$\begin{array}{c} {\rm NUEN~629} \\ {\rm Numerical~Methods~in~Reactor~Analysis} \\ {\rm Homework~4~\&~5~\&~Project} \end{array}$

Due on: Thursday, November 19, 2015 & & Thursday, December 3, 2015

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Homework 4 Problem Statement

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

1. (150 points + 50 points extra credit) In class we discussed the diamond-difference spatial discretization. Another discretization is the step discretization (this has several other names from other disciplines). It writes the discrete ordinates equations with isotropic scattering as, for $\mu_n > 0$ to

$$\mu_n \frac{\psi_{i,n} - \psi_{i-1,n}}{h_r} + \Sigma_t \psi_{i,n} = \frac{\Sigma_s}{2} \phi_i + \frac{Q}{2}$$
 (1)

and for $\mu_n < 0$

$$\mu_n \frac{\psi_{i+1,n} - \psi_{i,n}}{h_x} + \Sigma_t \psi_{i,n} = \frac{\Sigma_s}{2} \phi_i + \frac{Q}{2}$$
 (2)

The codes provided in class should be modified to implement this discretization.

- (a) (50 Points) Your task (should you choose to accept it) is to solve a problem with uniform source of Q = 0.01, $\Sigma_t = \Sigma_s = 100$ for a slab in vacuum of width 10 using step and diamond difference discretizations. Use, 10, 50, and 100 zones ($h_x = 1, 0.02, 0.01$) and your expert choice of angular quadratures. Discuss your results and how the two methods compare at each number of zones.
- (b) (10 points) Discuss why there is a different form of the discretization for the different signs of μ .
- (c) (40 points) Plot the error after each iteration using a 0 initial guess for the step discretization with source iteration and GMRES.
- (d) (50 points) Solve Reed's problem (see finite difference diffusion codes). Present convergence plots for the solution in space and angle to a "refined" solution in space and angle.
- (e) (50 points extra credit) Solve a time dependant problem for a slab surrounded by vacuum with $\Sigma_t = \Sigma_s = 1$ and initial condition given by $\psi(\mathbf{0}) = \mathbf{1}/h_x$ (original problem statement said $\phi(0) = 1/h_x$ and I'm not sure how to solve that). Plot the solution at t = 1 s, using step and diamond difference. The particles have a speed of 1 cm/s. Which discretization is better with a small time step? What do you see with a small number of ordinates compared to a really large number (100s)?

Homework 4 Problem Background

Due to the complicated nature of this course, I provided this background for the lay person (me), so that they might have some grounding for the solution and hopefully believe the results. It should be noted that most of this background information is copied from various points in Dr. McClarren's notes, and is in no way original. Anything intelligent in the following is due to this fact and for any errors, I blame myself.

Beginning with the weighty neutron transport equation.

$$\left(\frac{1}{v}\frac{\delta}{\delta t} + \hat{\Omega}\cdot\nabla + \Sigma_t\right)\psi = \int_0^\infty dE' \int_{4\pi} d\hat{\Omega}' K(\hat{\Omega}'\cdot\hat{\Omega}, v'\to v)\Sigma_s\psi + \frac{1}{4\pi}\chi\int_0^\infty dE' \bar{\nu}\Sigma_f\phi + q$$

Where $K(\hat{\Omega}' \cdot \hat{\Omega}, v' \to v)$ represents the probability of scattering from one angle and energy to another given a scattering event occurred and Σ_s is the macroscopic scattering cross section. The dependencies for the variables are shown below.

$$\Sigma_{t}(\vec{x}, v, t)$$

$$\psi(\vec{x}, \hat{\Omega}, v, t)$$

$$\Sigma_{s}(\vec{x}, v, t)$$

$$\chi(\vec{x}, v)$$

$$\Sigma_{f}(\vec{x}, v, t)$$

$$\phi(\vec{x}, v, t)$$

$$q(\vec{x}, \hat{\Omega}, v, t)$$

There are 7 free variables (three spatial $[\vec{x}]$, two angular $[\hat{\Omega}]$, one energy [v] and one time [t]) in this equation. In the steady state $\left(\frac{\delta\psi}{\delta t}=0\right)$, i.e. no time dependence, non fissioning $(\Sigma_f=0)$ case the transport equation reduces to,

$$\left(\hat{\Omega} \cdot \nabla + \Sigma_t\right) \psi = \int_0^\infty dE' \int_{4\pi} d\hat{\Omega}' K(\hat{\Omega}' \cdot \hat{\Omega}, v' \to v) \Sigma_s \psi + q.$$

In order to reduce this to a single energy the following definitions are helpful (remembering all time dependence is gone).

$$\begin{split} \psi(\vec{x}, \hat{\Omega}) &= \int_0^\infty dE \ \psi(\vec{x}, \hat{\Omega}, v(E)) \\ \Sigma_t(\vec{x}) &= \frac{\int_0^\infty dE \ \Sigma_t(\vec{x}, v(E)) \psi(\vec{x}, \hat{\Omega}, v(E))}{\psi(\vec{x}, \hat{\Omega})} \\ K(\hat{\Omega}' \cdot \hat{\Omega}, v' \to v) &= K(\hat{\Omega}' \cdot \hat{\Omega}) K(v' \to v) \\ \Sigma_s(\vec{x}) &= \frac{\int_0^\infty dE \int_0^\infty dE' \ \Sigma_s(\vec{x}, v(E)) K(v' \to v) \psi(\vec{x}, \hat{\Omega}, v(E))}{\psi(\vec{x}, \hat{\Omega})} \\ q(\vec{x}, \hat{\Omega}) &= \int_0^\infty dE \ q(\vec{x}, \hat{\Omega}, v(E)) \end{split}$$

Using these definitions, integrating the transport equation over all energy, and assuming cross sections and sources do not vary in space or angle, our transport equation reduces again to,

$$\left(\hat{\Omega} \cdot \nabla + \Sigma_t\right) \psi(\vec{x}, \hat{\Omega}) = \int_{A\pi} d\hat{\Omega}' \ K(\hat{\Omega}' \cdot \hat{\Omega}) \Sigma_s \psi(\vec{x}, \hat{\Omega}') + q.$$

Where the double differential was assumed to be separable in angle and energy. The final simplification for our problem will be in space. If we assume that our geometry is infinite in y $\left(\frac{\delta}{\delta y}=0\right)$ and x $\left(\frac{\delta}{\delta x}=0\right)$. This also means that ψ depends only on z and mu, and if we recall that

$$\hat{\Omega} = (\sqrt{1 - \mu^2} \cos(\rho), \sqrt{1 - \mu^2} \sin(\rho), \mu),$$

and

$$\nabla = \left(\frac{\delta}{\delta x}, \frac{\delta}{\delta y}, \frac{\delta}{\delta x}\right)$$

also assuming that

$$K(\hat{\Omega}' \cdot \hat{\Omega}) = \frac{1}{4\pi}$$
 Isotropic Scattering

then our transport equation, and the equation I think we are trying to solve for this homework is.

$$\left(\mu \frac{\delta}{\delta z} + \Sigma_t\right) \psi(z, \mu) = \Sigma_s \frac{2\pi}{4\pi} \int_{-1}^1 d\mu' \ \psi(z, \mu') + q.$$

Checking units,

$$\left(\mu \frac{\delta}{\delta z} + \Sigma_t\right) \left[\frac{1}{cm}\right] \psi(z,\mu) \left[\frac{n \cdot cm}{str \cdot cm^3 \cdot s}\right] = \Sigma_s \frac{1}{2} \left[\frac{1}{cm \cdot rad}\right] \int_{-1}^1 d\mu' \, \psi(z,\mu') \left[\frac{n \cdot cm}{rad \cdot cm^3 \cdot s}\right] + q \left[\frac{n}{str \cdot cm^3 \cdot s}\right].$$

 Σ_s was moved outside the integral because it has no angular dependence integration over the azimuthal angle occurred because $\psi(z,\hat{\Omega})$ is assumed to be uniform and not depend on that angle.

Using Gauss-Legendre Quadrature for the integration term

$$\phi = \int_{-1}^{1} d\mu' \psi(z, \mu') = \sum_{i=1}^{n} w_i \psi(z, \mu'_i)$$

where

$$w_i = \frac{2}{(1 - \mu_i^2)[P_n'(\mu_i)]^2}$$

 P'_n is the differential of the legendre polynomial n, and μ'_i are the roots of P_n . The weights of even n's of the legendre polynomials should sum to 2, the value of $\int_{-1}^{1} d\mu$, which they do.

Putting this all together with time dependence:

$$\left(\frac{1}{v}\frac{\delta}{\delta t} + \mu \frac{\delta}{\delta z} + \Sigma_t\right)\psi_n(z) = \sum_s \frac{1}{2} \sum_{n'=1}^N w_{n'}\psi_{n'}(z) + q$$

Where n and n' denote the direction being solved for and N is the total number of angles being solved for. Also unites of w are rad.

