Python On Resonance (PyOR)

Everybody can simulate NMR

Version: Jeener

Tutorial: Introduction to PyOR

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Define the Path to PyOR source code, PythonOnResonance.py

```
In [1]: pathSource = '/media/HD2/Vineeth/PostDoc_Simulations/Github/PyOR_V1/Source'
```

Load Python packages

```
In [2]:
    from IPython.display import display, HTML
    display(HTML("<style>.container { width:100% !important; }</style>"))
    import sys
    sys.path.append(pathSource)

    import PythonOnResonance as PyOR

    import time
    import numpy as np
    import matplotlib.pyplot as plt
    from matplotlib import rc
    %matplotlib notebook
    import sympy as sp
    from sympy import *
```

Define Spin System (Two Spin Half)

```
• For single spin half system
```

- Spin_list = [1/2]
- · For single spin one system
 - Spin_list = [1]
- · For two spin half system
 - Spin_list = [1/2, 1/2]
- · For spin half and spin one system
 - Spin_list = [1/2, 1]

```
In [3]: Spin_list = [1/2, 1/2]
```

Define the unit of Hamiltonian

hbarEQ1 is True, then unit of Hamiltonian is in angular frequency

```
In [4]: hbarEQ1 = True
```

Generate the spin operators: Sx, Sy and Sz

```
Sx [i], Sy [i] and Sz [i]
```

'i' is the index of the spin

```
In [5]: System = PyOR.Numerical_MR(Spin_list,hbarEQ1)
Sx,Sy,Sz = System.SpinOperator()
```

Matrix representation of the spin operators for the first spin (Sx, Sy and Sz)

Matrix representation of the spin operators for the second spin (Sx, Sy and Sz)

```
In [9]: Matrix(Sx[1])
Out[9]:
```

```
0.5
In [10]:
         Matrix(Sy[1])
                -0.5i
Out[10]:
                            -0.5i
In [11]:
         Matrix(Sz[1])
Out[11]:
               -0.5 0
        Generate the spin operators: S+ and S-
        Sp[i] and Sm[i]
        'i' is the index of the spin
In [12]:
         Sp, Sm = System.PMoperators(Sx,Sy)
        Matrix representation of the spin operators for the first spin (S+
        and S-)
In [13]:
         Matrix(Sp[0])
Out[13]:
                     1.0
In [14]:
         Matrix(Sm[0])
Out[14]:
```

0

0.5

Matrix representation of the spin operators for the second spin (S+ and S-)

```
Out[15]:
                      0
                  0 1.0
In [16]:
         Matrix(Sm[1])
Out[16]:
                       0.
                   0
           0
        Generating Zeeman Hamiltonian (Lab and Rotating Frame)
In [17]:
          # Gyromagnetic Ratio
          Gamma = [System.gammaH1, System.gammaH1]
In [18]:
          # BO Field in Tesla, Static Magnetic field (BO) along Z
          B0 = 9.4
In [19]:
          # Rotating Frame Frequency
          OmegaRF = [-System.gammaH1*B0, -System.gammaH1*B0]
In [20]:
          # Offset Frequency in rotating frame (Hz)
          Offset = [10.0, 20.0]
In [21]:
          # generate Larmor Frequencies
         LarmorF = System.LarmorFrequency(Gamma, B0, Offset)
         Larmor Frequency in MHz: [-400.22802765 -400.22803765]
In [22]:
         # Lab Frame Hamiltonian
         Hz_lab = System.Zeeman(LarmorF,Sz)
In [23]:
         # Rotating Frame Hamiltonian
         Hz = System.Zeeman_RotFrame(LarmorF, Sz, OmegaRF)
```

Matrix(Sp[1])

In [15]:

Out[24]:

Matrix representation of Zeeman Hamiltonian in Lab frame (Hilbert Space)

```
In [24]: Matrix(Hz_lab/(2.0*np.pi))
```

$\lceil -400228032.646802 \rceil$	0	0	0	
0	5.00000002530119	0	0	
0	0	-5.00000002530119	0	
0	0	0	400228032.646802	

