Sensitivity Analysis of Biochemical Systems Using Bond Graphs: Additional Material – Illustrative Examples

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1 Introduction

This notebook generates the figures for the paper: Sensitivity Analysis of Biochemical Systems Using Bond Graphs. The companion notebook Sensitivity_PPP.ipynb contains the Pentose Phosphate Pathway example.

2 Supporting software

2.1 Import packages

```
[1]: ## Some useful imports
  import BondGraphTools as bgt
  import numpy as np
  import sympy as sp
  import matplotlib.pyplot as plt
  import control as con
  import copy
  import importlib

## For reimporting: use imp.reload(module)
  import importlib as imp

## Stoichiometric analysis
  import stoich as st

## SVG
```

```
import svgBondGraph as sbg
## Stoichiometry to BG
import stoichBondGraph as stbg
## Modularity
# import modular as bgm
## Sloppy parameters
import sloppy as slp
## Display (eq disp.SVG(), disp.
import IPython.display as disp
##
quiet = True
Plotting = False
Titles = not Plotting
grid = False
fontsize=20
## Normalisation
Normalise = True
## Use lambda as system input or just perturb parameter
lambdaIn = True
```

Warning - scikit.odes not found. Simulations are disabled.

2.2 Transfer function properties

```
[2]: def tfProps(tf,method='truncate'):
         ## Steady-state gain
         g = con.dcgain(tf)
         if not con.issiso(tf):
             g = g[0][0]
         ## Time constant
         ## Check if direct link
         direct_link = np.any(tf.D)
         ## Only set tau=0 if siso
         if direct_link and con.issiso(tf):
             ## Instant response
             tau = 0
         else:
             ## Reduce to first-order to estimate time constant
             ## Note that method='matchdc' can give a kernel crash - use 'truncate'
             tf1 = con.balred(sys,orders=1,method=method)
```

```
poles = con.poles(tf1)
  realPoles = np.real(poles)
  tau = -1/min(realPoles)

return g,tau
```

2.3 Plotting and printing

```
[3]: ## Optional plotting
def Savefig(name):
    if Plotting:
        plt.rcParams.update({'font.size': fontsize})
        plt.tight_layout()
        plotname = 'Figs/'+name+'.pdf'
        print('Saving',plotname)
        plt.savefig(plotname)
```

```
[4]: def latex(m,name):
         lm = sp.latex(sp.Matrix(m),mat_delim="(")
         return name+' &= '+lm
     ## Optional print latex for the paper
     printing = True
     def printLatex(s,sc=None):
         if printing:
             ## System properties in LaTeX
             AA = Sys.A; print(latex(AA,'A'))
             BB = Sys.B; print(latex(BB, 'B'))
             CC = Sys.C; print(latex(CC, 'C'))
             DD = Sys.D; print(latex(DD, 'D'))
             for m in ['species', 'reaction', 'Nf', 'Nr', 'N']:
                 print(st.sprintl(s,m))
             if not (sc==None):
                 print('%% Nc matrix')
                 print(st.sprintl(sc,'N'))
             print(st.sprintrl(s,all=True,chemformula=True))
             print(st.sprintvl(s))
```

2.4 Steady-state by simulation

```
[5]: ## Steady-state by simulation
def SteadyState(s,sc,sf=None,parameter={},X0=None,t_ss=1000):
    t = np.linspace(0,t_ss)
    dat = st.

→sim(s,sc=sc,sf=sf,t=t,parameter=parameter,X0=X0,tol=1e-4,quiet=quiet)
    X_ss = dat['X'][-1]
```

```
V_ss = dat['V'][-1]
#dX_ss = sc['N']@V_ss
dX_ss = dat['dX'][-1]
dXc_ss = dat['dXc'][-1]
print(f'Steady-state finder error: {np.linalg.norm(dXc_ss):.2e}')
return X_ss,V_ss
```

2.5 Stoichiometry

```
[6]: def_
     →Stoichiometry(model,chemostats=[],flowstats=[],CommonSpecies=None,sensitivity=False):
         ## Stoichiometry
         s = st.stoich(model,quiet=quiet)
         ## Unify species
         if not (CommonSpecies is None):
             commonSpecies = st.merge(s,CommonSpecies=CommonSpecies)
               print(commonSpecies)
             st.unify(s,commonSpecies=commonSpecies)
         ## Sensitivity
         if sensitivity:
             extra = st.stoichSensitivity(s)
         else:
             extra = []
         print(chemostats+extra)
         ## Chemostats and flowstats
         sc = st.statify(s,chemostats=chemostats+extra)
         sf = st.statify(s,flowstats=flowstats)
         return s,sc,sf
```

2.6 Linearisation

2.7 Extract subsystem from linear system

```
[9]: def Index(A,a):
         I = \Gamma
         for aa in a:
             i = A.index(aa)
             I.append(i)
         return np.array(I)
     def zapSmall(x,tol=1e-10,quiet=True):
         xx = np.zeros(len(x))
         for i,val in enumerate(x):
             if abs(val)>tol:
                 xx[i] = x[i]
             else:
                 if not quiet:
                     print(f'Setting {i}th coefficient {val:.2} to zero')
         return xx
     def⊔
      →extractSubsystem(SYS,sc,sf,inp,outp,minreal=False,tol=None,order=None,quiet=False):
```

```
Sys = copy.copy(SYS)
chemostats = sc['chemostats']
if sf is None:
    flowstats = []
else:
    flowstats = sf['flowstats']
species = sc['species']
reaction = sc['reaction']
## Index of input and output
if inp[0] in chemostats:
    i_inp = Index(chemostats,inp)
      print('Input:',i_inp,chemostats[i_inp[0]])
else:
    i_inp = Index(flowstats,inp)+len(chemostats)
 print(i_inp)
if outp[0] in chemostats:
    i_outp = Index(chemostats,outp)
elif outp[0] in species:
    i_outp = Index(species,outp)
else:
    if outp[0] in reaction:
        i_outp = Index(reaction,outp)
    else:
        print(f'Output {outp} does not exist')
## Extract tf
n_y = len(i_outp)
n_u = len(i_inp)
nn = Sys.A.shape
n_x = nn[0]
 print(n_x)
sys = con.ss(Sys.A,
             Sys.B[:,i_inp].reshape(n_x,n_u),
             Sys.C[i_outp,:].reshape(n_y,n_x),
             Sys.D[i_outp][:,i_inp].reshape(n_y,n_u))
    sys = con.minreal(sys,tol=tol,verbose=False)
## Reduce order
if not (order is None):
    sys = con.balred(sys,order,method='matchdc')
return sys
```

Plotting

```
[10]: def_
       →plotSensitivity(dat,reactions=['r1','r2'],plotSim=True,name=None,grid=False):
          ## Folt
          plt.rcParams.update({'font.size': fontsize})
          if plotSim:
              plt.plot(t,y_lin,color='black',lw=3)
          else:
              plt.plot(t,y_lin,label=label,lw=3)
          for reac in reactions:
              i = s['reaction'].index(reac)
              if plotSim:
                  label = reac + r' (\frac{\pi}{1} (\frac{\pi}{1})'
                  pc = int(round(100*(lam-1)))
                  label = f'{reac} ({pc}%)'
                  plt.plot(t,(dat['V'][:,i]-V_ss[i])/(lam-1),
                           lw=5,ls='dashed',label=label)
          if grid:
             plt.grid()
          plt.legend()
         plt.xlabel(r'$t$')
         plt.ylabel(r'$\tilde{f}/\tilde{\lambda}$')
           plt.tight_layout()
[11]: def plotLines(lw=4,ls='dotted'):
              plt.hlines(g,min(t),max(t),color='black',ls=ls,lw=lw)
             plt.vlines(tau,min(y_lin.flatten()),max(y_lin.
       →flatten()),color='black',ls=ls,lw=lw)
```

Normalisation constants 2.9

```
[12]: T_human = 37 # Human body temperature
      K_0 = 273.15
      print(f'T_human = {T_human} degC = {T_human+K_0} K')
      mu_0 = RT = st.RT(T_cent=T_human)
      print(f'mu_0 = {mu_0*1e-3:0.3f} kJ/mol')
      F = st.F() # Faraday's constant
      print(f'F = \{F*1e-3:0.2f\} kC/mol')
      V_0 = RT/F
      print(f'V_0 = {V_0*1e3:0.2f} mV')
      P_0 = 1e-3
```

```
v_0 = P_0/mu_0
print(f'v_0 = {v_0*1e6:0.4f} micro mol /s')

i_0 = F*v_0
print(f'i_0 = {i_0*1e3:0.2f} mA')
```

```
T_human = 37 degC = 310.15 K

mu_0 = 2.579 kJ/mol

F = 96.49 kC/mol

V_0 = 26.73 mV

v_0 = 0.3878 micro mol /s

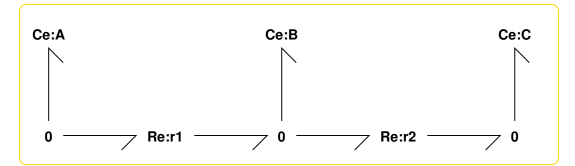
i_0 = 37.42 mA
```

3 Simple example A = B = C revisited.

3.1 Bond Graph

```
[13]: # Simple example A = B = C
sbg.model('ABC_abg.svg')
import ABC_abg
disp.SVG('ABC_abg.svg')
```

[13]:



```
[14]: ## BG generated equations
    # model = ABC_abg.model()
    # for cr in model.constitutive_relations:
    # print(cr)
    # #print(sp.diff(cr,'x_0'))
```

3.2 Parameters

```
[15]: ## Parameters
parameter = {}
parameter['K_A'] = 1
parameter['K_B'] = 1
parameter['kappa_r1'] = 1
parameter['kappa_r2'] = 9
print(parameter)
```

```
## Initial states
      X_A_0 = 2
     {'K_A': 1, 'K_B': 1, 'kappa_r1': 1, 'kappa_r2': 9}
          Stoichiometry & linearisation
[16]: ## Stoichiometry
      s,sc,sf = Stoichiometry(ABC_abg.model(),chemostats=['A','C'],flowstats=[])
[17]: ## Linearise
      Parameter = copy.copy(parameter)
      Parameter['K_A'] = X_A_0*parameter['K_A']
      Sys, X_ss, V_ss, dX_ss =
      →Linear(s,sc,parameter=parameter,X0=[X_A_0,1,1],quiet=quiet)
      i_A = s['species'].index('A')
      X_A_ss = X_ss[i_A]
      ## Show transfer function
      con.tf(Sys)
      ## Lambda for comparison
      lam = 1.1
     Steady-state finder error: 8.88e-16
     X_ss = [2. 1.11.]
     V_ss = [0.9 \ 0.9]
[18]: printLatex(s,sc=sc)
     A &= \left(\begin{matrix}-10.0\end{matrix}\right)
     B &= \left(\begin{matrix}1.0 & 9.0\end{matrix}\right)
     C &= \left(\begin{matrix}-1.0\\9.0\end{matrix}\right)
     D &= \left(\frac{matrix}{1.0 \& 0}\  \  -9.0\right)
     \begin{align}
     X&= \begin{pmatrix}
         X_{A}\
         X_{B}\
         X_{C}\
     \end{pmatrix}
     \end{align}
     \begin{align}
     V&= \begin{pmatrix}
         V_{r1}\\
         V_{r2}\\
     \end{pmatrix}
     \end{align}
     \begin{align}
```

```
\left(\left(\frac{n}{matrix}\right) \& 0\0 \& 1\0 \& 0\end{matrix}\right)
     \end{align}
     \begin{align}
     Nr &=
     \left(\left(\frac{matrix}{0 \& 0}\right) \& 1\right) 
     \end{align}
     \begin{align}
     N &=
     \left(\left(\frac{matrix}-1 \& 0\right)^2 \& -1\right)^2 \& 1\left(\frac{matrix}\right)^2
     \end{align}
     %% Nc matrix
     \begin{align}
     N &=
     \label{left(begin{matrix}0 & 0\\1 & -1\\0 & 0\\matrix\\right)
     \end{align}
     \begin{align}
     \ch{A & <> [ r1 ] B }\\
     \ch{B & <> [ r2 ] C }
     \end{align}
     \begin{align}
     v_{r1} \&= \kappa_{r1} \left(K_{A} x_{A} - K_{B} x_{B}\right)\
     v_{r2} &= \kappa_{r2} \left(K_{B} x_{B} - K_{C} x_{C}\right)
     \end{align}
[19]: | ## Show dc gain
      print('DC gain: \n', con.dcgain(Sys))
     DC gain:
      [[0.9 - 0.9]
      [0.9 - 0.9]
           Sensitivity Bond Graph – change chemostats
[20]: ## Extract sensitivity system
      inp = ['A']
      outp = ['r1','r2']
      sys = extractSubsystem(Sys,sc,sf,inp,outp)
      ## Include factor X_A_O into the linarised system
      sys.B = sys.B*X_A_0
      sys.D = sys.D*X_A_0
[21]: ## Show transfer function
      con.tf(sys)
```