Diamond difference discretization

$$\frac{1}{v}\frac{\psi_{n,i}^{\ell+1,j+1}-\psi_{n,i}^{L,j}}{\Delta t} + \mu_n\frac{\psi_{n,i+1/2}^{\ell+1,j+1}-\psi_{n,i-1/2}^{\ell+1,j+1}}{hz} + \Sigma_t\psi_{n,i}^{\ell+1,j+1} = \Sigma_s\frac{1}{2}\sum_{i=1}^Nw_i\psi_{n',i}^{\ell,j+1} + q.$$

Where n is for angle, i is the midplane of a spacial discretization, ℓ is the iteration index for spacial convergence, j is for a time step and

$$\psi_{n,i}^{\ell+1,j+1} = \frac{1}{2} (\psi_{n,i+1/2}^{\ell+1,j+1} + \psi_{n,i-1/2}^{\ell+1,j+1})$$

Writing this in terms of a steady state

$$\mu_n \frac{\psi_{n,i+1/2}^{\ell+1,j+1} - \psi_{n,i-1/2}^{\ell+1,j+1}}{hz} + \Sigma_t^* \psi_{n,i}^{\ell+1,j+1} = \Sigma_s \frac{1}{2} \sum_{n'=1}^N w_i \psi_{n',i}^{\ell,j+1} + q^*.$$

where

$$\Sigma_t^* = \Sigma_t + \frac{1}{v\Delta t}$$
$$q^* = q + \frac{\psi_{n,i}^{L,j}}{v\Delta t}$$

The above equation has L for the iteration index to indicate that its value was iteratively determined in the previous time step.

Step discretization

Writing this in terms of a steady state for $\mu > 0$

$$\mu_n \frac{\psi_{n,i}^{\ell+1,j+1} - \psi_{n,i-1}^{\ell+1,j+1}}{hz} + \Sigma_t^* \psi_{n,i}^{\ell+1,j+1} = \Sigma_s \frac{1}{2} \sum_{n'=1}^N w_i \psi_{n',i}^{\ell,j+1} + q^*.$$

and for $\mu < 0$

$$\mu_n \frac{\psi_{n,i+1}^{\ell+1,j+1} - \psi_{n,i}^{\ell+1,j+1}}{hz} + \Sigma_t^* \psi_{n,i}^{\ell+1,j+1} = \Sigma_s \frac{1}{2} \sum_{n'=1}^N w_i \psi_{n',i}^{\ell,j+1} + q^*.$$

where

$$\Sigma_t^* = \Sigma_t + \frac{1}{v\Delta t}$$
$$q^* = q + \frac{\psi_{n,i}^{L,j}}{v\Delta t}$$

GMRES

The generalized minimium residual (GMRES) method is an iterative method for solving linear systems of equations. The method approximates the solution by the vector in a Krylov subspace with a minimum residual (see wikipedia or Dr. McClarren's notes, I'm not really sure how this method works, but python has a solver for it).

The system $A\vec{\phi} = b$ is solved with GMRES, where for our situation,

$$A = \left(I - \sum_{n'=1}^{N} L^{-1} \Sigma_{s} \frac{1}{2}\right)$$

where L^{-1} is a sweep solve for our system and acts as an operator (I think), and

$$b = \sum_{n'=1}^{N} L^{-1} q^*$$

Reeds Problem

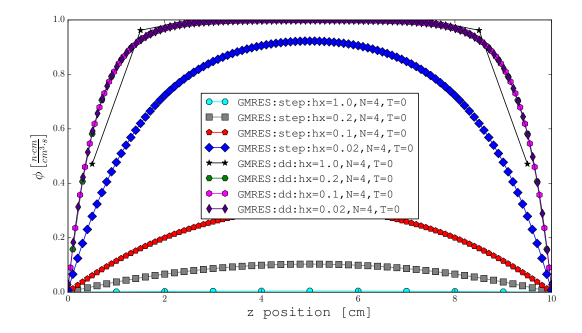
Reeds problem is a similar system as above, except the source and scattering and total cross sections are variable in z, and the width of z is 16.

Homework 4 Problem Solution

The code for this problem will be at the end of this section. The answers are below.

(a) (50 Points) Your task is to solve a problem with uniform source of Q = 0.01, $\Sigma_t = \Sigma_s = 100$ for a slab in vacuum of width 10 using step and diamond difference discretizations. Use, 10, 50, and 100 zones $(h_x = 1, 0.02, 0.01)$ and your expert choice of angular quadratures. Discuss your results and how the two methods compare at each number of zones.

The angular quadrature used was the Gauss-Legendre Quadrature because of the integration range. Its form was shown in the background section. The plot below was produced with the GMRES method, but the source iteration scheme produced the same results.



Both of the iterative solutions converged with max iterations of 100,000 and a slight modification on cross section ($\Sigma_t = \Sigma_t \cdot 1.0001$) to help the system converge. As the number of zones increased for the step solution, the flux magnitude kept increasing to match with the diamond difference and maintained a cosine(ish) shape. As the number of zones increased with the diamond difference, the shape started to converge towards the cosine, but maintained the proper magnitude.

Something else I would like to point out in the solution is that the step solution always had one more point plotted than the diamond difference. The reason for this is due to how each solution was solved. This is easier highlighted (for me) with an example, which is shown in the case where the number of zones is 10.

For the Diamond difference, the average locations (remember they were averaged), $\psi_{n,i}^{L,j+1}$, being solved for were,

$$z = [0.5, 1.5, 2.5, 3.5, 4.5, 5.5, 6.5, 7.5, 8.5, 9.5]$$

The points for $\psi_{n,i+1/2}^{L,j+1}$ and $\psi_{n,i-1/2}^{L,j+1}$ were at the points,

$$z = [0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10]$$

When sweeping to the right, $\psi_n(z=0)$ was set to zero, because the incoming flux is zero, and all points were solved for up to where z=10, and $\psi_{n,i}$ values were determined with averaging. This same thing occurred when sweeping to the left (except here $\psi_n(z=10)$ was set to zero). This would yield 10 values at the points [0.5, 1.5, ..., 9.5].

For the step discretization scheme, the locations (non averaged), $\psi_{n,i}^{L,j+1}$, being solved for were,

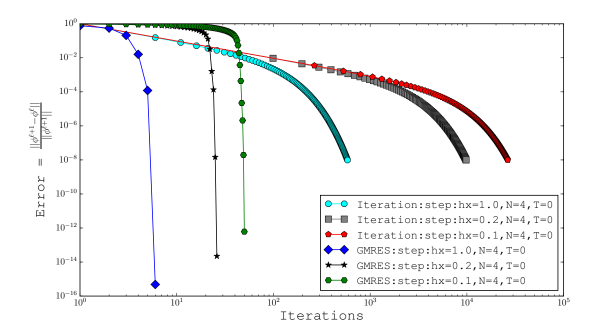
$$x = \begin{cases} [1, 2, 3, 4, 5, 6, 7, 8, 9, 10] & \mu > 0 \\ [0, 1, 2, 3, 4, 5, 6, 7, 8, 9], & \mu < 0 \end{cases}$$

When combining these two lists for ϕ , this was considered, and hence the step discretization scheme had one extra point (both lists have 10 points, but the location 10 is unquie in the first list, and 0 in the second).

- (b) (10 points) Discuss why there is a different form of the discretization for the different signs of μ .
 - The different forms are needed in the step discretization because in both the diamond and step approaches to the solution a value is needed from a previous zone. Our vacuum boundary condition states that the incoming neutrons are zero, which at the left side of the boundary, determines the angular flux moving to the right, and at the right side of the boundary, the angular flux moving to the left (these values are 0).
- (c) (40 points) Plot the error after each iteration using a 0 initial guess for the step discretization with source iteration and GMRES.

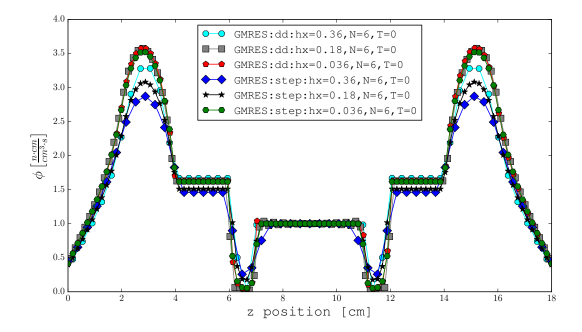
Error will be determined with the following:

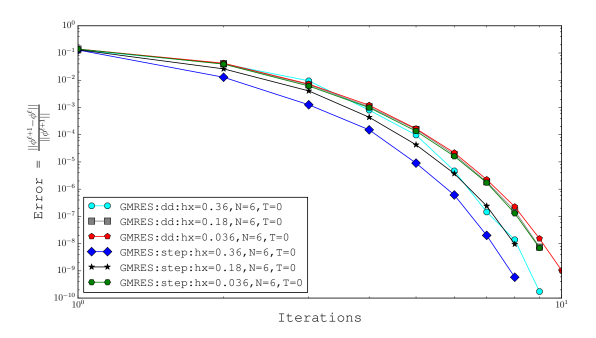
Error =
$$\frac{||\phi^{\ell+1} - \phi^{\ell}||}{||\phi^{\ell+1}||}$$

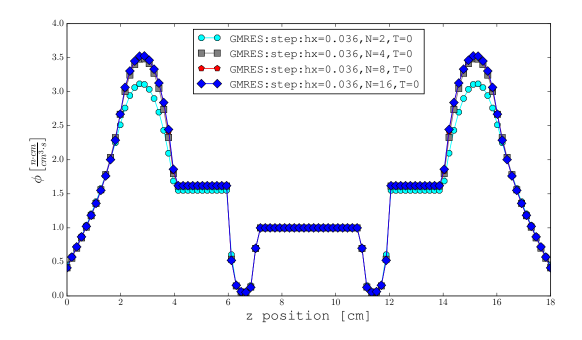


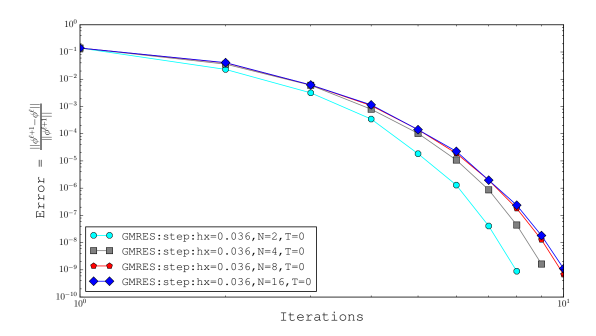
(d) (50 points) Solve Reed's problem (see finite difference diffusion codes). Present convergence plots for the solution in space and angle to a "refined" solution in space and angle.