Matrix representation of Zeeman Hamiltonian in Rotating frame (Hilbert Space)

```
In [25]:
          Matrix(Hz/(2.0*np.pi))
            -15.0000000000126
                                         0
                                                              0
                                                                                   0
Out[25]:
                                 5.00000002530119
                                                              0
                                                                                   0
                    0
                    0
                                         0
                                                     -5.00000002530119
                                         0
                                                                          15.0000000000126
                                                              0
```

Generating J Coupling Hamiltonian

Jlist [i][j], J coupling between 'i' th spin and 'j' th spin

```
In [26]:
    Jlist = np.zeros((len(Spin_list),len(Spin_list)))
    Jlist[0][1] = 1

    Jcoupling_Strong = True

    if Jcoupling_Strong:
        Hj = System.Jcoupling(Jlist,Sx,Sy,Sz)
    else:
        Hj = System.Jcoupling_Weak(Jlist,Sz)
```

Matrix representation of J coupling Hamiltonian (Hilbert Space)

Zeeman Basis Kets

Zeeman Basis Bras

```
In [29]:
         ['<1/2,1/2|<1/2,1/2|',
Out[29]:
          '<1/2,1/2|<1/2,-1/2|',
          '<1/2,-1/2|<1/2,1/2|',
          '<1/2, -1/2|<1/2, -1/2|']
        Zeeman Basis states
In [30]:
         Basis_Zeeman_state = System.ZBasis_H(Hz_lab)
        Matrix represenation of Zeeman Basis states
In [31]:
         Matrix(Basis_Zeeman_state[0])
Out[31]:
         \Gamma 1.0
           0
In [32]:
         Matrix(Basis_Zeeman_state[1])
Out[32]:
          1.0
           0
In [33]:
         Matrix(Basis_Zeeman_state[2])
Out[33]:
In [34]:
         Matrix(Basis_Zeeman_state[3])
Out[34]:
        Singlet Triplet Basis states
```

Bras = System.Basis_Bra()

In [35]:

Basis_ST_state = System.STBasis(Hz_lab)

Basis: $T_{-}, T_{0}, T_{+}, S_{0}$

Matrix representation of Singlet Triplet Basis states

```
In [36]:
         Matrix(Basis_ST_state[0])
Out[36]:
           0
           0
In [37]:
         Matrix(Basis_ST_state[1])
Out[37]:
          0.707106781186547
          0.707106781186547
In [38]:
         Matrix(Basis_ST_state[2])
Out[38]:
           0
In [39]:
         Matrix(Basis_ST_state[3])
Out[39]:
           0.707106781186547
           -0.707106781186547
        Transformation Between Zeeman state and Singlet-Triplet state
In [40]:
         U_Z_ST = System.Transform_StateBasis(Basis_Zeeman_state, Basis_ST_state)
        Matrix Representation of the unitary transformation matrix
In [41]:
         Matrix(U_Z_ST)
Out[41]:
                                  0
               0.707106781186547
                                       0.707106781186547
           0
                                  0
           0
               0.707106781186547
                                  0
                                       -0.707106781186547
                      0
                                 1.0
```

Transformation of Zeeman and J coupling Hamiltonian into Singlet-Triplet basis

```
In [42]: Hj_ST = System.Operator_BasisChange(Hj,U_Z_ST)
Hz_ST = System.Operator_BasisChange(Hz,U_Z_ST)
```

Matrix representation of Zeeman Hamiltonian in Singlet-Triplet basis

Matrix representaion of J coupling Hamiltonian in Singlet-Triplet basis

```
In [44]: Matrix(System.Matrix_Round(System.Matrix_Tol(Hj_ST/(2.0*np.pi), 1.0e-5), 2))  \begin{bmatrix} 0.25 & 0 & 0 & 0 \\ 0 & 0.25 & 0 & 0 \\ 0 & 0 & 0.25 & 0 \\ 0 & 0 & 0 & -0.75 \end{bmatrix}
```

Product Operator Basis: Zeeman (Hilbert Space)

```
In [45]: Basis_Zeeman, dic_Zeeman, coh_Zeeman, coh_Zeeman_array = System.ProductOperators_Zeeman(Sz
```

