

✓ BPL_STEM_AIR_Perfusion script with PyFMI

The key library PyFMI is installed.

After the installation a small application BPL_STEM_AIR_Pefusion 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 https://repo.anaconda.com/miniconda/Miniconda3-py310_23.1.0-1-Linux-x86_64.
!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/')
```

```
⇒ --2024-10-03 09:49:37-- https://repo.anaconda.com/miniconda/Miniconda3-py310
Resolving repo.anaconda.com (repo.anaconda.com)... 104.16.32.241, 104.16.191.1
Connecting to repo.anaconda.com (repo.anaconda.com)|104.16.32.241|:443... con
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 181MB/s in 0.4s
```

```
2024-10-03 09:49:37 (181 MB/s) - 'Miniconda3-py310_23.1.0-1-Linux-x86_64.sh' :
```

```
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
```

```
⇨
```

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

✓ BPL_STEM_AIR_Perfusion setup

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

- FMU - BPL_STEM_AIR_Perfusion_linux_om_me.fmu
- Setup-file - BPL_STEM_AIR_Perfusion_explore

```
# Filter out DeprecationWarnings for 'np.float as alias' is needed – wish I could  
import warnings  
warnings.filterwarnings("ignore")
```

```
%%bash  
git clone https://github.com/janpeter19/BPL_STEM_AIR_Perfusion
```

```
📁 Cloning into 'BPL_STEM_AIR_Perfusion'...
```

```
%cd BPL_STEM_AIR_Perfusion
```

```
📁 /content/BPL_STEM_AIR_Perfusion
```

✓ BPL_STEM_AIR_Perfusion - test

Author: Jan Peter Axelsson

Here we show simulations of stem cell cultivation in an aerated hollow fiber reactor. The reactor volume is kept constant and cells recycled, thus the setup is similar to perfusion cultivation.

The model combines rudimentary cell growth and metabolism combined with times series data of of the metabolic rates: q_{Nmax} , q_{Lc} , and q_{O2} , marked with red in the comprehensive plot.

Ref Greuel et al: "Online measurement of oxygen enables continuous noninvasive evalation of human-induced pluripotent stem cell (hiPSC) culture in a perfused 3D hollow-fober bioreactor", Biotech. Bioeng., 2019.

✓ Setup of the simulation model

```
run -i BPL_STEM_AIR_Perfusion_explore.py
```

Linux – run FMU pre-compiled OpenModelica

Model for bioreactor has been setup. Key commands:

- par() - change of parameters and initial values
- init() - change initial values only
- simu() - simulate and plot
- 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'] = [30/2.54, 24/2.54]
```