Plots are below, reduced the number of points so that figures wouldn't take so long to load.









The solution converges with more spatial slices. Increasing the number of angular slices helps upto when N=4, but beyond that it doesn't do much.

(e) (50 points extra credit) Solve a time dependant problem for a slab surrounded by vacuum with $\Sigma_t = \Sigma_s = 1$ and initial condition given by $\psi(\mathbf{0}) = \mathbf{1}/h_x$ (original problem statement said $\phi(0) = 1/h_x$ and I'm not sure how to solve that). Plot the solution at t = 1 s, using step and diamond difference. The particles have a speed of 1 cm/s. Which discretization is better with a small time step? What do you see with a small number of ordinates compared to a really large number (100s)?

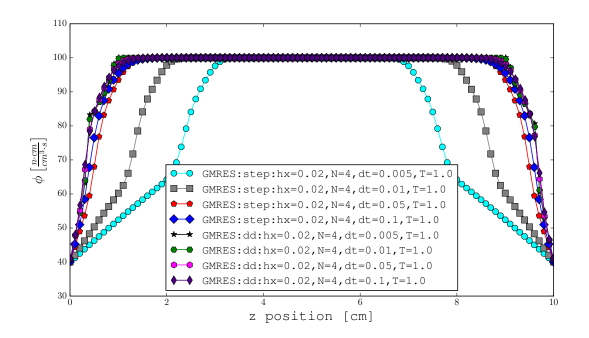


Figure 1: Q=0.01

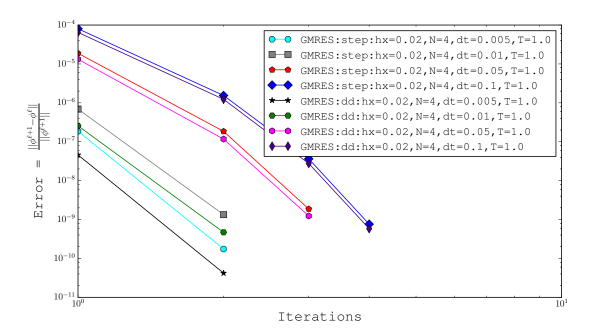
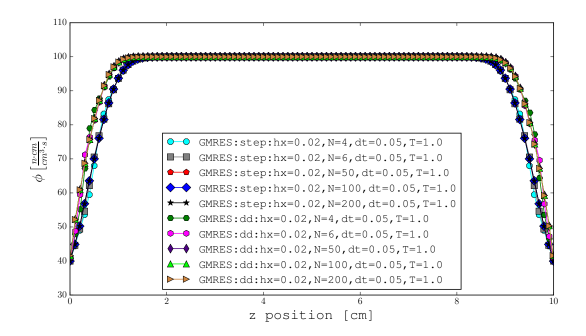
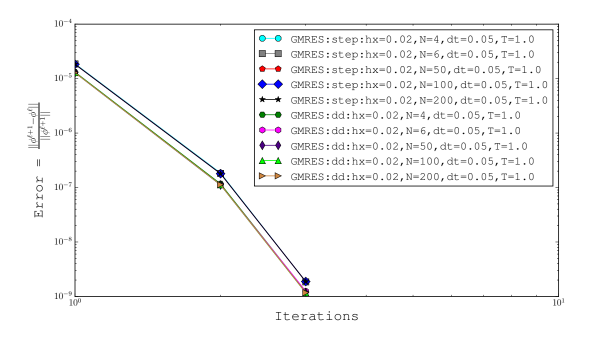


Figure 2: Q=0.01

Based on the above graphs, I am not sure which solution does better with a smaller step size. It depends on what the answer should be. I think the step solution, as the step size increases, look like

they have some nonphysical bends in the solution. This could be due to lots of things, but maybe its because of the smaller step size, which makes me think the diamond difference method is better with smaller step sizes.





As the number of ordinances increase in the problem there isn't much change in the solution, which is expected because with the Quadrature rule we used, the integral can usually be expressed within

around 6 terms. There was an increase in computational time though.

Homework 4 Code

Listing 1: Main Code For Parts a,b and c

```
#!/usr/bin/env python3
 import time
 start_time = time.time()
 import Functions as f
 # Constants
 0 = 0.01
 Sigma_t = 100;Sigma_s=100
 # Add adsorption to help converge
 if Sigma_t==Sigma_s:
   Sigma_t=Sigma_t * 1.0001
 # Geometry
                  # Width of slab
 L = 10.
                 # Number of cuts in slab (looped)
 slices=[10,50,100,500]
_{25} N = 4
                  # Number of angle slices
 BCs = f.np.zeros(N)
                  # Zero incoming flux
 #Time
 T=0
                  # total Time (A plot made at T)
                  # Time steps width
 dt=1
                  # Velocity
 v=1
 MAXITS=100000
                 # Max iterations for source iter
 loud=False
                  # Echo every Iteration?
 #Method
 Methods=['GMRES:step', # 'Iteration' or 'GM.
'GMRES:dd'] # Methods to solve with?
                  # 'Iteration' or 'GMRES'
                 # 'step' or 'dd'
 tol=1e-8
 PlotError=False # Do we plot the error?
 NumOfPoints=100
                 # Max Number of points for plots
 50
```

```
Check=0
 fig=f.plt.figure(figsize=f.FigureSize) # Plot all Methods
 ax=fig.add_subplot(111)
 if PlotError:
    erfig=f.plt.figure(figsize=f.FigureSize) # Err Plot
    erax=erfig.add_subplot(111)
  for Scheme in Methods:
    Method=Scheme.split(':')[1]
65
    for II in slices:
      if Method == 'step': #Step Dude needs one extra
       elif Method == 'dd':
         I = II
      #Width, ang lists for materials
      hx = L/II
      q = f.np.ones(I) *Q
      Sig_t_discr = f.np.ones(I) *Sigma_t
80
      Sig_s_discr = f.np.ones(I) *Sigma_s
      #Initialize psi (for time steps)
      if T==0:
         psi=f.np.zeros((N,I))
         Time=[0]
      else:
         psi=f.np.ones((N,I))*(1/hx)
         Time=f.Timevector(T, dt)
      label_tmp=Scheme+":hx="+str(hx)+",N="+str(N)+",T="
      for t in Time: #Loop over time
         label=label_tmp+str(t)
         #Determine phi (new psi is determined for time steps)
         x,phi,it,er,psi=f.solver(I,hx,q,Sig_t_discr,
         Sig_s_discr, N, psi, v, dt, Time, BCs, Scheme, tol, MAXITS, loud)
```

```
ax, fig=f.plot(x,phi,ax,label,fig,Check,NumOfPoints)
         if t==0 and PlotError:
            erfig
            erax,erfig=f.plotE(it,er,erax,label,erfig,
110
                       Check, NumOfPoints)
         Check=Check+1
  fiq
  f.Legend(ax)
  #f.plt.savefig('Plots/FluxPlot.pdf')
  if PlotError:
    erfia
    f.Legend(erax)
    #f.plt.savefig('Plots/ErrorPlot.pdf')
    f.plt.savefig('Plots/ErrorPlotTime.pdf')
125
    #f.plt.clf()
    f.plt.close()
  fiq
  f.plt.savefig('Plots/FluxPlotTime.pdf')
  #f.plt.show()
  #Why is tmp_psi in the GMRES going negative?
  print("--- %s seconds ---" % (time.time() - start_time))
```

Listing 2: Main Code For Part d

```
#!/usr/bin/env python3
 ##################### Import packages ###########################
 import time
 start_time = time.time()
 import Functions as f
 15 # Geometry
 L = 18.
              # Width of slab
 slices=[500] # Number of cuts in slab (looped)
            # Number of angle slices
 NN = [2, 4, 8, 16]
```

```
#Time
 T=0
                   # total Time (A plot made at T)
                    # Time steps width
 dt=1
 v=1
                   # Velocity
                 # Max iterations for source iter
 MAXITS=1000000
 loud=False
                   # Echo every Iteration?
 #Method
 Methods=['GMRES:step']#, # 'Iteration' or 'GMRI
# 'GMRES:step'] # Methods to solve with?
                    # 'Iteration' or 'GMRES'
                   # 'step' or 'dd'
 tol=1e-8
 PlotError=True # Do we plot the error?
 NumOfPoints=100
                  # Max Number of points for plots
  Check=0
 fig=f.plt.figure(figsize=f.FigureSize) # Plot all Methods
 ax=fig.add_subplot(111)
  if PlotError:
   erfig=f.plt.figure(figsize=f.FigureSize) # Err Plot
    erax=erfig.add_subplot(111)
                            # at T=0
  for Scheme in Methods:
    Method=Scheme.split(':')[1]
    60
    for II in slices:
      if Method == 'step': #Step Dude needs one extra
         I = II + 1
      elif Method == 'dd':
        T = T T
      #Width, ang lists for materials
      hx = L/II
      q = f.np.zeros(I)
70
      Sig_t_discr = f.np.zeros(I)
```

```
Sig_s_discr = f.np.zeros(I)
         if Method == 'step':
           x = f.np.linspace(0, (I-1)*hx, I)
         elif Method == 'dd':
           x = f.np.linspace(hx/2,I*hx-hx/2,I)
         for i in range (0, len(x)):
           q[i]=f.QReed(x[i])
           Sig_t_discr[i]=f.Sigma_tReed(x[i])
           Sig_s_discr[i]=Sig_t_discr[i]-f.Sigma_aReed(x[i])
        for N in NN:
           BCs = f.np.zeros(N)
                                  # Zero incoming flux
           #Initialize psi (for time steps)
           if T==0:
              psi=f.np.zeros((N,I))
              Time=[0]
           else:
              psi=f.np.ones((N,I))*(1/hx)
              Time=f.Timevector(T,dt)
           label tmp=Scheme+":hx="+str(hx)+", N="+str(N)+", T="
95
           for t in Time: #Loop over time
              label=label_tmp+str(t)
               #Determine phi (new psi is determined for time steps)
              x, phi, it, er, psi=f.solver(I, hx, q, Sig_t_discr,
              Sig_s_discr, N, psi, v, dt, Time, BCs, Scheme, tol, MAXITS, loud)
               ############# Plot Information ###################
               110
              ax, fig=f.plot(x,phi,ax,label,fig,Check,NumOfPoints)
               if t==0 and PlotError:
                 erfig
                 erax, erfig=f.plotE(it, er, erax, label, erfig,
115
                                Check, NumOfPoints)
              Check=Check+1
   ############## Legend/Save
                           ##########################
   fiq
  f.Legend(ax)
```