Matrix representation of Product Operators (Zeeman Basis)

Dictionary of Product Operators (Zeeman Basis)

```
In [47]: print(dic_Zeeman)

['|1/2,1/2>|1/2,1/2><1/2,1/2|<1/2,1/2|', '|1/2,1/2>|1/2,1/2><1/2,1/2|<1/2,-1/2|', '|1/2,1/2>|1/2,1/2><1/2,-1/2|', '|1/2,1/2>|1/2,1/2><1/2,-1/2|', '|1/2,1/2>|1/2,1/2><1/2,-1/2|', '|1/2,1/2>|1/2,-1/2|', '|1/2,1/2>|1/2,-1/2|<1/2,-1/2|', '|1/2,1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,-1/2><1/2,-1/2|', '|1/2,-1/2>|1/2,-1/2><1/2,-1/2|']
```

Coherence order of Product Operators (Zeeman Basis)

```
In [48]: print(coh_Zeeman)
[0.0, 1.0, 1.0, 2.0, -1.0, 0.0, 0.0, 1.0, -1.0, 0.0, 0.0, 1.0, -2.0, -1.0, -1.0, 0.0]
```

Coherence order of Product Operators as array (Zeeman Basis)

Product Operator Basis: PMZ (Hilbert Space)

```
In [50]:
    sort = 'negative to positive'
    Index = False
    Normal = True
    Basis_PMZ, coh_PMZ, dic_PMZ = System.ProductOperators_SpinHalf_PMZ(sort,Index,Normal)
```

Call Product Operator with string index

```
In [53]: Compare_with_SpinDynamica = True

if Compare_with_SpinDynamica:
    Basis_PMZ = [0pB_H['Im1Im2'], 0pB_H['Im1Iz2'], 0pB_H['Iz1Im2'], 0pB_H['Im1'], 0pB_H['Im2']
    dic_PMZ = ['Im1Im2', 'Im1Iz2', 'Iz1Im2', 'Im1', 'Im2', 'Im1Ip2', 'Ip1Im2', 'Iz1Iz2', 'Iz1', 'Iz
```

Matrix representation of Product Operators (PMZ Basis)

```
In [54]: Matrix(Basis_PMZ[0])
Out[54]:
```

Dictionary of Product Operators (PMZ Basis)

Coherence order of Product Operators (PMZ Basis)

```
In [56]: print(coh_PMZ)
[-2, -1, -1, -1, 0, 0, 0, 0, 0, 1, 1, 1, 1, 2]
```

Zeeman Hamiltonian in Liouville Space (Zeeman Basis)

```
In [57]: Hz_L = System.CommutationSuperoperator(Hz)
```

Matrix representation of Zeeman Hamiltonian in Liouville Space (Zeeman Basis)

```
In [58]:
            Matrix(System.Matrix_Round(Hz_L/(2.0*np.pi),2))
Out[58]:
             0
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                    0
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```

Convert Product Operators into Liouville Space (Zeeman Basis)

```
In [59]: Basis_Zeeman_L = System.ProductOperators_ConvertToLiouville(Basis_Zeeman)
```

Matrix representation of Product Operators into Liouville Space (Zeeman Basis)

```
In [60]:
           Matrix(Basis_Zeeman_L[0])
Out[60]:
            1.0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
             0
```

Convert Product Operators into Liouville Space (PMZ Basis)

```
In [61]: Basis_PMZ_L = System.ProductOperators_ConvertToLiouville(Basis_PMZ)
```

Matrix representation of Product Operators into Liouville Space (PMZ Basis)

```
In [62]: Matrix(Basis_PMZ_L[0])
```

Out[62]:

Transformation Between From Zeeman to PMZ Basis (Liouvillie Space)

```
In [63]: U_Z_PMZ = System.Transform_StateBasis(Basis_Zeeman_L, Basis_PMZ_L)
```

Matrix representation of transformation matrix between Zeeman to PMZ Basis (Liouvillie Space)

```
In [64]:
            Matrix(System.Matrix_Round(System.Matrix_Tol(U_Z_PMZ, 1.0e-5), 2))
Out[64]:
                      0
                               0
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```

