# Oscillation

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## Contents

1	1.1 Select system to be analysed	3
2	Submodels	13
3	Feedback system 3.1 Stoichiometry	28 29
4	Open-loop analysis 4.1 Derive linear closed-loop response	<b>30</b> 34
5	Investigate parameter dependencies	35
6	Closed-loop analysis 6.1 Compare two versions of closed-loop TF 6.2 Linear time response 6.3 Non-linear simulation 6.4 Phase plane 6.5 Signals at integrator 6.6 Period 6.7 Period 6.8 Compare two versions of closed-loop TF 6.9 Compare two versions of closed-loop TF 6.0 Compare two versions of closed-loop TF 6.1 Compare two versions of closed-loop TF 6.2 Compare two versions of closed-loop TF 6.3 Compare two versions of closed-loop TF 6.4 Compare two versions of closed-loop TF 6.5 Compare two versions of closed-loop TF 6.6 Compare two versions of closed-loop TF 6.7 Compare two ve	38 40 41 41 44 46 47
7	Power	48
1	7.1 Power	49
8	Split-loop analysis 8.1 Model 8.2 Stoichiometry 8.3 Steady-state analysis 8.4 Linearise 8.5 Active and passive loop gains 8.6 Bode plots 8.7 Nichols plots 8.8 Root Locus 8.9 Root Locus - active only 8.10 Sisotool 8.11 Sisotool - active only	544 545 555 566 588 599 600 611 622 633
9	Open-loop analysis of dynamic part of Toy and Goodwin examples	68
	Linear + saturation	69

10.1	Signals at integrator		•				•	•				 	•				74	

#### 1 Introduction

This Jupyter notebook (Oscillation.ipynb) contains the code used to generate the following examples for the paper "Analysis of Biochemical Oscillators Using Bond Graphs and Linear Control Theory" by Peter Gawthrop and Michael Pan:

- 3. Illustrative Example (system Toy)
- 4. The Sel'kov Oscillator (system Selkov)

The example - 5. The Repressilator is in the notebook Repressilator.ipynb

#### 1.1 Select system to be analysed

```
[1]: | ## Select system to be analysed
     ## NB Repressilator example is in a separate notebook:
     \rightarrow Repressilator, Repressilator. ipynb
     SystemName = 'Toy'
     ToyVersion = 'basic'
     # ToyVersion = 'varyAct'
     # ToyVersion = 'varyR'
     # SystemName = 'Toycc'
     # SystemName = 'Toy1'
     # SystemName = 'ToyTwo'
     # SystemName = 'ToyTwo3'
     # SystemName = 'ToyTwo4'
     # SystemName = 'Goodwin'
     # SystemName = 'Selkov'
     # SelkovVersion = 'basic'
     # # SelkovVersion = 'varyATP'
     # SelkovVersion = 'varyR'
     # SystemName = 'Selkov1'
     # SystemName = 'Selkov3'
```

```
[2]: if SystemName in ['Toy']:
    if ToyVersion in ['basic']:
        K_Act = 1
        kappa_rf = 10
        SysName = 'Toy'
    elif ToyVersion in ['varyAct']:
        K_Act = 0.2
        kappa_rf = 10
        SysName = 'Toymod'
    elif ToyVersion in ['varyR']:
        K_Act = 1
        kappa_rf = 25
        SysName = 'ToymodR'
    else:
        print('ToyVersion', ToyVersion, 'not known.')
```

```
print('K_Act=', K_Act)
elif SystemName in ['Selkov']:
    if SelkovVersion in ['basic']:
        v_ATP = 0.6
        kappa_rf = 10
        SysName = 'Selkov'
    elif SelkovVersion in ['varyATP']:
        v_ATP = 0.7
        kappa_rf = 10
        SysName = 'Selkovmod'
    elif SelkovVersion in ['varyR']:
        v_ATP = 0.6
        kappa_rf = 8
        SysName = 'SelkovmodR'
    else:
        print('SelkovVersion', SelkovVersion,'not known.')
else:
    SysName = SystemName
    v_ATP = 0.6
    K_Act = 1
   kappa_rf = 10
print(SysName)
```

Toy

```
[3]: ## Decide whether to compute parametric variation results - takes a long time!

ParametricVariation = False

## Save data for comparative plots and for printing equations and parameters.

SavingData = False

SaveData = SysName in

→['Toy','Toy1','Toycc','ToyTwo','ToyTwo3','ToyTwo4','Toymod','ToymodR',

'Selkov','Selkovmod','SelkovmodR','Selkov1','Selkov3']

SaveData = SaveData and SavingData

if SaveData:

SavedData = {}

print("Saving data = ",SaveData)
```

Saving data = False

#### 1.2 Import some python code

The bond graph analysis uses a number of Python modules:

```
[4]: ## For path etc: sys.path
import sys
sys.path.append("/home/peterg/WORK/Research/SystemsBiology/lib/python")
```

```
[5]: ## Some useful imports
import BondGraphTools as bgt
import numpy as np
import sympy as sym
```

```
import scipy.optimize as opt
     import scipy.integrate as integrate
     import matplotlib.pyplot as plt
     from cycler import cycler
     import IPython.display as disp
     import copy
     ## 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
     con.config.defaults['xferfcn.display_format'] = 'zpk'
     import slycot
     ## Stoichiometry to BG
     import stoichBondGraph as stbg
     ## Set slycot=True if slycot is installed (see control module)
     slycot=False
     ## For reimporting: use imp.reload(module)
     import importlib as imp
     # Allow output from within functions
     from IPython.core.interactiveshell import InteractiveShell
     InteractiveShell.ast_node_interactivity = "all"
     ## Saving data
     import pickle
     ## Set quiet=False for verbose output
     quiet = True
     ## Plot figure to folder Figs
     Plotting = False
[6]: def printSS(s,x_ss,parameter):
         print('\n## Steady state')
         for i,spec in enumerate(s['species']):
             # print(i,spec,x_ss[i])
             val = x_ss[i]
             if not (val==1):
                 print(f'x_{{spec}}) &= {val:.2g}///')
```

```
print('\n## Parameters')
          for par in parameter:
              val = parameter[par]
              if not (val==1):
                  print(f'{par} & {val:.2g}\\\')
 [7]: def zero_crossings(a):
          """Zero crossings from positive to negative
          return np.where(np.diff(np.sign(a))>0)[0]
 [8]: def SetPlot(fontsize=14,linewidth=5,RL=False):
              ## Sizes
              plt.rcParams.update({'font.size': fontsize})
              plt.rcParams.update({'lines.linewidth': linewidth})
              plt.rcParams.update({'lines.markersize': 6*linewidth})
              ## set up colour cycling for plot
              if RL:
                  ## Root locus colors
                  default_cycler = (cycler(color=['grey', 'grey', 'r', 'g', 'b']))
              else:
                  default_cycler = (cycler(color=['r', 'g', 'b']))
              plt.rc('axes', prop_cycle=default_cycler)
      SetPlot()
 [9]: def SaveFig(SystemName,PlotName,fontsize=14,linewidth=5,RL=False):
          if Plotting:
              SetPlot(RL=RL)
              plotname = f'Figs/{SystemName}_{PlotName}.pdf'
              plt.tight_layout()
              plt.savefig(plotname)
[10]: def zpk(tf,display_format='zpk'):
          p = np.real(con.poles(tf))
          z = np.real(con.zeros(tf))
          return con.zpk(z,p,1)
[11]: def balred(sys,n_red):
          if n_red<sys.nstates:</pre>
              return con.balred(sys,n_red)
          else:
              return sys
[12]: def printTF(tf):
          return tf
[13]: def Properties(s,sc):
          ## Pathways
          print('Paths')
```

```
print(st.sprintp(sc))
          ## Conserved Moieties (Pools)
          print('Pools:')
          disp.Latex(st.sprintml(s))
[14]: def SetChemostatsToy(N=3,Config='Closed',quiet=False):
          print(f'Setting feedback loop with configuration {Config}')
          chemostats = ['fb_Act','fb_E0']
          if Config in ['Closed']:
              chemostats = chemostats
          elif Config in ['Open']:
              chemostats.append('P')
          elif Config in ['Dynamic']:
              chemostats += ['E1']
          elif Config in ['SplitLoop']:
              chemostats += ['Pf','P']
          else.
              print(f'Config={Config} not recognised')
          for i in range(1,N+1):
              # for comp in ['A', 'Z', 'Zf']:
              for comp in ['A', 'Zf']:
                  chemostats.append(f'decr{i}_{comp}')
          return chemostats
[15]: def SetChemostatsGoodwin(N=3,Config='Closed',quiet=False):
          print(f'Setting feedback loop with configuration {Config}')
          chemostats = ['fb_Act','fb_E0']
          if Config in ['Closed']:
              chemostats = chemostats
          elif Config in ['Open']:
              chemostats.append('P')
          elif Config in ['Dynamic']:
              chemostats += ['E1']
          elif Config in ['SplitLoop']:
              chemostats += ['Pf','P']
          else:
              print(f'Config={Config} not recognised')
          for i in range(1,N+1):
              for comp in ['A', 'Z', 'Zf']:
                  chemostats.append(f'decr{i}_{comp}')
          return chemostats
[16]: def SetChemostatsSelkov(Config='Closed',quiet=False):
```

```
print(f'Setting feedback loop with configuration {Config}')
          chemostats = []
          if Config in ['Closed']:
              chemostats = chemostats
          elif Config in ['Open']:
              chemostats.append('P')
          elif Config in ['SplitLoop']:
              chemostats += ['Pf','P']
          else:
              print(f'Config={Config} not recognised')
          for comp in ['ATPO', 'Z', 'Zf']:
              chemostats.append(f'selkov_{comp}')
          return chemostats
[17]: | # def SetChemostatsRepressilator(Config='Closed', quiet=False):
            print(f'Setting feedback loop with configuration {Config}')
            chemostats = ['A', 'G1\_XM', 'G1\_XP', 'G2\_XM', 'G2\_XP', 'G3\_XM', 'G3\_XP']
            if Config in ['Closed']:
      #
                chemostats = chemostats
            elif Config in ['Open']:
      #
      #
                chemostats.append('P3')
            elif Config in ['SplitLoop']:
                chemostats += ['P3f', 'P3']
      #
      #
                print(f'Config={Config} not recognised')
            return chemostats
[18]: def SetChemostats(SystemName, Config='Closed', quiet=False):
          if SystemName in ['Toy', 'Toy1', 'Toycc']:
              chemostats = SetChemostatsToy(N=3,Config=Config)
          elif SystemName in ['ToyTwo','ToyTwo3','ToyTwo4']:
              chemostats = SetChemostatsToy(N=2,Config=Config)
          elif SystemName in ['Goodwin']:
              chemostats = SetChemostatsGoodwin(N=3,Config=Config)
          elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
              chemostats = SetChemostatsSelkov(Config=Config)
          # elif SystemName in ['Repressilator']:
                chemostats = SetChemostatsRepressilator(Config=Config)
          else:
              print('System Name', SystemName, 'is not known')
          return chemostats
[19]: def stoichiometry(abg,chemostats=[]):
            print('Comp s')
          s = st.stoich(abg,quiet=quiet)
          print('comp sc')
```

```
sc = st.statify(s,chemostats=chemostats)
          return s,sc
[20]: def SetParameterToy(N=3,kappa_rf=1e1,K_Act=1,K_Zf=1e-6): ## K_Zf = 0.2,1e-6
          parameter = {}
          parameter['K_fb_Act'] = K_Act
          for i in range(1,N+1):
              \# parameter[f'K_decr\{i\}_Z'] = 1 \# 1e-6
              parameter[f'K_decr{i}_Zf'] = K_Zf
              parameter[f'kappa_decr{i}_r'] = 1e0
              parameter[f'kappa_decr{i}_rf'] = kappa_rf
              parameter[f'K_decr{i}_A'] = 1e2
              parameter[f'K_E{i}'] = 1
          return parameter
[21]: def SetParameterGoodwin(N=3,kappa_rf=1e1,K_Act=1,K_Z=1e-6):
          parameter = {}
          parameter['K_fb_Act'] = K_Act
          for i in range(1,N+1):
              parameter[f'K_decr{i}_Z'] = parameter[f'K_decr{i}_Zf'] = K_Z
                  parameter[f'kappa_decr{i}_r'] = 1e0
              else:
                  parameter[f'kappa_decr{i}_r1'] = 1e0
                  parameter[f'kappa_decr{i}_r2'] = 1e6
                  parameter[f'K_decr{i}_C'] = 1e-3
              parameter[f'kappa_decr{i}_rf'] = kappa_rf
              parameter[f'K_decr{i}_A'] = 1e2
          return parameter
[22]: def SetParameterSelkov(subname='_selkov',v_ATP=0.0,kappa_rf=1e1):
          parameter = {}
           parameter['kappa_rd'] = 1
          parameter[f'K{subname}_Z'] = parameter[f'K{subname}_Zf'] = 1e-10
          parameter[f'kappa{subname}_r0'] = 1e3
          parameter[f'kappa{subname}_r1'] = 1e3
         parameter[f'kappa{subname}_r2'] = 1e3
          parameter[f'kappa{subname}_rf'] = kappa_rf
          parameter[f'K{subname}_PFK'] = 1
          parameter[f'K{subname}_C'] = 1
           Large = 1e2
         Large = 1e3
          parameter[f'K{subname}_ATPO'] = Large
          parameter[f'kappa{subname}_rs'] = v_ATP/Large
```

```
return parameter
[23]: def SetParameter():
          if SystemName in ['Toy', 'Toy1', 'Toycc']:
              parameter = SetParameterToy(N=3,K_Act=K_Act,kappa_rf=kappa_rf)
          elif SystemName in ['ToyTwo', 'ToyTwo3', 'ToyTwo4']:
              parameter = SetParameterToy(N=2,K_Act=K_Act,kappa_rf=kappa_rf)
          elif SystemName in ['Goodwin']:
              parameter = SetParameterGoodwin()
          elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
              parameter = SetParameterSelkov(v_ATP=v_ATP,kappa_rf=kappa_rf)
          elif SystemName in ['Repressilator']:
              parameter = SetParameterRepressilator()
          else:
              print('System Name',SystemName,'is not known')
          return parameter
[24]: def SetAll(SystemName, Config='Closed', quiet=False):
          if SystemName in ['Toy','Toy1','Toycc']:
              chemostats = SetChemostatsToy(N=3,Config=Config)
              parameter = SetParameterToy(N=3,K_Act=K_Act,kappa_rf=kappa_rf)
              InpVar = 'fb_Act'
              OutpVar = 'P'
              T = np.linspace(0,4,1000)
              T_{long} = np.linspace(0,10,5000)
              n_red = 2
```

```
elif SystemName in ['ToyTwo', 'ToyTwo3', 'ToyTwo4']:
    chemostats = SetChemostatsToy(N=2,Config=Config)
    parameter = SetParameterToy(N=2,K_Act=K_Act,kappa_rf=kappa_rf)
    InpVar = 'fb_Act'
    OutpVar = 'P'
    T = np.linspace(0,4,1000)
    T_{long} = np.linspace(0,10,5000)
    n_red = 2
elif SystemName in ['Goodwin']:
    chemostats = SetChemostatsGoodwin(N=3,Config=Config)
    parameter = SetParameterGoodwin(N=3)
    InpVar = 'fb_Act'
    OutpVar = 'P'
    T = np.linspace(0,5,1000)
    T_{long} = np.linspace(0,20,5000)
    n_red = 3
elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
    chemostats = SetChemostatsSelkov(Config=Config)
    parameter = SetParameterSelkov(v_ATP=v_ATP,kappa_rf=kappa_rf)
    InpVar = 'selkov_ATPO'
    OutpVar = 'P'
    T = np.linspace(0,5,1000)
```

```
T_long = np.linspace(0,50,5000)
    n_red = 2 ## Order reduction
# elif SystemName in ['Repressilator']:
# chemostats = SetChemostatsRepressilator(Config=Config)
# parameter = SetParameterRepressilator()
# InpVar = 'A'
# OutpVar = 'P3'
# T = np.linspace(0,5,100)
# T_long = np.linspace(0,50,5000)
else:
    print('System Name',SystemName,'is not known')

return chemostats,parameter,InpVar,OutpVar,T,T_long,n_red
```

```
[25]: def extractSysflow(Sys,s,chemo,chemostats,reac):
          ## Index of reaction
          reaction = s['reaction']
          i_v = reaction.index(reac)
          ## Index of input
          i = chemostats.index(chemo)
          sys = con.ss(Sys.A,Sys.B[:,i],Sys.C[i_v,:],Sys.D[i_v,i])
          return sys
      def extractSysdX(Sys,s,chemo,chemostats,outp,tol=None):
          ## Index of output
          species = s['species']
          i_v = species.index(outp)
          ## Index of input
          i = chemostats.index(chemo)
          sys = con.ss(Sys.A,Sys.B[:,i],Sys.C[i_v,:],Sys.D[i_v,i])
          return con.minreal(sys,tol=tol)
```

```
ld = len(den0)
              num = num0
              den = np.zeros(ld+1)
              den[:1d] = den0
          L = con.tf(num,den)
          return L
[27]: def Lin(s,sc,parameter=None,x_ss=None,outvar='dX',Inp=['P','E'],__
       →Outp=['P','E'],quiet=True):
          ## Linearise
          SYS = st.lin(s,sc,x_ss=x_ss,parameter=parameter,outvar='dX',quiet=quiet)
          # Extract individual transfer functions
          TF = \{\}
          Sys = {}
          for inp in Inp:
              for outp in Outp:
                  if not quiet:
                      print(inp,'-->',outp)
                  sys = extractSysdX(SYS,s,inp,chemostats,outp)
                  tf = con.tf(sys)
                  Sys[f'{inp}_{outp}'] = sys
                  TF[f'\{inp\}_{outp}'] = tf
                  if not quiet:
                      print(tf)
          return TF,Sys
[28]: def step_response(sys,T=None):
          resp = con.step_response(sys,T=T)
          t = resp.t
          y = np.array(resp.y).flatten()
          plt.plot(t,y)
          return y
[29]: def impulse_response(sys,T=None):
          resp = con.impulse_response(sys,T=T)
          t = resp.t
          y = np.array(resp.y).flatten()
          plt.plot(t,y)
          return y
[30]: def SteadyState(s,sc,parameter,x0,OutpVar='P',returnAll=False):
          t = np.linspace(0,1e4)
          ndat = st.sim(s,sc=sc,t=t,parameter=parameter,X0=x0,quiet=True)
          x_ss = ndat['X'][-1,:]
          ## Flow into P
```

```
species = s['species']
    v_ss = ndat['dX'][-1,species.index(OutpVar)]
    if returnAll:
        SS = \{\}
        ## Save up all steady-state data
        for key in ndat:
            if not key in ['t']:
                  print(key)
                SS[key] = ndat[key][-1,:]
        return x_ss,v_ss,SS
    else:
        return x_ss,v_ss
def func(x_P):
    x0 = np.ones(s['n_X'])
    x0[species.index(OutpVar)] = x_P
    x_ss,v_ss = SteadyState(s,sc,parameter,x0,OutpVar=OutpVar)
    return v_ss
def findSteadyState(s,sc,parameter,x0,OutpVar='P',returnAll=False):
    species = s['species']
    root = opt.fsolve(func,1)
    x_P_s = root[0]
    x0[species.index(OutpVar)] = x_P_ss
    if returnAll:
        x_s, v_s, S = 
 →SteadyState(s,sc,parameter,x0,OutpVar=OutpVar,returnAll=returnAll)
        return x_ss,SS
    else:
        x_ss,v_ss =
 →SteadyState(s,sc,parameter,x0,OutpVar=OutpVar,returnAll=returnAll)
        return x_ss,x_P_ss
```