Nf &=

```
[21]:
```

$$\begin{bmatrix} \frac{2s+18}{s+10} \\ \frac{18}{s+10} \end{bmatrix}$$

[22]: ## System matrix sys

[22]:

$$\begin{pmatrix} -10 & 2 \\ -1 & 2 \\ 9 & 0 \end{pmatrix}$$

[23]: ## Show dc gain print('DC gain: \n', con.dcgain(sys))

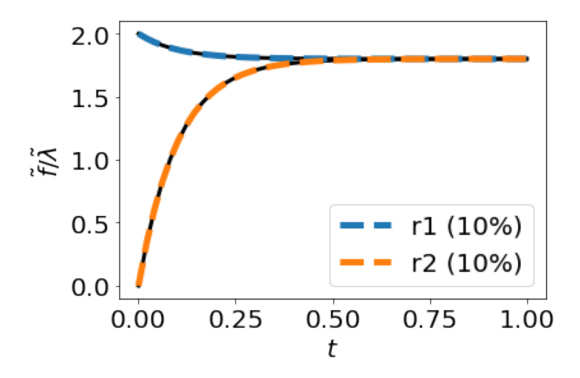
DC gain: [[1.8] [1.8]]

3.4.1 Compare sensitivity with exact simulation

```
[24]: ## Exact Simulate with changed x_A
X_ss_1 = copy.copy(X_ss)
X_ss_1[i_A] = lam*X_A_ss
print(X_ss)
t = np.linspace(0,1)
dat = st.sim(s,sc=sc,t=t,parameter=parameter,X0=X_ss_1,quiet=quiet)
print(parameter)
```

```
[2. 1.1 1.] {'K_A': 1, 'K_B': 1, 'kappa_r1': 1, 'kappa_r2': 9}
```

- [25]: ## Step response to change in A
 y_lin = linStep(sc,sys,T=t)
- [26]: ## Plot
 plotSensitivity(dat,)



3.5 Sensitivity Bond Graph – change K_B

```
[27]: # Simple example A = B = C: sensitivity
sbg.model('sKABC_abg.svg')
import sKABC_abg
disp.SVG('sKABC_abg.svg')
```

Ce:B

Ce:A

Ce:C

Re:r1

Ce:B

3.5.1 Stoichiometry & linearisation

```
[28]: ## Stoichiometry
     s,sc,sf = Stoichiometry(sKABC_abg.
      →model(),chemostats=['A','C','sB'],flowstats=[])
      ## Linearise
     Sys, X_ss, V_ss, dX_ss =
      →Linear(s,sc,parameter=parameter,X0=[2,1,1,1],quiet=quiet)
      ## Extract sensitivity system
     inp = ['sB']
     outp = ['r1', 'r2']
     sys = extractSubsystem(Sys,sc,sf,inp,outp)
     ## Species and reactions
     print(s['species'])
     print(s['reaction'])
     Steady-state finder error: 8.88e-16
     X_ss = [2. 1.11. 1.]
     V_ss = [0.9 \ 0.9]
     ['A', 'B', 'C', 'sB']
     ['r1', 'r2']
[29]: ## Show transfer function
     con.tf(sys)
[29]:
[30]:
[31]: | ## Show dc qain
     print('DC gain: \n', con.dcgain(sys))
     DC gain:
      [[0.]
      [0.]]
```

3.5.2 Compare sensitivity with exact simulation

```
[32]: ## Exact Simulate with changed K
     parameter1 = copy.copy(parameter)
     parameter1['K_B'] *= lam
     t = np.linspace(0,1)
     dat = st.sim(s,sc=sc,t=t,parameter=parameter1,X0=X_ss,quiet=quiet)
[33]: ## Step response to sB
     y_lin = linStep(sc,sys,T=t)
[34]: ## Plot
     plotSensitivity(dat,)
              10.0
                                                          r1 (10%)
                7.5
                                                          r2 (10%)
                5.0
                2.5
                0.0
```

3.6 Sensitivity Bond Graph – change κ_1

0.00

```
[35]: # Simple example A = B = C: sensitivity
sbg.model('sKapABC_abg.svg')
import sKapABC_abg
disp.SVG('sKapABC_abg.svg')
```

0.25

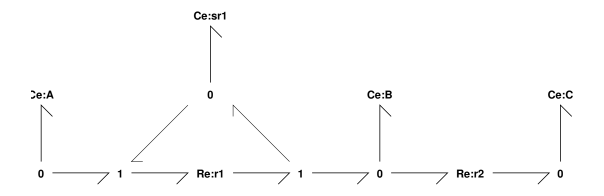
0.50

t

0.75

1.00

[35]:



```
[36]: ## BG generated equations
      model = sKapABC_abg.model()
      for cr in model.constitutive_relations:
          print(cr)
          #print(sp.diff(cr,'x_0'))
```

```
K_A*K_sr1*kappa_r1*x_0*x_3 - K_B*K_sr1*kappa_r1*x_1*x_3 + dx_0
-K_A*K_sr1*kappa_r1*x_0*x_3 + K_B*K_sr1*kappa_r1*x_1*x_3 + K_B*kappa_r2*x_1 -
K_C*kappa_r2*x_2 + dx_1
-K_B*kappa_r2*x_1 + K_C*kappa_r2*x_2 + dx_2
dx_3
```

Stoichiometry & linearisation 3.6.1

Steady-state finder error: 8.88e-16

 $X_ss = [2. 1.11. 1.]$

 $V_ss = [0.9 \ 0.9]$

[37]:

```
[37]: print(s['reaction'])
      ## Stoichiometry
      s,sc,sf = Stoichiometry(sKapABC_abg.model(),chemostats=['A','C','sr1'])
      print(sf['flowstats'])
      ## Linearise
      Sys, X_ss, V_ss, dX_ss =
      →Linear(s,sc,parameter=parameter,X0=[2,1,1,1],quiet=quiet)
      ## Extract sensitivity system
      inp = ['sr1']
      outp = ['r1','r2']
      sys = extractSubsystem(Sys,sc,sf,inp,outp)
      ## Show transfer function
      con.tf(sys)
     ['r1', 'r2']
```

$$\begin{bmatrix} \frac{0.9s + 8.1}{s + 10} \\ \frac{8.1}{s + 10} \end{bmatrix}$$

```
[38]: ## System matrix sys
```

[38]:

$$\begin{pmatrix} -10 & 0.9 \\ -1 & 0.9 \\ 9 & 0 \end{pmatrix}$$

```
[39]: ## Show dc gain
print('DC gain: \n', con.dcgain(sys))
```

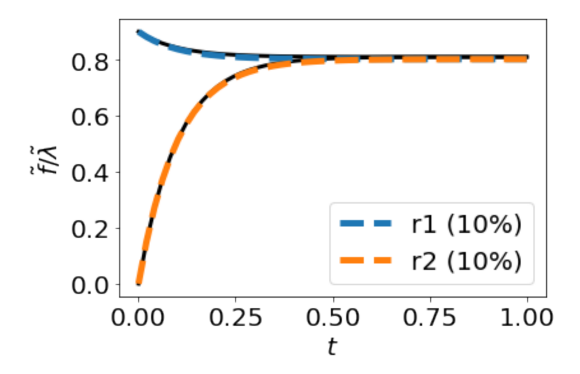
DC gain: [[0.81] [0.81]]

3.6.2 Compare sensitivity with exact simulation

```
[40]: ## Exact Simulate with changed K
lam = 1.1
parameter1 = copy.copy(parameter)
parameter1['kappa_r1'] *= lam
t = np.linspace(0,1)
dat = st.sim(s,sc=sc,sf=sf,t=t,parameter=parameter1,X0=X_ss,quiet=quiet)
```

```
[41]: ## Step response to sB
y_lin = linStep(sc,sys,T=t)
```

```
[42]: ## Plot plotSensitivity(dat,)
```



[43]: sys

[43]:

$$\begin{pmatrix} -10 & 0.9 \\ -1 & 0.9 \\ 9 & 0 \end{pmatrix}$$

[44]: disp.Latex(st.sprintvl(s))

[44]:

$$v_{r1} = K_{sr1}\kappa_{r1}x_{sr1} (K_A x_A - K_B x_B)$$
 (1)

$$v_{r2} = \kappa_{r2} \left(K_B x_B - K_C x_C \right) \tag{2}$$

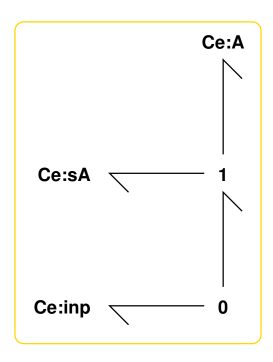
4 Sensitivity components

4.1 sCe

4.1.1 Bond Graph

```
[45]: # sCe
sbg.model('sCe_abg.svg')
import sCe_abg
disp.SVG('sCe_abg.svg')
```

[45]:

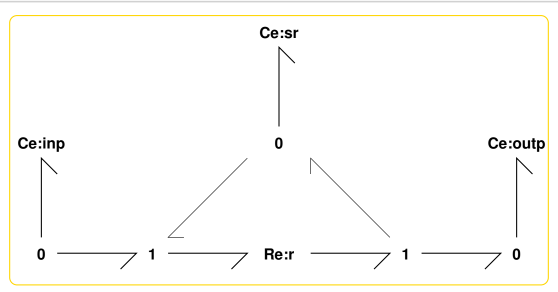


4.2 sRe

4.2.1 Bond Graph

```
[46]: # sRe
sbg.model('sRe_abg.svg')
import sRe_abg
disp.SVG('sRe_abg.svg')
```

[46]:

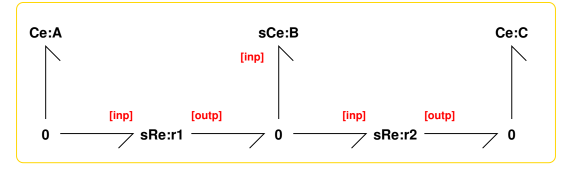


4.3 Simple system revisited

```
[47]: # Simple example A = B = C with sensitivity components
sbg.model('sABC_abg.svg')
import sABC_abg
disp.SVG('sABC_abg.svg')
```

Creating subsystem: sCe:B Creating subsystem: sRe:r1 Creating subsystem: sRe:r2

[47]:



4.3.1 Parameters

```
[48]: # ## Parameters
# parameter = {}
# parameter['K_A'] = 1
# parameter['K_B_A'] = 1
# parameter['kappa_r1'] = 0.1
# parameter['kappa_r2'] = 0.9
# print(parameter)

# ## Initial states
# X_A_0 = 2
```

```
[49]: ## Parameters
parameter = {}
parameter['K_A'] = 1
parameter['K_B_A'] = 1
parameter['kappa_r1_r'] = 1
parameter['kappa_r2_r'] = 9
print(parameter)

## Initial states
X_A_O = 2
```

```
{'K_A': 1, 'K_B_A': 1, 'kappa_r1_r': 1, 'kappa_r2_r': 9}
```

4.3.2 Stoichiometry & linearisation

```
[50]: ## Stoichiometry
      s,sc,sf = Stoichiometry(sABC_abg.
      →model(),chemostats=['A','C','B_sA','r1_sr','r2_sr'],flowstats=[])
      ## Species
      print(s['species'])
      ## Reactions
      print(s['reaction'])
      ## Linearise
      Sys, X_ss, V_ss, dX_ss =
      Linear(s,sc,parameter=parameter,X0=[X_A_0,1,1,1,1,1],quiet=quiet)
      ## Lambda for comparison
      lam = 1.1
     ['A', 'C', 'B_A', 'B_sA', 'r1_sr', 'r2_sr']
     ['r1_r', 'r2_r']
     Steady-state finder error: 8.88e-16
     X_ss = [2. 1. 1.11. 1. 1.]
     V_ss = [0.9 \ 0.9]
```

[51]: ## Show reactions disp.Latex(st.sprintrl(s,all=True,chemformula=True))

[51]:

$$A + r_{1s}r \xrightarrow{r_{1r}} B_A + B_sA + r_{1s}r$$
 (3)

$$B_A + B_s A + r_{2s} r \stackrel{r_{2r}}{\longleftarrow} C + r_{2s} r \tag{4}$$

[52]: ## Show system
Sys

[52]:

$$\begin{pmatrix} -10 & 1 & 9 & -11 & 0.9 & -0.9 \\ -1 & 1 & 0 & -1.1 & 0.9 & 0 \\ 9 & 0 & -9 & 9.9 & 0 & 0.9 \end{pmatrix}$$

[53]: ## Show transfer function con.tf(Sys)

[53]:

$$\begin{bmatrix} \frac{s+9}{s+10} & \frac{-9}{s+10} \frac{-1.1s}{s+10} \frac{0.9s+8.1}{s+10} \frac{0.9}{s+10} \\ \frac{9}{s+10} & \frac{-9s-9}{s+10} \frac{9.9s}{s+10} \frac{8.1}{s+10} \frac{0.9s+0.9}{s+10} \end{bmatrix}$$

4.3.3 Stoichiometric matrix

[54]: disp.Latex(st.sprintl(s,'N'))

[54]:

$$N = \begin{pmatrix} -1 & 0 \\ 0 & 1 \\ 1 & -1 \\ 1 & -1 \\ 0 & 0 \\ 0 & 0 \end{pmatrix} \tag{5}$$

