BPL_IEC_operation script with PyFMI

The key library PyFMI is installed.

After the installation a small application BPL_IEC_operation is loaded and run. You can continue with this example if you like.

!lsb_release -a # Actual VM Ubuntu version used by Google

No LSB modules are available. Distributor ID: Ubuntu

Description: Ubuntu 22.04.3 LTS

!conda update -n base -c defaults conda --yes

Release: 22.04 Codename: jammy

%env PYTHONPATH=



env: PYTHONPATH=

```
!wget $$ \underline{$https://repo.anaconda.com/miniconda/Miniconda3-py310\_23.1.0-1-Linux-x86\_64.sh} $$ \underline{$https://repo.anaconda.com/miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Miniconda/Minico
!chmod +x Miniconda3-py310_23.1.0-1-Linux-x86_64.sh
!bash ./Miniconda3-py310_23.1.0-1-Linux-x86_64.sh -b -f -p /usr/local
import sys
sys.path.append('/usr/local/lib/python3.10/site-packages/')
              --2024-01-18 11:47:29-- <a href="https://repo.anaconda.com/miniconda/Miniconda3-py310">https://repo.anaconda.com/miniconda/Miniconda3-py310</a> 23.1.0-1-Linux-x86 64.sh
              Resolving repo.anaconda.com (repo.anaconda.com)... 104.16.131.3, 104.16.130.3, 2606:4700::6810:8203, ...
              Connecting to repo.anaconda.com (repo.anaconda.com)|104.16.131.3|:443... connected.
              HTTP request sent, awaiting response... 200 OK
Length: 74403966 (71M) [application/x-sh]
              Saving to: 'Miniconda3-py310_23.1.0-1-Linux-x86_64.sh'
             Miniconda3-py310_23 100%[============] 70.96M
                                                                                                                                                                                               123MB/s
              2024-01-18 11:47:29 (123 MB/s) - 'Miniconda3-py310_23.1.0-1-Linux-x86_64.sh' saved [74403966/74403966]
              PREFIX=/usr/local
              Unpacking payload ...
              Installing base environment...
              Downloading and Extracting Packages
             Downloading and Extracting Packages
              Preparing transaction: done
              Executing transaction: done
              installation finished.
```

Preparing transaction: done Verifying transaction: done Executing transaction: done

!conda --version
!python --version

conda 23.11.0 Python 3.10.13

!conda install -c conda-forge pyfmi --yes # Install the key package

```
Preparing transaction: done
Verifying transaction: done
Executing transaction: done
```

Preparation of BPL_IEC_operation

Now specific installation and the run simulations. Start with connecting to Github. Then upload the two files:

- FMU BPL_IEC_Column_system_linux_om_me
- Setup-file BPL_IEC_explore

```
%bash
git clone https://github.com/janpeter19/BPL_IEC_operation
    Cloning into 'BPL_IEC_operation'...
%cd BPL_IEC_operation
    /content/BPL_IEC_operation
```

BPL_IEC_operation

Authors: Karl Johan Brink and Jan Peter Axelsson

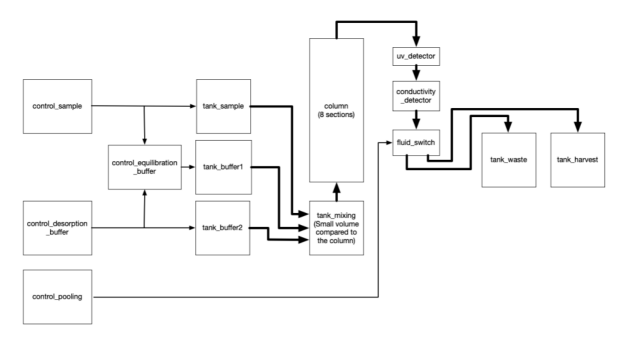
In this notebook we show operation of a typical ion-exchange chromatography step. The impact of pH is also illustrated.

