# Modules, Linearisation and Retroactivity

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Note: this is the Retroactivity.ipynb notebook. The PDF version "Modules, Linearisation and Retroactivity" is available here.

### 1 Introduction

As discussed by (Gawthrop and Crampin, 2016): "Computational modularity is a necessary condition for building physically correct computational models of biomolecular systems. However, computational modularity does not imply that module properties (such as ultrasensitivity) are retained when a module is incorporated into a larger system. In the context of engineering, modules often have buffer amplifiers at the interface so that they have unidirectional connections and may thus be represented and analysed on a block diagram or signal flow graph where the properties of each module are retained. This will be called behavioural modularity in this paper. However, biological networks do not usually have this unidirectional property, but rather display retroactivity; retroactivity modifies the properties of the interacting modules."

The concept of retroactivity is used by, for example: (Saez-Rodriguez et al., 2004), (Saez-Rodriguez et al., 2005), (Del Vecchio and Murray, 2014) and (Del Vecchio et al., 2016).

This note looks at retroactivity in a simple linear case: a chain of reactions. The analysis is based on (Gawthrop and Crampin, 2016). The analysis uses Bond Graph Tools.

### 1.1 Import some python code

The bond graph analysis uses a number of Python modules:

```
In [1]: ## Some useful imports
        import BondGraphTools as bgt
        import numpy as np
        import sympy as sp
        import matplotlib.pyplot as plt
        import IPython.display as disp
        ## Stoichiometric analysis
        import stoich as st
        ## SVG bg representation conversion
        import svgBondGraph as sbg
        ## Modularity
        import modularBondGraph as mbg
        ## Control systems package
        import control as con
        ## Set quiet=False for verbose output
        quiet = True
        ## Set slycot=True if slycot is installed (see control module)
        slycot=True
```

```
## Chemical equation formatting
chemformula = True

import importlib as imp

## Allow output from within functions
from IPython.core.interactiveshell import InteractiveShell
InteractiveShell.ast_node_interactivity = "all"
```

### **2** A simple reaction network: $A \Leftrightarrow B \Leftrightarrow C$

The simple reaction network  $A \Leftrightarrow B \Leftrightarrow C$  can be concatenated to form a chain of N such networks. There are two approaches to this: - the modular bond graph approach using the chain() function from modularBondGraph and - Concatenate linearised subsystem approach using the Python Control Systems Library.

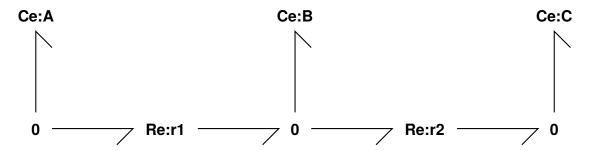
The species at each end of the chain are chemostats (Gawthrop and Crampin, 2016).

### 2.1 Bond graph model

```
In [2]: sbg.model('ABC_abg.svg')
    import ABC_abg
    disp.SVG('ABC_abg.svg')
    s = st.stoich(ABC_abg.model(),quiet=True)
    sc = st.statify(s,chemostats=['A','C'])

## Transfer function form (with chemostats)
    sys = st.lin(s,sc,outvar = 'port',quiet=quiet)
    con.ss2tf(sys)
```

#### Out[2]:



$$\begin{bmatrix} \frac{-s-1}{s+2} & \frac{1}{s+2} \\ \frac{1}{s+2} & \frac{-s-1}{s+2} \end{bmatrix}$$

### 2.2 Analysis of the chain of reactions

The function analyse() performs some basic analysis on the chain: - The poles and dc gain - The interaction loop-gain L from equation (86) of (Gawthrop and Crampin, 2016) - The step responses of the chain transfer function