Listing 3: Main Code For Part e

```
#!/usr/bin/env python3
 import time
 start_time = time.time()
 import Functions as f
 # Geometry
 L = 10
                 # Width of slab
 # Constants
 Q = 0.01
 Sigma_t = 1;Sigma_s=1
 # Add adsorption to help converge
 if Sigma_t==Sigma_s:
   Sigma_t=Sigma_t *1.0001
 slices=[500]  # Number of cuts in slab (looped)
NN = [4, 6, 50, 100, 200]
                     # Number of angle slices
 #Time
 T=1
                  # total Time (A plot made at T)
                   # Time steps width
 dtt=[0.05]
                   # Velocity
 v=1
 MAXITS=1000000
                  # Max iterations for source iter
 loud=False
                  # Echo every Iteration?
 #Method
 Methods=['GMRES:step', # 'Iteration' or 'GMRES'
```

```
'GMRES:dd']
                 # Methods to solve with?
                  # 'step' or 'dd'
 |tol=1e-8
 Ttol=1e-3
 PlotError=True
                 # Do we plot the error?
 NumOfPoints=100
                  # Max Number of points for plots
 Check=0
 fig=f.plt.figure(figsize=f.FigureSize) # Plot all Methods
 ax=fig.add_subplot(111)
 if PlotError:
   erfig=f.plt.figure(figsize=f.FigureSize) # Err Plot
    erax=erfig.add_subplot(111)
 for Scheme in Methods:
   Method=Scheme.split(':')[1]
    for II in slices:
      if Method == 'step': #Step Dude needs one extra
        I = II + 1
      elif Method == 'dd':
        I = II
      #Width, ang lists for materials
      hx = L/II
      q = f.np.ones(I) *Q
      Sig_t_discr = f.np.ones(I) *Sigma_t
      Sig_s_discr = f.np.ones(I) *Sigma_s
80
      for N in NN:
        BCs = f.np.zeros(N)
                         # Zero incoming flux
        for dt in dtt:
85
           #Initialize psi (for time steps)
           if T==0:
             psi=f.np.zeros((N,I))
             Time=[0]
```

```
else:
90
                 psi=f.np.ones((N,I))*(1/hx)
                 Time=f.Timevector(T,dt)
              label_tmp=Scheme+":hx="+str(hx)+",N="+str(N)+",dt="+
                      str(dt)+", T="
95
              ################## Determine phi #######################
              for t in Time: #Loop over time
                 label=label_tmp+str(round(t,3))
                 #Determine phi (new psi is determined for time steps)
                 x,phi,it,er,psi=f.solver(I,hx,q,Sig_t_discr,
105
                 Sig_s_discr, N, psi, v, dt, Time, BCs, Scheme, tol, MAXITS, loud)
                 ############## Plot Information ####################
                 110
                 PlotQuestion=abs(t-dt)<Ttol\
                 or abs(t-0.5)<Ttol or abs(t-1)<Ttol
                 PlotOuestion=abs(t-1)<Ttol
                 if PlotQuestion:
115
                    ax, fig=f.plot(x,phi,ax,label,fig,Check,NumOfPoints)
                    if PlotError:
                      erfig
                      erax, erfig=f.plotE(it, er, erax, label, erfig,
                                    Check, NumOfPoints)
120
                    Check=Check+1
   fig
  f.Legend(ax)
  #f.plt.savefig('Plots/FluxPlot.pdf')
  if PlotError:
     erfiq
     f.Legend(erax)
     #f.plt.savefig('Plots/ErrorPlot.pdf')
     f.plt.savefig('Plots/ErrorPlotTimeVaryN.pdf')
     #f.plt.clf()
135
     f.plt.close()
  fig
  f.plt.savefig('Plots/FluxPlotTimeVaryN.pdf')
  #f.plt.show()
140
```

```
|\operatorname{print}("---\ s seconds ---" % (time.time() - start_time))
```

