# Homework 5

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## NUEN 629, Homework 5

Due Date Dec. 3

Solve the following problem and submit a detailed report, including a justification of why a reader should believe your results.

## 1 Clean Fusion Energy ¬\\_('ソ)\_/¬

(100 points) Consider a thermonuclear fusion reactor producing neutrons of energy 14.1 and 2.45 MeV. The reactor is surrounded by FLiBe (a 2:1 mixture of LiF and BeF<sub>2</sub>, https://en.wikipedia.org/wiki/FLiBe) to convert the neutron energy into heat. All the constituents in the FLiBe have their natural abundances. Using data from Janis (https://www.oecd-nea.org/janis/). Assume the total neutron flux is  $10^{14}$  n/cm<sup>2</sup>·s. Perform the following analyses.

- 1. (25 points) Write out the depletion (or in this case activation) chains that will occur in the system.
- 2. (50 points) Over a two-year cycle compute the inventory of nuclides in the system using two methods discussed in class. What is the maximum concentration of tritium?
- 3. (25 points) After discharging the FLiBe blanket, how long will it take until the material is less radioactive than Brazil nuts (444 Bq/kg, http://www.orau.org/PTP/collection/consumer%20products/brazilnuts.htm).

### Solution 1-1:

I tracked all of the nuclides suggested in the document FLiBE use in Fusion Reactors: an Initial Safety Assessment by L.C Cadwallader and G.R. Longhurst. The decay paths from this document are given below.

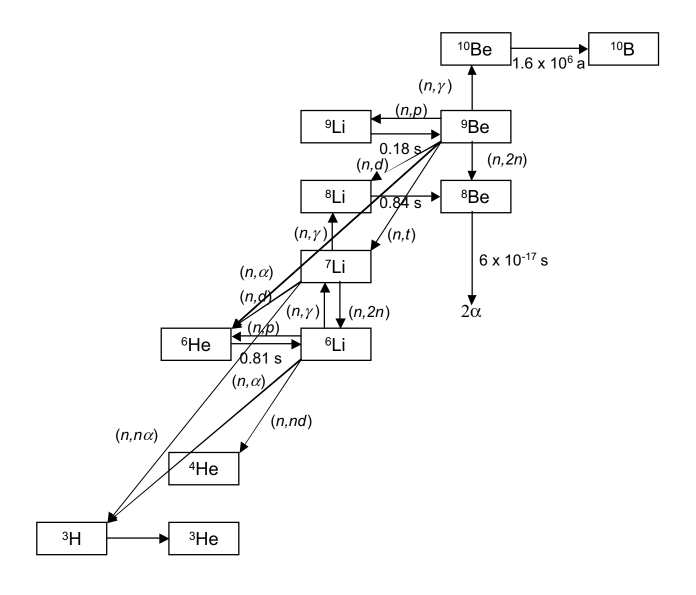
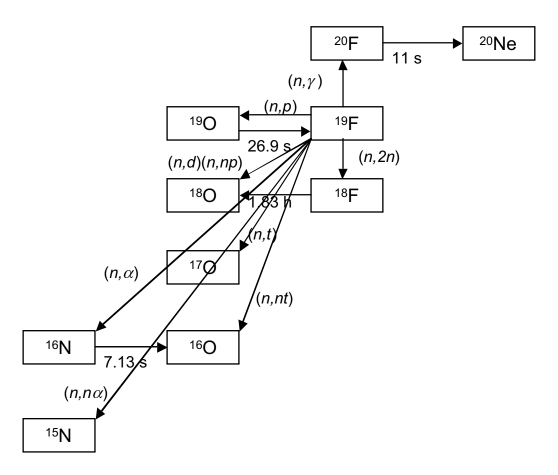


Figure 1. Activation and decay paths for lithium and beryllium in Flibe.[40]