Zeeman Hamiltonian in Liouville Space (PMZ Basis)

```
In [65]: Hz_L_PMZ = System.Operator_BasisChange(Hz_L,U_Z_PMZ)
```

Matrix representation of Zeeman Hamiltonian in Liouville Space (PMZ Basis)

```
In [66]:
            Matrix(System.Matrix_Round(System.Matrix_Tol(Hz_L_PMZ/(2.0*np.pi),1.0e-5),2))
Out[66]:
             30.0
                      0
                              0
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```

Initialize Density Matrix

```
In [67]:
          Thermal_DensMatrix = True
          if Thermal_DensMatrix:
              # Spin temperature of individual spins (initial) Kelvin
              Tin = [300.0, 300.0]
              # Spin temperature of individual spins (equlibrium) Kelvin
              Tfi = [300.0, 300.0]
              # High Temperature
              HT_approx = False
              # Initial Density Matrix
              rho_in = System.EqulibriumDensityMatrix_Advance(LarmorF, Sz, Tin, HT_approx)
              # Equlibrium Density Matrix
              rhoeq = System.EqulibriumDensityMatrix_Advance(LarmorF, Sz, Tfi, HT_approx)
          else:
              rho_in = np.sum(Sz,axis=0)
              rhoeq = np.sum(Sz, axis=0)
```

Trace of density metrix = 1.0Trace of density metrix = 1.0

Converting initial and equlibrium density matrix into Liouvillian

```
In [68]: rho_in_L = System.Vector_L(rho_in)
rhoeq_L = System.Vector_L(rhoeq)
```

Pulse (Hilbert Space)

```
In [69]: flip_angle1 = 0.0  # Flip angle Spin 1
flip_angle2 = 180.0 # Flip angle Spin 2

rho = System.Rotate_H(rho_in,flip_angle1,Sy[0])
rho = System.Rotate_H(rho,flip_angle2,Sy[1])
```

Pulse (Liouville Space)

```
In [70]:
    rho_L = System.Rotate_L(rho_in_L,flip_angle1,Sy[0])
    rho_L = System.Rotate_L(rho_L,flip_angle2,Sy[1])
```

Relaxation in Hilbert Space

```
In [71]: R1 = None
R2 = None

# Correlation Time
tau = [10.0e-12]

# Dipolar coupling constant (Hz)
bIS = [30.0e3]

Rprocess = "Auto-correlated Dipolar Homonuclear"

System.Relaxation_Constants(R1,R2)
System.Relaxation_Parameters(LarmorF, OmegaRF, tau, bIS)
```

Relaxation Rate Spin 1

```
In [72]:
R1_rate = System.RelaxationRate_H(Sz[0],Sz[0],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)
Rcross_rate = System.RelaxationRate_H(Sz[1],Sz[0],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)
Longit_relaxa = R1_rate + Rcross_rate
R2_rate = System.RelaxationRate_H(Sp[0],Sp[0],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)

print("T1 = %.5f and T2 = %.5f" % ((1.0/R1_rate).real, (1.0/R2_rate).real))
print("R1 = %.5f and R2 = %.5f" % ((R1_rate).real, (R2_rate).real))
print("Cross Relaxation rate = %.5f and time = %.5f" % (Rcross_rate.real, 1.0/Rcross_rate)
print("Longitudinal Relaxation rate = %.5f and time = %.5f" % (Longit_relaxa.real, 1.0/Lor

T1 = 5.63856 and T2 = 5.63482
R1 = 0.17735 and R2 = 0.17747
Cross Relaxation rate = 0.08856 and time = 11.29210
```

Relaxation Rate Spin 2

```
In [73]: R1_rate = System.RelaxationRate_H(Sz[1],Sz[1],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)
    Rcross_rate = System.RelaxationRate_H(Sz[0],Sz[1],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)
```

Longitudinal Relaxation rate = 0.26591 and time = 3.76070