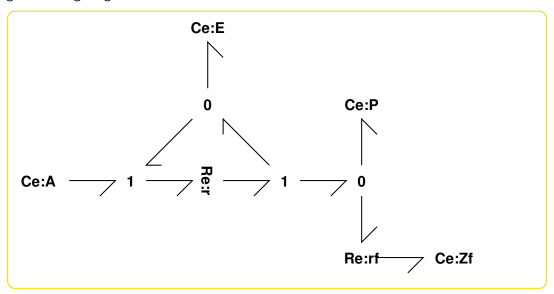
#### 2 Submodels

```
- decr: basic enzyme reaction with degredation
- decrc: basic enzyme reaction with degredation and cooperativity
- dECR: enzyme reaction with intermediate complex and degredation
- dECRc: enzyme reaction withintermediate complex and degredation and cooperativity
- ActInh: activation/inhibition module
- Selkov: The Sel'kov model of glycolitic oscillations from Keener and Sneyd
[31]: subTF = {}
V_ss = {}
```

```
for subname in ['decr_abg', 'decrc_abg', 'decrc3_abg', 'decrc4_abg',
                'dECR_abg', 'dECRc_abg', 'ActInh_abg', 'Selkov_abg']:
    svg = subname+'.svg'
   print('\n\nUsing',svg)
    disp.SVG(svg)
    sbg.model(svg,convertCe=True,convertR=True,quiet=quiet)
    exec(f'import {subname} as sys_abg')
    imp.reload(sys_abg)
    s = st.stoich(sys_abg.model(),quiet=quiet)
    species = s['species']
    print(species)
    disp.Latex(st.sprintrl(s,all=True))
    disp.Latex(st.sprintvl(s))
    print(st.sprintrl(s,all=True,chemformula=True))
    print(st.sprintvl(s))
    if subname not in ['ActInh_abg', 'Selkov_abg']:
        parameter={}
        chemostats = ['P','E','A','Zf']
        sc = st.statify(copy.deepcopy(s),chemostats=chemostats)
        parameter['K_A'] = 1
        # parameter['K_Z'] =
        parameter['K_Zf'] = 1e-6
        parameter['kappa_rf'] = 10
        if subname in ['dECR_abg','dECRc_abg']:
            parameter['kappa_r1'] = parameter['kappa_r2'] = 2*1000
        else:
            parameter['kappa_r'] = 1000
        Sys = st.lin(s,sc,outvar='dX',parameter=parameter)
        for inp in ['E', 'P']:
            for outp in ['E', 'P']:
                sys = extractSysdX(Sys,s,inp,chemostats,outp)
                print(inp, ' - ', outp)
                con.tf(sys)
        chemostats = ['A','Zf','P']
        sc = st.statify(copy.deepcopy(s),chemostats=chemostats)
        K = np.linspace(0.1,2)
        e0 = 0.1
        V_ss[subname] = []
        x0 = np.ones(s['n_X'])
        for K_A in K:
            parameter['K_A'] = K_A
            if subname in ['dECR_abg','dECRc_abg']:
                x0[species.index('E')] = x0[species.index('C')] = e0/2
```

Using decr\_abg.svg

[31]:



{}

[31]: <module 'decr\_abg' from '/home/peterg/WORK/Research/SystemsBiology/Notes/2024/Oscillation/decr\_abg. →py'>

['A', 'E', 'P', 'Zf']

[31]:

$$A + E \Leftrightarrow E + P \tag{1}$$

$$P \Leftrightarrow Zf$$
 (2)

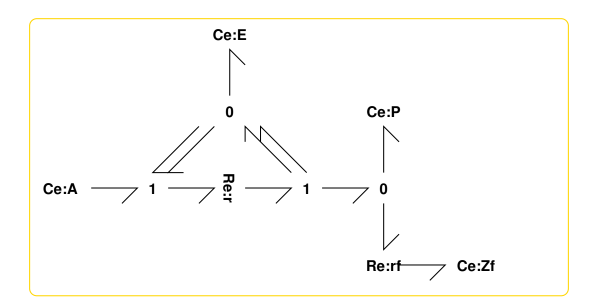
[31]:

```
v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right)
                                                                                       (4)
     \begin{align}
     \ch{A + E \& <> [r] E + P}
     \ch{P & <> [ rf ] Zf }
     \end{align}
     \begin{align}
     v_{r} &= K_{E} \times_{r} x_{E} \left(K_{A} x_{A} - K_{P} x_{P}\right) \right)
     v_{rf} &= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right)
     \end{align}
     Setting K_A to 1
     Setting K_Zf to 1e-06
     Setting kappa_r to 1000
     Setting kappa_rf to 10
     O states have been removed from the model
     E - E
     /home/peterg/anaconda3/envs/bgt/lib/python3.8/site-
     packages/scipy/signal/_filter_design.py:1746: BadCoefficients: Badly conditioned
     filter coefficients (numerator): the results may be meaningless
       warnings.warn("Badly conditioned filter coefficients (numerator): the "
     /home/peterg/anaconda3/envs/bgt/lib/python3.8/site-
     packages/scipy/signal/_filter_design.py:1091: RuntimeWarning: invalid value
     encountered in divide
       b /= b[0]
[31]:
     O states have been removed from the model
     E - P
[31]:
                                               0
                                               1
     O states have been removed from the model
     P - E
[31]:
                                               0
     O states have been removed from the model
     P - P
[31]:
                                             -1010
```

 $v_r = K_E \kappa_r x_E \left( K_A x_A - K_P x_P \right)$ 

(3)

Using decrc\_abg.svg



{}

[31]: <module 'decrc\_abg' from

 $\label{local_peterg_WORK_Research_SystemsBiology_Notes_2024_Oscillation_decrc_abg. $$ \hookrightarrow py'> $$$ 

['A', 'E', 'P', 'Zf']

[31]:

$$A + 2E \Leftrightarrow 2E + P \tag{5}$$

$$P \Leftrightarrow Zf$$
 (6)

[31]:

$$v_r = K_E^2 \kappa_r x_E^2 \left( K_A x_A - K_P x_P \right) \tag{7}$$

$$v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right) \tag{8}$$

\begin{align}

 $\ch{A + 2 E \& <> [r] 2 E + P}$ 

\ch{P & <> [ rf ] Zf }

\end{align}

\begin{align}

 $v_{r} &= K_{E}^{2} \left(K_{A} x_{A} - K_{P} x_{P}\right) \\ v_{rf} &= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right) \\ \end{align}$ 

Setting K\_A to 1

Setting K\_Zf to 1e-06

Setting kappa\_r to 1000

Setting kappa\_rf to 10

O states have been removed from the model

E - E

 $\frac{0}{1}$ 

O states have been removed from the model E  $\,$  -  $\,$  P  $\,$ 

[31]:

 $\frac{0}{1}$ 

0 states have been removed from the model  $\mathbf{p}$ 

[31]:

 $\frac{0}{1}$ 

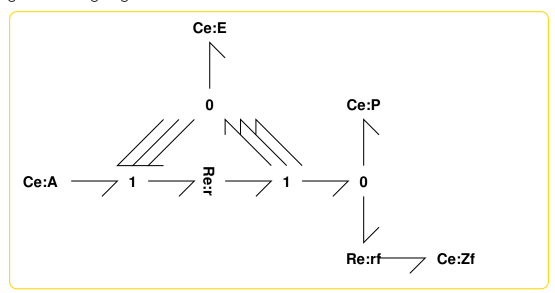
O states have been removed from the model P - P  $\,$ 

[31]:

 $\frac{-1010}{1}$ 

Using decrc3\_abg.svg

[31]:



{}

[31]: <module 'decrc3\_abg' from '/home/peterg/WORK/Research/SystemsBiology/Notes/  ${\hookrightarrow}2024/$ 

Oscillation/decrc3\_abg.py'>

['A', 'E', 'P', 'Zf']

[31]:

$$A + 3E \Leftrightarrow 3E + P \tag{9}$$

$$P \Leftrightarrow Zf \tag{10}$$

$$v_r = K_E^3 \kappa_r x_E^3 \left( K_A x_A - K_P x_P \right) \tag{11}$$

$$v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right) \tag{12}$$

\begin{align}
\ch{A + 3 E & <> [ r ] 3 E + P }\\
\ch{P & <> [ rf ] Zf }
\end{align}

\begin{align}

 $v_{r} &= K_{E}^{3} \left(K_{A} x_{A} - K_{P} x_{P}\right) \\ v_{rf} &= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right) \\ \end{align}$ 

Setting K\_A to 1
Setting K\_Zf to 1e-06
Setting kappa\_r to 1000
Setting kappa\_rf to 10
0 states have been removed from the model
E - E

[31]:

 $\frac{0}{1}$ 

O states have been removed from the model E -  $\mbox{P}$ 

[31]:

 $\frac{0}{1}$ 

O states have been removed from the model P  $\,$  -  $\,$  E

[31]:

 $\frac{0}{1}$ 

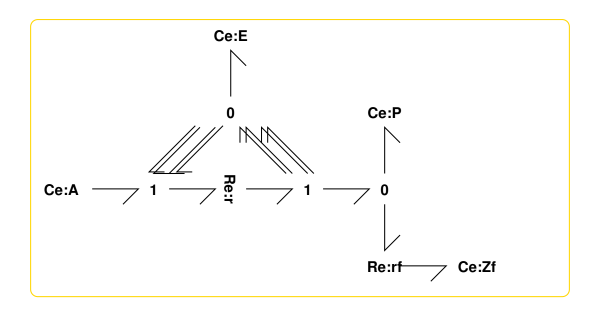
O states have been removed from the model  $P\ -\ P$ 

[31]:

 $\frac{-1010}{1}$ 

Using decrc4\_abg.svg

[31]:



{}

[31]: <module 'decrc4\_abg' from '/home/peterg/WORK/Research/SystemsBiology/Notes/  ${}_{\hookrightarrow}2024/$ 

Oscillation/decrc4\_abg.py'>

[31]:

$$A + 4E \Leftrightarrow 4E + P \tag{13}$$

$$P \Leftrightarrow Zf \tag{14}$$

[31]:

$$v_r = K_E^4 \kappa_r x_E^4 \left( K_A x_A - K_P x_P \right) \tag{15}$$

$$v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right) \tag{16}$$

\begin{align}
\ch{A + 4 E & <> [ r ] 4 E + P }\\
\ch{P & <> [ rf ] Zf }
\end{align}

\begin{align}

 $v_{r} &= K_{E}^{4} \left(K_{A} x_{A} - K_{P} x_{P}\right) \\ v_{rf} &= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right) \\ \end{align}$ 

Setting K\_A to 1
Setting K\_Zf to 1e-06
Setting kappa\_r to 1000
Setting kappa\_rf to 10
0 states have been removed from the model
E - E

 $\frac{0}{1}$ 

O states have been removed from the model E  $\,$  -  $\,$  P  $\,$ 

[31]:

 $\frac{0}{1}$ 

0 states have been removed from the model  $\mathbf{p}$   $\mathbf{r}$ 

[31]:

 $\frac{0}{1}$ 

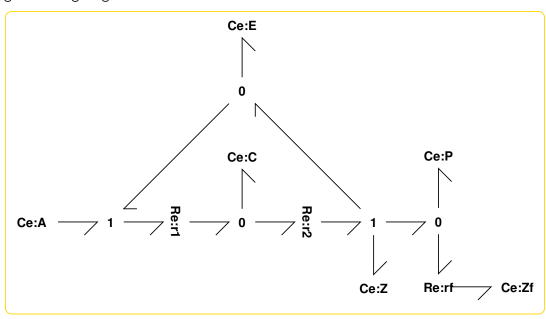
O states have been removed from the model P  $\mbox{-}$  P

[31]:

 $\frac{-1010}{1}$ 

Using dECR\_abg.svg

[31]:



{}

[31]: <module 'dECR\_abg' from

'/home/peterg/WORK/Research/SystemsBiology/Notes/2024/Oscillation/dECR\_abg.  $\leadsto$  py'>

['A', 'C', 'E', 'P', 'Z', 'Zf']

$$A + E \Leftrightarrow C \tag{17}$$

$$C \Leftrightarrow E + P + Z \tag{18}$$

$$P \Leftrightarrow Zf \tag{19}$$

[31]:

$$v_{r1} = \kappa_{r1} \left( K_A K_E x_A x_E - K_C x_C \right) \tag{20}$$

$$v_{r2} = \kappa_{r2} \left( K_C x_C - K_E K_P K_Z x_E x_P x_Z \right) \tag{21}$$

$$v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right) \tag{22}$$

\begin{align}

\ch{A + E & <> [ r1 ] C }\\

 $\ch{C \& <> [ r2 ] E + P + Z }\$ 

\ch{P & <> [ rf ] Zf }

\end{align}

\begin{align}

 $v_{r2} \&= \kappa_{r2} \ | \ v_{C} - K_{E} K_{P} K_{Z} x_{E} x_{P}$ 

 $x_{Z}\left( x_{z}\right)$ 

\end{align}

Setting K\_A to 1

Setting K\_Zf to 1e-06

Setting kappa\_r1 to 2000

Setting kappa\_r2 to 2000

Setting kappa\_rf to 10

O states have been removed from the model

E - E

[31]:

$$\frac{-4000(s - 1.431 \times 10^{-12})(s + 1000)}{(s + 763.9)(s + 5236)}$$

O states have been removed from the model

E - P

[31]:

$$\frac{-2000(s-4.367\times10^{-5})(s+4.367\times10^{-5})}{(s+763.9)(s+5236)}$$

O states have been removed from the model

P - E

[31]:

$$\frac{-2000(s - (4.657 \times 10^{-13} + 3.088 \times 10^{-5}j))(s - (4.657 \times 10^{-13} - 3.088 \times 10^{-5}j))}{(s + 763.9)(s + 5236)}$$

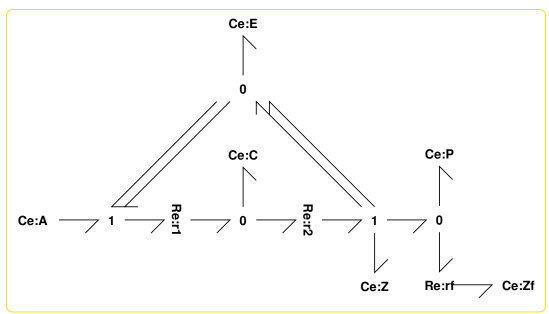
O states have been removed from the model

P - P

$$\frac{-2010(s+9.901)(s+2010)}{(s+763.9)(s+5236)}$$

Using dECRc\_abg.svg

[31]:



{}

[31]: <module 'dECRc\_abg' from

'/home/peterg/WORK/Research/SystemsBiology/Notes/2024/Oscillation/dECRc\_abg.  $\leadsto$  py'>

['A', 'C', 'E', 'P', 'Z', 'Zf']

[31]:

$$A + 2E \Leftrightarrow C \tag{23}$$

$$C \Leftrightarrow 2E + P + Z \tag{24}$$

$$P \Leftrightarrow Zf \tag{25}$$

[31]:

$$v_{r1} = \kappa_{r1} \left( K_A K_E^2 x_A x_E^2 - K_C x_C \right) \tag{26}$$

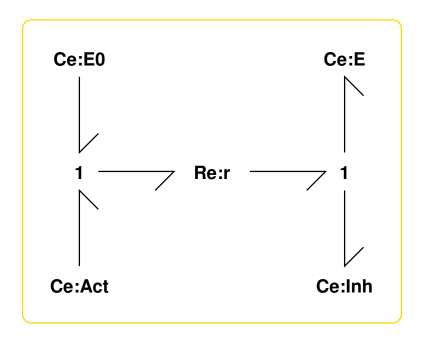
$$v_{r2} = \kappa_{r2} \left( K_C x_C - K_E^2 K_P K_Z x_E^2 x_P x_Z \right) \tag{27}$$

$$v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right) \tag{28}$$

\begin{align}
\ch{A + 2 E & <> [ r1 ] C }\\
\ch{C & <> [ r2 ] 2 E + P + Z }\\
\ch{P & <> [ rf ] Zf }
\end{align}

```
\begin{align}
                                                         v_{r1} &= \kappa_{r1} \left(K_{A} K_{E}^{2} x_{A} x_{E}^{2} - K_{C}\right)
                                                         x_{C}\rightarrow 
                                                         v_{r2} \&= \kappa_{r2} \ \text{k=ppa_{r2}} \ \text{k=ppa_{
                                                        x_{Z}\right)
                                                         v_{rf} \&= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right)
                                                         \end{align}
                                                         Setting K_A to 1
                                                         Setting K_Zf to 1e-06
                                                         Setting kappa_r1 to 2000
                                                         Setting kappa_r2 to 2000
                                                         Setting kappa_rf to 10
                                                         O states have been removed from the model
                                                        E - E
[31]:
                                                                                                                                                                                                                                                                                                                 \frac{-1.6 \times 10^4 (s - 1.431 \times 10^{-12})(s + 1000)}{(s + 763.9)(s + 5236)}
                                                        O states have been removed from the model
                                                        E - P
[31]:
                                                                                                                                                                                                                                                                                                     \frac{-4000(s - 4.367 \times 10^{-5})(s + 4.367 \times 10^{-5})}{(s + 763.9)(s + 5236)}
                                                        O states have been removed from the model
                                                        P - E
[31]:
                                                                                                                   \frac{-4000(s - (4.657 \times 10^{-13} + 3.088 \times 10^{-5}j))(s - (4.657 \times 10^{-13} - 3.088 \times 10^{-5}j))}{(s + 763.9)(s + 5236)}
                                                        O states have been removed from the model
                                                        P - P
[31]:
                                                                                                                                                                                                                                                                                                                                                                             \frac{-2010(s+9.901)(s+2010)}{(s+763.9)(s+5236)}
                                                        Using ActInh_abg.svg
```

24



{}

$$Act + E0 \Leftrightarrow E + Inh \tag{29}$$

[31]:

$$v_r = \kappa_r \left( K_{Act} K_{E0} x_{Act} x_{E0} - K_E K_{Inh} x_E x_{Inh} \right) \tag{30}$$

\begin{align}
\ch{Act + E0 & <> [ r ] E + Inh }
\end{align}
\begin{align}
v\_{r} &= \kappa\_{r} \left(K\_{Act} K\_{E0} x\_{Act} x\_{E0} - K\_{E} K\_{Inh} x\_{E} x\_{Inh}\rangle)
\end{align}

0 states have been removed from the model
Inh - Inh

[31]:

 $\frac{-1}{1}$ 

0 states have been removed from the model Inh -  $\mbox{E}$ 

$$\frac{-1}{1}$$

O states have been removed from the model E  $\,$  -  $\,$  Inh

[31]:

 $\frac{-1}{1}$ 

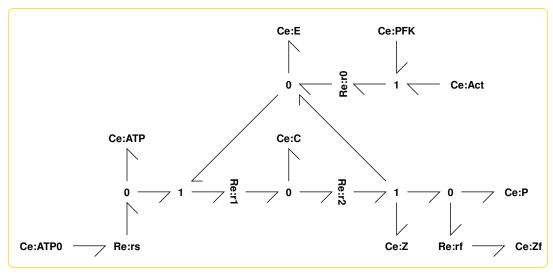
0 states have been removed from the model E - E

[31]:

 $\frac{-1}{1}$ 

Using Selkov\_abg.svg

[31]:



{}

[31]: <module 'Selkov\_abg' from '/home/peterg/WORK/Research/SystemsBiology/Notes/  ${}_{\hookrightarrow}2024/$ 

Oscillation/Selkov\_abg.py'>

['ATP', 'ATPO', 'Act', 'C', 'E', 'P', 'PFK', 'Z', 'Zf']

[31]:

$$Act + PFK \Leftrightarrow E \tag{31}$$

$$ATP + E \Leftrightarrow C \tag{32}$$

$$C \Leftrightarrow E + P + Z \tag{33}$$

$$P \Leftrightarrow Zf$$
 (34)

$$ATP0 \Leftrightarrow ATP$$
 (35)

[31]:

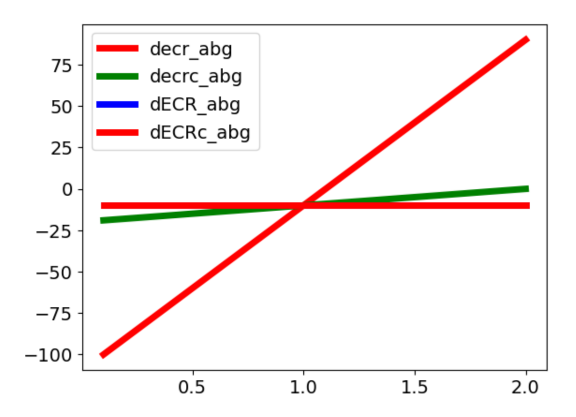
```
v_{r1} = \kappa_{r1} \left( K_{ATP} K_E x_{ATP} x_E - K_C x_C \right)
                                                                                                (37)
                                v_{r2} = \kappa_{r2} \left( K_C x_C - K_E K_P K_Z x_E x_P x_Z \right)
                                                                                                (38)
                                v_{rf} = \kappa_{rf} \left( K_P x_P - K_{Zf} x_{Zf} \right)
                                                                                                (39)
                                v_{rs} = \kappa_{rs} \left( -K_{ATP} x_{ATP} + K_{ATP0} x_{ATP0} \right)
                                                                                                (40)
      \begin{align}
      \ch{Act + PFK & <> [ r0 ] E }\\
      \ch{ATP + E & <> [ r1 ] C }\\
      \ch{C \& <> [ r2 ] E + P + Z }\
      \ch{P & <> [ rf ] Zf }\\
      \ch{ATPO & <> [ rs ] ATP }
      \end{align}
      \begin{align}
      v_{r0} &= \kappa_{r0} \ \left(K_{Act} K_{PFK} x_{Act} x_{PFK} - K_{E}\right)
      x_{E}\rangle
      v_{r1} \&= \kappa_{r1} \ (K_{ATP} K_{E} x_{ATP} x_{E} - K_{C} x_{C}\right)
      v_{r2} &= \kappa_{r2} \ \text{Left}(K_{C} x_{C} - K_{E} K_{P} K_{Z} x_{E} x_{P})
      x_{Z}\right)
      v_{rf} &= \kappa_{rf} \left(K_{P} x_{P} - K_{Zf} x_{Zf}\right)\
      v_{rs} \&= \kappa_{rs} \left( K_{ATP} x_{ATP} + K_{ATPO} x_{ATPO} \right)
      \end{align}
[32]: for subname in ['decr_abg', 'decrc_abg', 'dECR_abg', 'dECRc_abg']:
           plt.plot(K,V_ss[subname],label=subname)
       plt.legend()
[32]: [<matplotlib.lines.Line2D at 0x76d410e43430>]
[32]: [<matplotlib.lines.Line2D at 0x76d410e437c0>]
[32]: [<matplotlib.lines.Line2D at 0x76d410e439d0>]
```

 $v_{r0} = \kappa_{r0} \left( K_{Act} K_{PFK} x_{Act} x_{PFK} - K_{E} x_{E} \right)$ 

(36)

[32]: [<matplotlib.lines.Line2D at 0x76d410e43ac0>]

