2]'],'r-')
            ax2.plot([0, simulationTime], [model.get('monitor.S_min'), model.get('mo
            ax3.plot(sim_res['time'], sim_res['bioreactor.culture.q[1]'],'r-')
            ax4.step(sim_res['time'],sim_res['monitor.batch_evaluation'],where='post
plt.show()
```



Batches represented by blue lines are those that in end got accepted. The red ones failed.

```
In [8]: # Show end results
plt.figure()
```

```
ax1 = plt.subplot(1,2,1)
ax2 = plt.subplot(1,2,2)
for j in range(nY):
    for k in range(nqSmax):
        if data[j,k,4] > 0:
            ax1.scatter(data[j,k,0],data[j,k,1],c='b')
        else:
            ax1.scatter(data[j,k,0],data[j,k,1],c='r')
ax1.grid()
#plt.axis([0, 0.8, 0, 0.8])
ax1.set_ylabel('qSmax [g/g,h]')
ax1.set_xlabel('Y [g/g]')
ax1.set_title('Process parameter space')
for j in range(nY):
    for k in range(nqSmax):
        if data[j,k,4] > 0:
            ax2.scatter(data[j,k,2],data[j,k,3],c='b')
            ax2.scatter(data[j,k,2],data[j,k,3],c='r')
ax2.grid()
#plt.axis([0, 8, 0, 8])
ax2.set_xlabel('T_final [h]')
ax2.set_ylabel('X_final [g/L]')
ax2.set_title('Batch evaluation space')
plt.show()
```



Here we visualize the previous simulations results in a different way. Each dot in the left diagram (process parameter space) represent a simulation that give a result in the rigiht diagram (batch evaluation space). The blue dots are those batches that were accepted and the red ones those that failed.

The blue dots in the process parameter space show the "design space" for the acceptance criteria we have.

2 Batch end detection - with measurement noise

Here we load a system model with normal noise added to the sampled value of substrate concentration. The measurement of substrate conentration usually has a higher variation than measurement of cell concentration and therefore we focuse here on the impact on substrate conentrations.

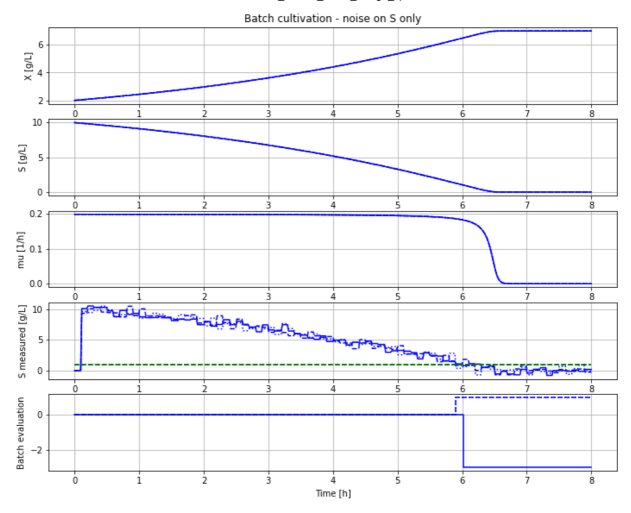
Thus detection of end of batch is now in discrete time with a give samplePeriod (default 0.1 hour). This discreteization also introduce an error in detection of the end point. By changing this sample intervall to shorter values you can see the impact of this error but not done here.

2.1 Batch evaluation under substrate measurement error

Here we see an example of how substrate measurement noise directly affect the evaluation of the batch from accetable to not acceptable.