Note that the chain inputs and outputs are determined by the chemostats at each end of the chain; thus the two inputs are the *potential* at each end and the two outputs are the coresponding flows.

```
In [3]: def analyse(sys,Sys,csys=None):
           """Analyse the linearised systems"""
           print('=======')
           print('Chain length N =',N)
           print('=======')
           ## Properties of the chain
           ## Note that poles are real; np.real removes spurious imag parts.
           print('Chain poles:',np.sort(np.real(con.pole(Sys))))
           print('Chain gain:',con.dcgain(Sys))
           if csys is not None:
               print('Analyse cys')
               csys_tf = con.ss2tf(csys)
               g_ef = csys_tf[0,0]
               g_fe = csys_tf[1,1]
               L = con.negate(con.series(g_ef,g_fe))
               print("L:",L)
           ## Step responses
           k = 0
           for i in range(2):
               for j in range(2):
                   k += 1
                   T = np.linspace(0,20,100)
                   tt,yy = con.step_response(Sys,input=j,output=i,T=T)
                   plt.subplot(2,2,k)
                   plt.plot(tt,yy)
                   plt.title('Step response '+str(i+1)+','+str(j+1))
                   plt.grid()
           plt.tight_layout()
           plt.show()
```

### 3 Modular bond graph approach

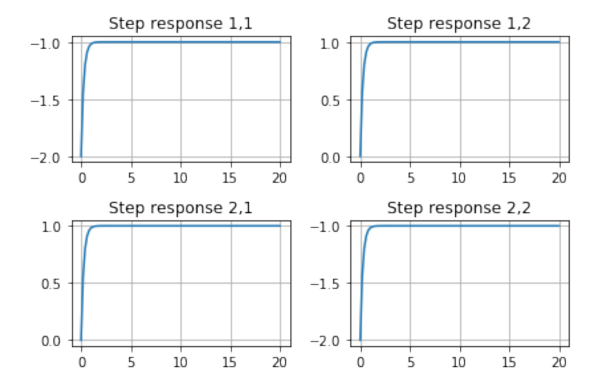
### 3.1 Create and analyse chain for various lengths N

```
In [4]: help(mbg.chain)
```

```
chain(model, inport='in', outport='out', N=2, quiet=False)
   Concatenate N instances of model via ports inport and outport
   The ports are represented in model as Ce components
   Ce:in in the first link of the chain and Ce:out in the last link remain as Ce components
   The method unifies Ce:out of link i and Ce:in of link i+1 by replacing them by ports
   and connecting them to a new Ce component with associated zero junction.
In [5]: NN = [1,2,5]
       for N in NN:
           ## Create chain
           NABC = mbg.chain(ABC_abg.model(),inport='A',outport='C',N=N,quiet=quiet)
           s = st.stoich(NABC,quiet=True)
           #print(s['reaction'])
           sc = st.statify(s,chemostats=['A','C'])
           #disp.Latex(st.sprintrl(sc,chemformula=chemformula))
           ## Set each Re component to have kappa = 2
           parameter = {}
           for r in s['reaction']:
               parameter['kappa_'+r] = 2
           ## Linearise the chain
           Sys = st.lin(s,sc,outvar = 'port',parameter=parameter,quiet=quiet)
           ## Analyse the properties
           analyse(sys,Sys)
           print('Chain transfer function:')
           con.ss2tf(Sys)
Chain length N = 1
_____
Chain poles: [-4.]
Chain gain: [[-1. 1.]
```

Help on function chain in module modularBondGraph:

[1. -1.]



Out[5]:

 $\begin{bmatrix} \frac{-2s-4}{s+4} & \frac{4}{s+4} \\ \frac{4}{s+4} & \frac{-2s-4}{s+4} \end{bmatrix}$ 

\_\_\_\_\_

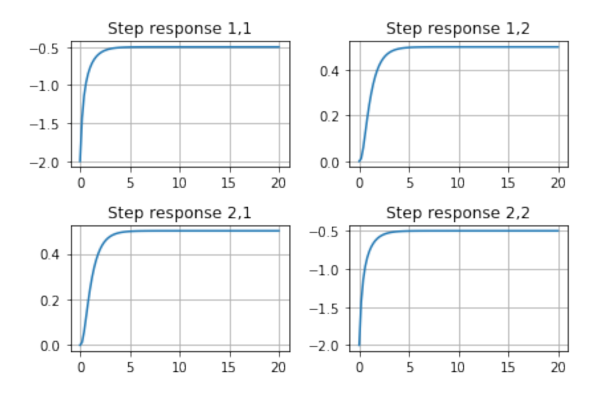
Chain length N = 2

\_\_\_\_\_

Chain poles: [-6.82842712 -4. -1.17157288]

