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BPL_TEST2_Perfusion script with PyFMI

The key library PyFMI is installed.

After the installation a small application BPL_TEST2_Perfusion 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
             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/')
          --2024-11-07 08:26:17-- <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)... 104.16.32.241, 104.16.191.158, 2606:4700::6810:bf9e, ...
             Connecting to repo.anaconda.com (repo.anaconda.com)|104.16.32.241|: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 80.3MB/s
             2024-11-07 08:26:18 (80.3 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.
!conda update -n base -c defaults conda --yes
```

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

!conda --version
!python --version

conda 23.1.0 Python 3.10.15

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

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

Notes of BPL_TEST2_Perfusion

This notebook explore perfusion cultivation in comparison with ordinary continuous cultivation (chemostat) and use comparable settings to earlier notebook. Further you see here examples of interaction with the simplified commands par(), init(), simu() etc as well as direct interaction with the FMU which is called "model" here. The last simulation is always available in the workspace and called "sim_res". Note that describe() brings mainly up from descriptive information from the Modelica code from the FMU but is complemented by some information given in the Python setup file.

Now specific installation run a simulation and notebook for that Start with connecting to Github. Then upload the two files:

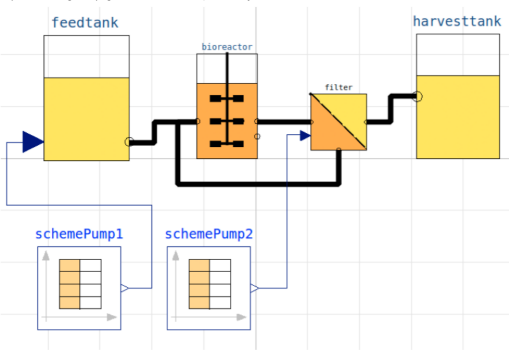
- FMU BPL_TEST2_Perfusion_linux_om_me.fmu
- Setup-file BPL_TEST2_Perfusion_explore.py

```
%bash
git clone https://github.com/janpeter19/BPL_TEST2_Perfusion
Cloning into 'BPL_TEST2_Perfusion'...
%cd BPL_TEST2_Perfusion
/content/BPL_TEST2_Perfusion
run -i BPL_TEST2_Perfusion_explore.py
→ Linux - run FMU pre-comiled OpenModelica
    Model for bioreactor has been setup. Key commands:
      - par()
                   - change of parameters and initial values
                   - change initial values only
     - init()
     - simu()

    simulate and plot

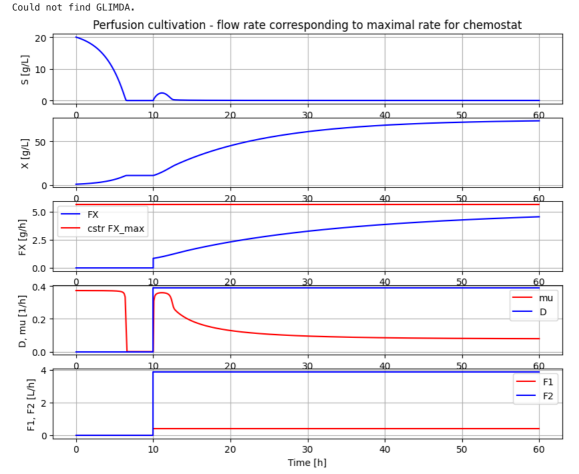
                  - make a new plot
     - newplot()
     - show()
                   - show plot from previous simulation
                   - display parameters and initial values from the last simulation
     - disp()
     - describe() - describe culture, broth, parameters, variables with values/units
    Note that 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()
# Filter out DepracationWarnings for 'np.float as alias' is needed - wish
import warnings
warnings.filterwarnings("ignore")
%matplotlib inline
plt.rcParams['figure.figsize'] = [25/2.54, 20/2.54]
process_diagram()
```

 \longrightarrow No processDiagram.png file in the FMU, but try the file on disk.