**Figure 2**. Activation and decay processes for fluorine in Flibe and oxygen impurity.[40]

Considering that stainless steel 316 with a Mn component (rather than nickel) might be present, erodants could be generated and activated [38], giving the following radioisotopes:

$$^{54}{
m Mn}$$
  $^{55}{
m Fe}$   $^{58}{
m Co}$   $^{60}{
m Co}$ 

Some of these radioisotopes may oxidize in air, but concentrations should be low since the impurity concentrations were in the ppm range. The actual curie inventory of these radioisotopes depends on fluence and the level of impurity or erosion. For HYLIFE-II, Tobin [38] showed site boundary doses between 1 and 26 rem for each of the impurity and erodant radioisotopes listed above. The <sup>18</sup>F from pure Flibe gave a site boundary dose of 340 rem for HYLIFE-II. Considering that workers would be closer to the more concentrated source of these radioisotopes, their doses would be higher than the values stated for the site boundary unless appropriate mitigative actions are taken. Specific calculations need to be performed for the magnetic fusion designs under consideration to determine the worker and the site boundary doses. While these calculations are important, it is also noted that the MSRE operated successfully from 1965 to 1969 without any extreme personnel safety events (i.e., no radiation exposure over 15 rem, and

#### Solution 1-2:

The relative abundances of each isotope is computed assuming natural abundances and the chemical makeup of Li<sub>2</sub>BeF<sub>4</sub>. The initial atom fractions are

$$\vec{n_0} = \begin{pmatrix} {}^{6}\text{Li} \\ {}^{7}\text{Li} \\ {}^{9}\text{Be} \\ {}^{19}\text{F} \end{pmatrix} = \begin{pmatrix} 0.0214 \\ 0.2643 \\ 0.1429 \\ 0.5714 \end{pmatrix} \tag{1}$$

with the rest of the tracked isotopes from part 1 set to zero. The neutron flux was assumed to be 90% at 14.1 MeV and 10% at 2.45 MeV. The blanket is assumed thin enough that scattering is negligible, so neutrons remain at these two energies, which is fairly inaccurate as Be is a strong moderator. The invetory of nuclides in the system was computed using a parabolic contour integral approximation to the exponential matrix and with backward Euler. The backward Euler was implemented by taking 100 time steps between each data point, using the previous data point as the initial conditon. With this approach the results were found to agree to with at least two digits of accuracy. All tritium (h1) and  $\alpha$  (he4) production are tracked as well. The plots of isotopic atom fractions are given below. The mass of tritium per initial kg of FLiBe after two years of irradiation is computed by multiplying the atom fraction by the ratio of atomic masses; this ratio was found to be  $1.04 \times 10^{-04}$  kg of tritium per initial kg of FLiBe.

#### Solution 1-3:

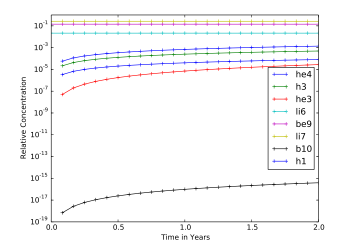
The activity, per unit mass of FLiBe, is computed as

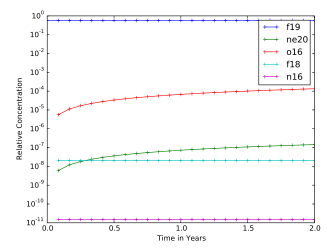
$$A(t) = 7 \sum_{j=1}^{N} \frac{\gamma_j(t) N_A}{\mathcal{A}_{\text{FLiBe}}} \lambda_j$$
 (2)

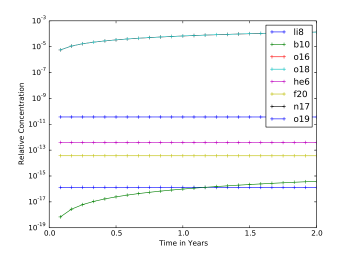
where  $\gamma_j(t)$  is the atom fraction for the j-th of N isotopes at time t after irradiation stops. The matrix exponential and backward Euler were found to be numerically inaccurate for computing the new activities of isotopes, resulting in negative answers, due to the large range of decay constants relative to the time scale. Instead, the atom fractions were computed ignoring secondary decay chains as

$$\gamma_j(t) = \gamma_j(t_0)e^{-\lambda_j t} \tag{3}$$

where  $\gamma_j(t_0)$  is the atom fractions after irradiation stops at 2 years. The factor of 7 is because my atom fractions are normalized to be atom of isotope j per any atom in the system, rather than per mol (which would not sum to 1). With this approach, it was found it will take  $395 \times 10^3$  years before the activity is below the threshold of 444 Bq/kg. This is completely dominated by the slow decay of <sup>10</sup>Be, although the majority of the radioactivity is removed within 100 years once the tritium has decayed away.