[32]: <matplotlib.legend.Legend at 0x76d411ba37c0>



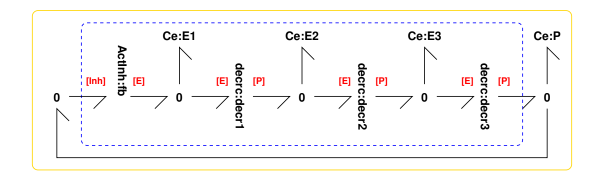
## 3 Feedback system

```
[33]: name = SystemName+'FB_abg'
    svg = name+'.svg'
    print('Using',svg)
    sbg.model(svg,convertCe=True,convertR=True,quiet=quiet)
    disp.SVG(svg)

    exec(f'import {name} as sys_abg')
    imp.reload(sys_abg)

Using ToyFB_abg.svg
    Creating subsystem: ActInh:fb
    Creating subsystem: decrc:decr1
    Creating subsystem: decrc:decr2
    Creating subsystem: decrc:decr3
    {}

[33]:
```



```
[33]: <module 'ToyFB_abg' from '/home/peterg/WORK/Research/SystemsBiology/Notes/2024/Oscillation/ToyFB_abg. →py'>
```

```
[34]: ## Set up system

# v_ATP=0.6

chemostats,parameter,InpVar,OutpVar,T,T_long,n_red = __

→SetAll(SystemName,Config='Open',quiet=False)

chemostats_open = chemostats

# print(parameter)
```

Setting feedback loop with configuration Open

#### 3.1 Stoichiometry

```
[35]: s,sc = stoichiometry(sys_abg.model(),chemostats=chemostats)
    species = s['species']
    species_open = copy.copy(species)
    reaction = s['reaction']
    print(species)
    print()
    print(reaction)
    print()
    print(parameter)

if SaveData:
        StoichData = {}
        StoichData['s'] = s
        StoichData['sc'] = sc
        StoichData['parameter'] = parameter
        SavedData['Stoich'] = StoichData
```

```
['fb_Act', 'fb_E0', 'E1', 'E2', 'E3', 'P', 'decr1_A', 'decr1_Zf', 'decr2_A',
'decr2_Zf', 'decr3_A', 'decr3_Zf']

['fb_r', 'decr1_r', 'decr1_rf', 'decr2_r', 'decr2_rf', 'decr3_r', 'decr3_rf']

{'K_fb_Act': 1, 'K_decr1_Zf': 1e-06, 'kappa_decr1_r': 1.0, 'kappa_decr1_rf': 10,
'K_decr1_A': 100.0, 'K_E1': 1, 'K_decr2_Zf': 1e-06, 'kappa_decr2_r': 1.0,
```

```
'kappa_decr2_rf': 10, 'K_decr2_A': 100.0, 'K_E2': 1, 'K_decr3_Zf': 1e-06,
      'kappa_decr3_r': 1.0, 'kappa_decr3_rf': 10, 'K_decr3_A': 100.0, 'K_E3': 1}
[36]: ## Reactions and properties
       disp.Latex(st.sprintrl(s,all=True,chemformula=False))
       # Properties(s,sc)
       print('Pools:')
       disp.Latex(st.sprintml(s))
       print('Paths')
       print(st.sprintp(sc))
[36]:
                                         fbAct + fbE0 \Leftrightarrow E1 + P
                                                                                                   (41)
                                         2E1 + decr1A \Leftrightarrow 2E1 + E2
                                                                                                   (42)
                                                    E2 \Leftrightarrow decr1Zf
                                                                                                   (43)
                                         2E2 + decr2A \Leftrightarrow 2E2 + E3
                                                                                                   (44)
                                                    E3 \Leftrightarrow decr2Zf
                                                                                                   (45)
                                         2E3 + decr3A \Leftrightarrow 2E3 + P
                                                                                                   (46)
                                                     P \Leftrightarrow decr3Zf
                                                                                                   (47)
      Pools:
[36]:
                                  fbAct \Leftrightarrow fbE0
                                                                                                   (48)
                                        \Leftrightarrow fbAct + E1
                                                                                                   (49)
                                        \Leftrightarrow E2 + decr1A + decr1Zf
                                                                                                   (50)
                                        \Leftrightarrow E3 + decr2A + decr2Zf
                                                                                                   (51)
                                        \Leftrightarrow fbAct + P + decr3A + decr3Zf
                                                                                                   (52)
      Paths
      4 pathways
      0: + decr1_r + decr1_rf
      1: + decr2_r + decr2_rf
      2: + decr3_r
      3: + decr3_rf
      4 Open-loop analysis
[37]: def OpenLoop(OutpVar='P',InpVar='fb_Act'):
            ## Steady-state analysis
            x0 = np.ones(s['n_X'])
```

→findSteadyState(s,sc,parameter,x0,OutpVar=OutpVar,returnAll=True)

 $x_ss,SS =$ 

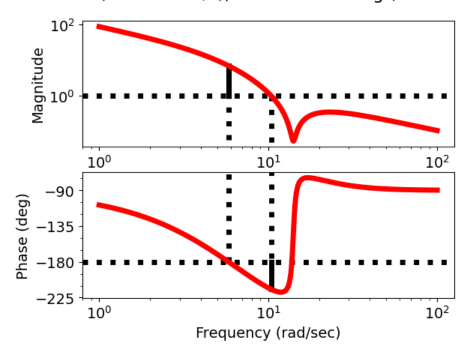
## Linearise

printSS(s,x\_ss,parameter)

```
TF,Sys_OL =
      →Lin(s,sc,parameter=parameter,x_ss=x_ss,outvar='dX',Inp=[InpVar,OutpVar],_
      →Outp=[OutpVar])
         ## Loop gain
         Lname = OutpVar+'_'+OutpVar
         L0 = -TF[Lname]
         L = IntegrateTF(L0)
         L0_{sys} = -Sys_{0L}[Lname]
         ## Forward gain
         F = TF[InpVar+'_'+OutpVar]
         return L,L0,L0_sys,F,x_ss,SS
     L,LO,LO_sys,F,x_ss,SS = OpenLoop(OutpVar=OutpVar,InpVar=InpVar)
     x_s_open = x_s
      # print(x_ss)
     ## Steady state
     x_{E1} &= 0.17 \
     x_{E2} \&= 0.28 \
    x_{E3} \&= 0.79
    x_{P} &= 5.9
     ## Parameters
    K_decr1_Zf & 1e-06\\
    kappa_decr1_rf & 10\\
    K_decr1_A & 1e+02\
    K_decr2_Zf & 1e-06\\
    kappa_decr2_rf & 10\\
    K_decr2_A & 1e+02\\
    K_decr3_Zf & 1e-06\\
    kappa_decr3_rf & 10\\
     K_decr3_A & 1e+02\\
     O states have been removed from the model
     O states have been removed from the model
 []:
[38]: print('L0:',L0)
     print(f'Gain: {con.dcgain(L0):.2f}')
     print('L:',L)
     print(f'Gain: {con.dcgain(L):.2f}')
     LO:
     10.8 (s + (0.4024-14.08j)) (s + (0.4024+14.08j)) (s + 25.15)
     _____
                (s + 5.936) (s + 10.03) (s + 10.08)
```

```
Gain: 89.77
     10.8 (s + (0.4024-14.08j)) (s + (0.4024+14.08j)) (s + 25.15)
              (s) (s + 5.936) (s + 10.03) (s + 10.08)
     Gain: inf
[39]: # print(L0)
     sysL0 = con.ss(L0)
     sysL0_r = balred(sysL0,n_red)
     L0_r = con.tf(sysL0_r)
     print('L0_r:',L0_r)
     print(f'Gain: {con.dcgain(L0_r):.2f}')
     print(f'Poles: {con.poles(L0_r)}')
     L0_r:
     10.8 (s + (1.934-14.22j)) (s + (1.934+14.22j))
     -----
       (s + (4.006-3.068j)) (s + (4.006+3.068j))
     Gain: 87.35
     Poles: [-4.00558944+3.06802235j -4.00558944-3.06802235j]
[40]: print('F:',F)
     print(f'Gain: {con.dcgain(F):.2f}')
     F:
     (s - (26.16+56.63j)) (s - (26.16-56.63j)) (s + 72.44)
              (s + 5.936) (s + 10.03) (s + 10.08)
     Gain: 469.79
[41]: print('Poles', (con.poles(L)))
     print('Zeros', (con.zeros(L)))
     Poles [-10.08007675+0.j -10.02837804+0.j -5.93620394+0.j
                                                                         +0.j]
                                                               0.
     Zeros [-25.14726161 +0.j
                                     -0.40240029+14.07807795j
       -0.40240029-14.07807795j]
[42]: mag,phase,omega=con.bode_plot(L,margins=True)
     SaveFig(SysName, 'Bode')
```

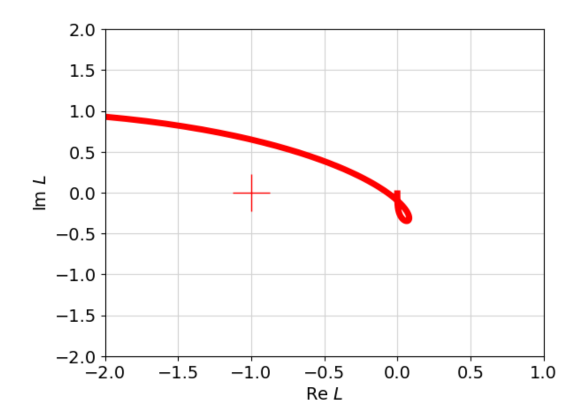
Gm = 0.14 (at 5.83 rad/s), Pm = -34.63 deg (at 10.44 rad/s)



```
[43]: # mag,phase,omega=con.nyquist_plot([L])
    con.nyquist_plot(L,mirror_style=False)
    plt.xlim(-2,1)
    plt.ylim(-2,2)
    plt.xlabel('Re $L$')
    plt.ylabel('Im $L$')
    SaveFig(SysName,'Nyquist')

[43]: 2
[43]: (-2.0, 1.0)
[43]: (-2.0, 2.0)
[43]: Text(0.5, 0, 'Re $L$')
```

[43]: Text(0, 0.5, 'Im \$L\$')



#### 4.1 Derive linear closed-loop response

- the closed-loop system is derived from the open loop system L(s)
- the input corresponds to an additive input to the product input  $f_P$ .
  - this is equivalent to the integrator 1/s in the forward path and  $L_0(s)$  in the feedback path
  - the feedback() function from the control toolbox is used.
  - a unit impulse on this input is equivalent to a unit perturbation in the *initial* product state  $x_P(0)$ .

```
[45]: ## Simple integrator 1/s
Integrator = con.tf(1,[1,0])

## Full model
```

```
CLL = con.feedback(Integrator,L0)
# CLL = con.feedback(con.series(Integrator,L0))
# CLL = con.feedback(L)
CLL_tf = con.tf(CLL)
print(CLL_tf)
print(con.poles(CLL_tf))

## Reduced model
CLL_r = con.feedback(Integrator,L0_r)
CLL_r_tf = con.tf(CLL_r)
print(CLL_r_tf)
print(CLL_r_tf)
```

### 5 Investigate parameter dependencies

```
[46]: if ParametricVariation:
         if SystemName in_
      RR = np.linspace(1,30,50)
            xlabel = r'$\kappa_{rf}$'
            ACT = [0.2, 1, 5]
            ActName = '$K_{Act}$'
         elif SystemName in ['Goodwin']:
            RR = np.linspace(1,10,10)
            xlabel = r'$\kappa_{rf}$'
            ACT = [0.5, 1, 10]
            ActName = '$K_{Act}$'
         elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
            RR = np.linspace(0.15, 1, 50)
            xlabel = '$v_{ATP}$'
            ACT = [8,10,12] # used as kappa_rf
            ActName = '$\kappa_{rf}$'
         ylabel = 'Pole'
```

```
\# RR = []
   \# ACT = []
   PPM = \{\}
   XX_P_ss = {}
   FREQ = \{\}
   POLES = \{\}
   MAXrpole = {}
   MAXipole= {}
   for Act in ACT:
       Freq = []
       PM = []
       Maxrpole = []
       Maxipole = []
       X_P_ss = []
       for rr in RR:
           print(f'rr = {rr:0.2f}')
           if SystemName in ['Toy', 'Toy1']:
               parameter = SetParameterToy(N=3,kappa_rf=rr,K_Act=Act)
           elif SystemName in ['ToyTwo','ToyTwo3','ToyTwo4']:
               parameter = SetParameterToy(N=2,kappa_rf=rr,K_Act=Act)
           elif SystemName in ['Goodwin']:
               parameter = SetParameterGoodwin(kappa_rf=rr,K_Act=Act)
           elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
               parameter = SetParameterSelkov(v_ATP=rr,kappa_rf=Act)
           else:
               print('SystemName', SystemName, 'not known')
           LL,LLO,LLO_sys,FF,xx_ss,SS =_
→OpenLoop(OutpVar=OutpVar,InpVar=InpVar)
           ## Closed-loop properties
           Integrator = con.tf(1,[1,0])
           CL = con.feedback(Integrator, LL0)
                 CL = con.minreal(CL)
           gmi, pmi, wcgi, wcpi = con.margin(LL)
           freq = wcpi/(2*np.pi)
           Freq.append(freq)
           PM.append(pmi)
           poles = con.poles(CL)
           if rr == min(RR):
               Poles = poles
           else:
               Poles = np.vstack((Poles,poles))
       #
                 print(poles)
           maxrpole = max(np.real(poles))
                 print(f'Max real pole = {maxpole:0.2f}')
           maxipole = max(np.imag(poles))
           Maxrpole.append(maxrpole)
           Maxipole.append(maxipole)
```

```
## Steady-state properties
                  x_P_ss = SS['X'][species.index('P')]
                  X_P_ss.append(x_P_ss)
              PPM[Act] = PM
              XX_P_ss[Act] = X_P_ss
              FREQ[Act] = Freq
              POLES[Act] = Poles
              MAXrpole[Act] = Maxrpole
              MAXipole[Act] = Maxipole
[47]: if ParametricVariation:
          ## CL poles
          for Act in ACT:
              plt.plot(RR,MAXrpole[Act],label=f'real: {ActName} = {Act}')
              plt.plot(RR,MAXipole[Act],label=f'imag: {ActName} = {Act}')
          plt.grid()
          plt.legend()
          plt.xlabel(xlabel)
          plt.ylabel(ylabel)
          SaveFig(SysName, 'RePole')
[48]: if ParametricVariation:
          ## CL poles - root -locus
          for Act in ACT:
              Poles = POLES[Act]
              plt.plot(np.real(Poles),np.imag(Poles),label=f'{ActName} = {Act}')
          if SystemName in ['Toy']:
              plt.xlim(-1,5)
          else:
              plt.xlim(-4,4)
          plt.ylim(bottom=0)
          plt.grid()
          # plt.legend()
          plt.xlabel('Real')
          plt.ylabel('Imag')
          SaveFig(SysName, 'RL')
[49]: if ParametricVariation:
          ## Steady states
          for Act in ACT:
              plt.semilogy(RR,XX_P_ss[Act],label=f'{ActName} = {Act}')
          plt.grid()
          plt.legend()
          plt.xlabel(xlabel)
          plt.ylabel(r'$\bar{x}_P$')
          SaveFig(SysName, 'Pss')
```

```
[50]: if ParametricVariation:
    ## Phase margin
    for Act in ACT:
        plt.plot(RR,PPM[Act],label=f'{ActName} = {Act}')
    plt.grid()
    plt.legend()
    plt.xlabel(xlabel)
    plt.ylabel('$\phi_m^\circ$')
    SaveFig(SysName,'PM')
[51]: if ParametricVariation:
### Phase margin for your and the property of the younger.
### Phase margin for your and the property of the younger.
### Phase margin for younger.
```

```
[51]: if ParametricVariation:
    ## Phase margin frequncy
    for Act in ACT:
        plt.plot(RR,FREQ[Act],label=f'{ActName} = {Act}')
    plt.grid()
    plt.legend()
    plt.xlabel(xlabel)
    plt.ylabel('$F_m$ Hz')
    SaveFig(SysName,'Fm')
```