Listing 4: Functions holder

```
#!/usr/bin/env python3
  ################### Import packages ##############################
  import sys
  import numpy as np
  import scipy.sparse.linalg as spla
  import scipy.special as sps
  import matplotlib.pyplot as plt
  plt.rcParams["font.family"] = "monospace"
  import matplotlib
matplotlib.rc('text', usetex=True)
  matplotlib.rcParams['text.latex.preamble']=[r"\usepackage{amsmath}"]
  import random as rn
  import matplotlib.mlab as mlab
  import copy
  import os
  # Basic information
  FigureSize = (11, 6) # Dimensions of the figure
TypeOfFamily='monospace' # This sets the type of font for text
  font = {'family' : TypeOfFamily} # This sets the type of font for text
30 LegendFontSize = 12
  Lfont = {'family' : TypeOfFamily} # This sets up legend font
  Lfont['size'] = LegendFontSize
  Title = ''
  TitleFontSize = 22
  TitleFontWeight = "bold" # "bold" or "normal"
  #Xlabel='E (eV)' # X label
  XFontSize=18 # X label font size
40 | XFontWeight="normal" # "bold" or "normal"
  XScale="linear"
                  # 'linear' or 'log'
  XScaleE='log' # Same but for error plot
  YFontSize=18
                           # Y label font size
 YFontWeight="normal"
                         # "bold" or "normal"
  YScale="linear"
                          # 'linear' or 'log'
  YScaleE='log'
  Check=0
```

```
Colors=["aqua", "gray", "red", "blue", "black",
               "green", "magenta", "indigo", "lime", "peru", "steelblue",
               "darkorange", "salmon", "yellow", "lime", "black"]
  # If you want to highlight a specific item
  # set its alpha value =1 and all others to 0.4
  # You can also change the MarkSize (or just use the highlight option below)
  # Can change all these to "." or "" for nothing "x" isn't that good
 MarkerType=["8", "s", "p", "D", "*", "H", "h", "d", "^", ">"]
  # LineStyles=["solid", "dashed", "dash_dot", "dotted", "."]
  LineStyles=["solid"]
 SquishGraph = 0.75
  BBOXX = 1.24
  BBOXY = 0.5
                # Set legend on right side of graph
  NumberOfLegendColumns=1
  Xlabel='z position [cm]'
  Ylabel="$\phi\left[\\frac{n\cdot cm}{cm^3\cdot s}\\right]$"
  XlabelE='Iterations'
  |YlabelE="Error = $\\frac{||\phi^{\ell+1}-\phi^\ell||}{||\phi^{\ell+1}||}$"
  85
  def Sigma_tReed(r):
     value = 0 + ((1.0*(r>=14) + 1.0*(r<=4)) +
                5.0 * ((np.abs(r-11.5)<0.5)  or (np.abs(r-6.5)<0.5)) +
                50.0 * (np.abs(r-9) <= 2))
     return value:
  def Sigma_aReed(r):
     value = 0 + (0.1*(r>=14) + 0.1*(r<=4) +
               5.0 * ((np.abs(r-11.5)<0.5)  or (np.abs(r-6.5)<0.5)) +
                50.0 * (np.abs(r-9) <= 2))
     return value;
  def QReed(r):
     value = 0 + 1.0 \times ((r<16) \times (r>14)) + 1.0 \times ((r>2) \times (r<4)) + 50.0 \times (np.abs(r-9) <=2)
     return value;
  def Timevector(T, dt):
     Time=[dt]
     while Time[-1]<T:
        Time.append(Time[-1]+dt)
```

```
return (Time)
105
   def diamond_sweep1D(I, hx, q, sigma_t, mu, boundary):
      """Compute a transport diamond difference sweep for a given
      Inputs:
       I:
                         number of zones
       hx:
                         size of each zone
110
                        source array
       q:
                      array of total cross-sections
       sigma_t:
       mu:
                         direction to sweep
       boundary:
                       value of angular flux on the boundary
      Outputs:
115
                         value of angular flux in each zone
       psi:
     assert(np.abs(mu) > 1e-10)
     psi = np.zeros(I)
     ihx = 1./hx
120
      if (mu > 0):
       psi_left = boundary
        for i in range(I):
          psi_right = (q[i] + (mu*ihx-0.5*sigma_t[i])*psi_left)
                      /(0.5*sigma_t[i] + mu*ihx)
125
          psi[i] = 0.5*(psi_right + psi_left)
          psi_left = psi_right
      else:
       psi_right = boundary
        for i in reversed(range(I)):
130
          psi_left = (q[i] + (-mu*ihx-0.5*sigma_t[i])*psi_right)
                     /(0.5*sigma_t[i] - mu*ihx)
          psi[i] = 0.5*(psi_right + psi_left)
          psi_right = psi_left
      return psi
135
   def step_sweep1D(I, hx, q, sigma_t, mu, boundary):
      """Compute a transport step sweep for a given
      Inputs:
       I:
                        number of zones
140
                         size of each zone
       hx:
                       source array
       sigma_t:
                       array of total cross-sections
       mu:
                         direction to sweep
                        value of angular flux on the boundary
       boundary:
145
      Outputs:
       psi:
                        value of angular flux in each zone
     assert (np.abs(mu) > 1e-10)
     psi = np.zeros(I)
150
      ihx = 1./hx
      if (mu > 0):
       psi_left = boundary
       psi[0] = 0
        for i in range (1, I):
155
          psi\_right = (q[i] + mu*ihx*psi\_left) / (mu*ihx + sigma\_t[i])
```

```
psi[i] = 0.5*(psi_right + psi_left)
          psi_left = psi_right
      else:
160
       psi_right = boundary
       psi[-1] = 0
        for i in reversed(range(0, I-1)):
          psi_left = (q[i] - mu*ihx*psi_right)/(sigma_t[i] - mu*ihx)
          psi[i] = 0.5*(psi\_right + psi\_left)
          psi_right = psi_left
165
      return psi
    def source_iteration(I, hx, q, sigma_t, sigma_s, N, psiprevioustime,
                          v, dt, Time, BCs, sweep_type,
170
                          tolerance = 1.0e-8, maxits = 100, LOUD=False ):
      """Perform source iteration for single-group steady state problem
      Inputs:
       I:
                         number of zones
                        size of each zone
       hx:
175
        q:
                         source array
                       array of total cross-sections array of scattering cross-sections
        sigma_t:
        sigma_s:
       N:
                         number of angles
                         Boundary conditions for each angle
       BCs:
180
                       type of 1D sweep to perform solution
        sweep_type:
       tolerance:
                       the relative convergence tolerance for the iterations
                         the maximum number of iterations
        maxits:
                         boolean to print out iteration stats
        LOUD:
      Outputs:
185
                         value of center of each zone
       X:
       phi:
                         value of scalar flux in each zone
     iterations = []
190
      Errors = []
     phi = np.zeros(I)
     phi_old = phi.copy()
     converged = False
     MU, W = np.polynomial.legendre.leggauss(N)
     iteration = 1
195
      tmp_psi=psiprevioustime.copy()
      if len(Time) == 1:
          sigma_ts=sigma_t
      else:
          sigma_ts=sigma_t+1/(v*dt)
200
      while not (converged):
       phi = np.zeros(I)
        #sweep over each direction
        for n in range(N):
205
          \#qs = (q*W[n])/2 + (phi_old*sigma_s)/2 + psiprevioustime[n,:]/(v*dt)
          qs=(q)/2+(phi_old*sigma_s)/2+psiprevioustime[n,:]/(v*dt)
          if sweep_type == 'dd':
            tmp_psi[n,:] = diamond_sweep1D(I,hx,qs,sigma_ts,MU[n],BCs[n])
```

```
elif sweep_type == 'step':
210
            tmp_psi[n,:] = step_sweep1D(I,hx,qs,sigma_ts,MU[n],BCs[n])
          else:
            sys.exit("Sweep method specified not defined in SnMethods")
          phi = phi+tmp_psi[n,:]*W[n]
        #check convergence
215
        change = np.linalg.norm(phi-phi_old)/np.linalg.norm(phi)
        iterations.append(iteration)
        Errors.append(change)
        #iterations.append(iteration)
        #Errors.append(change)
        converged = (change < tolerance) or (iteration > maxits)
        if (LOUD>0) or (converged and LOUD<0):</pre>
          print("Iteration", iteration, ": Relative Change =", change)
        if (iteration > maxits):
          print("Warning: Source Iteration did not converge: "+\
225
                sweep_type+", I : "+str(I)+", Diff : %.2e" % change)
        #Prepare for next iteration
        iteration += 1
        phi_old = phi.copy()
      if sweep_type == 'step':
230
          x = np.linspace(0, (I-1)*hx, I)
      elif sweep_type == 'dd':
          x = np.linspace(hx/2, I*hx-hx/2, I)
      return x, phi, iterations, Errors, tmp_psi
235
    def gmres_solve(I, hx, q, sigma_t, sigma_s, N, psiprevioustime,
                    v, dt, Time, BCs, sweep_type,
                    tolerance = 1.0e-8, maxits = 100, LOUD=False,
                    restart = 20):
240
      """Solve, via GMRES, a single-group steady state problem
      Inputs:
        I:
                          number of zones
        hx:
                         size of each zone
245
        q:
                         source array
                       array of total cross-sections
        sigma_t:
                         array of scattering cross-sections
        sigma_s:
        N:
                        number of angles
                        Boundary conditions for each angle
        BCs:
        sweep_type: type of 1D sweep to perform solution
tolerance: the relative convergence tolerance for the iterations
        maxits:
                        the maximum number of iterations
        LOUD:
                          boolean to print out iteration stats
      Outputs:
       X:
                         value of center of each zone
255
                        value of scalar flux in each zone
       phi:
      iterations = []
      Errors = []
260
      #compute RHS side
      RHS = np.zeros(I)
```

```
MU, W = np.polynomial.legendre.leggauss(N)
      tmp_psi=psiprevioustime.copy()
265
      if len(Time) == 1:
          sigma_ts=sigma_t
      else:
          sigma_ts=sigma_t+1/(v*dt)
270
      for n in range(N):
        qs=q/2+psiprevioustime[n,:]/(v*dt)
        if sweep_type == 'dd':
          tmp_psi[n,:] = diamond_sweep1D(I,hx,qs,sigma_ts,MU[n],BCs[n])
        elif sweep_type == 'step':
          tmp_psi[n,:] = step_sweep1D(I,hx,qs,sigma_ts,MU[n],BCs[n])
        \#tmp\_psi = sweep1D(I,hx,q,sigma\_t,MU[n],BCs[n])
        RHS += tmp_psi[n,:]*W[n]
      #define linear operator for gmres
      def linop(phi):
        tmp = phi * 0
        #sweep over each direction
        for n in range(N):
          if sweep_type == 'dd':
            tmp_psi[n,:] = diamond_sweep1D(I,hx,(phi*sigma_s)/2,
                                       sigma_ts,MU[n],BCs[n])
          elif sweep_type == 'step':
            tmp_psi[n,:] = step_sweep1D(I,hx,(phi*sigma_s)/2,
                                         sigma_ts, MU[n], BCs[n])
290
          tmp += tmp_psi[n,:]*W[n]
        return phi-tmp
      A = spla.LinearOperator((I,I), matvec = linop, dtype='d')
295
      #define a little function to call when the iteration is called
      iteration = np.zeros(1)
      def callback(rk, iteration=iteration):
        iteration += 1
300
        if (LOUD>0):
          print("Iteration", iteration[0], "norm of residual", np.linalg.norm(rk))
        iterations.append(iteration[0])
        Errors.append(np.linalg.norm(rk))
      #Do the GMRES Solve
      phi, info = spla.gmres(A, RHS, x0=RHS, tol=tolerance,
                             restart=int(restart), callback=callback)
      #Print important information
      if (LOUD):
310
        print("Finished in", iteration[0], "iterations.")
      if (info >0):
        print("Warning, convergence not achieved :"+str(sweep_type)+" "+str(hx))
      if sweep_type == 'step':
          x = np.linspace(0,(I-1)*hx,I)
315
```

```
elif sweep_type == 'dd':
         x = np.linspace(hx/2, I*hx-hx/2, I)
     #Calculate Psi for time iterations
     phi2 = np.zeros(I)
320
     #sweep over each direction
     for n in range(N):
         \#qs = (q*W[n])/2 + (phi_old*sigma_s)/2 + psiprevioustime[n,:]/(v*dt)
         qs=(q)/2+(phi*sigma_s)/2+psiprevioustime[n,:]/(v*dt)
         if sweep_type == 'dd':
325
             tmp_psi[n,:] = diamond_sweep1D(I,hx,qs,sigma_ts,MU[n],BCs[n])
         elif sweep_type == 'step':
             tmp_psi[n,:] = step_sweep1D(I,hx,qs,siqma_ts,MU[n],BCs[n])
         else:
             sys.exit("Sweep method specified not defined in SnMethods")
330
         phi2 = phi2+tmp_psi[n,:]*W[n]
     return x, phi, iterations, Errors, tmp_psi
   def solver(I, hx,q, Sig_t, Sig_s, N, psi, v, dt, Time, BCs, Scheme, tol, MAXITS, loud):
       Method=Scheme.split(':')[1]
       if "Iteration" in Scheme:
           x, phi, iterations, errors, psi =source_iteration(I,
               hx, q, Sig_t, Sig_s, N, psi, v, dt, Time, BCs,
340
               Method, tolerance=tol, maxits=MAXITS, LOUD=loud)
       elif "GMRES" in Scheme:
           x, phi, iterations, errors, psi =gmres_solve(I,
               hx,q,Sig_t,Sig_s,N,psi,v,dt,Time,BCs,
               Method, tolerance=tol, maxits=MAXITS, LOUD=loud, restart=MAXITS)
345
       else:
           print("Improper sweep selected")
           quit()
       return x, phi, iterations, errors, psi
    #################### Plotting Function #########################
    def reduceList(List, N):
       List2=[List[0]]
355
       Div=int(len(List)/N)
       for i in range(1,len(List)-1):
           if i % Div == 0:
               List2.append(List[i])
       List2.append(List[-1])
       return (List2)
    def loop_values(list1,index):
       This function will loop through values in list even if
       outside range (in the positive sense not negative)
       while True:
```

```
try:
                 list1[index]
                 break
            except IndexError:
                 index=index-len(list1)
        return(list1[index])
375
    def plot(x, y, ax, label, fig, Check, NumOfPoints):
        if len(x) > 300:
            x=reduceList(x, NumOfPoints)
            y=reduceList(y,NumOfPoints)
        \#Plot\ X\ and\ Y
380
        ax.plot(x,y,
                 linestyle=loop_values(LineStyles, Check),
                 marker=loop_values(MarkerType, Check),
                 color=loop_values(Colors, Check),
                 markersize=loop_values(MarkSize,Check),
385
                 alpha=loop_values(Alpha_Value, Check),
                 label=label)
        #Log or linear scale?
        ax.set_xscale(XScale)
390
        ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                      fontweight=TitleFontWeight, fontdict=font,
                                                                  ha='center')
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        ax.set_ylabel(Ylabel,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
        return (ax, fig)
405
    def plotE(x, y, erax, label, erfig, Check, NumOfPoints):
        if len(x) > 300:
            x=reduceList(x, NumOfPoints)
            y=reduceList(y,NumOfPoints)
410
        \#Plot\ X\ and\ Y
        erax.plot(x,y,
                 linestyle=loop_values(LineStyles, Check),
                 marker=loop_values(MarkerType, Check),
                 color=loop_values(Colors, Check),
415
                 markersize=loop_values(MarkSize,Check),
                 alpha=loop_values(Alpha_Value,Check),
                 label=label)
        #Log or linear scale?
420
        erax.set_xscale(XScaleE)
```

```
erax.set_yscale(YScaleE)
        #Set Title
        erfig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font,
425
                                                                 ha='center')
        #Set X and y labels
        erax.set_xlabel(XlabelE,
                      fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
430
        erax.set_ylabel(YlabelE,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
        return (erax, erfig)
435
    def Legend(ax):
        handles, labels=ax.get_legend_handles_labels()
        ax.legend(handles, labels, loc='best',
                  fontsize=LegendFontSize,prop=font)
440
        return (ax)
    # def Legend(ax):
              handles, labels=ax.get_legend_handles_labels()
              box=ax.get_position()
445
              ax.set_position([box.x0, box.y0, box.width*SquishGraph,
                                box.height])
              ax.legend(handles, labels, loc='center',
                        bbox_to_anchor=(BBOXX,BBOXY),
                         fontsize=LegendFontSize, prop=font,
450
                         ncol=NumberOfLegendColumns)
    #
              return(ax)
```