[55]: disp.Latex(st.sprintl(s,'Nf'))

[55]:

$$Nf = \begin{pmatrix} 1 & 0 \\ 0 & 0 \\ 0 & 1 \\ 0 & 1 \\ 1 & 0 \\ 0 & 1 \end{pmatrix} \tag{6}$$

[56]: disp.Latex(st.sprintl(s,'Nr'))

[56]:

$$Nr = \begin{pmatrix} 0 & 0 \\ 0 & 1 \\ 1 & 0 \\ 1 & 0 \\ 1 & 0 \\ 0 & 1 \end{pmatrix} \tag{7}$$

```
[57]: ## Show dc gain print('DC gain: \n', con.dcgain(Sys))
```

DC gain:

```
[[ 0.9 -0.9 0. 0.81 0.09]
[ 0.9 -0.9 0. 0.81 0.09]]
```

4.4 Sensitivity Bond Graph – change B

```
[58]: ## Extract sensitivity system
inp = ['B_sA']
outp = ['r1_r','r2_r']
sys = extractSubsystem(Sys,sc,sf,inp,outp)
```

```
[59]:
```

$$\begin{bmatrix} \frac{-1.1s}{s+10} \\ \frac{9.9s}{s+10} \end{bmatrix}$$

[60]: ## System matrix sys

[60]:

$$\begin{pmatrix} -10 & -11 \\ -1 & -1.1 \\ 9 & 9.9 \end{pmatrix}$$

[61]: ## Show dc gain print('DC gain: \n', con.dcgain(sys))

DC gain:

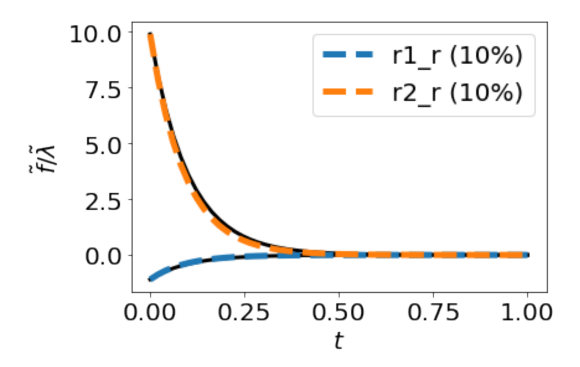
[[0.]

[0.]]

4.4.1 Compare sensitivity with exact simulation

```
[62]: ## Exact Simulate with changed K
parameter1 = copy.copy(parameter)
parameter1['K_B_A'] *= lam
t = np.linspace(0,1)
dat = st.sim(s,sc=sc,t=t,parameter=parameter1,X0=X_ss,quiet=quiet)
```

- [63]: ## Step response to sB y_lin = linStep(sc,sys,T=t)
- [64]: ## Plot plotSensitivity(dat,reactions=outp)



4.5 Sensitivity Bond Graph – change r1

```
[65]: ## Extract sensitivity system
inp = ['r1_sr']
outp = ['r1_r', 'r2_r']
sys = extractSubsystem(Sys,sc,sf,inp,outp)
```

[66]: ## Show transfer function con.tf(sys)

[66]:

$$\begin{bmatrix} \frac{0.9s + 8.1}{s + 10} \\ \frac{8.1}{s + 10} \end{bmatrix}$$

[67]: ## System matrix

[67]:

$$\begin{pmatrix} -10 & 0.9 \\ -1 & 0.9 \\ 9 & 0 \end{pmatrix}$$

DC gain: [[0.81] [0.81]]

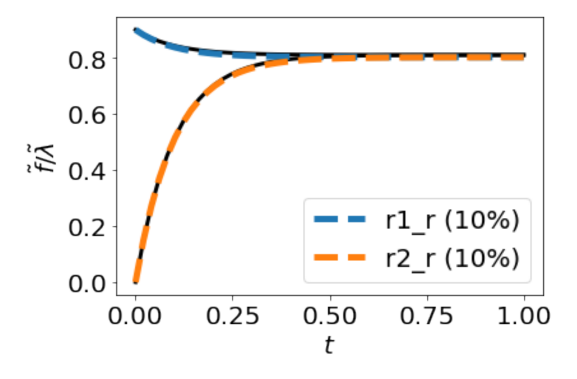
4.5.1 Compare sensitivity with exact simulation

```
[69]: ## Exact Simulate with changed r1
parameter1 = copy.copy(parameter)
parameter1['kappa_r1_r'] *= lam
t = np.linspace(0,1)
dat = st.sim(s,sc=sc,t=t,parameter=parameter1,X0=X_ss,quiet=quiet)
print(parameter1)
```

{'K_A': 1, 'K_B_A': 1, 'kappa_r1_r': 1.1, 'kappa_r2_r': 9}

```
[70]: ## Step response to sr1
y_lin = linStep(sc,sys,T=t)
```

[71]: ## Plot plotSensitivity(dat,reactions=outp)



4.6 Sensitivity Bond Graph – change r2

```
[72]: ## Extract sensitivity system
inp = ['r2_sr']
outp = ['r1_r','r2_r']
sys = extractSubsystem(Sys,sc,sf,inp,outp)
[73]: ## Show transfer function
```

[73]:

con.tf(sys)

$$\begin{bmatrix} \frac{0.9}{s+10} \\ \frac{0.9s+0.9}{s+10} \end{bmatrix}$$

```
[74]: ## System matrix sys
```

[74]:

$$\begin{pmatrix} -10 & -0.9 \\ -1 & 0 \\ 9 & 0.9 \end{pmatrix}$$

```
[75]: ## Show dc gain
print('DC gain: \n', con.dcgain(sys))
```

DC gain: [[0.09] [0.09]]

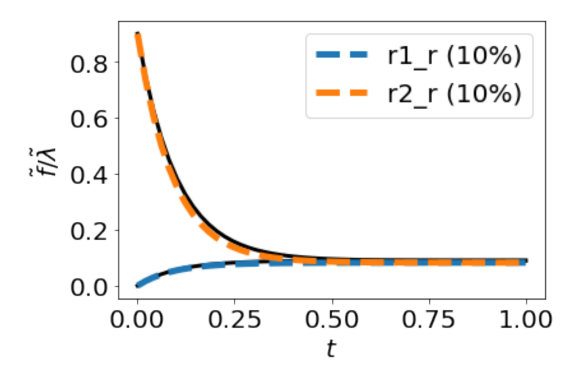
4.6.1 Compare sensitivity with exact simulation

```
[76]: ## Exact Simulate with changed r2
parameter1 = copy.copy(parameter)
parameter1['kappa_r2_r'] *= lam
t = np.linspace(0,1)
dat = st.sim(s,sc=sc,t=t,parameter=parameter1,X0=X_ss,quiet=quiet)
print(parameter1)
print(X_ss)
```

```
{'K_A': 1, 'K_B_A': 1, 'kappa_r1_r': 1, 'kappa_r2_r': 9.9}
[2. 1. 1.1 1. 1. ]
```

```
[77]: ## Step response to sr2
y_lin = linStep(sc,sys,T=t)
```

```
[78]: ## Plot plotSensitivity(dat,reactions=outp)
```



5 Stoichiometric approach

5.1 Supporting software

```
[79]: def stoichSensitivity(model,chemostats=['A','C'],
       →parameter={},X0=None,CommonSpecies=None):
          """Sensitivity analysis via stoichiometric approach """
          ## Stoichiometry
          s,sc,sf = 
       →Stoichiometry(model,chemostats=chemostats,CommonSpecies=CommonSpecies,sensitivity=True)
          ## Linearise
          if XO is None:
              X0 = np.ones(s['n_X'])
              i_chem = s['species'].index(chemostats[0])
              X0[i\_chem] = 2
          Sys,X_ss,V_ss,dX_ss = Linear(s,sc,parameter=parameter,X0=X0,quiet=quiet)
          return s,sc,sf,Sys,X_ss,V_ss,dX_ss
      def simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,chemostats=['A','C'],
                         inp=['sr1'],outp=['r1','r2'],t_last=2,parameter={},
                         lam=1.2,order=None,sX0=None,lambdaIn=True):
```

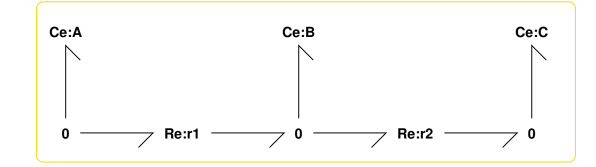
```
## Extract sensitivity system
   sys = extractSubsystem(Sys,sc,sf,inp=inp,outp=outp,order=order)
   ## Time
   t = np.linspace(0,t_last,500)
   ## Sensitivity step response
    step = con.step_response(sys, T=t)
     y_lin = step.y[:, 0, :].T
   y_lin = linStep(sc,sys,T=t,X0=sX0)
   ## Exact simulation with changed parameter or state
   X_s_1 = copy.copy(X_s)
   parameter1 = copy.copy(parameter)
   inComp = inp[0][1:]
   if sXO is None:
       ## Parameter or chemostat perturbation
       X_{chemo} = \{\}
       if inComp in chemostats:
           ## Perturb state
           iComp = s['species'].index(inComp)
           X_ss_1[iComp] = lam*X_ss_1[iComp]
             y_lin *= X_ss_1[iComp]
   #
       else:
           if lambdaIn:
               print("Use lambda as input")
               ## Perturb lambda chemostat initial state
               iComp = s['species'].index('s'+inComp)
               X_ss_1[iComp] = lam*X_ss_1[iComp]
           else:
           ## Perturb parameter
               print('Perturb parameter')
               if inComp[0].isupper():
                   parname = 'K_'+inComp
               else:
                   parname = 'kappa_'+inComp
               if parname in list(parameter.keys()):
                   parameter1[parname] = lam*parameter[parname]
                   parameter1[parname] = lam
       dat = st.
⇒sim(s,sc=sc,sf=sf,t=t,parameter=parameter1,X0=X_ss_1,X_chemo=X_chemo,
                    quiet=quiet,Normalise=Normalise)
   else:
       ## Initial condition perturbation.
       X_s_1 += (lam-1)*sX0
       dat = st.
→sim(s,sc=sc,sf=sf,t=t,parameter=parameter1,X0=X_ss_1,quiet=quiet,Normalise=Normalise)
```

```
return dat,y_lin,t,sys
```

5.2 Simple example A = B = C

```
[80]: # Simple example A = B = C
sbg.model('ABC_abg.svg')
import ABC_abg
disp.SVG('ABC_abg.svg')
```

: [08]



```
[81]: ## Parameters
parameter = {}
parameter['K_A'] = 1
parameter['K_B'] = 1
parameter['kappa_r1'] = 1
parameter['kappa_r2'] = 9
print(parameter)

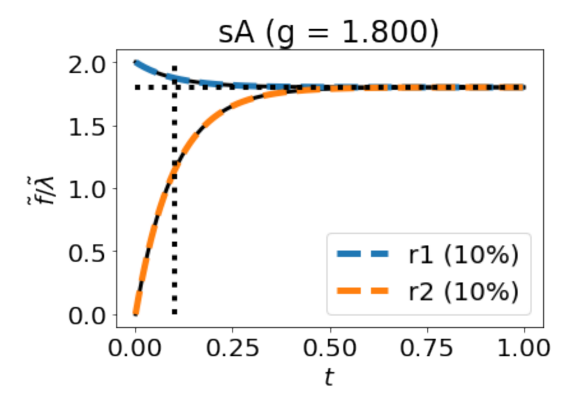
## Initial states
X_A_0 = 2
```

{'K_A': 1, 'K_B': 1, 'kappa_r1': 1, 'kappa_r2': 9}

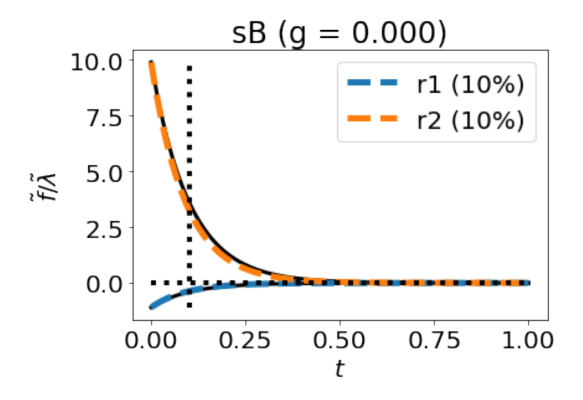
```
plt.title(f'{inp} (g = {g:.3f})')
plotSensitivity(dat,reactions=outp)
plotLines()

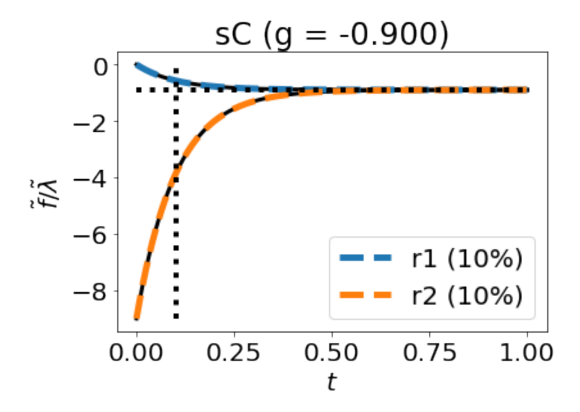
# plt.hlines(g,min(t),max(t),color='black',ls='dashed')
Savefig('ABC_'+inp)
plt.show()
```

```
Steady-state finder error: 8.88e-16
X_ss = [2. 1.1 1. 1. 1. 1. 1. 1.]
V_ss = [0.9 0.9]
```