```
In [10]: # Nominal parameters
    par(S_min=1.0, time_final_max=6.0, X_final_min=5.0)
    init(VX_0=2, VS_0=10)
    par(Y=0.5, qSmax=0.5, Ks=0.1)
    par(sigma=0.48, samplePeriod=0.1)

In [11]: # Simulation of nominal parameters that gives a batch that meed the end criteria
    newplot(plotType='TimeSeries_2')
    par(Y=0.5, qSmax=0.4);
    for value in [2,3,5]: par(seed=value); simu(8)
```

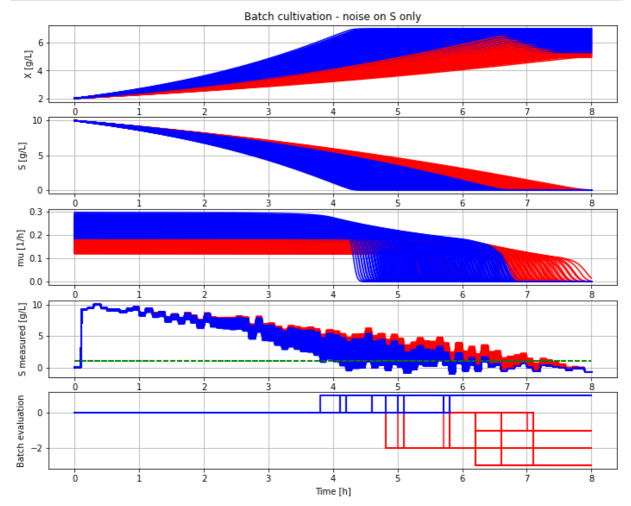


2.2 Batch evaluation under process variation and measurement error - parameter sweep

Now let us again systematically sweep through a number of combinations of process parameters Y and qSmax and evaluate the batches and visualise their result.

```
In [12]:
          # Define sweep ranges and storage of final data
          nY = 20
          nqSmax = 20
          Y range = np.linspace(0.3,0.5,nY)
          qSmax_range = np.linspace(0.4,0.6,nqSmax)
          data = np.zeros([nY,nqSmax,5])
In [13]:
          # Run parameter sweep - takes a few minuts
          newplot(plotType='TimeSeries 2 diagrams')
          par(sigma=0.48, seed=1, samplePeriod=0.1)
          for j in range(nY):
              for k in range(nqSmax):
                  par(Y=Y_range[j])
                  par(qSmax=qSmax_range[k])
                  simu(8)
                  # Store final results
                  data[j,k,0] = Y_range[j]
                  data[j,k,1] = qSmax_range[k]
                  data[j,k,2] = sim_res['monitor.time_final'][-1]
                  data[j,k,3] = sim_res['monitor.X_final'][-1]
```

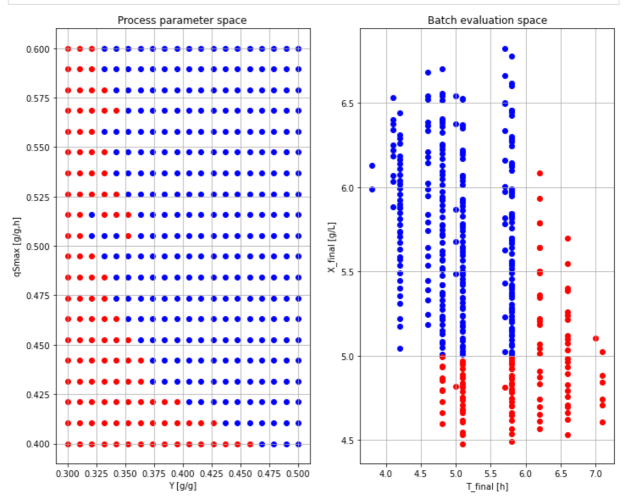
```
data[j,k,4] = sim_res['monitor.batch_evaluation'][-1]
        # Plot simulation results
        if sim_res['monitor.batch_evaluation'][-1] > 0:
            ax1.plot(sim_res['time'], sim_res['bioreactor.c[1]'],'b-')
            ax2.plot(sim_res['time'], sim_res['bioreactor.c[2]'],'b-')
            ax3.plot(sim_res['time'], sim_res['bioreactor.culture.q[1]'],'b-')
            ax4.plot(sim_res['time'], sim_res['sensor.out.c[2]'],'b-')
            ax4.plot([0, simulationTime], [model.get('monitor.S_min'), model.get('mo
            ax5.step(sim_res['time'],sim_res['monitor.batch_evaluation'],where='post
        else:
            ax1.plot(sim_res['time'], sim_res['bioreactor.c[1]'],'r-')
            ax2.plot(sim_res['time'], sim_res['bioreactor.c[2]'],'r-')
            ax3.plot(sim_res['time'], sim_res['bioreactor.culture.q[1]'],'r-')
            ax4.plot(sim_res['time'], sim_res['sensor.out.c[2]'],'r-')
            ax4.plot([0, simulationTime], [model.get('monitor.S_min'), model.get('mo
            ax5.step(sim_res['time'],sim_res['monitor.batch_evaluation'],where='post
plt.show()
```