Homework 5 Problem Statement

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

Clean Fusion Energy

(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 BeF₂) to convert the neutron energy into heat. All the constituents in the FLiBe have their natural abundances. Using data from JANIS, and assuming the total neutron flux is 10^{14} n/cm²·s. Perform the following analyses.

- (a) (25 points) Write out the depletion (or in thise case activation) chains that will occur in the system.
- (b) (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?
- (c) (25 points) After discharging the FLiBe blanket, how long will it take until the material is less radioactive than Brazil Nuts? (444 Bq/kg)

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Homework 5 Background

Please note, that most of this background is copied directly from Dr. McClarren's notes, but are reproduced here.

The production of an isotope is dictated by production and loss

$$\frac{dn_i}{dt} = -\lambda_i^{eff} n_i + \sum_{j=1}^{N} b_{j \to i}^{eff} \lambda_j^{eff} n_j$$

Where,

$$\lambda_i^{eff} = \lambda_i + \phi \sum_{j=1}^N \sigma_{i \to j}$$

and

$$b_{j \to i}^{eff} = \frac{b_{j \to i} \lambda_j + \sigma_{j \to i} \phi}{\lambda_i^{eff}}$$

For a system of isotopes, this can be reduced to:

$$\frac{d\vec{n}}{dt} = \mathbf{A}\vec{n}(t)$$

Where \boldsymbol{A} is a matrix whose diagonal elements are $[-\lambda_1^{eff}, -\lambda_2^{eff}, ..., -\lambda_N^{eff}]$, all off diagonal elements are $b_{j \to i}^{eff} \lambda_j^{eff}$ (i for the diagonal, and j is for the off diagonal position) and $\vec{n}(t) = [n_1, n_2, ..., n_N]$.

The solution to this system is obvious (it wasn't to me at first - but that's because I'm a newb)

$$\vec{n} = e^{\mathbf{A}t}\vec{n}_0$$

Determing $e^{\mathbf{A}t}\vec{n}_0$ will be done 3 different ways,

Matrix Exponential

Analytic Solution, unstable with large N.

$$\vec{n}(t) = e^{\mathbf{A}t} \vec{n}_0 \approx \left[\sum_{m=0}^{\infty} \frac{1}{m!} \mathbf{A}^m t^m \right] \vec{n}_0$$

Backward Euler

Unstable for large Δt , but can take time steps.

$$\frac{d\vec{n}}{dt} \approx \frac{\vec{n}(\Delta t) - \vec{n}_0}{\Delta t} \approx \mathbf{A}\vec{n}(\Delta t)$$
$$\vec{n}(\Delta t) \approx (\mathbf{I} - \mathbf{A}\Delta t)^{-1}\vec{n}_0$$

Rational Approximation

$$\vec{n}(t) = e^{\mathbf{A}t} \vec{n}_0 \approx -2\Re \sum_{k=1}^{N/2} c_k (z_k \mathbf{I} - \mathbf{A}t)^{-1} \vec{n}_0$$

The \Re symbol means taking the real part of the solution. Further,

$$c_k = \frac{i}{N} e^{z_k} w_k$$

where z_k and w_k are both scalers defined as

$$z_k = \phi(\theta_k)$$
$$w_k = \phi'(\theta_k)$$

with

$$\begin{split} \phi(\theta) = & N[0.1309 - 0.1194\theta^2 + 0.2500i\theta] \\ \text{or} \\ \phi(\theta) = & 2.246N \left[1 - sin(1.1721 - 0.3443i\theta] \right] \\ \text{or} \\ \phi(\theta) = & N[0.5071\theta cot(0.6407\theta) - 0.6122 + 0.2645i\theta] \\ \text{or} \\ \phi(\theta) = & \text{Best Possible} \end{split}$$

and

$$\theta_k = \pm \frac{\pi}{N} (1 + 2k)$$
 k from 0 to N-1

Where N doesn't have to go much higher than 10 to have low errors (for the best Rational Approximation). Also both plus and minus terms were written here, but the first equation in this solution method only uses the positive terms. This is because using the negative β 's yields the same real part as the positive β 's, with opposite complex parts (the complex cancels). Thats why the \sum only goes to N/2 and the solution is multiplied by 2.

The solution utilized the assumption that half the neutron flux was 14 MeV and the other half was 2.45 MeV. The A matrix was built in terms of days and atoms per kg of initial fuel.

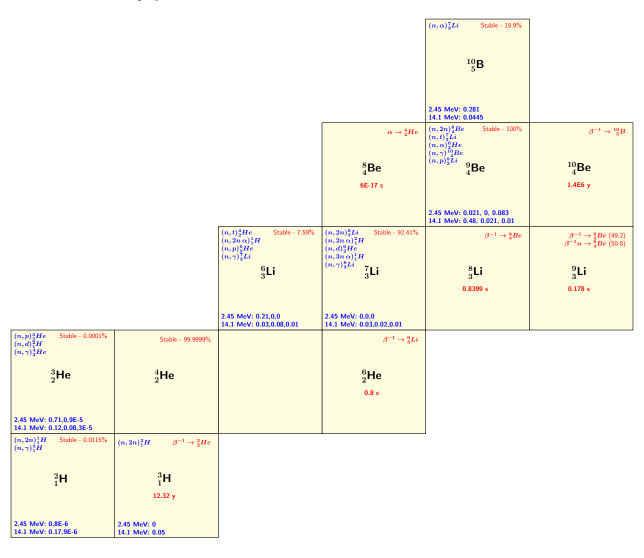
1 Kg FLiBe
$$\cdot \frac{6.022E23}{98.9 \text{ g}} = 6.09E24 \text{ atoms of FLiBe}$$

This number was the starting condition for ⁹Be. Twice this number (with natural abundancy considerations) for ⁶Li and ⁷Li, and 4 times this number for ¹⁹F.

Homework 5 Solution

(a) (25 points) Write out the depletion (or in thise case activation) chains that will occur in the system.

The chains were written out in a form so that the diagram would be uncluttered, and not all possible reactions are displayed.