## 6 Closed-loop analysis

- The linearised closed-loop system is obtained by linearising the non-linear system from the bond graph
- the transfer function is from the activation (Act) to the product (P)
- the transfer function is compared with that deduced from the loop gain L(s)
  - the denominators are the same
  - the numerators are different as the inputs are different

```
[52]: ## Stoichiometry
      chemostats,parameter,InpVar,OutpVar,T,T_long,n_red =_

→SetAll(SystemName, Config='Closed', quiet=False)
      s,sc = stoichiometry(sys_abg.model(),chemostats=chemostats)
      ## Linearise
      print(InpVar)
      X_ss = copy.copy(x_ss)
      TF,Sys = Lin(s,sc,parameter=parameter,x_ss=X_ss,outvar='dX',Inp=[InpVar],_
      →Outp=['P'])
      ## Extract closed-loop transfer-function
      CL0 = TF[InpVar+'_P']
      print(CL0)
      CL = IntegrateTF(CL0,crite=0.1)
      print(CL)
            return CL,s,sc
      print(chemostats)
```

Setting feedback loop with configuration Closed

```
fb_Act
     O states have been removed from the model
             (s - (26.16+56.63j)) (s - (26.16-56.63j)) (s + 4.22e-15) (s + 72.44)
      ______
      (s - (1.326+10.32j)) (s - (1.326-10.32j)) (s + (19.75-10.35j)) (s +
      (19.75+10.35j))
                      (s - (26.16+56.63j)) (s - (26.16-56.63j)) (s + 72.44)
      (s - (1.326+10.32j)) (s - (1.326-10.32j)) (s + (19.75-10.35j)) (s +
      (19.75+10.35j)
      ['fb_Act', 'fb_E0', 'decr1_A', 'decr1_Zf', 'decr2_A', 'decr2_Zf', 'decr3_A',
      'decr3_Zf']
[53]: ## Reactions and properties
      disp.Latex(st.sprintrl(s,all=True,chemformula=False))
      # Properties(s,sc)
      print('Pools:')
      disp.Latex(st.sprintml(s))
      print('Paths')
      print(st.sprintp(sc))
[53]:
                                      fbAct + fbE0 \Leftrightarrow E1 + P
                                                                                            (53)
                                      2E1 + decr1A \Leftrightarrow 2E1 + E2
                                                                                            (54)
                                                E2 \Leftrightarrow decr1Zf
                                                                                            (55)
                                     2E2 + decr2A \Leftrightarrow 2E2 + E3
                                                                                            (56)
                                                E3 \Leftrightarrow decr2Zf
                                                                                            (57)
                                      2E3 + decr3A \Leftrightarrow 2E3 + P
                                                                                            (58)
                                                 P \Leftrightarrow decr3Zf
                                                                                            (59)
     Pools:
[53]:
                               fbAct \Leftrightarrow fbE0
                                                                                            (60)
                                     \Leftrightarrow fbAct + E1
                                                                                            (61)
                                     \Leftrightarrow E2 + decr1A + decr1Zf
                                                                                            (62)
                                     \Leftrightarrow E3 + decr2A + decr2Zf
                                                                                            (63)
                                     \Leftrightarrow fbAct + P + decr3A + decr3Zf
                                                                                            (64)
```

Paths

3 pathways

0: + decr1\_r + decr1\_rf

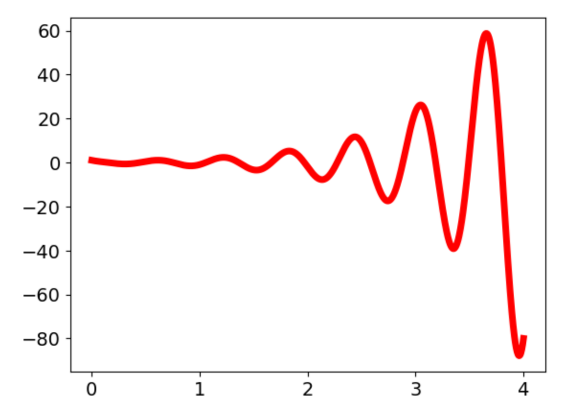
```
1: + decr2_r + decr2_rf
2: + decr3_r + decr3_rf
```

#### 6.1 Compare two versions of closed-loop TF

```
[54]: # CL,s,sc = ClosedLoopTF(SystemName,sys_abq.model(),InpVar,parameter,x_ss)
     print('CLL:',CLL)
     print('CL:',CL)
     print(f'Gain: {con.dcgain(CL):.2f}')
    CLL:
                          (s + 5.936) (s + 10.03) (s + 10.08)
    (s - (1.326+10.32j)) (s - (1.326-10.32j)) (s + (19.75-10.35j)) (s +
    (19.75+10.35j))
    CL:
                  (s - (26.16+56.63j)) (s - (26.16-56.63j)) (s + 72.44)
    ______
    (s - (1.326+10.32j)) (s - (1.326-10.32j)) (s + (19.75-10.35j)) (s +
    (19.75+10.35j))
    Gain: 5.23
[55]: print('Poles:', (con.poles(CL)))
    Poles: [-19.74856549+10.35123185j -19.74856549-10.35123185j
       1.3264659 +10.3244805j 1.3264659 -10.3244805j ]
[56]: CLr = balred(con.ss(CL),3)
     print(con.tf(CLr))
     poles = con.poles(CLr)
     print('Poles:', poles)
       3.994 (s - (30.61+39.16j)) (s - (30.61-39.16j))
    _____
    (s - (1.326+10.32j)) (s - (1.326-10.32j)) (s + 16.97)
    Poles: [ 1.3264659 +10.3244805j 1.3264659 -10.3244805j
     -16.96616984 +0.j
[57]: print(abs(poles[0]))
     Omega = abs(np.imag(poles[0]))
     Freq = Omega/(2*np.pi)
     print(Freq,1/Freq)
    10.40934241269315
    1.6431921071431235 0.6085715697226747
```

#### 6.2 Linear time response

```
[58]: # y_cl = impulse_response(CL,T=T)
y_cll = impulse_response(CLL,T=T)
#y_cll_r = impulse_response(CLL_r,T=T)
```



```
[59]: | # print(CL.state_labels)
```

#### 6.3 Non-linear simulation

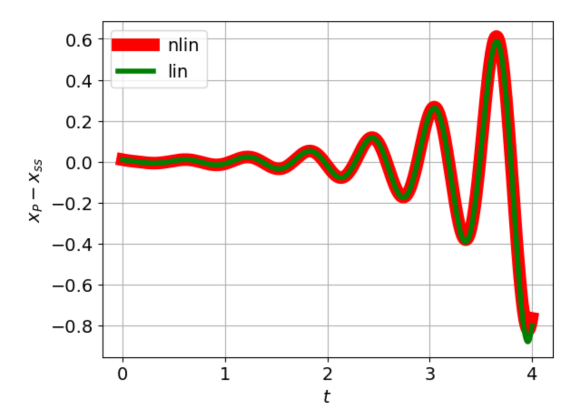
- the non-linear simulation has the initial state the same as the steady state except that the initial value  $x_P(0)$  of the product state  $x_P$  is perturbed by pert.
- the trajectory  $x_P(t)$  from the nonlinear system is compared with the impulse response of the linearised system
  - − the input to the linearised system is pert\*unit\_impulse.

```
[60]: ## Simulate
    print(InpVar)
    species = s['species']
    print(species)
    reaction = s['reaction']
    X0 = copy.copy(x_ss)
    pert = 1e-2
    # X0[species.index(InpVar)] *= 1+pert
    X0[species.index('P')] += pert
    ndat = st.sim(s,sc=sc,t=T,X0=X0,parameter=parameter,quiet=False)
```

```
fb_Act
     ['fb_Act', 'fb_E0', 'E1', 'E2', 'E3', 'P', 'decr1_A', 'decr1_Zf', 'decr2_A',
     'decr2_Zf', 'decr3_A', 'decr3_Zf']
     Setting K_fb_Act to 1
     Setting K_E1 to 1
     Setting K_E2 to 1
     Setting K_E3 to 1
     Setting K_decr1_A to 100.0
     Setting K_decr1_Zf to 1e-06
     Setting K_decr2_A to 100.0
     Setting K_decr2_Zf to 1e-06
     Setting K_decr3_A to 100.0
     Setting K_decr3_Zf to 1e-06
     Setting kappa_decr1_r to 1.0
     Setting kappa_decr1_rf to 10
     Setting kappa_decr2_r to 1.0
     Setting kappa_decr2_rf to 10
     Setting kappa_decr3_r to 1.0
     Setting kappa_decr3_rf to 10
[61]: if SystemName in_
      →['Toy','Toy1','Toycc','ToyTwo','ToyTwo3','ToyTwo4','Goodwin']:
          Yname = 'P'
          Xlabel = 't'
          Ylabel = 'P'
      elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
          Yname = 'P'
          Xlabel = 't'
          Ylabel = 'ADP'
      else:
          print('System Name',SystemName,'is not known')
      y_n = (ndat['X'][:,species.index('P')] - x_ss[species.index('P')] )
      plt.plot(T,y_n, label='nlin',lw=10)
      plt.plot(T,pert*y_cll,label='lin',lw=4)
      # plt.plot(T,pert*y_cll_r,label='lin (reduced)',lw=4,ls='dashed')
      plt.grid()
      plt.legend()
      plt.xlabel('$t$')
      plt.ylabel('x_{\text{'+Ylabel+'}} - x_{\text{ss}}')
      if SysName == 'Selkov3':
          plt.xlim((0,0.1))
          plt.ylim((0,0.1))
      SaveFig(SysName, 'Simulation')
[61]: [<matplotlib.lines.Line2D at 0x76d4107ffc10>]
[61]: [<matplotlib.lines.Line2D at 0x76d410f93af0>]
[61]: <matplotlib.legend.Legend at 0x76d411025820>
```

```
[61]: Text(0.5, 0, '$t$')
```

```
[61]: Text(0, 0.5, '$x_{P} - x_{ss}$')
```



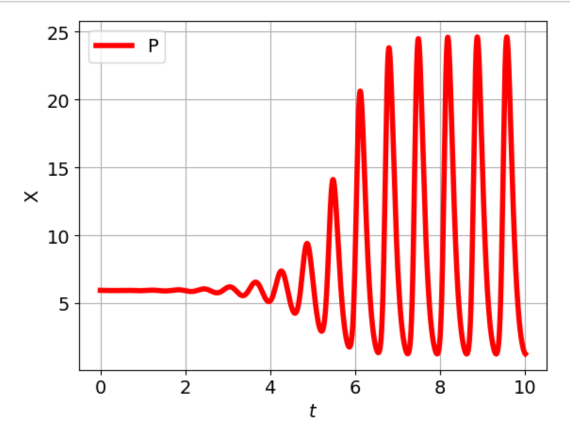
```
[62]: # st.plot(s,ndat,species=['P'],reaction=[])
# st.plot(s,ndat,species=['E1'],reaction=[])
# st.plot(s,ndat,species=['E2'],reaction=[])
# st.plot(s,ndat,species=['E3'],reaction=[])
```

#### [63]: ndat = st.sim(s,sc=sc,t=T\_long,X0=X0,parameter=parameter,quiet=False)

```
Setting K_E1 to 1
Setting K_E2 to 1
Setting K_E3 to 1
Setting K_decr1_A to 100.0
Setting K_decr1_Zf to 1e-06
Setting K_decr2_A to 100.0
Setting K_decr2_Zf to 1e-06
Setting K_decr3_A to 100.0
Setting K_decr3_Zf to 1e-06
Setting K_decr3_Zf to 1e-06
Setting kappa_decr1_r to 1.0
Setting kappa_decr1_rf to 10
Setting kappa_decr2_r to 1.0
Setting kappa_decr2_rf to 10
Setting kappa_decr3_r to 1.0
```

Setting K\_fb\_Act to 1

```
[64]: st.plot(s,ndat,species=['P'],reaction=[])
# st.plot(s,ndat,species=['E1'],reaction=[])
# st.plot(s,ndat,species=['E2'],reaction=[])
# st.plot(s,ndat,species=['E3'],reaction=[])
```



#### 6.4 Phase plane

```
[65]: print(species)
     if SystemName in⊔
      Xname = 'E1'
        Yname = 'P'
        Xlabel = Xname
        Ylabel = Yname
     elif SystemName in ['Selkov', 'Selkov1', 'Selkov3']:
        Xname = 'selkov_ATP'
        Yname = 'P'
        Xlabel = 'ADP'
        Ylabel = 'ATP'
     else:
        print('System Name',SystemName,'is not known')
     i_X = species.index(Xname)
     x_X = ndat['X'][:,i_X]
```

```
i_Y = species.index(Yname)
x_Y = ndat['X'][:,i_Y]
plt.plot(x_X,x_Y,lw=1,color='black')

plt.plot(x_ss[i_X],x_ss[i_Y],marker='+',color='black')
plt.locator_params(nbins=4)

# plt.plot(X[:,0],X[:,2])
plt.xlabel(f'$x_{{Xlabel}}$')
plt.ylabel(f'$x_{{Ylabel}}$')
# plt.xlim(left=0)
# plt.ylim(bottom=0)
plt.grid()
SaveFig(SysName, 'PhasePlane')
```