#### Code

```
import numpy as np
    import scipy as sp
    import scipy.sparse as sparse
    import scipy.sparse.linalg as splinalg
    import matplotlib.pyplot as plt
6
   import re
   # In [107]:
9
10
   z = [5.623151880088747 + 1.194068309420004j,
11
          5.089353593691644 + 3.588821962583661j,
          3.993376923428209 + 6.004828584136945j,
13
          2.269789514323265 + 8.461734043748510j,
14
          -0.208754946413353 + 10.991254996068200j,
         -3.703276086329081 + 13.656363257468552j
16
         -8.897786521056833 + 16.630973240336562j
17
18
    c = \begin{bmatrix} -0.278754565727894 - 1.021482174078080j, \end{bmatrix}
          0.469337296545605 \; + \; 0.456439548888464 \mathtt{j} \; ,
19
20
          -0.234984158551045 - 0.058083433458861j,
          0.048071353323537 - 0.013210030313639j
21
          -0.003763599179114 + 0.003351864962866j
22
          0.000094388997996 - 0.000171848578807j,
23
24
          -0.000000715408253 + 0.000001436094999j,
25
26
   ## Depletion Example
27
28
   # In [84]:
29
30
   #cross-sections in barns
31
   33
34
    pos = \{\}
    for i in range(len(ids)):
35
        pos [ ids [ i ]] = i
36
37
38
    stoa = 1./31557600
39
40
    siga = \{ 'h1' : [0.0, 0.0] ,
41
             ^{\prime}h3 ^{\prime}: [0.0, 0.98287 - 0.93317],
42
             'he4':[0.0,0.0],
43
             'he3': [3.090702 - 2.364785, 1.17 - 0.97],
44
             'he6':[0.0,0.0],
45
             'li6': [1.5421 - 1.27645, 1.448249 - 0.861308]
46
             'li7': [1.920451 - 1.733316, 1.44098 - 1.01017],
47
             'li8':[0.0,0.0],
'li9':[0.0,0.0],
48
49
             'be9': [2.510747-2.401078, 1.52753-1.10746],
50
             be10': [1.0e-5, 1.0e-5],
51
             'b10': [2.05-1.6767, 1.4676-0.907458],
52
             \frac{19}{19}: [3.040797 - 2.134332, 1.76351 - 0.9528]
53
54
55
   #Missing ones to zero
    for i in ids:
58
        if i not in siga:
             siga[i] = [0.0, 0.0]
59
60
    {\tt cap} \ = \ \left\{ \ {\tt 'li6':[\ 'li7',1.106851e-5,1.017352e-5]} \right.,
61
            'li7':['li8',4.670126e-6,4.1075e-6],
'be9':['be10',1.e-6,1.e-6],
62
63
            'f19': ['f20', 8.641807e - 5, 3.495e - 5]
64
                                                             }
65
```

```
n2n = \{ 'li7' : [ 'li6', 0.0, 0.03174] ,
66
             'be9':['he4-he4',0.0255,0.486034],
'f19':['f18',0,0.04162] }
67
68
69
     nalpha = \{ 'li7' : ['h3-he4', 0.0, 0.30215], \}
70
                 'li6':['h3-he4',0.206155,0.025975],
71
                 'be9': ['he6-he4',0.090833,0.0105],
'f19': ['he4-n16',2.551667e-5,0.028393]
72
73
74
     ntrit = \{'be9': ['li7-h3', 0.0, 0.02025],
75
                 'f19':['o16-h3',0.0,0.01303],}
76
77
    78
79
               'f19':['o19-h1',0.0,0.018438]}
80
81
82
    decays = { 'h3':['he3',12.5], 'he6':['li6',0.81*stoa],
83
84
                 'li8':['he4-he4',0.84*stoa],
85
                 'li9':['be9',0.5*0.18*stoa, 'he4-he4',0.5*0.18*stoa],
86
                 'be10':['b10',1.6E6],
'f18':['o18',6586*stoa],
87
88
                'f20':['ne20',11.16*stoa],
89
                'o19': ['f19',26.9*stoa],
90
                'n16':['o16',7.13*stoa]}
91
92
    #Convert halflifes to decays
93
     for i in decays:
94
         for j in range(0,len(decays[i]),2):
95
              decays[i][j+1] = np.log(2)/decays[i][j+1]
96
97
    A = np.zeros((len(ids), len(ids)))
98
99