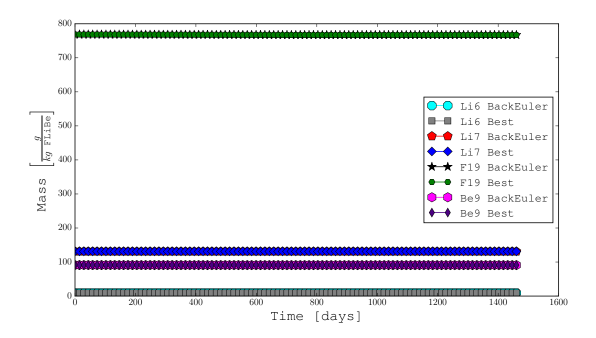
		Stable - 90.48%	
		$^{20}_{10}Ne$	
	$EC ightarrow {}^{18}_{8}O$ ${}^{18}_{9}$ F 110 min	$\begin{array}{c} (n,n\alpha)^{15}N & \text{Stable - } 100\% \\ (n,np)^{18}O \\ (n,2n)^{18}F \\ (n,\alpha)^{16}F \\ (n,\alpha)^{18}O \\ (n,p)^{18}O \\ (n,t)^{18}O \\ (n,\gamma)^{29}F \end{array}$	$eta^{-1} ightarrow rac{20}{10} Ne$ $rac{20}{9} extsf{F}$ 11.1 s
$(n,\alpha)^{13}_{6}C$ Stable - 99.757%	$(n,\alpha)^{14}_{6}C$ Stable - 0.038%	14.1 MeV: 0.4,0.06,0.04 Stable - 0.21%	$eta^{-1} ightarrow ^{19}_{9}F$
$(n,p)_{10}^{10}N$ $(n,d)_{15}^{15}N$ $(n,\gamma)_{18}^{17}O$	$\begin{array}{c} (n,\alpha)^{14}_{\ 6}C \\ (n,2n)^{16}_{\ 6}O \\ (n,n\alpha)^{13}_{\ 6}O \\ (n,n\alpha)^{13}_{\ 7}O \\ (n,p)^{17}_{\ 7}N \\ (n,d)^{16}_{\ 7}N \\ (n,\gamma)^{18}_{\ 8}O \end{array} \text{Stable - 0.038\%}$	¹⁸ ₈ O	¹⁹ 80 26.9 s
2.45 MeV: 0,0,0 14.1 MeV: 0.14,0.04,0.02	2.45 MeV: 0.12,0,0 14.1 MeV: 0.3,0.1,0.04		
$(n,2n)_{5}^{14}N$ Stable - 0.364% $(n,\alpha)_{5}^{12}B$ $(n,np)_{6}^{14}C$ $(n,t)_{6}^{13}C$ $(n,p)_{6}^{15}C$ $(n,p)_{6}^{15}C$ $(n,d)_{6}^{14}C$ 15	$EC ightarrow {}^{16}_{8}O$ $^{16}_{7}$ N $^{7.13}$ s		
2.45 MeV: 0,0,0 14.1 MeV: 0.11,0.07,0.04			

(b) (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?

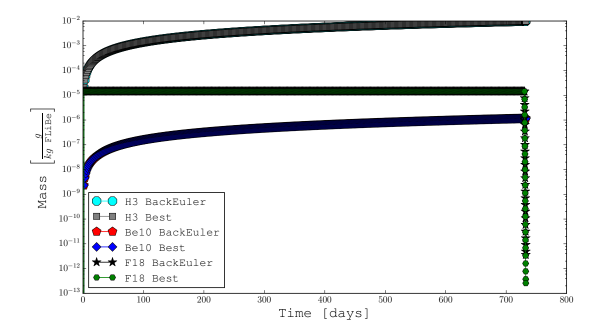
The matrix exponential method did not converge. The Backwards Euler and the Rational approximation solutions did converge. Their answers are basically the same (all plots will have both solutions, but their answers are on top of one another). It should be noted that I changed the solver from in the notes from a GMRES solver to a normal matrix inverter for the rational approach to speed up the algorithm and give better results.

Plots were split up into two groups, with initial nuclides, and product nuclides which would have appreciable activity after a couple of days. Below plots are shown in grams.

The rational approximation used the "Best" method with 10 quadrature points,



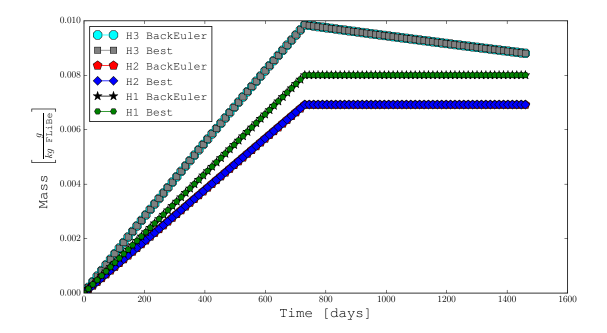
This figure shows that there wasn't sizeable amount of depletion in the coolant. Which is expected because the cross sections are less than a barn.



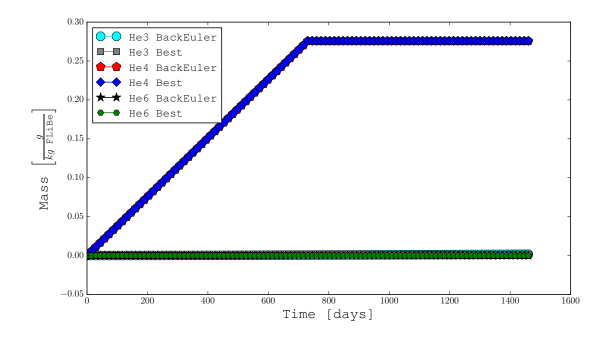
The above figure shows that the high activity elements with large half lives (all other radioactive elements have half lives less than a minute and would decay away and less than 7 minutes) are not produces in a sizeable quantity except for ³H, which has a 12 year half life. The sharp decrease in ¹⁸F

is because the plot has about a day of decay included to show that activity is primarily from ³H (Beta emitter) and ¹⁰Be (alpha emitter).

Below it is shown that the hydrogen is fairly enriched in $^3{\rm H}\approx 40\%$. If the hydrogen were removed, the dose due to the coolant would decrease dramatically but would still have a small amount of activity due to $^{10}{\rm Be}$.

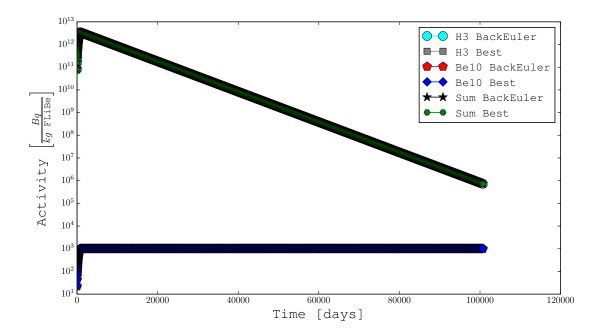


Below we look at the enrichment of ³He. Which is mostly ⁴He.



(c) (25 points) After discharging the FLiBe blanket, how long will it take until the material is less radioactive than Brazil Nuts? (444 Bq/kg)

In order to use the code to do decay, the A matrix was remade with $\phi = 0$. Looking into this problem without ¹⁸F. Also the reason why this plot looks linear is because the y axis has a log scale.



It will take a long time for the $^{10}{\rm Be}$ to decay to a point where its less radioactive than nuts. I decided to do this analytically with

$$t = \frac{-1}{\lambda_Y} ln\left(\frac{444}{N_0 \lambda_s}\right)$$

For the 3 H to decay to 444 Bq, it will take about 400 years, and the 10 Be it will take 1.8E6 years, which is a long time, and slightly over one half life for the 10 Be.

Homework 5 Code

In order to build the A matrix I utilized code from James Tompkins that he let me see a while back. I wanted to give him credit because building the A matrix is probably one of the harder parts of this homework. I had my own experience in building a much larger A matrix for the next problem.

Listing 5: Main Code

```
#!/usr/bin/env python3
##################### Import packages #############################
###################################
import time
start_time = time.time()
import Functions as f
NumOfPoints=200
                # Max Number of points for plots
PlottingG1=["Li6", "Li7", "F19", "Be9"] #List elements you want to Plot
CompareG1=["Li6","Li7","F19","Be9"]
PlottingG2=["H3", "Be10"] #List elements you want to Plot
CompareG2=["H3", "Be10"]
#List elements you want to compare between methods
########### Initialize Matrix #########################
high_flux_fraction=0.5
phi=1.0e14
A, n0=f.MakeAb (high_flux_fraction, phi)
if not A.shape[0] == A.shape[1] or not A.shape[0] == len(n0):
   print("A is not a square matrix")
   quit()
t=730.5; #Two years in days
Nt=1000;
      #Number of Time Steps
dt=t/Nt;
Time=f.np.linspace(dt,t,Nt) #Time steps
tDecay=100000; #No time of decay
NtDecay=10000; #Number of Decay Time Steps
dtDecay=tDecay/NtDecay
TimeDecay=f.np.linspace(t+dtDecay,t+tDecay,NtDecay)
```

```
############## And Time How long it takes ####################
  ############## And Plot at Each Time Step ####################
  #Matrix Exp - Unstable
  #Current_Time=time.time()
  #maxits=20
  # nt_Mat=f.MatExp(A, n0, t, maxits)
  # Mat_Time=time.time()-Current_Time
 #nt_Mat=n0.copy();TIMEOLD=0
  #for TIME in Time[1:len(Time)]: #Unstable
    #nt_Mat=f.MatExp(A,n0,TIME,maxits) No Converge
    #Step through
    #nt_Mat=f.MatExp(A, nt_Mat, TIME-TIMEOLD, maxits)
    #TIMEOLD=TIME.copy()
  #nt_Mat=f.RationalApprox(A, n0, t, maxits) #one Step
  #Mat_Time=time.time()-Current_Time
  Current_Time=time.time()
  File=f.PrepFile('BackEuler.csv',n0) #Prep File
nt_Back=n0.copy();
  #Irradiation Time
  for TIME in Time:
    nt_Back=f.BackEuler(A,nt_Back,dt)
    File.write(str(TIME)+","+f.ListToStr(nt_Back))
  #Decay Time
  phi=0
  A, n0=f.MakeAb(high_flux_fraction, phi)
  for TIME in TimeDecay:
    nt_Back=f.BackEuler(A, nt_Back, dtDecay)
    File.write(str(TIME)+","+f.ListToStr(nt_Back))
  File.close()
  Back_Time=time.time()-Current_Time
 ############# Rational Approx #########################
  #Reset A and n0
 high_flux_fraction=0.5
  phi=1.0e14
  A, n0=f.MakeAb (high_flux_fraction, phi)
  #Irradiation Time
```

```
Current_Time=time.time()
   N=10;
   Method="Best" #Parabola, Cotangent, Hyperbola, Best
   File=f.PrepFile(Method+".csv", n0)
   ck, zk=f.RationalPrep(N, Method); nt_Rational=n0.copy(); TIMEOLD=0
   for TIME in Time:
      nt_Rational=f.RationalApprox(A, n0, TIME, N, ck, zk)
      File.write(str(TIME)+","+f.ListToStr(nt_Rational))
      #Step through
      #nt_Rational=f.RationalApprox(A,nt_Rational,TIME-TIMEOLD,N,ck,zk)
      #TIMEOLD=TIME.copy()
   #nt_Rational=f.RationalApprox(A,n0,t,N,ck,zk) #one Step
   #Non Irradiation Time
   0=idq
A, n0=f.MakeAb (high_flux_fraction, phi)
   n0=nt_Rational
   for TIME in TimeDecay:
      nt_Rational=f.RationalApprox(A, n0, TIME-t, N, ck, zk)
      File.write(str(TIME) +", "+f.ListToStr(nt_Rational))
120
   File.close()
   Rational_Time=time.time()-Current_Time
   dfBack = f.pd.read_csv('BackEuler.csv',index_col=False)
   dfRational = f.pd.read_csv(Method+".csv",index_col=False)
   # #Plot group 1 dudes Back Euler method
   #f.plot(dfBack,PlottingG1,'BackEulerG1',NumOfPoints)
  # f.plot(dfRational,PlottingG1,Method+'G1',NumOfPoints)
   # #Plot group 2 dudes Rational method
   #f.plot(dfBack,PlottingG2,'BackEulerG2',NumOfPoints)
   f.plot(dfRational, PlottingG2, Method+'G2_Check', NumOfPoints)
   # #Plot group 1 dudes, compare both methods
   # Name='BackEuler_'+Method+"_G1"
   # f.plots2(dfBack, dfRational, CompareG1, Name,
           NumOfPoints, 'BackEuler', Method)
145
   #Plot group 2 dudes, compare both methods
   # Name='BackEuler_'+Method+"_G2_NoF"
   # f.plots2(dfBack, dfRational, CompareG2, Name,
           NumOfPoints, 'BackEuler', Method)
150
```