Chain gain: [[-0.5 0.5]

[ 0.5 -0.5]]



### Out[5]:

$$\begin{bmatrix} \frac{-2s^3 - 20s^2 - 48s - 16}{s^3 + 12s^2 + 40s + 32} & \frac{16}{s^3 + 12s^2 + 40s + 32} \\ \frac{16}{s^3 + 12s^2 + 40s + 32} & \frac{-2s^3 - 20s^2 - 48s - 16}{s^3 + 12s^2 + 40s + 32} \end{bmatrix}$$

\_\_\_\_\_

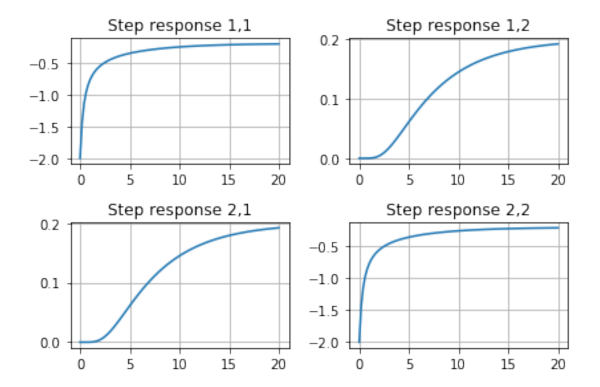
Chain length N = 5

\_\_\_\_\_

-1.64885899 -0.76393202 -0.19577393]

Chain gain: [[-0.2 0.2]

[ 0.2 -0.2]]



### Out[5]:

```
\begin{bmatrix} \frac{-2s^9 - 68s^8 - 960s^7 - 7280s^6 - 3.203e + 04s^5 - 8.237e + 04s^4 - 1.183e + 05s^3 - 8.448e + 04s^2 - 2.304e + 04s - 1024 \\ \frac{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120 \\ 1024 \\ \frac{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120}{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120} \\ \frac{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120}{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120} \\ \frac{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120}{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120} \\ \frac{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 1.098e + 05s^3 + 1.014e + 05s^2 + 4.224e + 04s + 5120}{s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406e + 04s^4 + 0.084e + 0.
```

#### 3.2 Discussion

- The overall step response from the start to the end of the chain (step response 2,1 from  $u_1$  to  $v_2$ ) becomes higher-order and slower as N increases.
- Because of retroactivity, the steady-state (DC) gain of the chain is *not* the product of the gain of each link it is 1/N in this case.
- Because of retroactivity, the poles of the chain are (except for the pole at s=4) different for each N

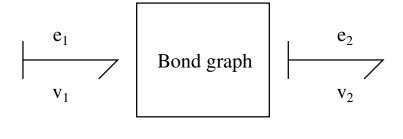
### 4 Concatenate linearised subsystem approach

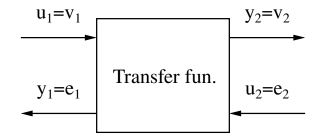
As discussed by (Gawthrop and Crampin, 2016), and as outlined in the following figure: a bond graph representing a linear system can be represented as a transfer function when causality has been asigned. Thus, in the example in the figure, a bond graph with two impinging bonds and causality assigned as shown, corresponds to a  $2 \times 2$  transfer function matrix so that:

$$\begin{pmatrix} y_1 \\ y_2 \end{pmatrix} = \begin{pmatrix} g_{11} & g_{12} \\ g_{21} & g_{22} \end{pmatrix} \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} \tag{1}$$

As in the modular bond-graph approach, a Ce component is put beween each link in the chain to unify the appropriate Ce components: - The system *sys* corresponds to the bond graph ABG\_abg above with **Ce:A** and **Ce:C** replaced by ports - The system *csys* is *sys* with a **Ce** component prepended.