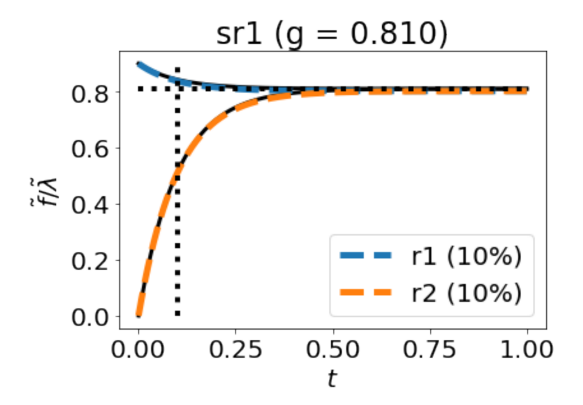


Use lambda as input

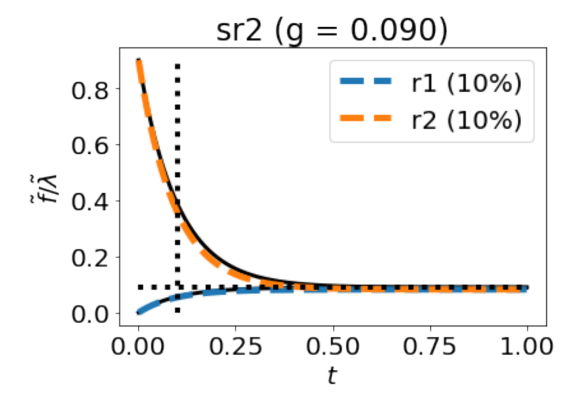




Use lambda as input



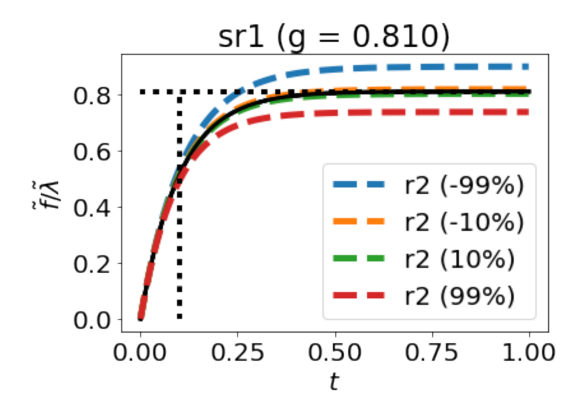
Use lambda as input



```
[83]: printLatex(s,sc=sc)
                A &= \left(\begin{matrix}-10.0\end{matrix}\right)
                B &= \left(\begin{matrix}1.0 & 9.0 & 2.0 & -11.0 & 9.0 & 0.9 &
                -0.90000000000001\end{matrix}\right)
                C &= \left(\begin{matrix}-1.0\\9.0\end{matrix}\right)
                D &= \left(\begin{matrix}1.0 & 0 & 2.0 & -1.1 & 0 & 0.9 & 0\\0 & -9.0 & 0 & 9.9
                & -9.0 & 0 & 0.9000000000001\end{matrix}\right)
                \begin{align}
                X&= \begin{pmatrix}
                           X_{A}\
                           X_{B}
                           X_{C}\
                           X_{sA}
                           X_{sB}
                           X_{sC}\
                           X_{sr1}\
                           X_{sr2}\
                \end{pmatrix}
                \end{align}
                \begin{align}
                V&= \begin{pmatrix}
                           V_{r1}\\
                           V_{r2}\\
                \end{pmatrix}
                \end{align}
                \begin{align}
                Nf &=
                \left(\begin{matrix}1.0 & 0\\0 & 1.0\\0 & 0\\1.0 & 0\\0 & 1.0\\0 & 0\\1.0
                & 1.0\end{matrix}\right)
                \end{align}
                \begin{align}
                \left(\begin{matrix}0 & 0\\1.0 & 0\\0 & 1.0\\0 & 0\\1.0 & 0\\0 & 1.0\\1.0 & 0\\0
                & 1.0\end{matrix}\right)
                \end{align}
                \begin{align}
                N &=
                1.0\\0 & 0\\0 & 0\end{matrix}\right)
                \end{align}
                %% Nc matrix
                \begin{align}
                \label{left(begin{matrix}0 & 0\\1.0 & -1.0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 0\\0 & 
                0\end{matrix}\right)
```

5.2.1 Vary λ

Use lambda as input Use lambda as input Use lambda as input Use lambda as input



```
[85]: ## Show system
Sys
```

[85]:

$$\begin{pmatrix}
-10 & 1 & 9 & 2 & -11 & 9 & 0.9 & -0.9 \\
-1 & 1 & 0 & 2 & -1.1 & 0 & 0.9 & 0 \\
9 & 0 & -9 & 0 & 9.9 & -9 & 0 & 0.9
\end{pmatrix}$$

```
[86]: ## Show system TF
print(sc['chemostats'])
con.tf(Sys)
```

['A', 'C', 'sA', 'sB', 'sC', 'sr1', 'sr2']

[86]:

$$\begin{bmatrix} \frac{s+9}{s+10} & \frac{-9}{s+10} \frac{2s+18}{s+10} \frac{-1.1s}{s+10} \frac{-9}{s+10} \frac{0.9s+8.1}{s+10} \frac{0.9}{s+10} \\ \frac{9}{s+10} & \frac{-9s-9}{s+10} \frac{18}{s+10} \frac{9.9s}{s+10} \frac{-9s-9}{s+10} \frac{8.1}{s+10} \frac{0.9s+0.9}{s+10} \end{bmatrix}$$

```
[87]: ## Show DC gain con.dcgain(Sys)
```

[88]: ## Show initial reponse print(Sys.D)

```
[[ 1.  0.  2.  -1.1  0.  0.9  0. ]
[ 0.  -9.  0.  9.9  -9.  0.  0.9]]
```

5.2.2 Sloppy parameters

```
[89]: imp.reload(slp)
      def sloppy(Sys,inp,outp,t=None,GainOnly=False):
          sys = extractSubsystem(Sys,sc,sf,inp,outp)
          #print(sys)
          H,eig,eigv,t = slp.Sloppy(sys,t=t,GainOnly=GainOnly)
          print(H)
          slp.SloppyPlot(eig,eigv,inp)
          slp.SloppyPrint(eig,eigv,inp,min_eig=0)
            if not GainOnly:
                slp.SloppyPlotData(t,y,inp,outp)
      def sloppyBoth(Sys,inp,outp,t=1):
          sys = extractSubsystem(Sys,sc,sf,inp,outp)
          SysName = sc['name']
          print(SysName)
          if (\max(t)<1e2):
              name = f'{SysName}_sloppy_{inp[0]}'
              name = f'{SysName}_sloppy_{inp[0]}_long'
          #print(sys)
          H,eig,eigv,t = slp.Sloppy(sys,t=t,GainOnly=False)
          print(f' n t_f = \{max(t): 0.2f\}')
          print('H:')
          slp.SloppyPrint(eig,eigv,inp,min_eig=0)
          H,Eig,Eigv,t = slp.Sloppy(sys,t=t,GainOnly=True)
          print('H_ss:')
          slp.SloppyPrint(Eig,Eigv,inp,min_eig=0)
          ## Direct computation
          print('Direct:')
          gain = con.dcgain(sys)[0]
          norm = np.sum(gain*gain)
          ngain = gain/np.sqrt(norm)
          slp.SloppyPrint([norm],np.array([ngain]).T,inp,min_eig=0.0,min_eigv=0.
       \rightarrow05,max_eigs=2)
          ## Plot
          slp.SloppyPlot(eig,eigv,inp,Eig=Eig,Eigv=Eigv)
          Savefig(name)
          plt.show()
```

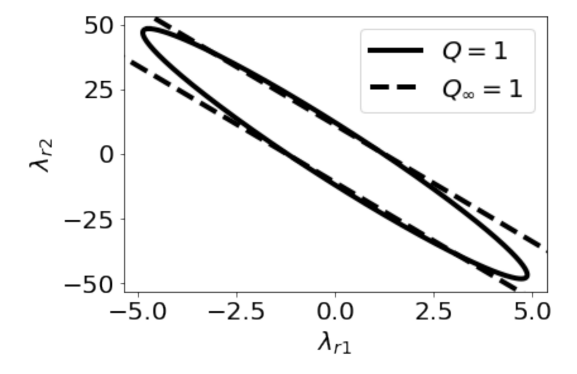
```
for t_last in [1,1e3]:
    if t_last==0:
        t = None
    else:
        t = np.linspace(0,t_last,100)
    sloppyBoth(Sys,['sr1','sr2'],['r1'],t=t)
    sloppyBoth(Sys,['sA','sC'],['r1'],t=t)
```

ABC

 $t_f = 1.00$

Η:

 $\sqrt sqrt \frac{1}{e} 0.81 \& V_1 \& e + 0.994 \ada_{r1} + 0.110 \ada_{r2}$

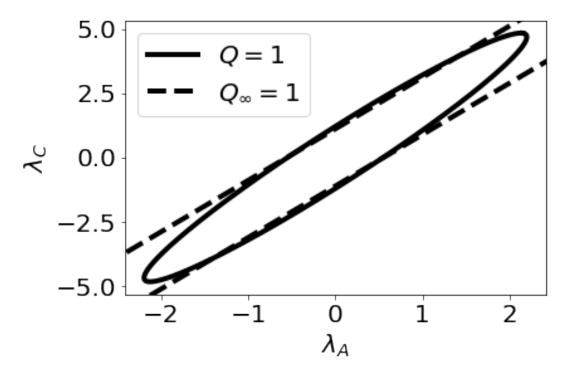


ABC

 $t_f = 1.00$

Η:

 $\label{lambda_A} $$ \sqrt 1 \end{array} $$ \sqrt 1 \end{a$



ABC

 $t_f = 1000.00$

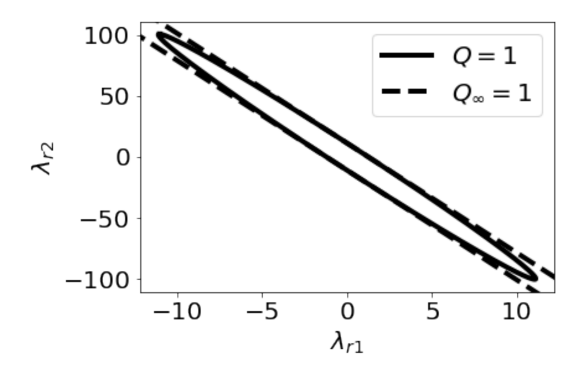
н:

 $\label{lambda_{r1}} $$ \sqrt 1 \le 0.82 & V_1\ \& = 0.994 \ambda_{r1} + 0.109 \ambda_{r2} \ambda_2 & 0.0099 & V_2\ \& = 0.994 \ambda_{r2} - 0.109 \ambda_{r1}$$

Н сс

\sqrt\sigma_1 &= 0.81 & V_1\Lambda &= + 0.994 \lambda_{r1} + 0.110 \lambda_{r2} \sqrt\sigma_2 &= 1e-06 & V_2\Lambda &= + 0.994 \lambda_{r2} - 0.110 \lambda_{r1} \Direct:

 $\sqrt\sigma_1 \&= 0.81 \& V_1\Lambda \&= + 0.994 \lambda_{r1} + 0.110 \lambda_{r2}$

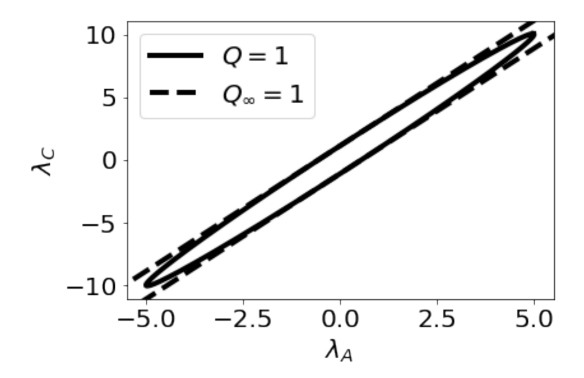


ABC

 $t_f = 1000.00$

Η:

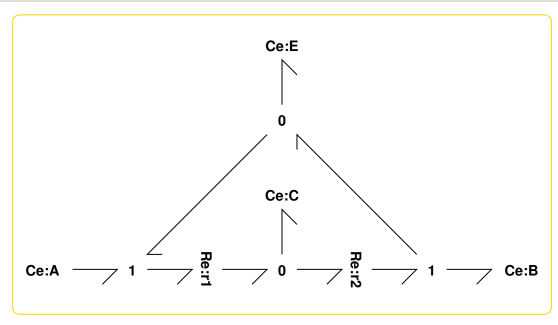
\sqrt\sigma_1 &= 2 & V_1\Lambda &= + 0.894 \lambda_{A} - 0.447 \lambda_{C} \sqrt\sigma_2 &= 1e-06 & V_2\Lambda &= + 0.894 \lambda_{C} + 0.447 \lambda_{A} \Direct:



5.3 Example: Enzyme-catalysed reaction

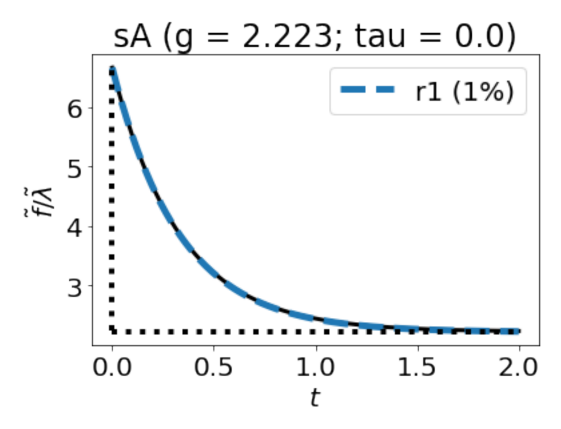
```
[90]: sbg.model('ECR_abg.svg')
  import ECR_abg
  imp.reload(ECR_abg)
  disp.SVG('ECR_abg.svg')
```

[90]:

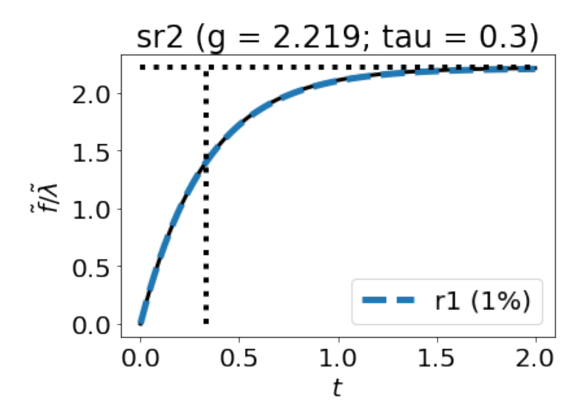


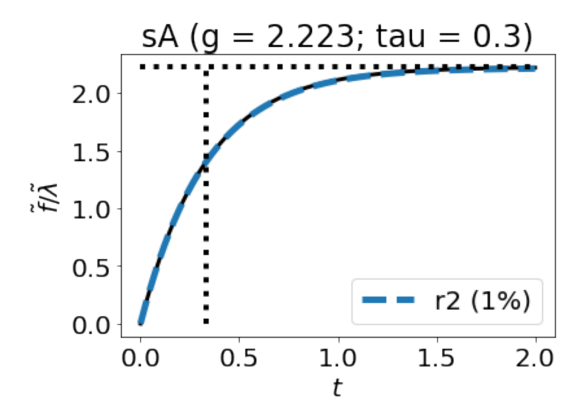
```
[91]: ## Stoichiometry
      s0 = st.stoich(ECR_abg.model(),quiet=quiet)
      species = s0['species']
      reaction=s0['reaction']
      n_X = s0['n_X']
      n_V = s0['n_V']
[92]: | ## Initial states (for sensitivity system)
      eO = 10 ## Total bound and unbound enzyme
      XXO = \{\}
      \# XXO['A'] = 1
      XX0['E'] = e0/2
      XX0['C'] = e0/2
      X0 = np.ones(2*n_X+n_V)
      for spec in XXO:
          X0[species.index(spec)] = XX0[spec]
[93]: ## Parameters
      kappa_1 = 1
      K_B = 1e-3
      parameter = {}
      parameter['K_A'] = 1
      parameter['K_B'] = K_B
      parameter['kappa_r1'] = kappa_1
      parameter['kappa_r2'] = 1
      print(parameter)
     {'K_A': 1, 'K_B': 0.001, 'kappa_r1': 1, 'kappa_r2': 1}
[94]: ## Chemostats
      chemostats = ['A','B']
[95]: dcgain = {}
      syss = {}
      s,sc,sf,Sys,X_ss,V_ss,dX_ss = stoichSensitivity(ECR_abg.
      →model(),parameter=parameter,chemostats=chemostats,X0=X0)
      lam = 1.01
      Outp = ['r1', 'r2']
      Inp = ['sA','sr2']
      t_last = 2/kappa_1
      for outp in Outp:
          for inp in Inp:
              dat,y_lin,t,sys = simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,
       \rightarrowparameter=parameter,inp=[inp],outp=[outp],
       →lam=lam,t_last=t_last,lambdaIn=lambdaIn)
              syss[inp] = sys
                g = con.dcgain(sys)
```

```
g,tau = tfProps(sys)
dcgain[inp] = g
    #print(g)
if Titles:
    plt.title(f'{inp} (g = {g:.3f}; tau = {tau:0.1f})')
plotSensitivity(dat,reactions=[outp])
plotLines()
Savefig('ECR_'+inp+'_'+outp)
plt.show()
```

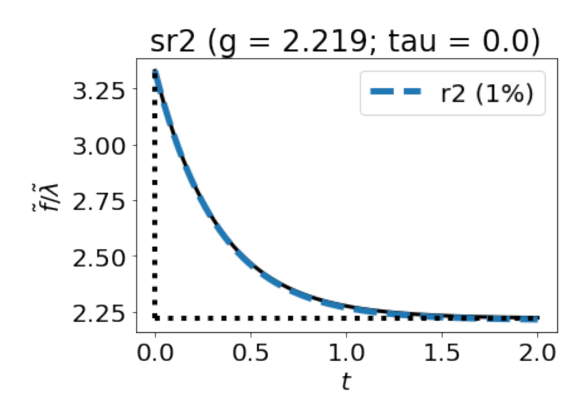


Use lambda as input

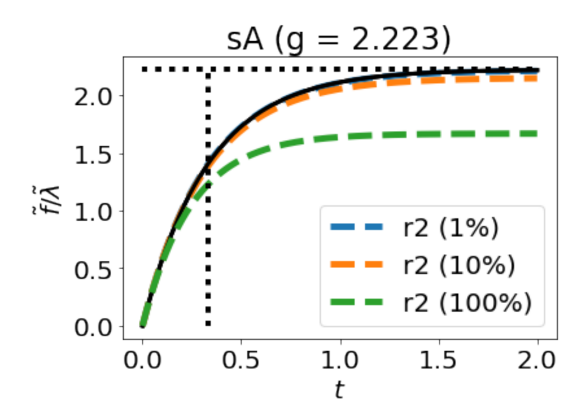




Use lambda as input



5.3.1 Vary λ



```
[97]: ## Show system #Sys  
[98]: ## Show system TFs  
con.tf(syss['sA'])  
[98]: \frac{6.671}{s+3.001}  
[99]: con.tf(syss['sr2'])  
[99]: \frac{3.329s+6.658}{s+3.001}
```

5.3.2 Simulate over flow range - quasi steady-state. Supporting software

```
[100]: def QuasiSteadyState(Inp=['sAct','sInh'],points=10,x_max=100,grid=False):
    ## Extract info
    species = s['species']
    reaction = s['reaction']

    ### Simulate over flow range.
    ## Slow ramp for x_A
    X_chemo = {}
```

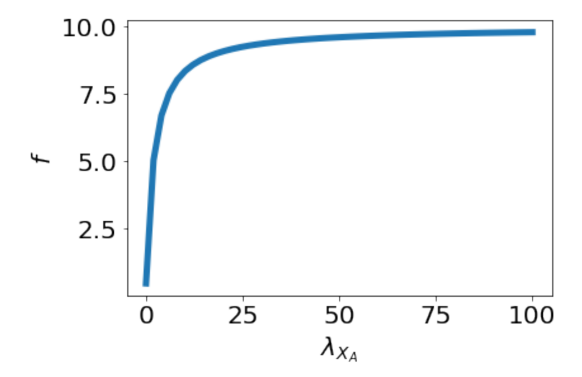
```
t_d = 1e3
   t_{last} = 1e6
   points = 50
   t = np.linspace(0,t_last,points)
   x_0 = 0.1
   X0[species.index('A')] = x_0
   slope = (x_max-x_0)/(t_last-t_d)
   X_{\text{chemo}}['A'] = f'(\{x_0\} + \{slope\}*(t_{t_d})*(1*(t_{t_d})))'
   print(X_chemo)
   ## Steady-state
   X_ss,V_ss = SteadyState(s,sc,parameter=parameter,X0=X0)
   ## Simulate
   ndat = st.
→sim(s,sc=sc,t=t,parameter=parameter,X0=X_ss,X_chemo=X_chemo,quiet=quiet)
   ## Plot
    st.plot(s,ndat,species=['E0','E','C'],reaction=['r2'])
   ## Plot flow v X_A
   ylabel = r'$\tilde {f}_2/\tilde{\lambda}$'
   X_A = ndat['X'][:,species.index('A')]
   V_2 = ndat['V'][:,reaction.index('r2')]
   plt.plot(X_A,V_2,lw=5)
   if grid:
       plt.grid()
   plt.xlabel('$\lambda_{X_A}$')
   plt.ylabel('$f$')
   sysname = s['name']
   plotname = f'{sysname}_flow'
   Savefig(plotname)
   plt.show()
   plt.clf()
   ## Compute sensitivity gain for each steady-state
   X = ndat['X']
   DCgain = {}
   Tau = \{\}
   for inp in Inp:
       DCgain[inp] = []
       Tau[inp] = []
       for x in X:
           ## Linearise about this steady-state
           Sys = st.lin(s,sc,x_ss=x,parameter=parameter,
                         quiet=quiet)
           ## Extract relevant subsystems
           outp = ['r1']
```

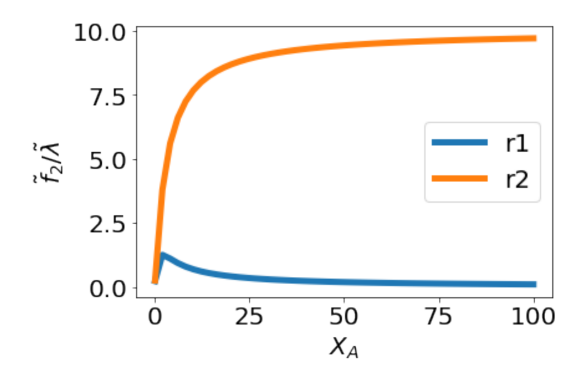
```
sys = extractSubsystem(Sys,sc,sf,inp=[inp],outp=outp)
          dcgain = con.dcgain(sys)
        dcgain,tau = tfProps(sys)
          print(dcgain)
        DCgain[inp].append(dcgain)
        Tau[inp].append(tau)
# plt.plot(X_A, V_2, lw=6, label='flow')
INP = ''
for inp in Inp:
    plt.plot(X_A,DCgain[inp],label=inp[1:],lw=5)
    INP += inp+'_'
plt.legend()
if grid:
    plt.grid()
plt.xlabel('$X_A$')
plt.ylabel(ylabel)
plotname = f'{sysname}_{INP}X'
Savefig(plotname)
plt.show()
# plt.plot(X_A, V_2, lw=6, label='flow')
## Plot DC gain v flow
for inp in Inp:
    plt.plot(V_2,DCgain[inp],label=inp[1:],lw=5)
plt.legend()
if grid:
    plt.grid()
plt.xlabel('$f$')
ylabel = '$g_\infty$'
plt.ylabel(ylabel)
plotname = f'{sysname}_{INP}f'
Savefig(plotname)
plt.show()
## Plot time-constant v flow
for inp in Inp:
    plt.plot(V_2,Tau[inp],label=inp[1:],lw=5)
plt.legend()
if grid:
    plt.grid()
plt.xlabel('$f$')
plt.ylabel(r'$\tau$')
plotname = f'{sysname}_{INP}f_tau'
Savefig(plotname)
plt.show()
```

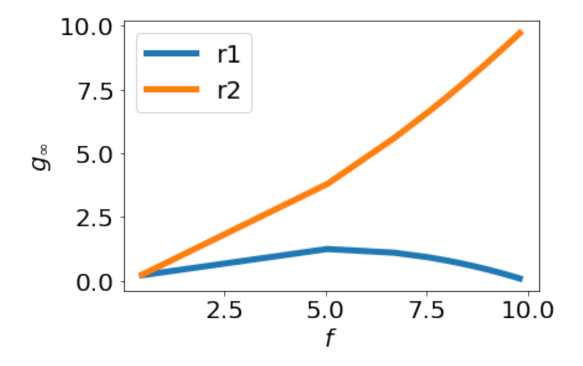
5.3.3 Simulate over flow range - quasi steady-state. Vary Re components.