```
Longit_relaxa = R1_rate + Rcross_rate
R2_rate = System.RelaxationRate_H(Sp[1],Sp[1],Rprocess,R1,R2,Sx,Sy,Sz,Sp,Sm)

print("T1 = %.5f and T2 = %.5f" % ((1.0/R1_rate).real, (1.0/R2_rate).real))
print("R1 = %.5f and R2 = %.5f" % ((R1_rate).real, (R2_rate).real))
print("Cross Relaxation rate = %.5f and time = %.5f" % (Rcross_rate.real, 1.0/Rcross_rate.print("Longitudinal Relaxation rate = %.5f and time = %.5f" % (Longit_relaxa.real, 1.0/Lor
```

```
T1 = 5.63856 and T2 = 5.63482 R1 = 0.17735 and R2 = 0.17747 Cross Relaxation rate = 0.08856 and time = 11.29210 Longitudinal Relaxation rate = 0.26591 and time = 3.76070
```

Evolution of Density Matrix in Hilbert Space

```
In [74]:
    dt = 0.0001
    AQ = 50.0
    Npoints = int(AQ/dt)

method = "ODE Solver"
    ode_solver = 'DOP853'
    System.ODE_Method(ode_solver)

start_time = time.time()
    t, rho_t = System.Evolution_H(rhoeq,rho,Sx,Sy,Sz,Sp,Sm,Hz+Hj,dt,Npoints,method,Rprocess)
    end_time = time.time()
    timetaken = end_time - start_time
    print("Total time = %s seconds " % (timetaken))
```

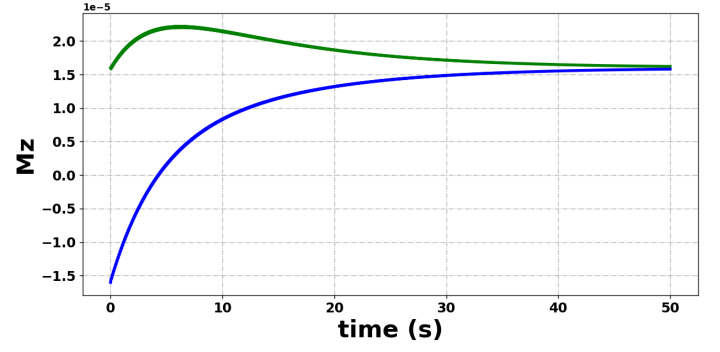
Total time = 9.728602886199951 seconds

Expectation Value (Hilbert Space)

```
In [75]: det_Z1 = Sz[0]
  det_Z2 = Sz[1]

  t, signal_Z1 = System.Expectation_H(rho_t, det_Z1, dt, Npoints)
  t, signal_Z2 = System.Expectation_H(rho_t, det_Z2, dt, Npoints)
```

```
In [76]: System.PlottingMulti(1,[t,t],[signal_Z1,signal_Z2],"time (s)","Mz",["green","blue"])
```



/opt/anaconda3/lib/python3.9/site-packages/numpy/core/_asarray.py:102: ComplexWarning: Cas ting complex values to real discards the imaginary part return array(a, dtype, copy=False, order=order)
/opt/anaconda3/lib/python3.9/site-packages/numpy/core/_asarray.py:102: ComplexWarning: Cas ting complex values to real discards the imaginary part return array(a, dtype, copy=False, order=order)
No handles with labels found to put in legend.

Relaxation in Liouville Space

```
In [77]: R = None
    Rprocess = "Auto-correlated Dipolar Homonuclear"
    tau = [10.0e-12]
    bIS = [30.0e3]
    System.Relaxation_Parameters(LarmorF, OmegaRF, tau, bIS)
    R_L = System.Relaxation_L(Rprocess, R, Sx, Sy, Sz, Sp, Sm)
```

Relaxation Rate Spin 1

```
In [78]:
R1_rate = System.RelaxationRate_L(Sz[0],Sz[0],R_L)
Rcross_rate = System.RelaxationRate_L(Sz[1],Sz[0],R_L)
Longit_relaxa = R1_rate + Rcross_rate
R2_rate = System.RelaxationRate_L(Sp[0],Sp[0],R_L)

print("T1 = %.5f and T2 = %.5f" % ((1.0/R1_rate).real, (1.0/R2_rate).real))
print("R1 = %.5f and R2 = %.5f" % ((R1_rate).real, (R2_rate).real))
print("Cross Relaxation rate = %.5f and time = %.5f" % (Rcross_rate.real, 1.0/Rcross_rate.
print("Longitudinal Relaxation rate = %.5f and time = %.5f" % (Longit_relaxa.real, 1.0/Lor