In [6]: disp.SVG('notation.svg')
Out[6]:





### 4.1 Linearise the subsystem

### 4.2 Create linear C component in control toolbox form

### 4.3 Create chain of linearised modules

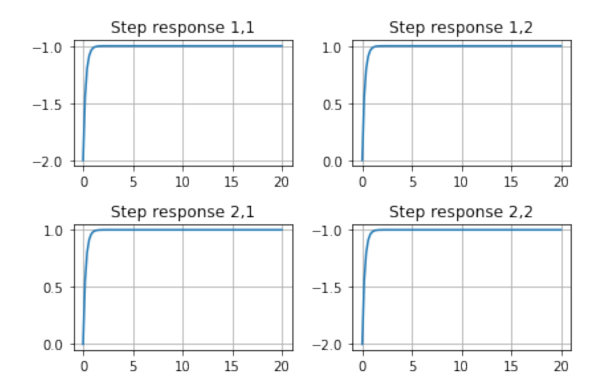
```
In [11]: def makeChain(sys,C,N=2):
    """Make the chain of modules"""

## Prepend C component to sys
    CSYS = con.append(C,sys)
    Q = np.array([[2,3],[3,2]])
    inout = np.array([1,4])
    csys = con.connect(CSYS,Q,inout,inout)

## Recursively create chain
    Sys = sys
    for i in range(N-1):
        SYS = con.append(Sys,csys)
        Sys = con.connect(SYS,Q,inout,inout)

return Sys,csys
```

```
Sys,csys = makeChain(sys,makeC(Retroactive=True),N=2)
       con.ss2tf(csys)
       con.ss2tf(Sys)
       con.pole(Sys)
  Out[11]:
  Out[11]:
Out[11]: array([-6.82842712, -4.
                                   , -1.17157288])
4.4 Analyse with and without retroactivity
In [12]: for Retroactive in [True,False]:
           for N in NN:
              print('======"')
              print('Retroactive =',Retroactive)
              print('=======')
              C = makeC(Retroactive=Retroactive)
              Sys,csys = makeChain(sys,C,N=N)
              analyse(sys,Sys,csys=csys)
              print('Chain transfer function:')
              con.ss2tf(Sys)
_____
Retroactive = True
_____
Chain length N = 1
Chain poles: [-4.]
Chain gain: [[-1. 1.]
[1. -1.]
Analyse cys
L:
     2 s^3 + 16 s^2 + 32 s
s^4 + 12 s^3 + 44 s^2 + 48 s + 16
```



### Out[12]:

 $\begin{bmatrix} \frac{-2s-4}{s+4} & \frac{4}{s+4} \\ \frac{4}{s+4} & \frac{-2s-4}{s+4} \end{bmatrix}$ 

\_\_\_\_\_

Retroactive = True

\_\_\_\_\_

Chain length N = 2

Chain poles: [-6.82842712 -4. -1.17157288]

Chain gain: [[-0.5 0.5]

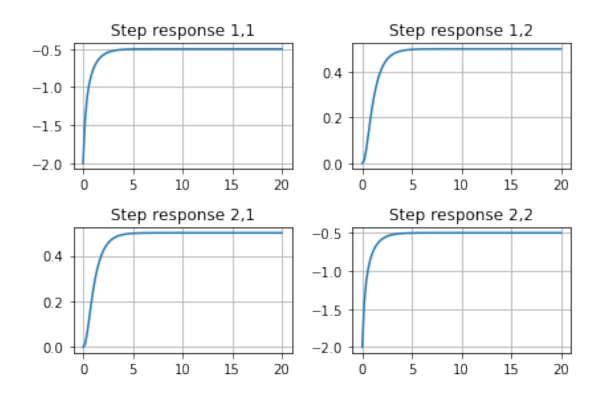
[ 0.5 -0.5]] Analyse cys

т.