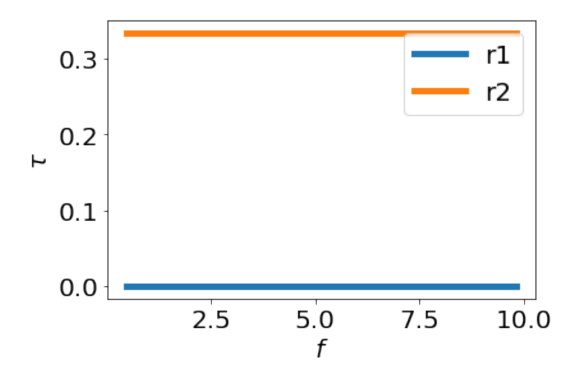
[101]: QuasiSteadyState(Inp=['sr1','sr2'],points=50)

 ${'A': '(0.1 + 0.0001*(t-1000.0)*(1*(t>1000.0)))'}$ Steady-state finder error: 7.85e-17





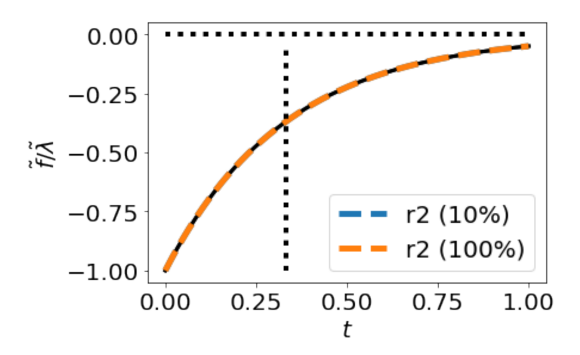




5.3.4 Initial condition sensitivity.

```
[102]: ## Set up deviation state for full system
       n_X = sc['n_X']
       sX0 = np.zeros(n_X)
       spec = sc['species']
       i_C = spec.index('C')
       i_E = spec.index('E')
       sX0[i_E] = 1
       sX0[i_C] = -1
       ## Test conserved moiety compatability
       G_c = sc['G']
       error = np.linalg.norm(G_c@sX0)
       print('Compatibility error =',error)
       ## Compare linear and non-linear
       inp = 'sr1'
       outp = ['r2']
       for lam in [1.1,2,]:
           dat,y_lin,t,sys = simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,
                                             sX0=sX0,
       →parameter=parameter,inp=[inp],outp=outp,lam=lam,t_last=1)
            print(con.dcgain(sys))
           g = con.dcgain(sys)
            if Titles:
                 plt.title(f'{inp} (g = {g:.3f})')
           plotSensitivity(dat,reactions=outp)
           g=0;plotLines()
       Savefig('ECR_IC')
```

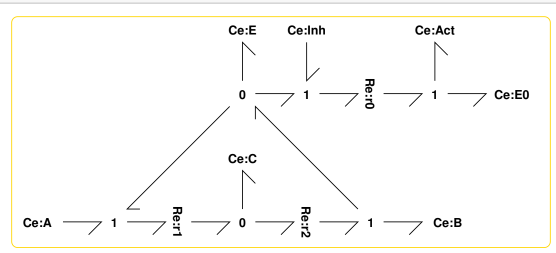
Compatibility error = 0.0



5.4 Example: Modulated Enzyme-catalysed reaction

```
[103]: sbg.model('ecr_abg.svg')
   import ecr_abg
   imp.reload(ecr_abg)
   disp.SVG('ecr_abg.svg')
```

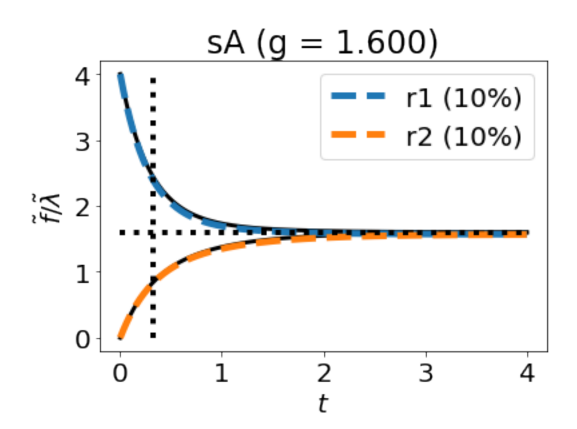
[103]:



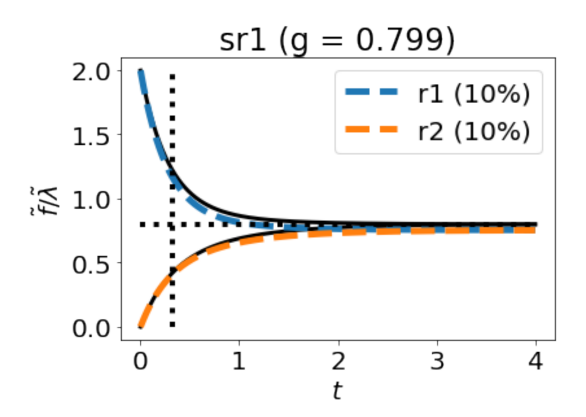
```
[104]: ## Stoichiometry
s0 = st.stoich(ecr_abg.model(),quiet=quiet)
species = s0['species']
n_X = s0['n_X']
n_V = s0['n_V']
```

```
[105]: | ## Initial states (for sensitivity system)
       \# e0 = 10
       XXO = \{\}
       \# XXO['A'] = 1
       XX0['E'] = e0/3
       XX0['E0'] = e0/3
       XX0['C'] = e0/3
       XXO['Inh'] = 1
       X0 = np.ones(2*n_X+n_V)
       for spec in XXO:
           X0[species.index(spec)] = XX0[spec]
[106]: ## Parameters
       parameter = {}
       parameter['K_A'] = 1
       parameter['K_B'] = K_B
       \# parameter['K_F'] = 1
       \# parameter['K_G'] = 1
       parameter['kappa_r0'] = 1
       parameter['kappa_r1'] = kappa_1
       parameter['kappa_r2'] = 1
       parameter['K_E0'] = 1
       print(parameter)
      {'K_A': 1, 'K_B': 0.001, 'kappa_r0': 1, 'kappa_r1': 1, 'kappa_r2': 1, 'K_E0': 1}
[107]: ## Chemostats
       chemostats = ['A','B','Act','Inh']
[108]: imp.reload(st)
       dcgain = {}
       syss = {}
       print(s['species'])
       s,sc,sf,Sys,X_ss,V_ss,dX_ss = stoichSensitivity(ecr_abg.
       →model(),parameter=parameter,chemostats=chemostats,X0=X0)
       lam = 1.1
       outp = ['r1', 'r2']
       Inp = ['sA','sr1','sr2','sAct','sInh']
       #Inp = ['sAct', 'sInh', 'sA', 'sB']
       t_{last} = 4
       for inp in Inp:
           dat,y_lin,t,sys = simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,
                                              parameter=parameter,inp=[inp],outp=outp,
                                             lam=lam,t_last=t_last,lambdaIn=lambdaIn)
           syss[inp] = sys
           print(sys)
             g = con.dcgain(sys)[0][0]
           g,tau = tfProps(sys)
           dcgain[inp] = g
           print(f'g = \{g:0.2f\}, tau = \{tau:0.2f\}')
```

```
if Titles:
        plt.title(f'{inp} (g = \{g:.3f\})')
    plotSensitivity(dat,reactions=outp)
    plotLines()
      plt.hlines(g,min(t),max(t),color='black',ls='dashed')
    Savefig('ecr_'+inp)
    plt.show()
['A', 'B', 'C', 'E', 'sA', 'sB', 'sC', 'sE', 'sr1', 'sr2']
Steady-state finder error: 1.66e-15
X_ss = [1.
            1.
                  1.
                        2.002 3.999 3.999 1. 1.
                                                      1.
                                                          1. 1. 1.
1. 1.
            1.
                  1.
                        1.
                           ]
V_ss = [-8.882e-16 \ 1.998e+00 \ 1.998e+00]
<LinearIOSystem>: sys[316]
Inputs (1): ['u[0]']
Outputs (2): ['y[0]', 'y[1]']
States (2): ['x[0]', 'x[1]']
A = [[-2.
            1.001]
    [ 1.
            -3.001]]
B = [[3.999]]
    [-3.999]]
C = [[-1.
            1. ]
    [ 1.
            -0.001]]
D = [[3.999]]
     [0. ]]
g = 1.60, tau = 0.32
```



```
Use lambda as input
<LinearIOSystem>: sys[318]
Inputs (1): ['u[0]']
Outputs (2): ['y[0]', 'y[1]']
States (2): ['x[0]', 'x[1]']
A = [[-2.
              1.001]
     [ 1.
             -3.001]]
B = [[1.998]]
     [-1.998]]
C = [[-1.
              1. ]
     [ 1.
             -0.001]]
D = [[1.998]]
     [0.
           ]]
g = 0.80, tau = 0.32
```



```
Use lambda as input 
<LinearIOSystem>: sys[320]
```

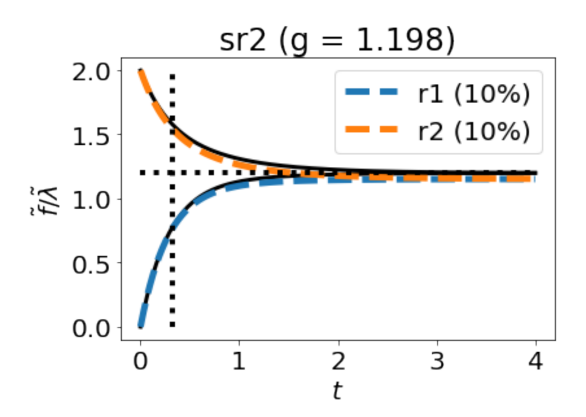
Inputs (1): ['u[0]']

Outputs (2): ['y[0]', 'y[1]']
States (2): ['x[0]', 'x[1]']

$$B = [[-1.998]$$
 [1.998]]

$$D = [[0.]]$$
 [1.998]]

$$g = 1.20$$
, tau = 0.32



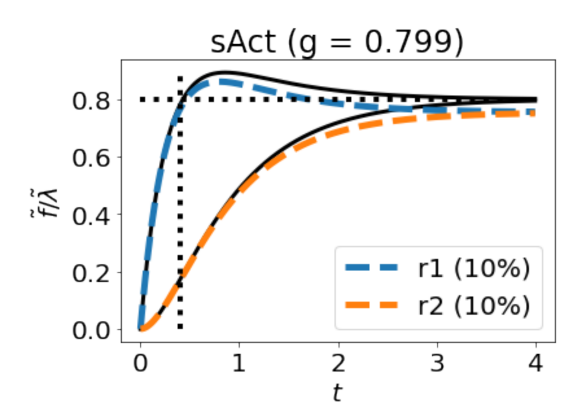
Use lambda as input
<LinearIOSystem>: sys[322]
Inputs (1): ['u[0]']

Outputs (2): ['y[0]', 'y[1]'] States (2): ['x[0]', 'x[1]']

B = [[0.]] [3.999]]

D = [[0.]]

g = 0.80, tau = 0.40



Use lambda as input <LinearIOSystem>: sys[324]

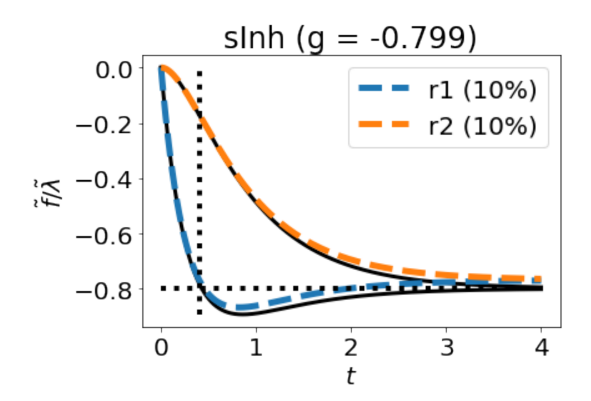
Inputs (1): ['u[0]']

Outputs (2): ['y[0]', 'y[1]'] States (2): ['x[0]', 'x[1]']

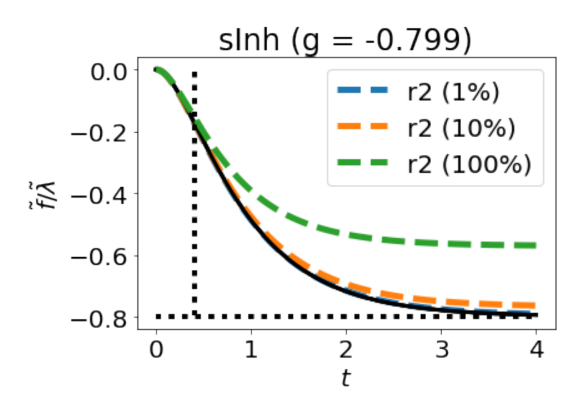
$$B = [[0.] \\ [-3.999]]$$

$$D = [[0.]]$$

$$g = -0.80$$
, tau = 0.40

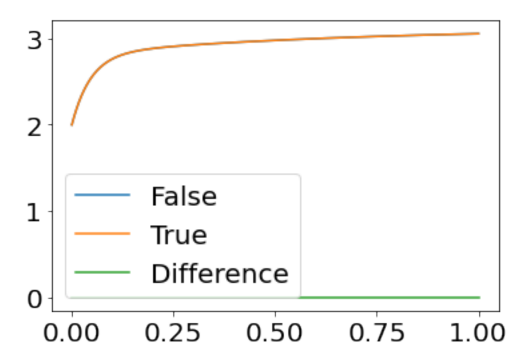


```
5.4.1
            Vary \lambda
[109]: tau
[109]: 0.39992001599680055
[110]: | inp = 'sInh'
       outp = ['r2']
       for lam in [1.01,1.1,2]:
           dat,y_lin,t,sys = simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,
                                              parameter=parameter,inp=[inp],outp=outp,
                                             lam=lam,t_last=t_last,lambdaIn=lambdaIn)
             print(con.dcgain(sys))
           g = con.dcgain(sys)
           if Titles:
               plt.title(f'{inp} (g = {g:.3f})')
           plotSensitivity(dat,reactions=outp)
           plotLines()
       Savefig('ecr_lambda')
      Use lambda as input
      Use lambda as input
      Use lambda as input
```