T1 = 5.63856 and T2 = 5.63482
R1 = 0.17735 and R2 = 0.17747
Cross Relaxation rate = 0.08856 and time = 11.29210
```

Relaxation Rate Spin 2

```
In [79]: R1_rate = System.RelaxationRate_L(Sz[1],Sz[1],R_L)
    Rcross_rate = System.RelaxationRate_L(Sz[0],Sz[1],R_L)
```

Longitudinal Relaxation rate = 0.26591 and time = 3.76070

```
Longit_relaxa = R1_rate + Rcross_rate
R2_rate = System.RelaxationRate_L(Sp[1],Sp[1],R_L)

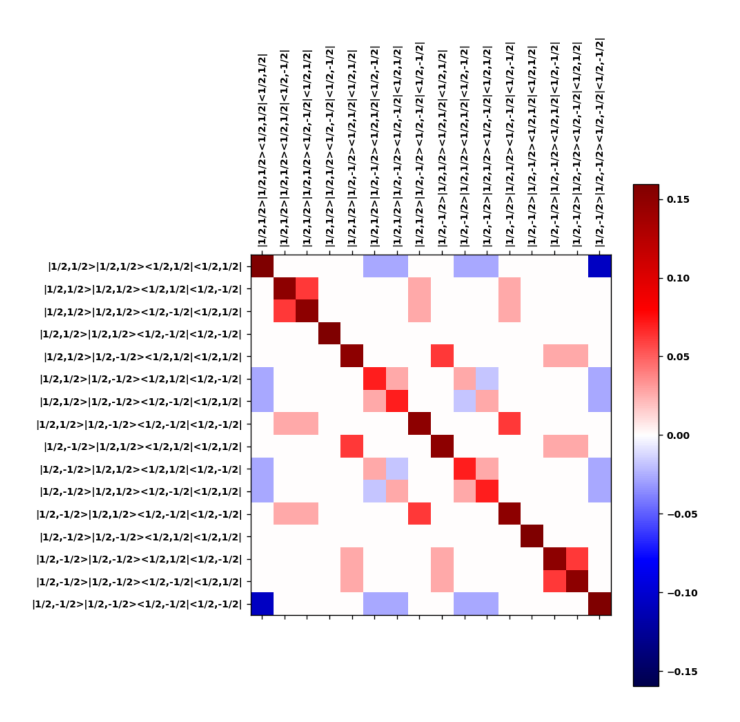
print("T1 = %.5f and T2 = %.5f" % ((1.0/R1_rate).real, (1.0/R2_rate).real))
print("R1 = %.5f and R2 = %.5f" % ((R1_rate).real, (R2_rate).real))
print("Cross Relaxation rate = %.5f and time = %.5f" % (Rcross_rate.real, 1.0/Rcross_rate.print("Longitudinal Relaxation rate = %.5f and time = %.5f" % (Longit_relaxa.real, 1.0/Lor
```