2 s^3 + 16 s^2 + 32 s

-----

 $s^4 + 12 s^3 + 44 s^2 + 48 s + 16$ 



### Out[12]:

$$\begin{bmatrix} \frac{-2s^3 - 20s^2 - 48s - 16}{s^3 + 12s^2 + 40s + 32} & \frac{16}{s^3 + 12s^2 + 40s + 32} \\ \frac{16}{s^3 + 12s^2 + 40s + 32} & \frac{-2s^3 - 20s^2 - 48s - 16}{s^3 + 12s^2 + 40s + 32} \end{bmatrix}$$

Retroactive = True

\_\_\_\_\_

\_\_\_\_\_

Chain length N = 5

Chain poles: [-7.80422607 -7.23606798 -6.35114101 -5.23606798 -4. -2.76393202

-1.64885899 -0.76393202 -0.19577393]

Chain gain: [[-0.2 0.2]

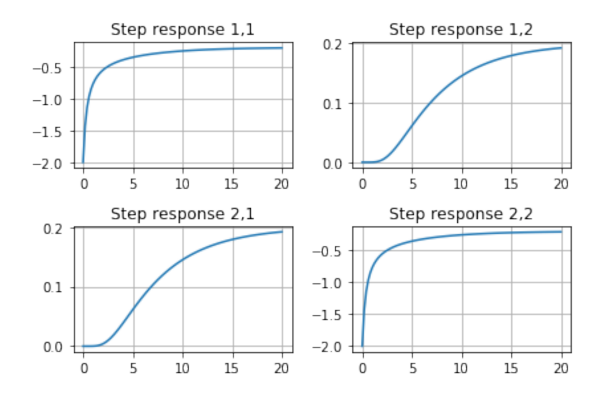
[ 0.2 -0.2]]

Analyse cys

L:

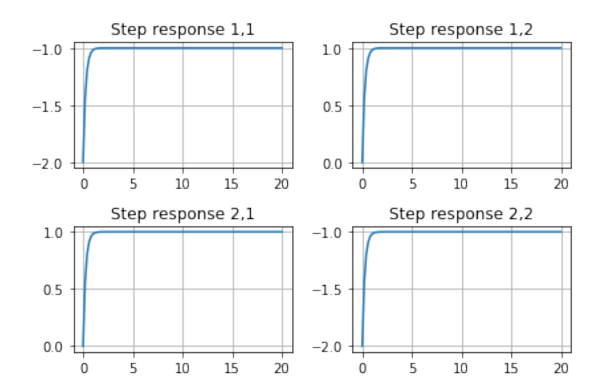
$$2 s^3 + 16 s^2 + 32 s$$

s^4 + 12 s^3 + 44 s^2 + 48 s + 16



### Out[12]:

 $\begin{array}{l} s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406 \\ -2s^9 - 68s^8 - 960s^7 - 7280s^6 - 3.203e + 04s^5 - 8.2 \\ s^9 + 36s^8 + 544s^7 + 4480s^6 + 2.184e + 04s^5 + 6.406 \end{array}$ 



### Out[12]:

 $\begin{bmatrix} \frac{-2s-4}{s+4} & \frac{4}{s+4} \\ \frac{4}{s+4} & \frac{-2s-4}{s+4} \end{bmatrix}$ 

\_\_\_\_\_

Retroactive = False

\_\_\_\_\_

Chain length N = 2

Chain poles: [-5.23606798 -4. -0.76393202]

Chain gain: [[-1. 0.]

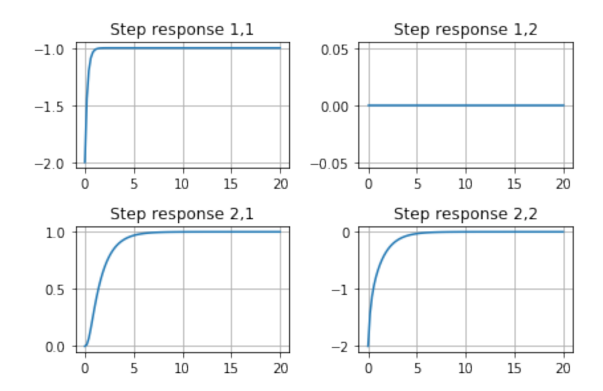
[ 1. 0.]]