The model is based on the simplified model [1].

```
run -i BPL_IEC_explore.py
    Linux - run FMU pre-comiled OpenModelica 1.21.0
    Model for bioreactor has been setup. Key commands:
                   - change of parameters and initial values
     - par()
                   - change initial values only
     - init()

    simulate and plot

     - simu()
     - newplot()
                  – make a new plot
     - show()
                   - show plot from previous simulation
                   - display parameters and initial values from the last simulation
     - describe() - describe culture, broth, parameters, variables with values/units
    Note that both disp() and describe() takes values from the last simulation
    and the command process_diagram() brings up the main configuration
    Brief information about a command by help(), eg help(simu)
    Key system information is listed with the command system\_info()
%matplotlib inline
plt.rcParams['figure.figsize'] = [36/2.54, 30/2.54]
# The process diagram is made outside Modelica for illustration of the configuration
process_diagram()
```



1 Typical parameters for a pilot scale ion exchange chromatography column process setup

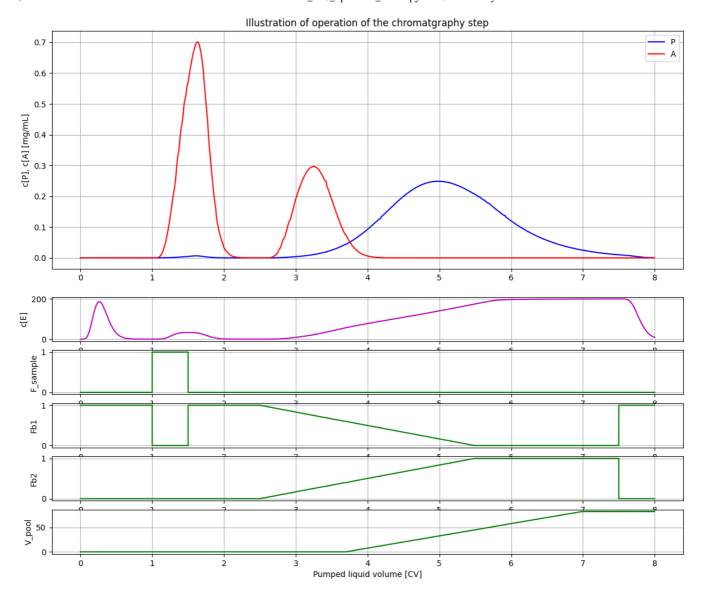
```
# From given column height (h) diameter (d) and linear flow rate (lfr)
# actual column volume (V) and volume flow rate (VFR) are calculated below.

from numpy import pi
h = 20.0
d = 1.261
a = pi*(d/2)**2
V = h*a
print('V =', np.round(V,1), '[mL]')

lfr = 48
VFR = a*lfr/60
print('VFR =', np.round(VFR,1), '[mL/min]')  # Pump schedule parameter

V = 25.0 [mL]
VFR = 1.0 [mL/min]
```

```
# Sample concentration product P_in and antagonist A_in
par(P_in = 1.0)
par(A in = 1.0)
par(E_in = 0.0)
# Column properties are described by the size and binding capacity of the resin Q_av
par(height = h)
par(diameter = d)
par(Q_av = 6.0)
# Remaining salt koncentration in the column from prvious batch and eliminated during the initial equilibration period
init(E_0 = 50)
# Salt koncentration of the desorption buffer
par(E_in_desorption_buffer = 8.0)
# Flow rate rate through the
par(LFR=lfr)
# Switching points during operation are conveniently described in terms of multiples of the column volume V
CV_ekv = 1.0
CV_ads = 0.5
CV_wash = 1.0
CV_desorb = 3.0
CV_start_pool = 1.2
CV_stop_pool = 4.5
CV_ekv2 = 2.5
\verb|par(scale_volume=True, start_adsorption=CV_ekv*V, stop_adsorption=(CV_ekv+CV_ads)*V)|
par(start_desorption=(CV_ekv+CV_ads+CV_wash)*V, stationary_desorption=(CV_ekv+CV_ads+CV_wash+CV_desorb)*V)
par(stop_desorption=7.5*V)
par(start_pooling=(CV_ekv+CV_ads+CV_wash+CV_start_pool)*V, stop_pooling=(CV_ekv+CV_ads+CV_wash+CV_stop_pool)*V)
# Simulation and plot of results
newplot(title='Illustration of operation of the chromatgraphy step', plotType='Elution-conductivity-vs-CV-combined-all')
\verb|simu|(CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR|
```



Comments of steps of operations:

- 1) Time: 0-1 hours equilibration. Just to illustrate the equilibration process the first part of the column is given an initial value of salt concentration.
- 2) Time: 1-1.5 hours sample is loaded on the column. The product P is adsorbed to the columne and just a small amount passes through and goes to the waste. The antagonist A is much less adsrobed.
- 3) Time: 1.5-2.5 hours washing 1. The column comes to equilibrium and both antagonist and product comes down to low levels.
- 4) Time: 2.5-5.5 hours desorption. A linear gradient of increaseing salt concentration is applied. First the antagonist and later the product comes out
- $5) \ Time: 5.5-7.5 \ hours washing \ 2 \ The \ The \ column \ has \ constant \ salt \ concentration \ and \ stationary \ desorption.$
- 6) Time: 3.7-7.0 hours pooling of product. The start- and stop of pooling are chosen with trade-off between maximizing the product pooled and minimize the amount of antagonist in the pooling.
- 7) Time: 7.5-8.0 hours desorption stopped and salt is washed out and preparation of the next batch to come.

