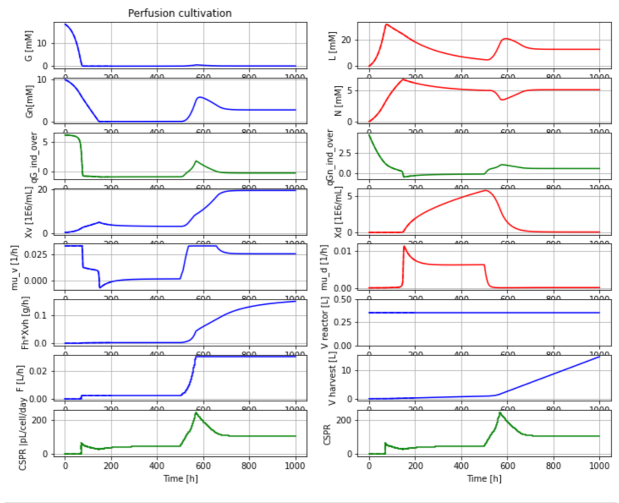
Notes: application CHO Perfusion cspr openloop

Here we run CHO Perfusion with cspr openloop for change of perfusion rate.

```
In [1]:
         run -i BPL_CHO_perfusion_cspr_openloop_explore.py
        Windows - run FMU pre-compiled JModelica 2.14
        Model for bioreactor has been setup. Key commands:
         - par() - change of parameters and initial values

    change initial values only

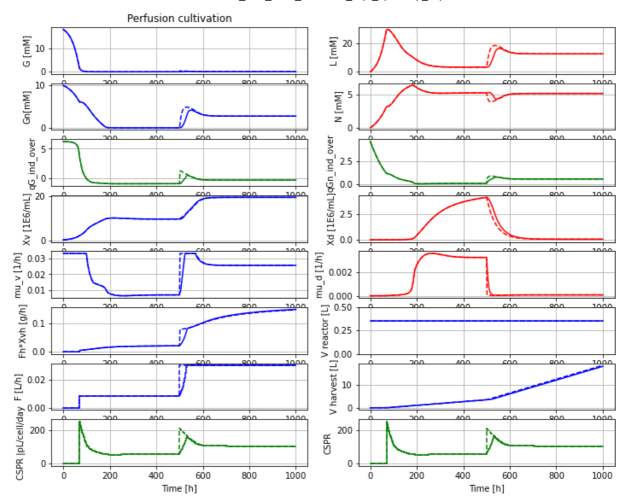
         - init()
         - simu() - simulate and plot
         - newplot() - make a new plot
         - show() - show plot from previous simulation - display parameters and initial value.
         - disp()
                      - display parameters and initial values from the last simulation
         - describe() - describe culture, broth, parameters, variables with values / units
        Note that both disp() and describe() takes values from the last simulation
        Brief information about a command by help(), eg help(simu)
        Key system information is listed with the command system_info()
In [2]:
         plt.rcParams['figure.figsize'] = [30/2.54, 24/2.54]
In [3]:
         describe('culture'); print(); describe('broth')
        Reactor culture CHO-MAb - cell line HB-58 American Culture Collection ATCC
        Reactor broth substances included in the model
        Cells viable index = 1 molecular weight = 24.6 Da
        Cells dead index = 2 molecular weight = 24.6 Da
        Glucose index = 3 molecular weight = 180.0 Da
        Glutamine index = 4 molecular weight = 146.1 Da
        Lactate index = 5 molecular weight = 90.1 Da
        Ammonia
                   index = 6 molecular weight = 17.0 Da
        Protein
                     index = 7 molecular weight = 150000.0 Da
In [4]:
         newplot('Perfusion cultivation', plotType='Extended')
         par(samplePeriod=1); par(F1=0.0020)
                                                            # General parameters
         par(mu_ref=0.04); simu()
                                                            # First simulation (solid)
         par(mu ref=1); simu(200)
                                                               # Second simulation (dashed)
```



```
In [5]: model.get('CSPR')
Out[5]: array([37.68370282])
```

We see here the shorter settling time for cell conc Xv when the slower exponential increase of perfusion rate is used, compared to an abrupt change.