Analyse cys

L: 0

\_

1



### Out[12]:

$$\begin{bmatrix} \frac{-2s^3 - 4s^2}{s^3 + 4s^2} & 0\\ \frac{16}{s^3 + 10s^2 + 28s + 16} & \frac{-2s^3 - 16s^2 - 32s}{s^3 + 10s^2 + 28s + 16} \end{bmatrix}$$

Retroactive = False

\_\_\_\_\_

\_\_\_\_\_

Chain length N = 5

\_\_\_\_\_

 $\hbox{Chain poles: $[-5.23607422 -5.23607422 -5.23606798 -5.23605549 -4. } \\ -0.76393527$ 

-0.76393527 -0.76393202 -0.76392553]

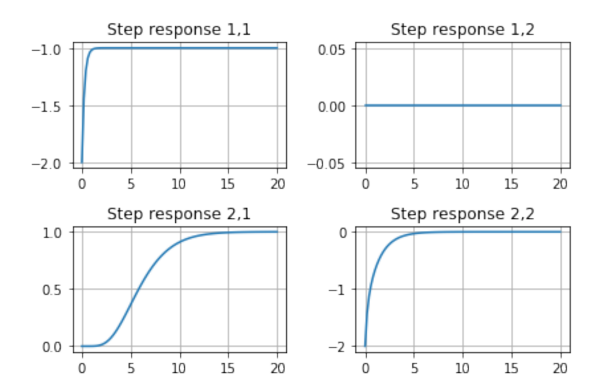
Chain gain: [[-1. 0.]

[ 1. 0.]]

Analyse cys

L: 0

1



### Out[12]:

```
\begin{bmatrix} \frac{-2s^9-4s^8}{s^9+4s^8} & 0 \\ \frac{1024}{s^9+28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e+04s^2+6400s+1024} \\ \frac{-2s^9-52s^8-560s^7-3216s^6-1.056e+04s^5-1.978e+04s^4-2.018e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e+04s^2+6400s+1024} \\ \frac{-2s^9-52s^8-560s^7-3216s^6-1.056e+04s^5-1.978e+04s^4-2.018e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^7+2080s^6+7728s^5+1.709e+04s^4+2.214e+04s^3+1.638e^{-28s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8+328s^8
```

### 4.5 Compute product of transfer functions

```
4
s + 4
 _____
Chain length N = 2
                                          16
_____
s^3 + 10 s^2 + 28 s + 16
_____
Chain length N = 3
_____
                                                                                         64
s^5 + 16 s^4 + 92 s^3 + 224 s^2 + 208 s + 64
_____
Chain length N = 4
_____
                                                                                                                                              256
s^7 + 22 s^6 + 192 s^5 + 840 s^4 + 1920 s^3 + 2208 s^2 + 1216 s + 256
_____
Chain length N = 5
_____
                                                                                                                                                                                                                                1024
s^9 + 28 s^8 + 328 s^7 + 2080 s^6 + 7728 s^5 + 1.709e + 04 s^4 + 2.214e + 04 s^3 + 1.638e + 04 s^2 + 68 to 100 t
```

#### 4.6 Discussion

- When the linear modules are concatenated using the standard **C** component, the properties of the chain are, as expected, the same as those from the modular BG approach
- When the non-standard **C** component is used to avoid retroactivity, chain shows behavioral modularity
- the steady-state (DC) gain of the chain is the product of the gain of each link
- the poles of the chain are the same for each N
- this is verified by directly computing the product of transfer functions
- But the use of this non-standard C implies the use of an amplifier and therfore consumes

power.

• The *interaction loop gain L* is discussed by (Gawthrop and Crampin, 2016). If L = 0 there no retroactive interaction between the links of the chain.

### References

- Domitilla Del Vecchio and Richard M Murray. *Biomolecular Feedback Systems*. Princeton University Press, 2014. ISBN 0691161534.
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