Note that step 4 and 5 is parallel to step 6.

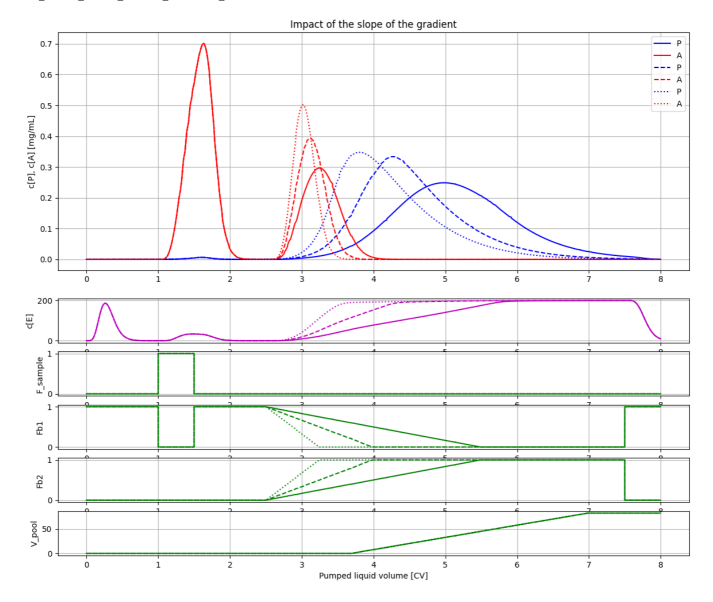
```
# Check mass-balance of P and A
P_mass = model.get('tank_harvest.m[1]') + model.get('tank_waste.m[1]')
A_mass = model.get('tank_harvest.m[2]') + model.get('tank_waste.m[2]')
print('P_mass [mg] =', P_mass)
print('A_mass [mg] =', A_mass)
```

```
P_{mass} [mg] = [12.42212162]
A_{mass} [mg] = [12.48878113]
```

These values should be compared with the expected value 12.5 mg, i.e. half a column volume with sample concentration 1 mg/L. The difference is due to numerical errors during simulation.

2 The impact of the slope of the desorption gradient

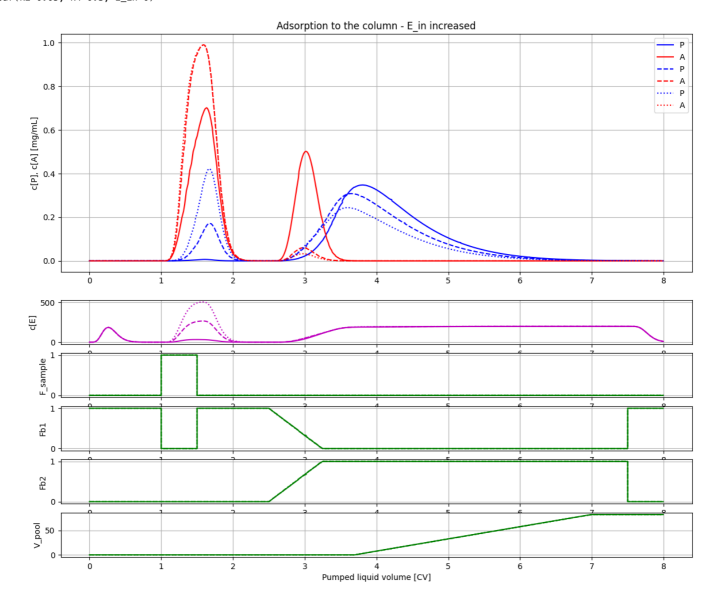
```
# Simulations showing the impact of change of slope of the desorption gradient
newplot(title='Impact of the slope of the gradient', plotType='Elution-conductivity-vs-CV-combined-all')
# Same gradienet as before
par(start_desorption=(CV_ekv+CV_ads+CV_wash)*V, stationary_desorption=(CV_ekv+ CV_ads+CV_wash+CV_desorb)*V)
par(stop_desorption=7.5*V)
simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)
# Gradeint finishes after 0.5 of the volume
par(stationary_desorption = (CV_ekv + CV_ads + CV_wash + 0.5*CV_desorb)*V)
simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)
# Fradient finishes after 0.25 of the volume
par(stationary_desorption = (CV_ekv + CV_ads + CV_wash + 0.25*CV_desorb)*V)
simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)
```



Note the pens shift style for each simulation in the order: solid, dashed, dotted, dash-dotted. The actual simulations done you see in the preceeding cell.