    phi \, = \, 1.0\,e14 \, * \, 60 \, * \, 60 \, * \, 24 \, * \, 365 \quad \#10\,^{\hat{}}14 \, 1/cm\,^{\hat{}}2/s \, in \, 1/cm\,^{\hat{}}2/year
100
    phi = phi *1.0e-24 # neutrons/barns-year
    phi1 = 0.1*phi
102
    phi2 = 0.9*phi
103
     for i in ids:
104
105
         row = pos[i]
106
         A[row, row] = - phi1*siga[i][0] - phi2*siga[i][1]
107
         if i in decays:
              #sum over branching ratios
108
              A[row, row] = sum(decays[i][j+1] \text{ for } j \text{ in } range(0, len(decays[i]), 2))
109
110
         #Loop over all reaction types
111
         for r in [cap, n2n, nalpha, ntrit, nprot]:
              if i in r:
113
                   target = r[i][0]. split("-")
114
                   for t in target: #from i to target
115
                       A[pos[t], row] += phi1*r[i][1] + phi2*r[i][2]
116
117
         #Loop over decays
118
         if i in decays:
119
              for j in range(0,len(decays[i]),2): #in sets of 2
120
                   #the first member is what it decays to, second is decay constant
121
                   target = decays[i][0+j].split("-") #if goes to two things, hyphen
122
                   for t in target:
123
124
                       A[pos[t], row] += decays[i][1+j] #A[target, src] = decay
126
127
     for j in ids:
         print ("Row: h3, column: ",j,"value", A[pos['h3'], pos[j]]/365)
128
     plt.spy(A)
129
130
131
    #Initial condition
    n0 = np.zeros(len(ids))
132
    abund = \{'1i6': (0.075*2),
```

```
134
               'li7':(0.925*2),
               'be9':1.,
135
136
               'f19':4.}
     for i in ids:
137
         if i in abund:
138
             n0[pos[i]] = abund[i]
139
140
141
    n0 /= sum(n0)
142
143
    from scipy.linalg import expm
144
145
    Npoints = (12,) #(2,4,6,8,10,12,14,16,18,20,22,24,26,28,30,32)
146
    times = np.linspace (0,2,num=25) #in years
147
    conc_{exp} = np.zeros((times.shape[0], n0.shape[0]))
148
149
    conc_be = np.zeros((times.shape[0], n0.shape[0]))
150
     for ti in range (times.shape [0]):
         t = times [ti]
152
153
         n = n0.copy()
         for N in Npoints:
154
155
             pos1 = 0
156
             theta = np.pi*np.arange(1,N,2)/N
             z = N*(.1309 - 0.1194*theta**2 + .2500j*theta)
157
             w = N*(-2*0.1194*theta + .2500j)
158
             c = 1.0 j/N*np.exp(z)*w
159
160
             #plt.plot(np.real(z),np.imag(z),'o-')
161
             #plt.show()
             u = np.zeros(len(n))
162
              for k in range (int(N/2)):
163
                  n, code = splinalg.gmres(z[k]*sparse.identity(len(n)) - A*t, n0, tol=1e-12,
164
                           maxiter=20000)
165
                  if (code):
166
                      print(code)
167
168
                  u -= c[k]*n
             u = 2*np.real(u)
169
         conc_exp[ti,:] = u
170
171
172
         #Backward euler
173
         T = 100
         if ti == 0:
174
             dt = 0.0
             n = n0.copy()
176
177
         else:
             dt = (t - times [ti -1])/T
178
             n = conc_be[ti-1,:].copy()
179
180
         I = sparse.identity(len(n0))
181
         for i in range(T):
182
    #
          print ("Iteration", i)
183
             n = splinalg.gmres(I - A*dt, n, tol=1e-12)[0]
184
         \verb|conc_be[ti|,:]| = n
185
186
    plot1 = ['he4', 'h3', 'he3', 'li6', 'be9', 'li7', 'b10', 'h1']
187
    plot2 = ['f19', 'ne20', 'o16', 'f18', 'n16']