```
In [14]:  # Show end results
plt.figure()
ax1 = plt.subplot(1,2,1)
ax2 = plt.subplot(1,2,2)

for j in range(nY):
    for k in range(nqSmax):
        if data[j,k,4] > 0:
            ax1.scatter(data[j,k,0],data[j,k,1],c='b')
        else:
            ax1.scatter(data[j,k,0],data[j,k,1],c='r')
ax1.grid()
```

```
#plt.axis([0, 0.8, 0, 0.8])
ax1.set_ylabel('qSmax [g/g,h]')
ax1.set_xlabel('Y [g/g]')
ax1.set_title('Process parameter space')
for j in range(nY):
    for k in range(nqSmax):
        if data[j,k,4] > 0:
            ax2.scatter(data[j,k,2],data[j,k,3],c='b')
        else:
            ax2.scatter(data[j,k,2],data[j,k,3],c='r')
ax2.grid()
#plt.axis([0, 8, 0, 8])
ax2.set_xlabel('T_final [h]')
ax2.set_ylabel('X_final [g/L]')
ax2.set_title('Batch evaluation space')
plt.show()
```



We see that we get somwehat different results in the parameter space. The acceptable region with blue dots (design space) get a more rounded corner. The vertical left line is also more rugged.

With much more simulations we could get a better idea of the probablity that a batch is accepted and determine the design space in proabilistic sense.

3 Summary

We have worked through a simple example of evaluation of batch with given acceptance criteria and how that criteria can be translated to acceptable variation in process parameters, i.e. the

design space.

In the deterministic case we get a rather clear cut design space.

In the more realistic case with subsrate measurment noise included we get a more complicated design space, but still similar.

The stochastic model introduce erorrs both due to the added normal noise in the substrate concentration, and due to the fact that we use time discrete system for the noise. The impact of the time discrete check when batch has ended can be made smaller by chosing a smaller sample intervall. This was not studied here and is left for the interested reader.

References

[1] Axelsson J.P. and A. Elsheikh: "An example of sensitivity analysis of a bioprocess using Bioprocess Library for Modelica", Proceedings MODPROD, Linköping, Sweden 2019, see presentation here.

Appendix

```
In [15]:
         describe('parts')
         ['bioreactor', 'bioreactor.culture', 'liquidphase', 'monitor', 'MSL', 'sensor']
In [16]:
          describe('MSL')
         MSL: 3.2.2 build 3 - used components: Noise.NormalNoise
In [17]:
          system_info()
         System information
          -OS: Windows
          -Python: 3.9.5
          -PyFMI: 2.9.5
          -FMU by: JModelica.org
          -FMI: 2.0
          -Type: FMUModelCS2
          -Name: BPL_TEST2.BatchWithNoise
          -Generated: 2022-09-04T14:31:31
          -MSL: 3.2.2 build 3
          -Description: Bioprocess Library version 2.1.0 beta
          -Interaction: FMU-explore ver 0.9.3
 In [ ]:
```