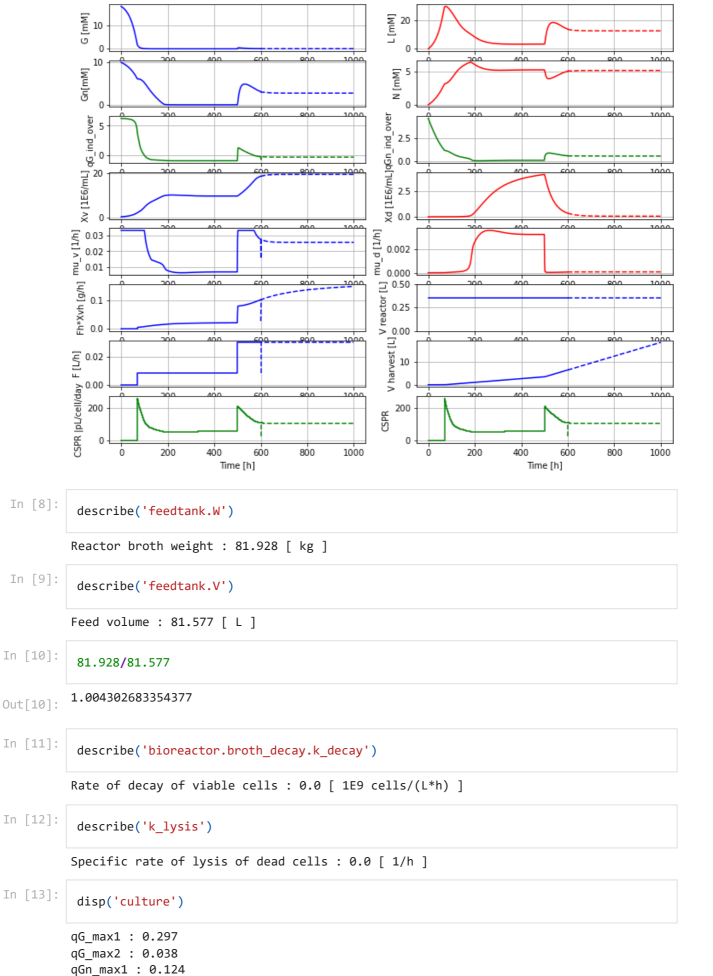
```
In [6]:
    newplot('Perfusion cultivation', plotType='Extended')
    par(samplePeriod=1); par(F1=0.0080)  # General parameters
    par(mu_ref=0.04); simu()  # First simulation (solid)
    par(mu_ref=1); simu()  # Second simulation (dashed)
```



We see that a somewhat smaller change in perfusion rate make the differnce in settling time of Xv for the two strategies to almost disappear.

```
newplot('Perfusion cultivation', plotType='Extended')
simu(600)
simu(400,'cont')
```

Perfusion cultivation



qGn_max2 : 0.022 mu_d_max : 0.13

```
In [14]: | describe('mu')
         Specific cell growth rate variable : 0.026 [ 1/h ]
In [15]:
          describe('parts')
         ['bioreactor', 'bioreactor.broth_decay', 'bioreactor.culture', 'CSPR', 'cspr_openloo
         p', 'D', 'feedtank', 'filter', 'harvesttank', 'liquidphase', 'MSL']
In [16]:
          describe('MSL')
         MSL: 3.2.2 build 3 - used components: RealInput, RealOutput
In [17]:
          system_info()
         System information
          -OS: Windows
          -Python: 3.9.5
          -Scipy: not installed in the notebook
          -PyFMI: 2.9.5
          -FMU by: JModelica.org
          -FMI: 2.0
          -Type: FMUModelCS2
          -Name: BPL_CHO.Perfusion_cspr_openloop
          -Generated: 2022-10-10T10:03:56
          -MSL: 3.2.2 build 3
          -Description: Bioprocess Library version 2.1.0
          -Interaction: FMU-explore ver 0.9.5
In [18]:
          model.get('Customer.name')
         ['Bioprocess Library']
Out[18]:
In [ ]:
```