## BPL\_TEST2\_Chemostat script with PyFMI

The key library PyFMI is installed.

After the installation a small application BPL\_TEST2\_Chemostat 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
                     Ubuntu 22.04.3 LTS
    Description:
    Release:
                     22.04
    Codename:
                     jammy
%env PYTHONPATH=
    env: PYTHONPATH=
!python --version
    Python 3.10.12
!wget https://repo.anaconda.com/miniconda/Miniconda3-py310_23.1.0-1-Linux-x86_64.sh
!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
sys.path.append('/usr/local/lib/python3.10/site-packages/')
Show hidden output
!conda update -n base -c defaults conda --yes
Show hidden output
!conda --version
!pvthon --version
    conda 23.11.0
    Python 3.10.13
!conda install -c conda-forge pyfmi --yes # Install the key package
Show hidden output
```

## Now specific installation run a simulation and notebook for that

Start with connecting to Github. Then upload the two files:

```
• FMU - BPL_TEST2_Chemostat_linux_om_me.fmu
```

```
    Setup-file - BPL_TEST2_Chemostat_explore.py
```

```
# Filter out DepracationWarnings for 'np.float as alias' is needed — wish I could make filter more narrow
import warnings
warnings.filterwarnings("ignore")
git clone https://github.com/janpeter19/BPL_TEST2_Chemostat
    Cloning into 'BPL_TEST2_Chemostat'...
%cd BPL_TEST2_Chemostat
    /content/BPL_TEST2_Chemostat
run -i BPL_TEST2_Chemostat_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
- disp()- display parameters and initial values from the last simulation- describe()- describe culture, broth, parameters, variables with values/units

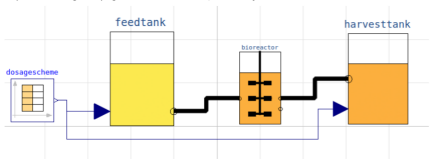
Note that both disp() and describe() takes values from the last simulation and the command process\_diagram() brings up the main configuration

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'] = [25/2.54, 20/2.54]
```

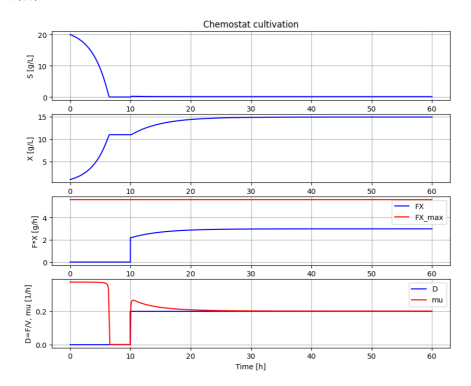
process\_diagram()

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



describe('culture')

Simplified text book model - only substrate S and cell concentration X



```
\# The maximal biomass productivity FX_max [g/h] marked red in the diagram above
# can be calculated for CSTR from the FMU and is
cstrProdMax(model)
    5.625
describe('cstrProdMax')
    Calculate from the model maximal chemostat productivity FX_max : 5.625 [ g/h ]
describe('parts')
     ['bioreactor', 'bioreactor.culture', 'D', 'dosagescheme', 'feedtank', 'harvesttank']
system_info()
    System information
     -OS: Linux
     -Python: 3.10.12
     -Scipy: not installed in the notebook
     -PyFMI: 2.11.0
     -FMU by: OpenModelica Compiler OpenModelica 1.21.0
     -FMI: 2.0
     -Type: FMUModelME2
     -Name: BPL_TEST2.Chemostat
      -Generated: 2023-04-20T12:24:50Z
     -Description: Bioprocess Library version 2.1.1
     -Interaction: FMU-explore version 0.9.8
                                                      + Code -
                                                                + Text
```