✓ About the process model

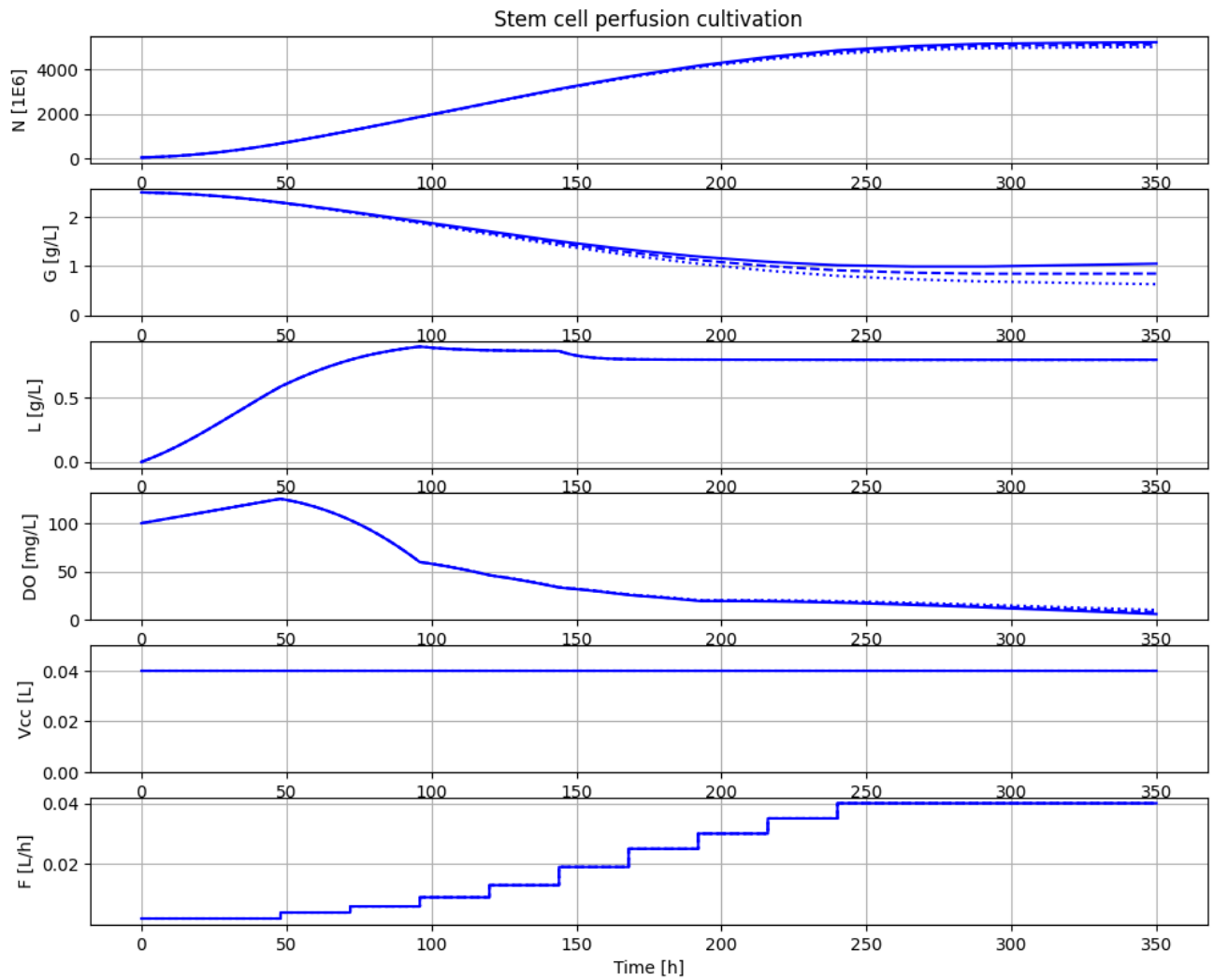
Here a process diagram is shown of the process. Further information about the culture stored in the model code is extracted.

```
#process_diagram()
```

✓ Simulations

```
# Process parameters to mimic stem cell example
# – study impact of different values maintenance qm
```