5.4.2 Sanity check on sensitivity system

Perturb parameter Use lambda as input Error:, 5.72e-12



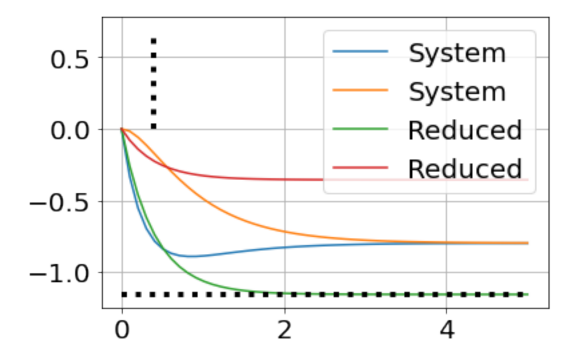
5.4.3 Show system transfer functions

5.4.4 Test model reduction

```
[114]: ## Test model reduction - truncate
method = 'truncate'
sys = syss['sInh']
g,tau = tfProps(sys)
print(f'g = {g:.2f}, tau = {tau:.2f}')
sys1 = con.balred(sys,orders=1,method=method)
# print(con.dcgain(sys))
# print(con.dcgain(sys1))
con.tf(sys1)
g,tau = tfProps(sys1)
print(f'g = {g:.2f}, tau = {tau:.2f}')
```

```
t = np.linspace(0,5)
step = con.step_response(sys,T=t)
step1 = con.step_response(sys1,T=t)
plt.plot(t,step.outputs[:,0,:].T,label='System')
plt.plot(t,step1.outputs[:,0,:].T,label='Reduced')
plt.legend()
plt.grid()
plotLines()
```

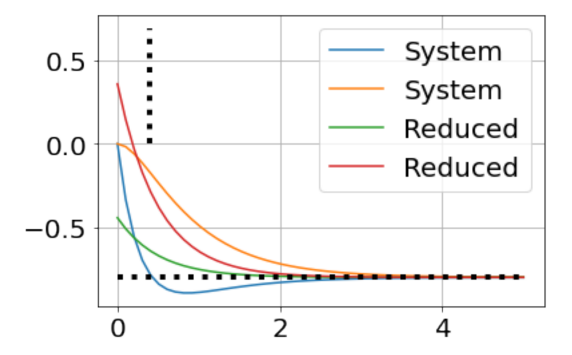
```
g = -0.80, tau = 0.40 g = -1.16, tau = 0.40
```



```
[115]: ## Test model reduction - matchdc
       method = 'matchdc'
       sys = syss['sInh']
       g,tau = tfProps(sys)
       print(f'g = {g:.2f}, tau = {tau:.2f}')
       sys1 = con.balred(sys,orders=1,method=method)
       # print(con.dcgain(sys))
       # print(con.dcgain(sys1))
       con.tf(sys1)
       g,tau = tfProps(sys1)
       print(f'g = \{g:.2f\}, tau = \{tau:.2f\}')
       t = np.linspace(0,5)
       step = con.step_response(sys,T=t)
       step1 = con.step_response(sys1,T=t)
       plt.plot(t,step.outputs[:,0,:].T,label='System')
       plt.plot(t,step1.outputs[:,0,:].T,label='Reduced')
```

```
plt.legend()
plt.grid()
plotLines()
```

$$g = -0.80$$
, tau = 0.40 $g = -0.80$, tau = 0.40

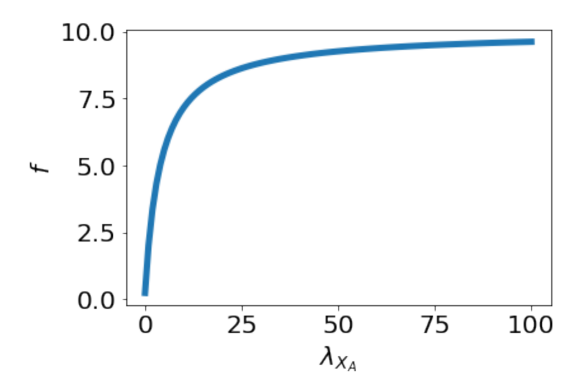


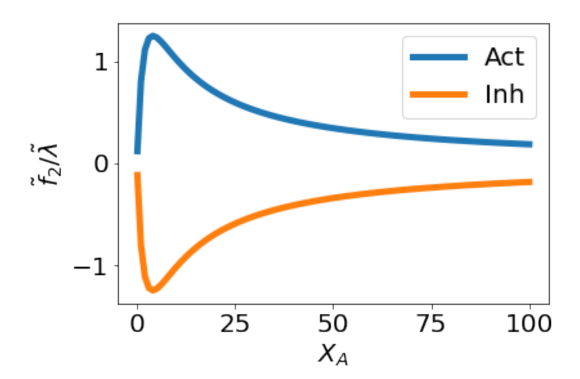
5.4.5 Simulate over flow range - quasi steady-state. Vary Activation and Inhibition.

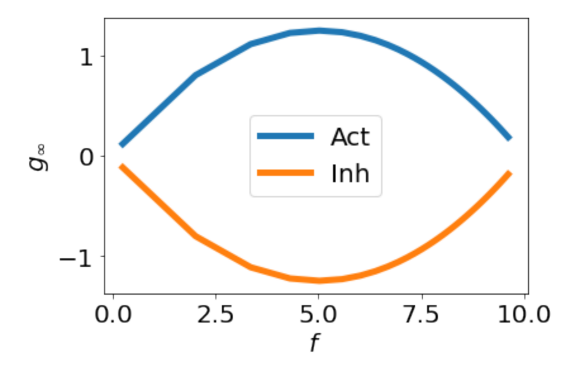
[116]: QuasiSteadyState(Inp=['sAct','sInh'],points=100)

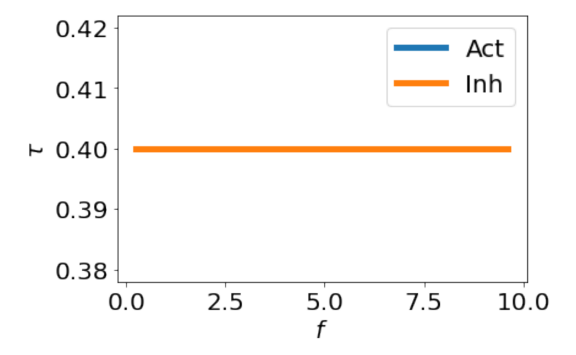
 ${'A': '(0.1 + 0.0001*(t-1000.0)*(1*(t>1000.0)))'}$

Steady-state finder error: 7.85e-17







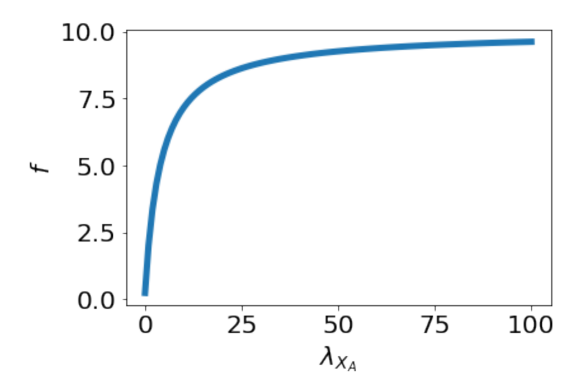


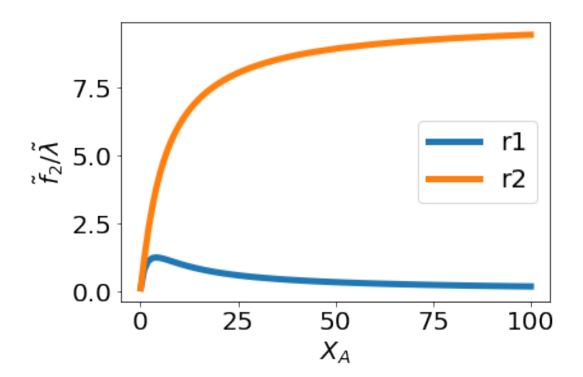
5.4.6 Simulate over flow range - quasi steady-state. Vary Re components

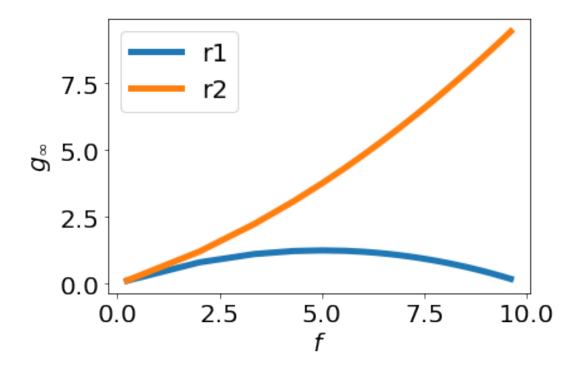
```
[117]: QuasiSteadyState(Inp=['sr1','sr2'],points=100)

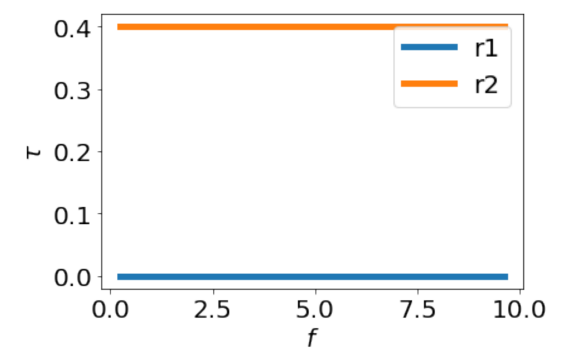
{'A': '(0.1 + 0.0001*(t-1000.0)*(1*(t>1000.0)))'}

Steady-state finder error: 7.85e-17
```







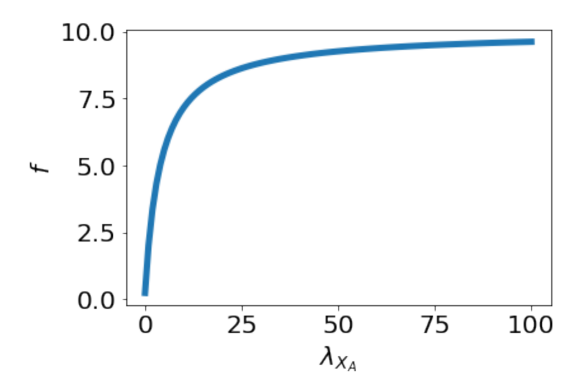


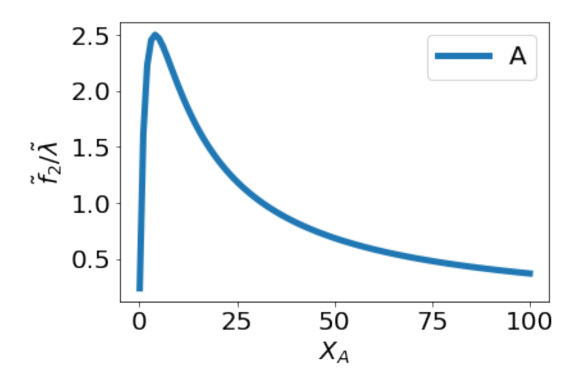
5.4.7 Simulate over flow range - quasi steady-state. Vary substrate concentration.