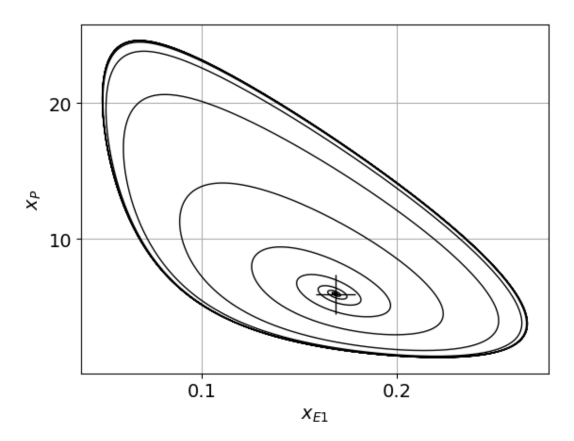
['fb\_Act', 'fb\_E0', 'E1', 'E2', 'E3', 'P', 'decr1\_A', 'decr1\_Zf', 'decr2\_A', 'decr2\_Zf', 'decr3\_A', 'decr3\_Zf']

[65]: [<matplotlib.lines.Line2D at 0x76d410730850>]

[65]: [<matplotlib.lines.Line2D at 0x76d410730be0>]

[65]: Text(0.5, 0, '\$x\_{E1}\$')

[65]: Text(0, 0.5, '\$x\_{P}\$')



## 6.5 Signals at integrator

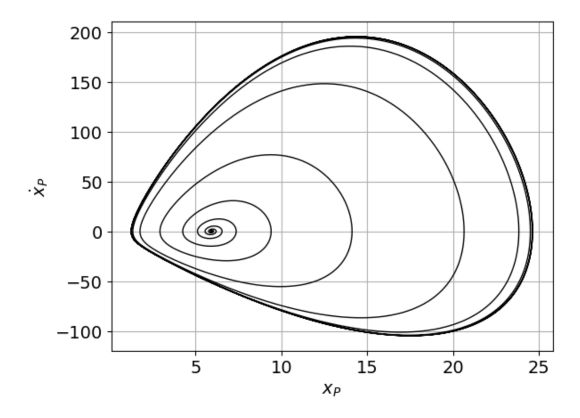
```
[66]: i_P = species.index('P')
    x_X = ndat['X'][10:,i_P]
    x_Y = ndat['dX'][10:,i_P]
    tt = ndat['t'][10:]
    plt.plot(x_X,x_Y,lw=1,color='black')
    plt.xlabel('$x_P$')
    plt.ylabel('$x_P$')
    # plt.xlim(left=0)
    # plt.ylim(bottom=0)
    plt.grid()

SaveFig(SysName,'PhasePlaneP')
```

[66]: [<matplotlib.lines.Line2D at 0x76d41070d280>]

[66]: Text(0.5, 0, '\$x\_P\$')

[66]:  $Text(0, 0.5, '\$\dot{x}_P$')$ 

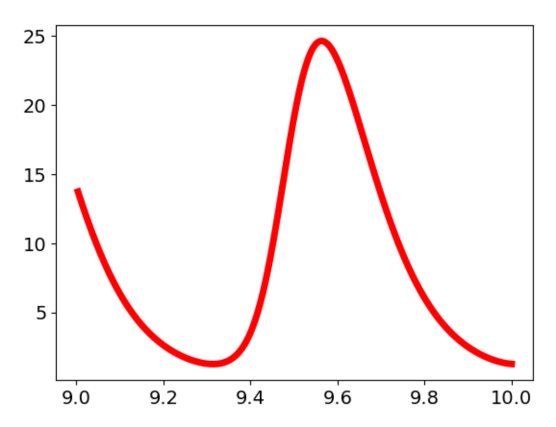


```
[67]: i0 = int(0.9*len(tt)) print(i0)
```

```
plt.plot(tt[i0:],x_X[i0:])
```

4491

[67]: [<matplotlib.lines.Line2D at 0x76d410692670>]



## 6.6 Period

```
[68]: x_Y_ss = x_ss[i_Y]
# print(x_Y_ss)
i_zc = zero_crossings(x_Y-x_Y_ss)
t = ndat['t']
t_zc = t[i_zc]
T_zc = np.diff(t_zc)

plt.hlines([1/Freq],min(t),max(t),ls='dashed',color='grey',label='linear')
plt.plot(t_zc[1:],T_zc, label='actual')

plt.grid()
plt.legend()
plt.xlabel('$t$')
plt.ylabel('$t$')
SaveFig(SysName,'Period')

# print(zc)
# print(1/Freq)
```

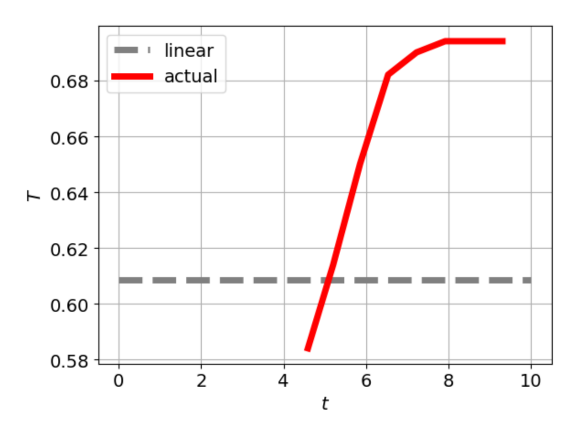
[68]: <matplotlib.collections.LineCollection at 0x76d410646460>

[68]: [<matplotlib.lines.Line2D at 0x76d4105f7610>]

[68]: <matplotlib.legend.Legend at 0x76d4106e0cd0>

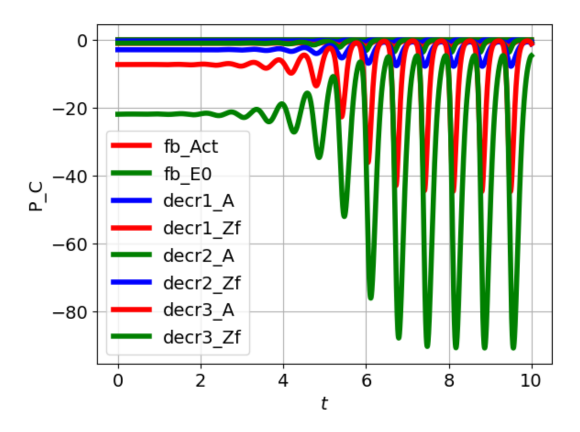
[68]: Text(0.5, 0, '\$t\$')

[68]: Text(0, 0.5, '\$T\$')



## 7 Power

[69]: st.plot(s,ndat,plotPower=True,species=chemostats,reaction=[])



```
[70]: # st.plot(s,ndat,plotPower=True,species=[],reaction=['decr3_r','decr3_rf'])
```

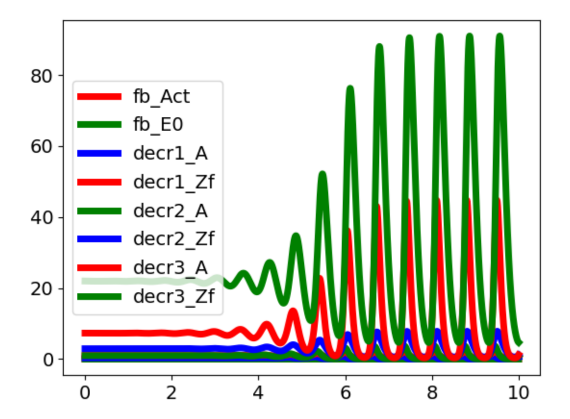
#### 7.1 Power

```
[71]: P_Re = ndat['P_Re']
      P_C = -ndat['P_C']
      i_{chemo} = []
      for chemo in chemostats:
          i_chemo.append(species.index(chemo))
      free = list(set(species)-set(chemostats))
      print(free)
      i_free = []
      for fr in free:
          i_free.append(species.index(fr))
      P_chemo = P_C[:,i_chemo]
      P_free = P_C[:,i_free]
      ## Total power
      PP_Re = np.sum(P_Re,axis=1)
      PP_C = np.sum(P_C,axis=1)
      PP_chemo = np.sum(P_chemo,axis=1)
      PP_free = np.sum(P_free,axis=1)
```

['E2', 'E1', 'E3', 'P']

```
[72]: t = ndat['t']
  plt.plot(t,P_chemo)
  plt.legend(chemostats,loc='center left')
```

[72]: <matplotlib.legend.Legend at 0x76d4104f4940>

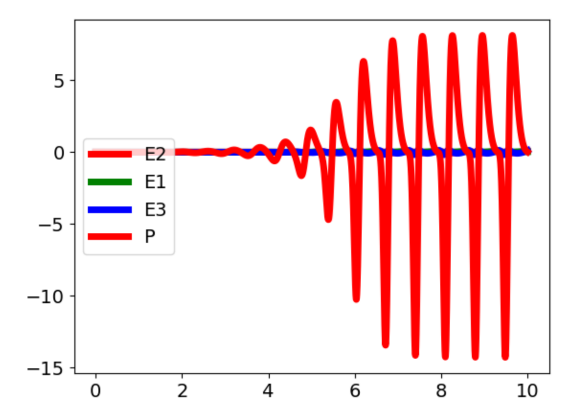


```
[73]: t = ndat['t']
plt.plot(t,P_free)
plt.legend(free,loc='center left')

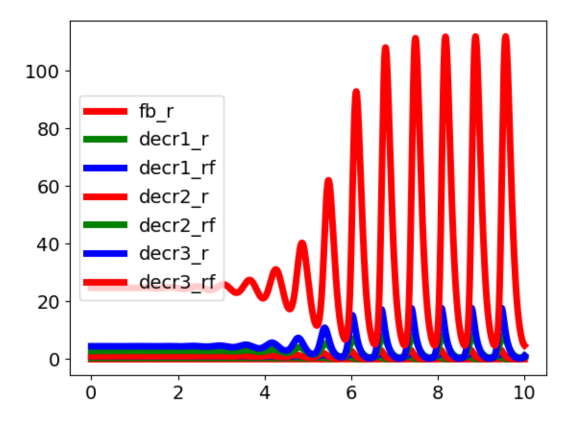
[73]: [cmatplotlib lines Line2D at 0x76d41052c490>
```

[73]: [<matplotlib.lines.Line2D at 0x76d41052c490>, <matplotlib.lines.Line2D at 0x76d41044c610>, <matplotlib.lines.Line2D at 0x76d41044c640>, <matplotlib.lines.Line2D at 0x76d41044c730>]

[73]: <matplotlib.legend.Legend at 0x76d4104e46d0>

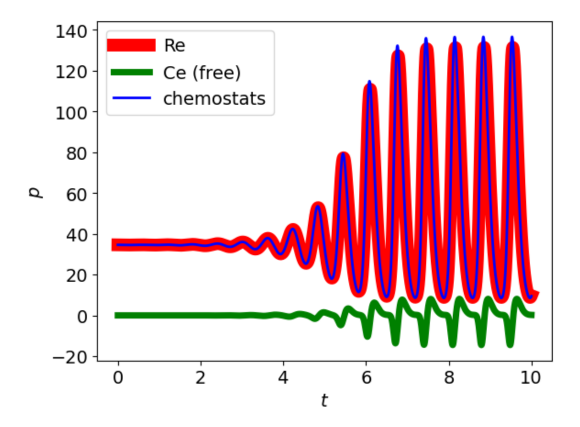


[74]: <matplotlib.legend.Legend at 0x76d4103ff550>

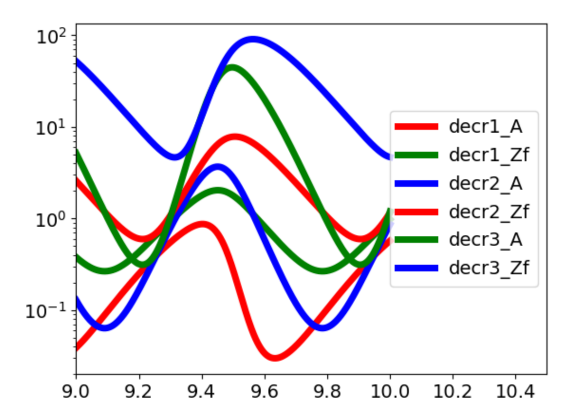


```
[75]: t = ndat['t']
    plt.plot(t,PP_Re,label='Re',lw=10)
    plt.plot(t,PP_free,label='Ce (free)',lw=5)
    plt.plot(t,PP_chemo,label='chemostats',lw=2)
    plt.legend()
    plt.xlabel('$t$')
    plt.ylabel('$p$')
    SaveFig(SysName,'Power')

[75]: [<matplotlib.lines.Line2D at 0x76d4102f6940>]
[75]: [<matplotlib.lines.Line2D at 0x76d4102f6d00>]
[75]: [<matplotlib.lines.Line2D at 0x76d41034fcd0>]
[75]: <matplotlib.legend.Legend at 0x76d41034fcd0>]
[75]: Text(0.5, 0, '$t$')
```



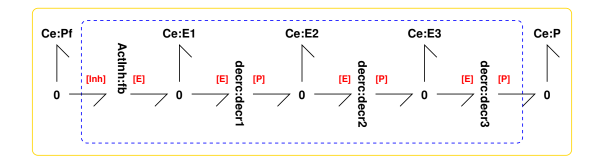
[76]: <matplotlib.legend.Legend at 0x76d410224f70>



# 8 Split-loop analysis

#### 8.1 Model

```
[77]: name = SystemName+'SL_abg'
      svg = name+'.svg'
      print('Using',svg)
      sbg.model(svg,convertCe=True,convertR=True,quiet=quiet)
      exec(f'import {name} as sys_abg')
      imp.reload(sys_abg)
      disp.SVG(svg)
      SplitVar = 'Pf'
     Using ToySL_abg.svg
     Creating subsystem: ActInh:fb
     Creating subsystem: decrc:decr1
     Creating subsystem: decrc:decr2
     Creating subsystem: decrc:decr3
     {}
[77]: <module 'ToySL_abg' from
      '/home/peterg/WORK/Research/SystemsBiology/Notes/2024/Oscillation/ToySL_abg.
      →py '>
[77]:
```