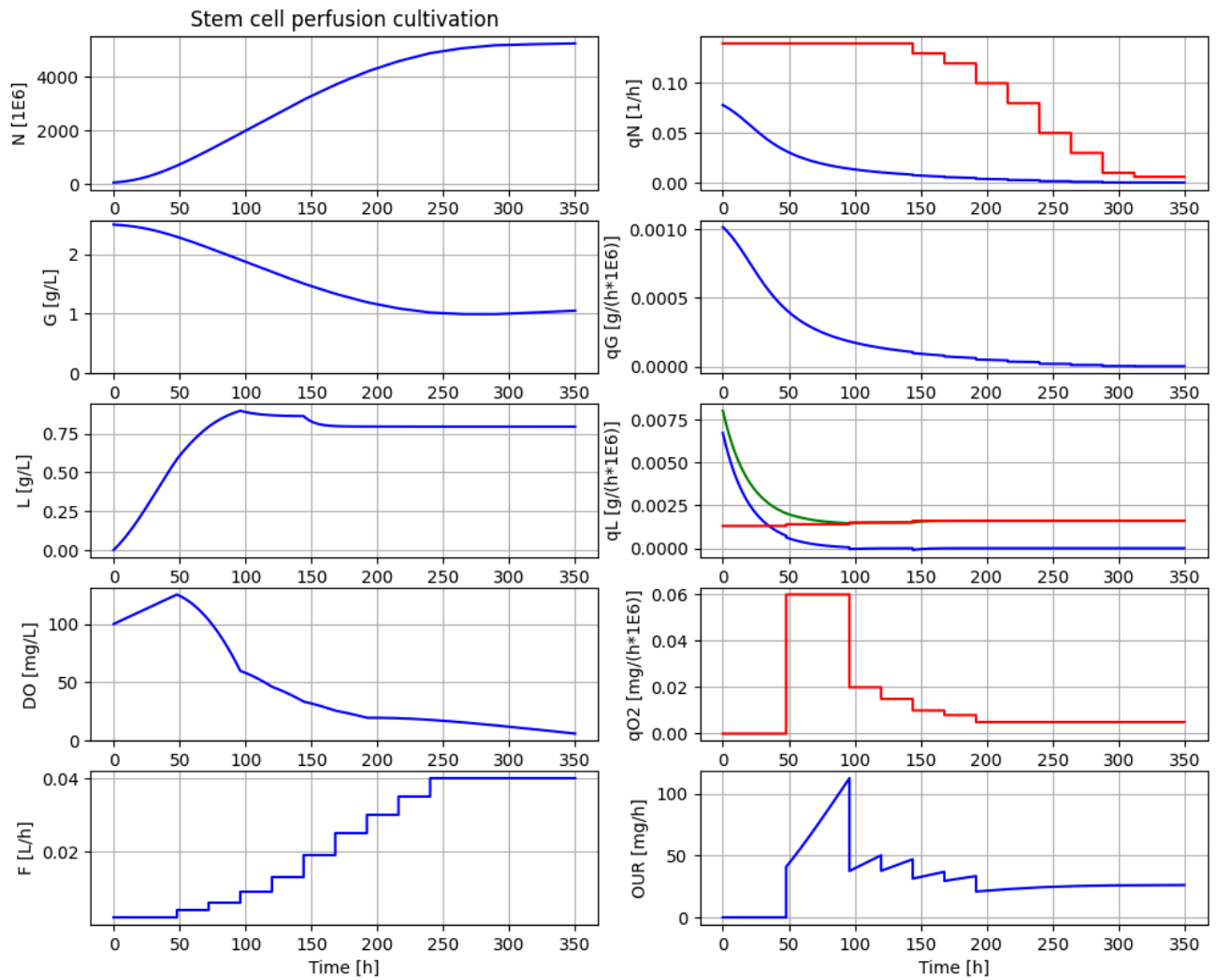
```
newplot(plotType='Basic')
for value in [1.0e-6, 10.0e-6, 20e-6]:
    par(qm=value)
    simu(350)
```



```
# Process parameters to mimic stem cell example
# . comprehensvie plot including metabolic rates
```

```
par(qm=1e-6)
```

```
newplot(plotType='Comprehensive')
simu(350)
```



```
describe('culture')
```



```
Reactor culture human-induced pluripotent stem cells - hiPSCs
```

```
describe('Vcc')
```



```
Volume of the reactor : 0.04 [ L ]
```

```
describe('N_start')
```

➦ Number of cells at start : 500 [1E6]

```
describe('N')
```

➦ Number of cells : 5223.752 [1E6]

```
describe('G_start')
```

➦ Glucose conc at start : 2.5 [g/L]

```
describe('G_in')
```

➦ Glucose feed conc : 2.5 [g/L]

```
describe('L')
```

➦ Lactate conc : 0.796 [g/L]

```
describe('D0')
```

➦ Dissolved oxygen conc : 6.044 [mg/L]

[+ Code](#)[+ Text](#)

✓ Appendix

```
system_info()
```



System information

-OS: Linux

-Python: 3.10.12

-Scipy: not installed in the notebook

-PyFMI: 2.14.0

-FMU by: OpenModelica Compiler OpenModelica 1.25.0~dev-51-ge672d09

-FMI: 2.0

-Type: FMUModelME2

-Name: BPL_STEM.Reactor

-Generated: 2024-10-03T11:45:51Z

-MSL: 3.2.3

-Description: BPL - not used

-Interaction: FMU-explore version 1.0.0

Start coding or [generate](#) with AI.