3 The impact of salt concentration in the sample

```
# Let us investigate the impact of increasing salt concetration in the sample E_in
# Simulate and plot the results
newplot(title='Adsorption to the column - E_in increased', plotType='Elution-conductivity-vs-CV-combined-all')
for value in [0, 10, 20]:
    par(E_in=value)
    simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)
# Restore default values
par(k2=0.05, k4=0.3, E_in=0)
```



Note, that increased salt concentration in the sample affect binding of both proteins. During adsorption less is bound. During desoprtion less product P can be harvested but the fraction of antagonist A may be lowered. Thus, some product is lost but the quality in terms of purity is improved.

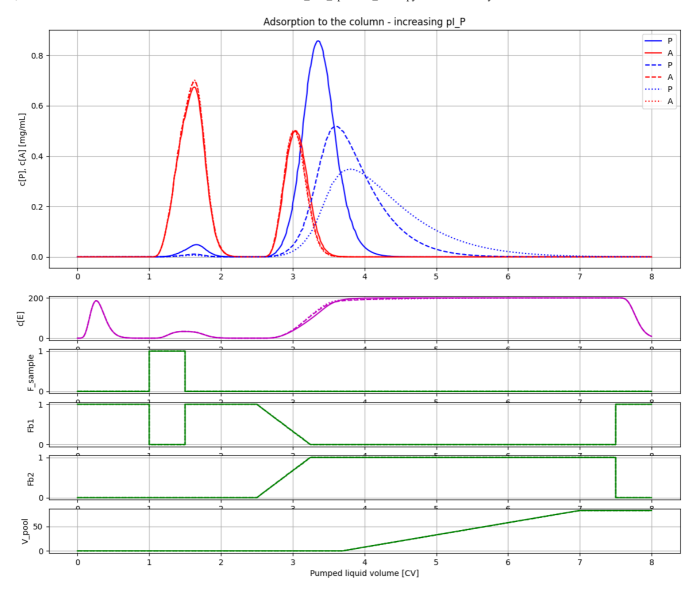
4 The impact of change of binding strength due to pH

There are many factors that contribute to the binding strength. A most important factor is the pH-value of the resin and the characteristic iso-electric point of the protein. The binding strength can be seen as proportional to the difference.

The binding strength of the resin is described by the quotient KP=k1/k2 for the protein P and similarly KA=k3/k4 for the protein A.

Below a few help-functions that describe this idea of the pH difference and its impact on binding strength in terms of the parameters k1, k2, k3, and k4 of the protein-resin interaction.

```
# Define function that describe the proportionality of binding strength ot
# the pH difference of the iso-electric point and the resin
def KP_pH_sensitivity(pI_P=8.0, pH_resin=7.0):
   coeff_pH = 6.0
    return coeff_pH*(pI_P-pH_resin)
def KA_pH_sensitivity(pI_A=7.1667, pH_resin=7.0):
    coeff_pH = 1.0
    return coeff_pH*(pI_A-pH_resin)
def par_pH(pI_P=8.0, pI_A=7.1667, pH_resin=7.0, TP=3.33, TA=20.0):
    if (pI_P > pH_resin) & (pI_A > pH_resin):
       par(k2 = 1/(TP*KP_pH_sensitivity(pI_P=pI_P, pH_resin=pH_resin)))
       par(k4 = 1/(TA*KA_pH_sensitivity(pI_A=pI_A, pH_resin=pH_resin)))
   else:
       print('Both pI_P > pH_resin and pI_A > pH_resin must hold - no parameter change made')
# The default parameters of the column
disp('column')
    diameter: 1.261
    height: 20.0
    x_m : 0.3
    k1 : 0.3
    k2 : 0.05
    k3: 0.05
    k4: 0.3
    Q_av : 6.0
    E_0 : 50.0
# Let us investigate the impact of change of the iso-electric pH for protein P
# Simulate and plot the results
newplot(title='Adsorption to the column - increasing pI_P', plotType='Elution-conductivity-vs-CV-combined-all')
for value in [7.2, 7.6, 8.0]:
   par_pH(pI_P=value, pI_A=7.1667, pH_resin=7.0)
   simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)
# Restore default values
par(k2 = 0.05, k4 = 0.3)
```