#### 8.2 Stoichiometry

```
[78]: chemostats = SetChemostats(SystemName,Config='SplitLoop',quiet=False)
```

Setting feedback loop with configuration SplitLoop

```
[79]: print(chemostats)
s,sc = stoichiometry(sys_abg.model(),chemostats=chemostats)
species_sl = s['species']
print(species_sl)
```

```
['fb_Act', 'fb_E0', 'Pf', 'P', 'decr1_A', 'decr1_Zf', 'decr2_A', 'decr2_Zf',
'decr3_A', 'decr3_Zf']
['fb_Act', 'fb_E0', 'E1', 'E2', 'E3', 'P', 'Pf', 'decr1_A', 'decr1_Zf',
'decr2_A', 'decr2_Zf', 'decr3_A', 'decr3_Zf']
```

#### 8.3 Steady-state analysis

```
[80]: ## Create the steady state corresponding to open loop with x_inh=x_P:
    parameter = SetParameter()
    x_sl_ss = np.ones(s['n_X'])
    X_ss = copy.copy(x_ss)
    for i,spec in enumerate(species):
        x_sl_ss[species_sl.index(spec)] = X_ss[i]
    x_sl_ss[species_sl.index(SplitVar)] = X_ss[species.index('P')]
```

#### 8.4 Linearise

```
[81]: # ttff = con.zpk([-1,-2],[-3,-4,-5],123,display_format='zpk')
# ttff
```

```
[82]: Inp = [SplitVar,OutpVar]
TF,Sys = L

→Lin(s,sc,parameter=parameter,x_ss=x_sl_ss,outvar='dX',Inp=Inp,Outp=Inp)
```

```
2 states have been removed from the model
0 states have been removed from the model
3 states have been removed from the model
3 states have been removed from the model
```

```
[83]: for name in TF:
            print(name)
            TFr = con.minreal(TF[name])
            # zTF = zpk(TF[name])
            # zTF
            print(con.poles(TF[name]))
      O states have been removed from the model
[83]:
                                                   \frac{-0.1685(s)}{s + 5.936}
      [-5.93620394+0.j]
      O states have been removed from the model
[83]:
                                       \frac{-1.639 \times 10^{-12} (s + 2.897 \times 10^{16})}{(s + 5.936)(s + 10.03)(s + 10.08)}
      [-10.08007675+0.j -10.02837804+0.j -5.93620394+0.j]
      P_Pf
      O states have been removed from the model
[83]:
                                                        0
                                                        \frac{-}{1}
      []
      O states have been removed from the model
[83]:
                                                     \frac{-10.63}{1}
```

#### 8.5 Active and passive loop gains

```
[84]: LLO = con.tf(0,1)
L_pas_0 = con.tf(0,1)
for index in TF:
    if not index in ['Pf_P']:
        L_pas_0 = con.minreal(con.parallel(L_pas_0,-TF[index]))

    LLO = con.minreal(con.parallel(LLO,-TF[index]))

# LLO = con.minreal(LLO)
print('LO')
LLO
L_act_0 = -TF[SplitVar+'_P']
```

```
print('L0_act')
      L_act_0
      \# L_pas_0 = con.minreal(L_pas_0)
      print('L0_pas')
      L_pas_0
      LL = IntegrateTF(LL0)
      # LL = con.minreal(LL)
      # LL = con.tf(balred(con.ss(LL),3))
      print('L')
      print('LL')
      LL
      L_act = IntegrateTF(L_act_0)
      print('L_act')
      L_act
      \# L_pas_0 = con.parallel(-TF['P_P'], -TF[SplitVar+'_'+SplitVar])
      L_pas = IntegrateTF(L_pas_0)
      print('L_pas')
      L_pas
      \# L_pas_P = IntegrateTF(-TF['P_P'])
      # print('L_pas_P', L_pas_P)
      # L_pas_Inh = IntegrateTF(-TF[SplitVar+'_'+SplitVar])
      # print('L_pas_Inh', L_pas_Inh)
     O states have been removed from the model
     O states have been removed from the model
     1 states have been removed from the model
     O states have been removed from the model
     LO
[84]:
                     10.8(s + (0.4024 - 14.08j))(s + (0.4024 + 14.08j))(s + 25.15)
                                  (s+5.936)(s+10.03)(s+10.08)
     L0_act
[84]:
                                  1.639 \times 10^{-12} (s + 2.897 \times 10^{16})
                                 \overline{(s+5.936)(s+10.03)(s+10.08)}
     L0_pas
[84]:
```

$$\frac{10.8(s+5.844)}{s+5.936}$$

L

[84]:

$$\frac{10.8(s + (0.4024 - 14.08j))(s + (0.4024 + 14.08j))(s + 25.15)}{(s)(s + 5.936)(s + 10.03)(s + 10.08)}$$

LL

[84]:

$$\frac{10.8(s + (0.4024 - 14.08j))(s + (0.4024 + 14.08j))(s + 25.15)}{(s)(s + 5.936)(s + 10.03)(s + 10.08)}$$

L\_act

[84]:

$$\frac{1.639 \times 10^{-12} (s + 2.897 \times 10^{16})}{(s)(s + 5.936)(s + 10.03)(s + 10.08)}$$

L\_pas

[84]:

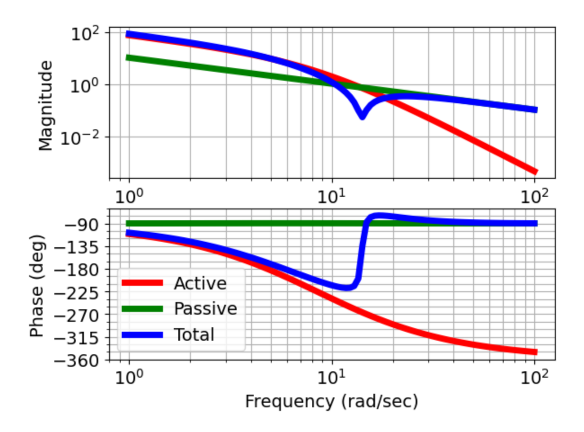
$$\frac{10.8(s+5.844)}{(s)(s+5.936)}$$

```
[85]: # print(f'Gain L0 = {con.dcgain(L0):.2f}')
# L0_pas_P = TF['P_P']
# K = parameter['K_decr3_A']*parameter['kappa_decr3_r']
# K_f = parameter['kappa_decr3_rf']
# print(f'Gain L0_pas_P = {con.dcgain(L0_pas_P):0.2f} ({K_f:0.2f})')
```

#### 8.6 Bode plots

```
[86]: ## Bode
L_list = [L_act,L_pas,LL]
  omega = np.logspace(0,2,100)
  mag,phase,om=con.bode_plot(L_list,omega)
  plt.legend(['Active','Passive','Total'])
  SaveFig(SysName,'SplitBode')
```

[86]: <matplotlib.legend.Legend at 0x76d410842070>

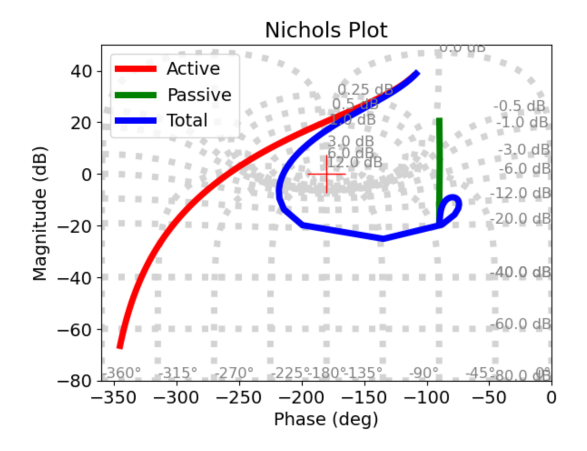


```
[87]: if SaveData:
    BodeData = {}
    BodeData['L_list'] = L_list
    BodeData['mag'] = mag
    BodeData['phase'] = phase
    BodeData['omega'] = om
    BodeData['wcp'] = wcp
    BodeData['pm'] = pm
SavedData['Bode'] = BodeData
```

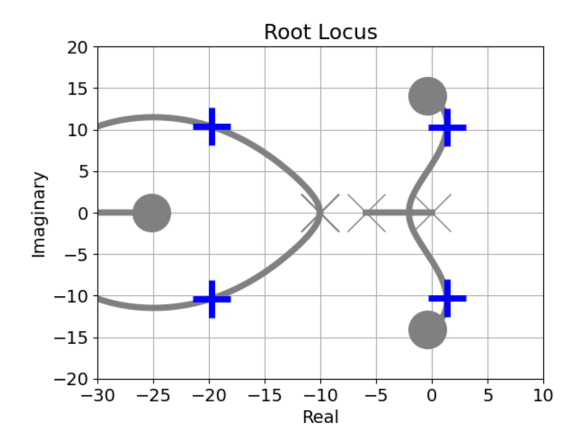
## 8.7 Nichols plots

```
[88]: ## Nichols
    con.nichols_plot(L_list,omega)
    plt.legend(['Active','Passive','Total'])
    SaveFig(SysName,'SplitNichols')
```

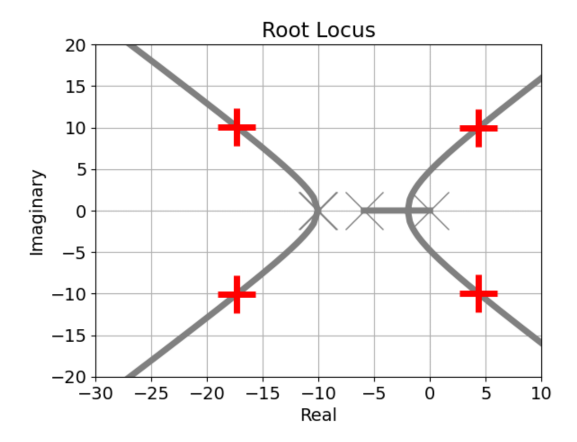
[88]: <matplotlib.legend.Legend at 0x76d410ba01f0>



#### 8.8 Root Locus

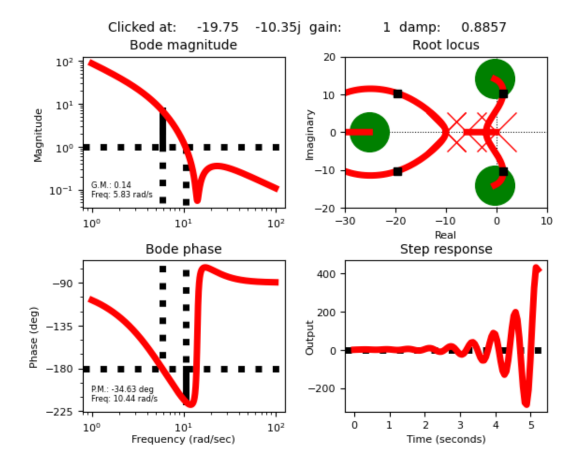


#### 8.9 Root Locus - active only



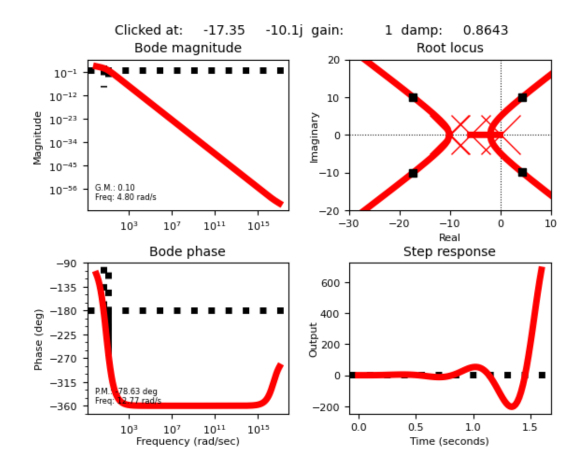
## 8.10 Sisotool

[91]: con.sisotool(L,xlim\_rlocus=xlim,ylim\_rlocus=ylim)
SaveFig(SysName,'SplitSisoTool')



## 8.11 Sisotool - active only

```
[92]: con.sisotool(L_act,xlim_rlocus=xlim,ylim_rlocus=ylim)
SaveFig(SysName,'SplitSisoTool')
```



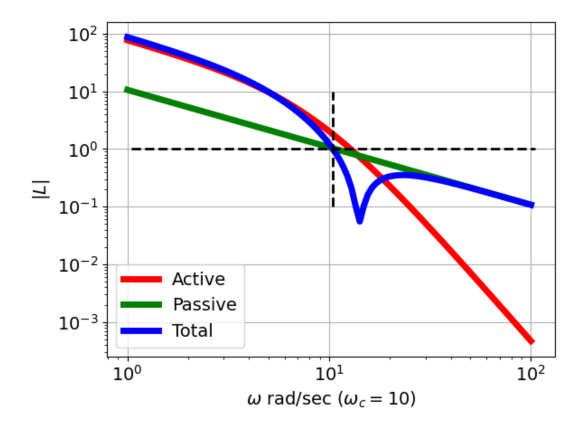
[93]: [<matplotlib.lines.Line2D at 0x76d3e4024790>]

[93]: <matplotlib.legend.Legend at 0x76d3e4024a30>

[93]: <matplotlib.collections.LineCollection at 0x76d3e40b3f70>

[93]: <matplotlib.collections.LineCollection at 0x76d3e40a5760>

```
[93]: Text(0.5, 0, '\$\setminus sample rad/sec (\$\setminus c = 10\$)')
[93]: Text(0, 0.5, '$|L|$')
```



```
[94]: for i,l in enumerate(L_list):
            print(i, l)
          mag,phase,om = con.bode_plot(1,omega,plot=False)
          phase_deg = phase*180/np.pi
          plt.semilogx(omega,phase_deg,label=Name[i])
      plt.legend()
      plt.hlines(-180,wcp/10,wcp*10,ls='dashed',color='black',lw=2)
      plt.vlines(wcp,-230,-130,ls='dashed',color='black',lw=2)
      # plt.xlabel(r'$\omega$')
      plt.xlabel(f'$\\omega$ rad/sec ($\\theta_{{pm}} = {int(round(pm))}^\circ$)')
      plt.ylabel(r'$\angle{L}$')
      plt.grid()
      SaveFig(SysName, 'SplitBodePha')
```