```
T1 = 5.63856 and T2 = 5.63482
R1 = 0.17735 and R2 = 0.17747
Cross Relaxation rate = 0.08856 and time = 11.29210
Longitudinal Relaxation rate = 0.26591 and time = 3.76070
```

Matrix Representation of Relaxation Superoperator in Liouville Space (Zeeman basis)

In [80]:

System.MatrixPlot(2, R_L.real, dic_Zeeman, dic_Zeeman)

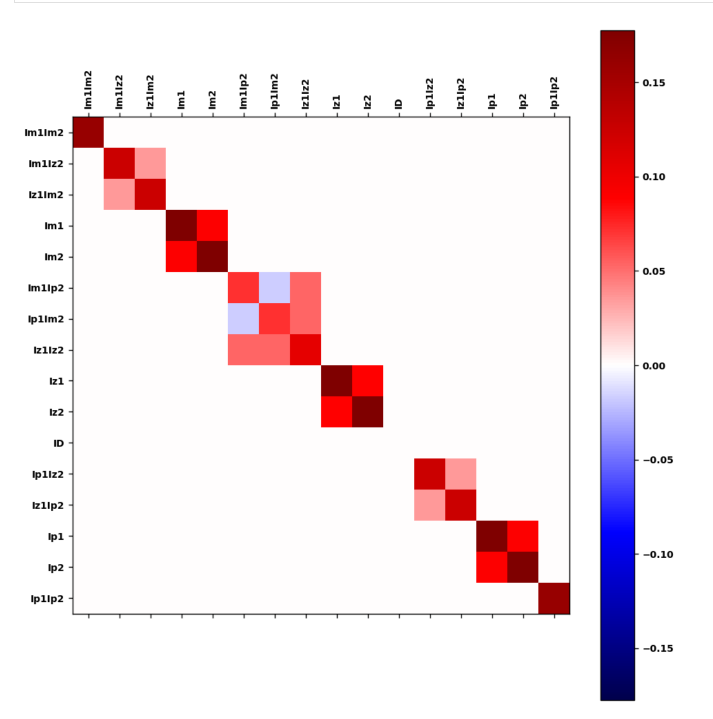


Converting Relaxation Superoperator in Liouville Space from Zeeman to PMZ basis

```
In [81]: R_L_PMZ = System.Operator_BasisChange(R_L.real,U_Z_PMZ).real
```

Matrix Representation of Relaxation Superoperator in Liouville Space (PMZ basis)

In [82]: System.MatrixPlot(3,R_L_PMZ,dic_PMZ,dic_PMZ)



Evolution of Density Matrix Liouville Space

```
In [83]: method = "ODE Solver"
System.ODE_Method('DOP853')
```

```
start_time = time.time()
t, rho_t = System.Evolution_L(rhoeq_L, rho_L, Sx, Sy, Hz_L, R_L, dt, Npoints, method)
end_time = time.time()
timetaken = end_time - start_time
print("Total time = %s seconds" % (timetaken))
```

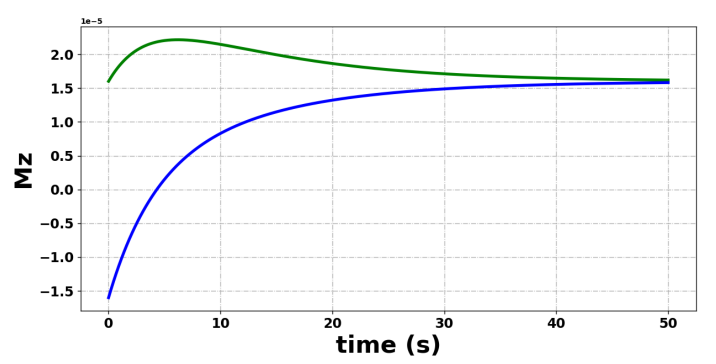
(16, 500000) Total time = 3.5411036014556885 seconds

Expectation value (Liouville Space)

```
In [84]: LEXP_Z1 = System.Detection_L(det_Z1)
    LEXP_Z2 = System.Detection_L(det_Z2)

In [85]: t, MZ_1 = System.Expectation_L(rho_t, LEXP_Z1, dt, Npoints)
    t, MZ_2 = System.Expectation_L(rho_t, LEXP_Z2, dt, Npoints)

In [86]: System.PlottingMulti(4,[t,t],[MZ_1,MZ_2],"time (s)","Mz",["green","blue"])
```



/opt/anaconda3/lib/python3.9/site-packages/numpy/core/_asarray.py:102: ComplexWarning: Cas
ting complex values to real discards the imaginary part
 return array(a, dtype, copy=False, order=order)
/opt/anaconda3/lib/python3.9/site-packages/numpy/core/_asarray.py:102: ComplexWarning: Cas
ting complex values to real discards the imaginary part
 return array(a, dtype, copy=False, order=order)
No handles with labels found to put in legend.

Remarks

If you see something is wrong, please write to me.

Any suggestion? write to me

vineethfrancis.physics@gmail.com