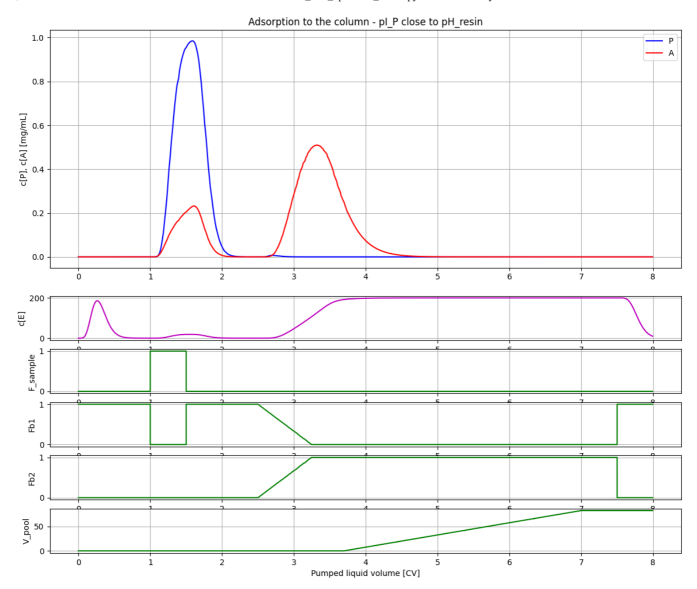
Note, with increasing pl_P the binding of P increase which leads less loss of product during adsorption. During desorption the peak height is lower with increasing binding strength, but the total amoiunt of product P that can be harvested is higher, due to the smaller loss during adsorption.

```
# Let us investigate the impact of pI_P close to pH_resin

# Simulate and plot the results
newplot(title='Adsorption to the column - pI_P close to pH_resin', plotType='Elution-conductivity-vs-CV-combined-all')

for value in [7.0001]:
    par_pH(pI_P=value, pI_A=8)
    simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)

# Restore default values
par(k2=0.05, k4=0.3)
```

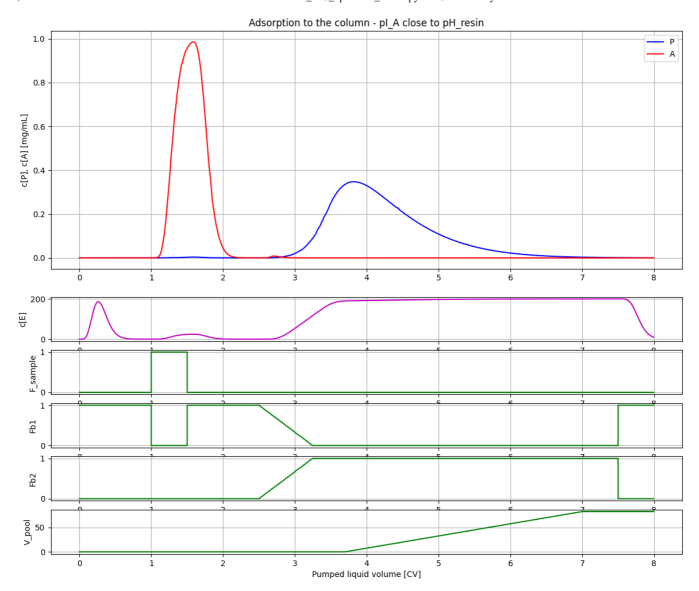


```
# Let us investigate the impact of pI_A close to pH_resin

# Simulate and plot the results
newplot(title='Adsorption to the column - pI_A close to pH_resin', plotType='Elution-conductivity-vs-CV-combined-all')

for value in [7.001]:
    par_pH(pI_P=8.0, pI_A=value)
    simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)

# Restore default values
par(k2=0.05, k4=0.3)
```



```
# Let us also investigate the impact of salt concentration of the desorptions buffer
# Simulate and plot the results
newplot(title='Adsorption to the column - desorption buffer salt conc varied', plotType='Elution-conductivity-vs-CV-combinec

for value in [8.0, 16.0]:
    par(E_in_desorption_buffer=value)
    par_pH(pI_P=8.0, pI_A=7.001, pH_resin=7.0)
    simu((CV_ekv+CV_ads+CV_wash+CV_desorb+CV_ekv2)*V/VFR)

# Restore default values
par(E_in_desorption_buffer=8.0)
par(k2=0.05, k4=0.3)
```