plot3 = ['li8', 'b10', 'o16', 'o18', 'he6', 'f20', 'n17', 'o19']
188
189
190
    A_{trit} = 3.0160492
191
    A_{\text{-flibe}} = 18.99*4 + 6.94 * 2 + 9.01
    print ("Mass \ ratio \ of \ tritium", \ conc\_exp[-1,pos['h3']]*7*A\_trit/A\_flibe)
193
194
195
    #Plot concentrations
196
    plt.figure()
197
    for i in plot1:
198
199
         plt.semilogy(times,conc_be[:,pos[i]],"-+",label=i)
    plt.legend(loc='best')
200
    plt.xlabel("Time in Years")
201
```

```
plt.ylabel("Relative Concentration")
     plt.savefig('pl.pdf',bbox_inches='tight')
203
204
205
     plt.figure()
     for i in plot2:
206
          plt.semilogy(times,conc_be[:,pos[i]],"-+",label=i)
207
     plt.legend(loc='best')
208
     plt.xlabel("Time in Years")
plt.ylabel("Relative Concentration")
209
210
     plt.savefig('p2.pdf',bbox_inches='tight')
211
212
     plt.figure()
213
     for i in plot3:
214
          plt.semilogy\,(\,times\,,conc\_be\,[:\,,pos\,[\,i\,]\,]\,\,,"-\!+"\,\,,labe\,l\!=\!i\,)
215
     plt.legend(loc='best')
216
     plt.xlabel("Time in Years")
217
     plt.ylabel("Relative Concentration")
218
     plt.savefig('p3.pdf',bbox_inches='tight')
219
220
221
     #Compute activities
222
223
     n0 = conc_be[-1,:].copy()
224
     n = n0.copy()
225
    #Rebuild A with no flux
226
     A = np.zeros((len(ids), len(ids)))
227
228
     for i in ids:
229
          row = pos[i]
          if i in decays:
230
              #sum over branching ratios
231
              A[row, row] = sum(decays[i][j+1] \text{ for } j \text{ in } range(0, len(decays[i]), 2))
232
233
          #Loop over decays
234
          if i in decays:
235
               for j in range(0,len(decays[i]),2): #in sets of 2
236
                   #the first member is what it decays to, second is decay constant
237
                   target = decays[i][0+j].split("-") #if goes to two things, hyphen
238
                   for t in target:
239
240
                        A[pos[t], row] += decays[i][1+j] #A[target, src] = decay
241
242
     #Use BE to advance in time the concentrations
     dt = 100
243
     t = 0.
244
     conc_decay = []
245
246
     times = []
     Na = 6.0221E23
247
248
     #Convert n to Atom densities
249
     n0 = Na/A_flibe*1000*7
250
     n *= Na/A_flibe*1000*7
251
     decay\_const = \{\}
252
253
     for i in ids:
          if i in decays:
254
               \operatorname{decay-const}[i] = \operatorname{sum}(\operatorname{decays}[i][j+1] \text{ for } j \text{ in } \operatorname{range}(0, \operatorname{len}(\operatorname{decays}[i]), 2)) * \operatorname{stoa}
255
256
          else:
              decay\_const[i] = 0.
257
258
     while True:
259
260
          t += dt
261
262
263
          #Compute based on exponential decay
          activity_new_v = [n0[pos[i]]*decay_const[i]*np.exp(-1.*decay_const[i]*t/stoa) for i in ids]
264
265
          N_{\text{new}} = [n0[pos[i]]*np.exp(-1.*decay\_const[i]*t/stoa)  for i in ids]
          activity_new = sum(activity_new_v)
266
           print("Activity", activity_new_v)
print("Nuces", N_new)
267
    #
268
     #
    #
           print ("pos", pos)
269
```

```
270 #
271 #
272
   #
          input()
    ##
273
274
         times.append(t)
275
        conc_decay.append(activity_new)
print(activity_new_v[pos['h3']],activity_new_v[pos['be10']])
print("Current activity: ",activity_new ,"time: ",t)
276
277
278
279
         if activity_new < 444 or t > 10000000:
280
             break
281
```