[94]: [<matplotlib.lines.Line2D at 0x76d3dfdb5280>]

[94]: [<matplotlib.lines.Line2D at 0x76d3dfdb5550>]

[94]: [<matplotlib.lines.Line2D at 0x76d3dfdb5820>]

```
[94]: <matplotlib.legend.Legend at 0x76d3dfd99250>
```

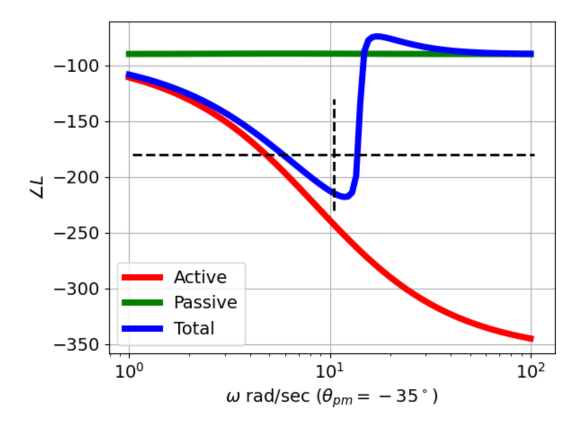
[94]: <matplotlib.collections.LineCollection at 0x76d3dfe27610>

[94]: <matplotlib.collections.LineCollection at 0x76d3dfd42bb0>

[94]: Text(0.5, 0, ' $\$  rad/sec ( $\$  = -35^\\circ\$)')

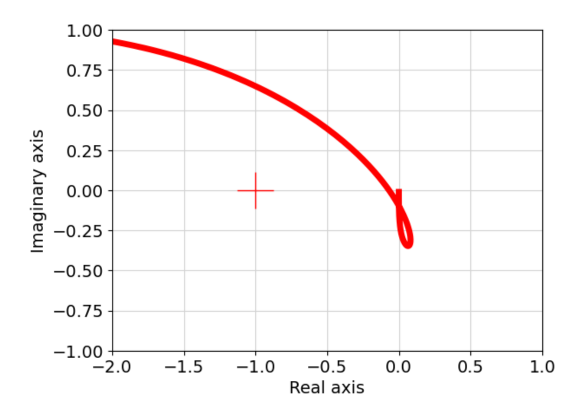
[94]: Text(0, 0.5, '\$\\angle{L}\$')

[95]: (-1.0, 1.0)



```
[ ]:
[95]: con.nyquist_plot([LL],mirror_style=False)
    plt.xlim(-2,1)
    plt.ylim(-1,1)

[95]: 2
[95]: (-2.0, 1.0)
```

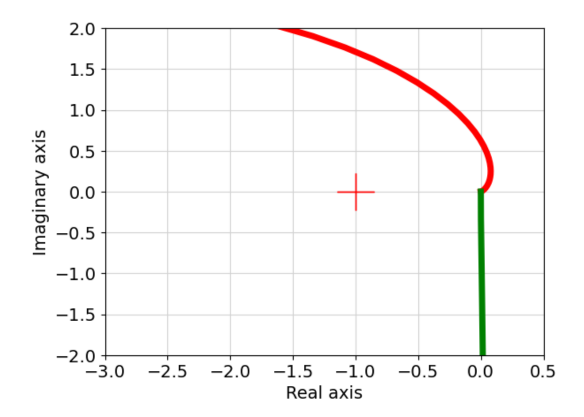


```
[96]: con.nyquist_plot([L_act,L_pas],mirror_style=False)
   plt.xlim(-3,0.5)
   plt.ylim(-2,2)
```

[96]: [2, 0]

[96]: (-3.0, 0.5)

[96]: (-2.0, 2.0)



# 9 Open-loop analysis of dynamic part of Toy and Goodwin examples

```
[97]: if SystemName in ['Toy', 'Goodwin']:
          # Stoichiometry
          chemostats,parameter,InpVar,OutpVar,T,T_long,n_red =_
       →SetAll(SystemName, Config='Dynamic', quiet=False)
          s,sc = stoichiometry(sys_abg.model(),chemostats=chemostats)
          species = s['species']
          ## Linearise
          X_ss = copy.deepcopy(x_sl_ss)
          TF,Sys = Lin(s,sc,parameter=parameter,x_ss=X_ss,outvar='dX',Inp=['E1'],__

Outp=['P','Pf'])
          ## Steady state values of the 3 enzymes - and product.
          x_s_E = 1
          for i in range(3):
              Ei = 'E' + str(i+1)
              x_ss_i = X_ss[species.index(Ei)]
              print(f'{Ei} steady state: {x_ss_i:.3f}')
```

```
x_s_E *= x_s_i
          print(f'P steady-state: {X_ss[species.index("P")]:.3f}')
          ## Extract transfer-functions
          for tfName in ['E1_P','E1_Pf']:
              print(tfName)
              tf = TF[tfName]
              0L_d_0 = tf
              # print(OL_d_0)
              OL_d = IntegrateTF(OL_d_0)
              print(OL_d)
              print(con.poles(OL_d))
              g = con.dcgain(OL_d)
              print(f'Gain: \{g:.2e\} (\{g/x\_ss\_E:.3e\})')
     Setting feedback loop with configuration Dynamic
     1 states have been removed from the model
     3 states have been removed from the model
     E1 steady state: 0.168
     E2 steady state: 0.283
     E3 steady state: 0.794
     P steady-state: 5.936
     E1_P
                  2.819e+05
     (s + 10.03) (s + 10.08) (s + 10.63)
     [-10.63108265+0.j -10.08007675+0.j -10.02837804+0.j]
     Gain: 2.62e+02 (6.927e+03)
     E1_Pf
       -5.936
     -----
     s + 0.1685
     [-0.16845782+0.j]
     Gain: -3.52e+01 (-9.305e+02)
[98]: con.config.defaults['xferfcn.display_format'] = 'zpk'
```

#### 10 Linear + saturation

```
[99]: def linpos_fun(tt,x):
        global _A_MATRIX_
        global _MIN_STATE_
        dx = _A_MATRIX_@x
        # if x[0]<-10:
        # dx[0] = 0</pre>
```

```
# print(dx.shape)
           for i,xx in enumerate(x):
               min = _MIN_STATE_[i]
               if xx<min:</pre>
                   # print(i)
                   x[i] = min
           return dx
       def linpos(A,x0,x_ss_0,t_span):
           global _A_MATRIX_
           global _MIN_STATE_
           _A_MATRIX_ = A
           _{MIN\_STATE\_} = -1*np.array(x_ss_0)
           ret = integrate.solve_ivp(linpos_fun, t_span, x0, max_step=0.01)
           t = ret['t']
           x = ret['y']
           dx = _A_MATRIX_{0x}
           return t,x.T,dx.T
[100]: ## Extract variable states
       x_s_0 = []
       species_0 = []
       for spec in species_open:
           if spec not in chemostats_open:
               species_0.append(spec)
       ## con.feedback puts P state first - so prepend
       species_0 = ['P'] + species_0
       for spec in species_0:
           x = x_ss_open[species.index(spec)]
           print(spec,x)
           x_ss_0.append(x)
       print(x_ss_0)
       print(species_0)
      P 5.93620393877866
      E1 0.16845782427848063
      E2 0.28297834855767506
      E3 0.7944071039555537
      [5.93620393877866, 0.16845782427848063, 0.28297834855767506, 0.7944071039555537]
      ['P', 'E1', 'E2', 'E3']
[101]: if SystemName in ['Toy']:
           print(x_ss_0)
           timespan = [0,10]
           x0 = np.zeros(len(x_ss_0))
           # Pert = 1e-2
           x0[0] = pert
```

```
Intsys = con.ss(0,1,1,0)
    con.tf(Intsys)
    linsys = con.feedback(Intsys,L0_sys)
    ## Show systems
    L0_sys
    Intsys
    linsys
    linsys.state_labels
    A = copy.copy(linsys.A)
    A.shape
    x0.shape
    tt,x,dx = linpos(A,x0,x_ss_0,timespan)
    X = x + x_s_0
    print(x[1,3])
    plt.plot(tt,X)
    plt.grid()
    plt.legend(species_0)
    plt.show()
    plt.plot(tt,x[:,0],label='sim')
    plt.plot(T,y_cll*pert,label='impulse',lw=2)
    # plt.xlim(0,2)
    plt.legend()
    plt.show()
    plt.plot(X[:,1],X[:,0])
    plt.grid()
[5.93620393877866, 0.16845782427848063, 0.28297834855767506, 0.7944071039555537]
```

[101]:

1

[101]:

$$\begin{pmatrix} -5.84 & -0.01 & -0.0169 & -16.8 \\ -33.6 & -10.1 & 0.133 & -0.0494 \\ 0 & -5.61 & -10.1 & 0.0669 \\ \hline -0 & -0 & -14.9 & 10.8 \end{pmatrix}$$

[101]:

$$\begin{pmatrix} 0 & 1 \\ \hline 1 & 0 \end{pmatrix}$$

[101]:

$$\begin{pmatrix}
-10.8 & 0 & 0 & 14.9 & 1 \\
-16.8 & -5.84 & -0.01 & -0.0169 & 0 \\
-0.0494 & -33.6 & -10.1 & 0.133 & 0 \\
0.0669 & 0 & -5.61 & -10.1 & 0 \\
\hline
1 & 0 & 0 & 0 & 0
\end{pmatrix}$$

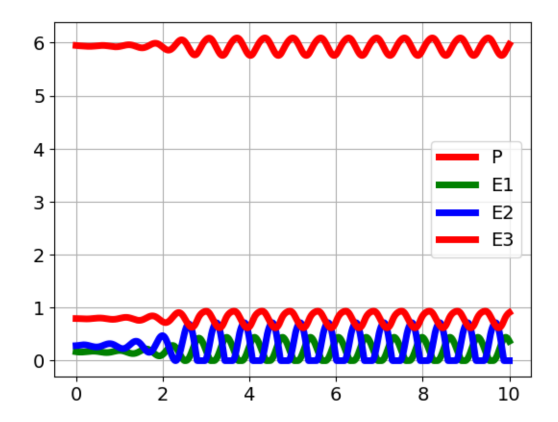
[101]: ['sys[227]\_x[0]', 'sys[82]\_x[0]', 'sys[82]\_x[1]', 'sys[82]\_x[2]']

[101]: (4, 4)

[101]: (4,)

#### 2.657990146241061e-06

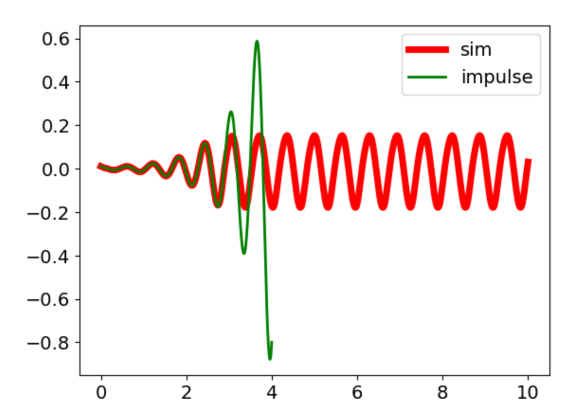
[101]: <matplotlib.legend.Legend at 0x76d3dfc31940>



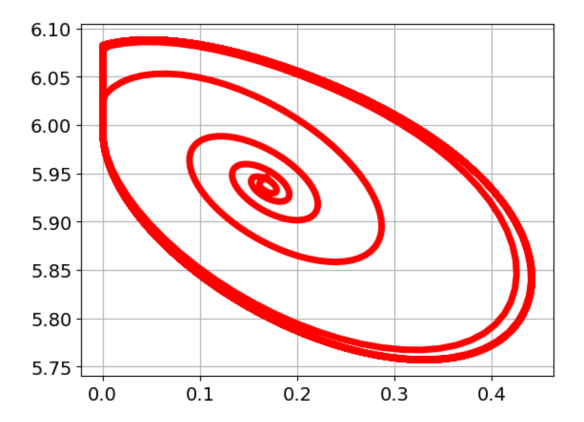
[101]: [<matplotlib.lines.Line2D at 0x76d3dfa07910>]

[101]: [<matplotlib.lines.Line2D at 0x76d3dfa07ca0>]

[101]: <matplotlib.legend.Legend at 0x76d3dfa07e50>



[101]: [<matplotlib.lines.Line2D at 0x76d3dfb1bee0>]



#### 10.1 Signals at integrator

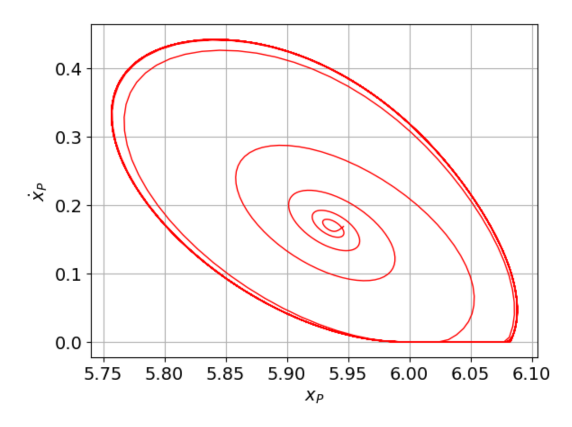
```
[102]: if SystemName in ['Toy']:
    print(species)
    i_P = 0 # See above
    i_E1 = 1
    x_X = X[:,i_P]
    x_Y = X[:,i_E1]
    plt.plot(x_X,x_Y,lw=1)
    plt.xlabel('$x_P$')
    plt.ylabel('$\dot{x}_P$')
    # plt.xlim(left=0)
    # plt.ylim(bottom=0)
    plt.grid()
    SaveFig(SysName, 'PhasePlaneP')
```

['fb\_Act', 'fb\_E0', 'E1', 'E2', 'E3', 'P', 'Pf', 'decr1\_A', 'decr1\_Zf', 'decr2\_A', 'decr2\_Zf', 'decr3\_A', 'decr3\_Zf']

[102]: [<matplotlib.lines.Line2D at 0x76d3df9f6730>]

[102]: Text(0.5, 0, '\$x\_P\$')

[102]: Text(0, 0.5, '\$\\dot{x}\_P\$')



```
[103]: ## Optionally save data
print(SysName)
if SaveData:
    file = open(f'{SysName}.dat', 'wb')
    pickle.dump(SavedData, file)
    file.close()

Toy
[]:
```

## References