```
# Process parameters used throughout
par(Y=0.5, qSmax=0.75, Ks=0.1)
                                                                    # Culture
par(filter_eps=0.10, filter_alpha_X=0.02, filter_alpha_S=0.10)
                                                                    # Filter
par(S_in=30.0)
                                                                    # Inlet substrate conc
init(V_start=1.0, VX_start=1.0)
                                                                    # Process initial conditions that are common
                                                                    # Pump schedule parameter
eps = parDict['filter_eps']
# Simulation of process with flow rate close to wash-out for chemostat
init(VS_start=20)
                                                         # Process initial
par(pump1_t1=10, pump2_t1=10)
                                                         # Pump schedule - recycle flow 10 times perfusion flow
par(pump1_F1=2.5*0.155, pump2_F1=2.5*0.155/eps)
par(pump1_t2=940, pump2_t2=940, pump1_t3=950, pump2_t3=950, pump1_t4=960, pump2_t4=960)
newplot(title='Perfusion cultivation - flow rate corresponding to maximal rate for chemostat')
simu(60)
```

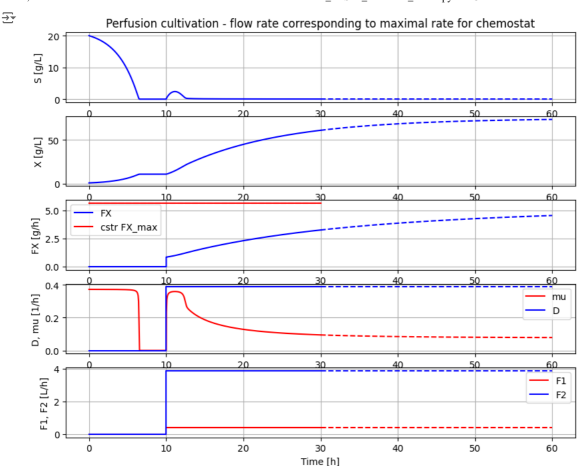
Could not find cannot import name 'dopri5' from 'assimulo.lib' (/usr/local/lib/python3.10/site-packages/assimulo/lib/_i Could not find cannot import name 'rodas' from 'assimulo.lib' (/usr/local/lib/python3.10/site-packages/assimulo/lib/_in Could not find cannot import name 'odassl' from 'assimulo.lib' (/usr/local/lib/python3.10/site-packages/assimulo/lib/_i Could not find ODEPACK functions. Could not find RADAR5



Simulation of process with flow rate close to wash-out for chemostat

```
init(VS_start=20)  # Process initial
par(pump1_t1=10, pump2_t1=10)  # Pump schedule - recycle flow 10 times perfusion flow
par(pump1_F1=2.5*0.155, pump2_F1=2.5*0.155/eps)
par(pump1_t2=940, pump2_t2=940, pump1_t3=950, pump2_t3=950, pump1_t4=960, pump2_t4=960)

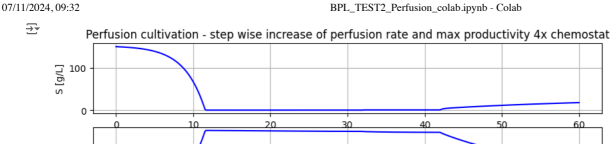
newplot(title='Perfusion cultivation - flow rate corresponding to maximal rate for chemostat')
simu(30)
simu(30,'cont')
```

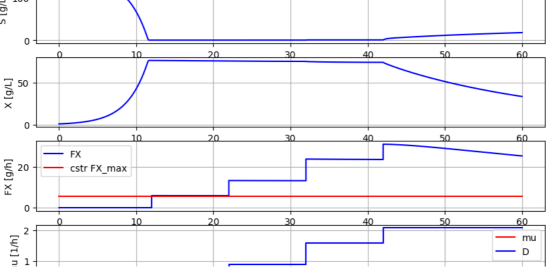


Note the inability of the OpenModelica FMU to handle simu('cont') properly.

```
# Concentration factor of the filter
c=model.get('filter.retentate.c[1]')[0]/model.get('filter.inlet.c[1]')[0]
print('Conc factor of perfusion filter =', np.round(c,3))
→ Conc factor of perfusion filter = 1.179
c_data=sim_res['filter.retentate.c[1]']/sim_res['filter.inlet.c[1]']
print('Conc factor variation', np.round(min(c_data[151:]), 3), np.round(max(c_data[151:]),3))

→ Conc factor variation 1.179 1.179
# Simulation of process with step-wise increase of pefusion rate until wash-out.
# This means that re-circulation rate change at the same time as the perfusion rate.
                                                         # Process initial varied
init(VS_start=150)
par(pump1_t1=12, pump2_t1=12)
                                                         # Pump schedule - recycle flow 10 times perfusion flow
par(pump1_F1=2.5*0.155, pump2_F1=2.5*0.155/eps)
par(pump1_t2=22, pump2_t2=22)
par(pump1_F2=2.5*0.35, pump2_F2=2.5*0.35/eps)
par(pump1_t3=32, pump2_t3=32)
par(pump1_F3=2.5*0.63, pump2_F3=2.5*0.63/eps)
par(pump1_t4=42, pump2_t4=42)
par(pump1_F4=2.5*0.83, pump2_F4=2.5*0.83/eps)
newplot(title='Perfusion cultivation - step wise increase of perfusion rate and max productivity 4x chemostat')
simu(60)
```





Simulation without a plot and just to check typical values at high production rate

c_data=sim_res['filter.retentate.c[1]']/sim_res['filter.inlet.c[1]'] print('Conc factor variation', np.round(min(c_data[190:]), 3), 'to', np.round(max(c_data[190:]),3))

1 1 #describe('cstrProdMax')

The maximal biomass productivity before washout is obtained aroudn 40 hours np.round (model.get('harvesttank.inlet.F')[0]*model.get('harvesttank.inlet.c[1]')[0], 1)

→ 23.5

Thus perfusion (with this filter) brings a productivity improvement of about np.round(23.5/5.6,1)

→ 4.2

Finally we check the filter flow rates at time 40 hour - note the negative sign for outlfow model.get('filter.inlet.F')[0]

15 740000000000000