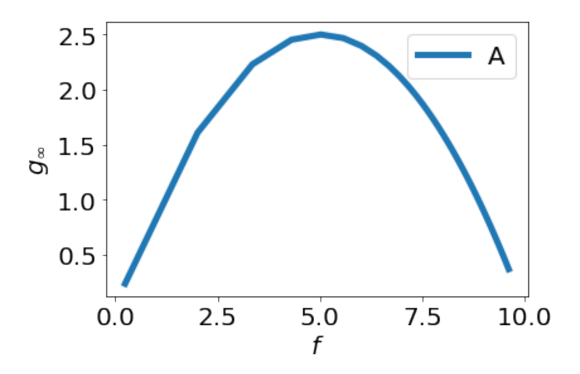
```
[118]: QuasiSteadyState(Inp=['sA'],points=100)

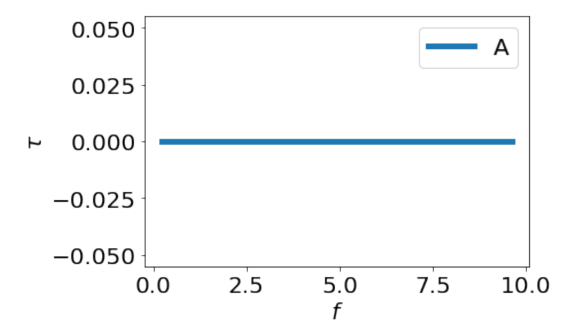
{'A': '(0.1 + 0.0001*(t-1000.0)*(1*(t>1000.0)))'}

Steady-state finder error: 7.85e-17
```









5.4.8 Sloppy parameters

```
[119]: imp.reload(slp)

for t_last in [4,1e2]:
    if t_last==0:
        t = None
```

```
else:
    t = np.linspace(0,t_last,100)

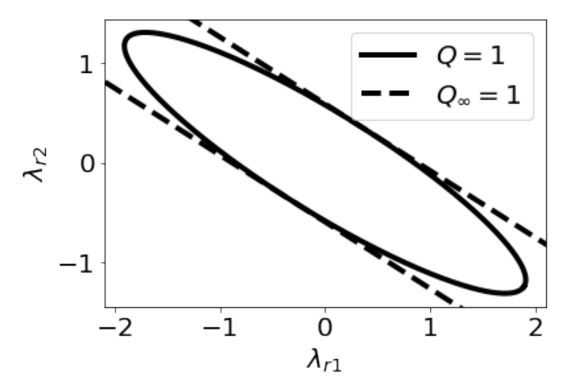
sloppyBoth(Sys,['sr1','sr2'],['r1','r2'],t=t)
sloppyBoth(Sys,['sE','sC'],['r1','r2'],t=t)
sloppyBoth(Sys,['sA','sB'],['r1','r2'],t=t)
```

 $t_f = 4.00$

Η:

 $\label{lambda_{r2} + 0.555 \ambda_{r1} &= 2 & V_1\ambda &= + 0.832 \ambda_{r2} + 0.555 \ambda_{r1} \\ \ambda_{r2} &= 1e-06 & V_2\ambda &= + 0.832 \ambda_{r1} - 0.555 \ambda_{r2} \\ \noalign{ \end{cases} Direct:$

 $\sqrt sqrt \frac{1.4 \& V_1\Delta \& = + 0.832 \ada_{r2} + 0.555 \ada_{r1}}$



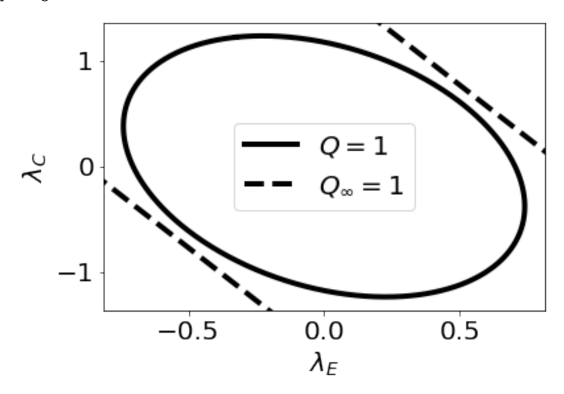
ecr

 $t_f = 4.00$

Η:

 $\label{eq:continuous} $$ \left(\frac{E} + 0.257 \right. & V_1\Delta_{E} + 0.966 \ad_{E} + 0.966 \ad_{E} + 0.257 \ad_{E} + 0.966 \ad_{E} + 0.9$

\sqrt\sigma_1 &= 0.89 & V_1\Lambda &= + 0.894 \lambda_{E} + 0.448 \lambda_{C}

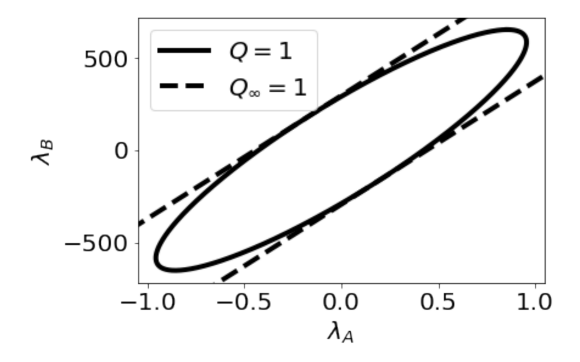


ecr

 $t_f = 4.00$

H:

\sqrt\sigma_1 &= 1.6 & V_1\Lambda &= + 1.000 \lambda_{A}



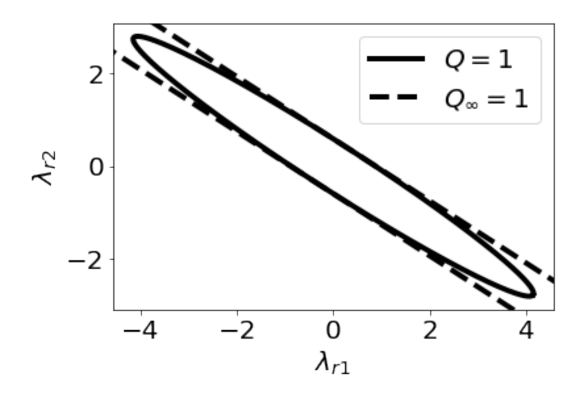
 $t_f = 100.00$

Η:

\sqrt\sigma_1 &= 2 & V_1\Lambda &= + 0.832 \lambda_{r2} + 0.555 \lambda_{r1} \sqrt\sigma_2 &= 0.2 & V_2\Lambda &= + 0.832 \lambda_{r1} - 0.555 \lambda_{r2} H ss:

 $\label{lambda_{r2} + 0.555 \ambda_{r1} &= 2 & V_1\ambda &= + 0.832 \ambda_{r2} + 0.555 \ambda_{r1} \\ \ambda_{r2} &= 1e-06 & V_2\ambda &= + 0.832 \ambda_{r1} - 0.555 \ambda_{r2} \\ \noalign{ \end{cases} Direct:$

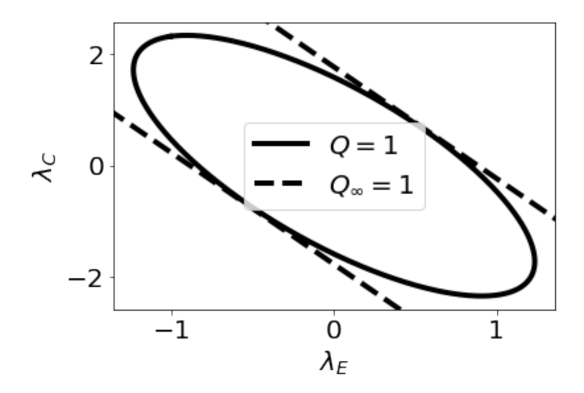
 $\sqrt\sigma_1 \&= 1.4 \& V_1\Lambda \&= + 0.832 \lambda_{r2} + 0.555 \lambda_{r1}$



 $t_f = 100.00$

Η:

\sqrt\sigma_1 &= 0.89 & V_1\Lambda &= + 0.894 \lambda_{E} + 0.448 \lambda_{C}

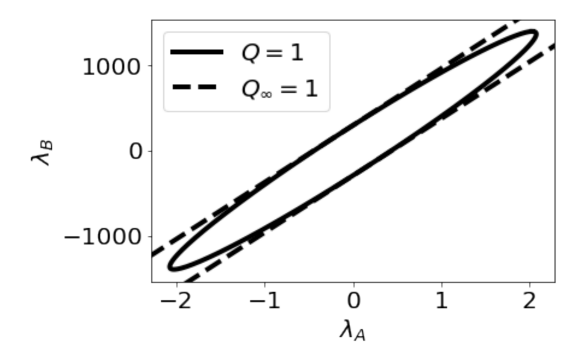


 $t_f = 100.00$

H:

\sqrt\sigma_1 &= 2.3 & V_1\Lambda &= + 1.000 \lambda_{A} - 0.001 \lambda_{B} \sqrt\sigma_2 &= 0.00072 & V_2\Lambda &= + 1.000 \lambda_{B} + 0.001 \lambda_{A} H_ss:

 $\label{eq:lambda_A} $$ \left(A - 0.001 \right) \ \end{a_{B}} \ \end{a_{B}} - 0.001 \ \end{a_{B}} \ \end{a_{B}} $$ \left(A - 0.001 \right) \ \end{a_{A}} $$ \left(A - 0.001 \right) \ \end{a_{A}} $$ \ \end{a_{A}} $$ \ \end{a_{B}} $$ \ \end{a_{A}} $$ \$



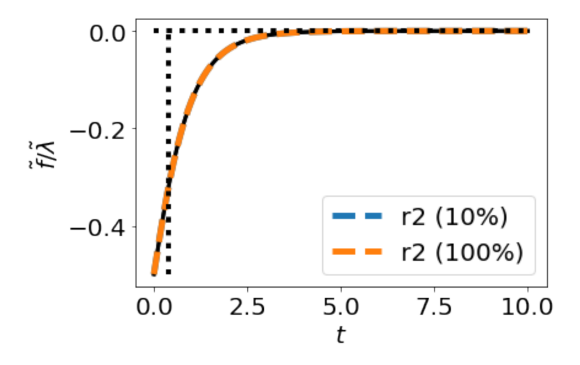
```
[120]: imp.reload(slp)
       Outp = ['r1']
       Inp_reac = ['sr1','sr2']
       def PrintSloppy(Inp,Outp,GainOnly=True,tf=None):
           blurb = '\n******\n'
           if tf is None:
               t = None
           else:
               t = np.linspace(0,tf)
           for outp in Outp:
               print(blurb,outp,blurb)
               sys = extractSubsystem(Sys,sc,sf,Inp,[outp])
                 print(con.dcgain(sys))
               gain = con.dcgain(sys)[0]
               norm = np.sum(gain*gain)
               ngain = gain/np.sqrt(norm)
                 print(norm, gain/np.sqrt(norm))
               H,eig,eigv,t = slp.Sloppy(sys,GainOnly=GainOnly,small=1e-10,t=tf)
               slp.SloppyPrint(eig,eigv,Inp,min_eig=0.0,min_eigv=0.05,max_eigs=2)
               if GainOnly:
                   print('Direct')
                   slp.SloppyPrint([norm],np.array([ngain]).T,Inp,min_eig=0.
        \rightarrow0,min_eigv=0.05,max_eigs=2)
                 print(eigv[:,0])
                 print(ngain)
       ## Reactions
```

```
for tf in [None, 1e2]:
             for GainOnly in [True,False]:
                        print('\nGainOnly =',GainOnly,'; tf =',tf)
                        PrintSloppy(Inp_reac,Outp,GainOnly=GainOnly,tf=tf)
GainOnly = True ; tf = None
*****
  r1
*****
\qquad \ensuremath{\mbox{ }} \ \sqrt\sigma_2 &= 1e-05 & V_2\Lambda &= + 0.832 \lambda_{r1} - 0.555 \lambda_{r2}
Direct
GainOnly = False ; tf = None
*****
*****
\qquad \ \sqrt\sigma_1 &= 1.4 & V_1\Lambda &= + 0.787 \lambda_{r2} + 0.617 \lambda_{r1}
\sqrt\sigma_2 \&= 0.29 \& V_2\Lambda \&= + 0.787 \lambda_{r1} - 0.617 \lambda_{r2}
GainOnly = True ; tf = 100.0
*****
  r1
******
\qquad \ \sqrt\sigma_1 &= 1.4 & V_1\Lambda &= + 0.832 \lambda_{r2} + 0.555 \lambda_{r1}
\qquad \ensuremath{\mbox{\mbox{\mbox{$\sim$}}} \ensuremath{\mbox{\mbox{$\sim$}}} \ensuremath{\mbox{$\sim$}} \ensur
Direct
\sqrt sqrt \frac{1.4 \& V_1\Delta \& = + 0.832 \ada_{r2} + 0.555 \ada_{r1}}
GainOnly = False ; tf = 100.0
*****
  r1
*****
\qquad \ \sqrt\sigma_1 &= 1.4 & V_1\Lambda &= + 0.830 \lambda_{r2} + 0.558 \lambda_{r1}
\sqrt\sigma_2 \&= 0.069 \& V_2\Lambda \&= + 0.830 \lambda_{r1} - 0.558 \lambda_{r2}
```

5.4.9 Initial condition sensitivity.

```
[121]: ## Set up deviation state for full system
       n_X = sc['n_X']
       sX0 = np.zeros(n_X)
       spec = sc['species']
       i_C = spec.index('C')
       i_E = spec.index('E')
       i_E0 = spec.index('E0')
       sX0[i_E0] = 1
       sX0[i_E] = -0.5
       sX0[i_C] = -0.5
       ## Test conserved moiety compatability
       G_c = sc['G']
       error = np.linalg.norm(G_c@sX0)
       print('Compatibility error =',error)
       ## Compare linear and non-linear
       inp = 'sr1'
       outp = ['r2']
       for lam in [1.1,2,]:
           dat,y_lin,t,sys = simSensitivity(s,sc,sf,Sys,X_ss,V_ss,dX_ss,
                                              sX0=sX0,
                                             parameter=parameter,inp=[inp],outp=outp,
                                             lam=lam,t_last=10,lambdaIn=lambdaIn)
            print(con.dcgain(sys))
           g = con.dcgain(sys)
            if Titles:
                 plt.title(f'{inp} (g = {g:.3f})')
           plotSensitivity(dat,reactions=outp)
           g=0;plotLines()
       Savefig('ecr_IC')
```

Compatibility error = 0.0



[]:

References