Listing 6: Functions holder

```
#!/usr/bin/env python3
  import sys
  import numpy as np
  import scipy.sparse as sparse
  import scipy.sparse.linalg as spla
  import scipy.linalg as scil
  import scipy.special as sps
  import matplotlib.pyplot as plt
  plt.rcParams["font.family"] = "monospace"
 import matplotlib
  matplotlib.rc('text', usetex=True)
  matplotlib.rcParams['text.latex.preamble']=[r"\usepackage{amsmath}"]
  import random as rn
  import matplotlib.mlab as mlab
20 | import copy
  import os
  import pandas as pd
  ####################### Variables ##########################
  # Basic information
  FigureSize = (11, 6)
FigureSize = (11, 6) # Dimensions of the figure
TypeOfFamily='monospace' # This sets the type of font for text
```

```
font = {'family' : TypeOfFamily} # This sets the type of font for text
  LegendFontSize = 12
  Lfont = {'family' : TypeOfFamily} # This sets up legend font
  Lfont['size'] = LegendFontSize
  Title = ''
  TitleFontSize = 22
  TitleFontWeight = "bold" # "bold" or "normal"
XFontSize=18
                    # X label font size
  XFontWeight="normal" # "bold" or "normal"
  XScale="linear" # 'linear' or 'log'
45 YFontSize=18
                             # Y label font size
  YFontWeight="normal"
YScale="linear"
                            # "bold" or "normal"
  YScale="linear"
                             # 'linear' or 'log'
  Check=0
  Colors=["aqua", "gray", "red", "blue", "black",
                "green", "magenta", "indigo", "lime", "peru", "steelblue",
                "darkorange", "salmon", "yellow", "lime", "black"]
  # If you want to highlight a specific item
  # set its alpha value =1 and all others to 0.4
  # You can also change the MarkSize (or just use the highlight option below)
  # Can change all these to "." or "" for nothing "x" isn't that good
65 | MarkerType=["8", "s", "p", "D", "*", "H", "h", "d", "^", ">"]
  # LineStyles=["solid", "dashed", "dash_dot", "dotted", "."]
  LineStyles=["solid"]
_{70} | SquishGraph = 0.75
  BBOXX = 1.24
  BBOXY = 0.5 # Set legend on right side of graph
  NumberOfLegendColumns=1
  Xlabel='Time [days]'
  Ylabel="Mass $\\left[\\frac{g}{kg \\text{ FLiBe}}\\right]$"
80 | nuclides = { 'H1':0, 'H2':1, 'H3':2, 'He3':3, 'He4':4,
              'He6':5, 'Li6':6, 'Li7':7, 'Li8':8, 'Be8':9,
              'Be9':10, 'Be10':11, 'Be11':12, 'B10':13, 'B11':14,
              'B12':15, 'C12':16, 'C13':17, 'C14':18, 'C15':19,
```

```
'N13':20, 'N14':21, 'N15':22, 'N16':23, 'N17':24,
                '016':25, '017':26, '018':27, '019':28, 'F18':29,
                'F19':30, 'F20':31, 'Ne20':32}
   atom_mass = np.array([1.007825032,2.014101778,3.0160492779,
                                                           #2
                       3.016029320, 4.002603254, 6.151228874,
                                                           #5
                       6.015122887, 7.0160034366, 8.022486246,
                                                           #8
                       8.005305102, 9.012183065, 10.013534695, #11
                       11.02166108, 10.01293695, 11.00930536,
                                                           #14
                       12.0269221, 12, 13.003354835,
                                                           #17
                       14.003241988, 15.01059926,13.00573861, #20
                       14.003074004, 15.000108898, 16.0061019, #23
                       17.008449, 15.994914619, 16.999131756, #26
                       17.999159612, 19.0035780,17.99915961286,#29
                       18.998403162, 19.999981252, 19.992440176])
   nuclide_names = ('H1', 'H2', 'H3', 'He3', 'He4', 'He6', 'Li6',
                   'Li7', 'Li8', 'Be8', 'Be9', 'Be10',
                   'Bell', 'Bl0', 'Bll', 'Bl2', 'Cl2', 'Cl3',
                   'C14', 'C15', 'N13', 'N14', 'N15', 'N16',
                   'N17', '016', '017', '018', '019', 'F18',
                   'F19', 'F20', 'Ne20')
105
   decay\_consts = np.array([0., 0., np.log(2)/3.887896E8, #H1 H2 H3
                          0., 0., np.log(2)/0.807, #He3 He4 He6
                          0.,0., np.log(2)/0.840, #Li6 #Li7 #Li8
                                              #Be8 #Be9
                          np.log(2)/6E-17,0.,
                          np.log(2)/4.73364E13,np.log(2)/13.8, # Be10,11
                          0., 0., np.log(2)/0.0202, #B10 B11 B12
                          0., 0., np.log(2)/1.803517E11, #C12 C13 C14
                          np.log(2)/2.45, np.log(2)/598.2, \#C15 N13
                          0., 0., np.log(2)/7.13, # N14 N15 N16
115
                          np.log(2)/4.174, 0., 0., 0., #N17 016 017 018
                          np.log(2)/26.9, np.log(2)/6586.2, #019 F18
                          0., np.log(2)/11.1, 0.]) #F19 F20 Ne20
   Na=6.0221409E23
120
   def MatExp(A, n0, t, maxits, tolerance=1e-12, LOUD=False):
     converged = False
     sum_old=n0.copy()*0
130
     while not (converged):
       if m==0:
          APowerm=np.identity(A.shape[0])
          Factorial=1
135
       else:
```

```
APowerm=np.dot(APowerm, A)
            Factorial=Factorial*m
        Sum=sum_old+(1/Factorial)*np.dot((APowerm)*(t**m),n0)
140
        #Avoid dividing by zero
        if sum(Sum) ==0: m+=1; sum_old=Sum.copy(); continue
        change = np.linalg.norm(Sum-sum_old)/np.linalg.norm(Sum)
        converged = (change < tolerance) or (m > maxits)
145
        if (LOUD>0) or (converged and LOUD<0):</pre>
          print("Iteration", m, ": Relative Change =", change)
        if (m > maxits):
          print("Warning: Source Iteration did not converge: "+\
150
                " m : "+str(m)+", Diff : %.2e" % change)
        #Prepare for next iteration
        m += 1
        sum_old = Sum.copy()
155
      return (Sum)
    def BackEuler(A, no, dt):
        I=np.identity(A.shape[0])
        return (np.dot (np.linalg.inv(I-A*dt), no))
160
    def DeterminePolesNResidues(n):
        This program takes the algorithm from the reference
        and converts to a python script... I know its janky
165
        but it works
        .....
        def Append(List1, List2):
            for item in List2:
                for item2 in item:
                    List1=np.append(List1,item2)
            return (List1)
        def absG(List):
            List2=[]
175
            for item in List:
                if abs(item)>1:
                    List2.append(item)
            return(List2)
        #function [zk, ck] = cf(n);
        K = 75;
                                              # no of Cheb coeffs
180
        nf = 1024;
                                              # no of pts for FFT
        #Roots correct?
        roots=np.arange(0,nf,1)/nf
        #w = np.exp(2i*pi*(0:nf-1)/nf);
                                              # roots of unity
185
        w=np.exp(2j*np.pi*roots)
        t = np.real(w);
                                              # Cheb pts (twice over)
                                              # scale factor for stability
        \#F = np.exp(scl*(t-1)./(t+1+1e-16)); \# exp(x) transpl. to [-1,1]
```

```
F = np.exp(scl*(t-1)/(t+1+1e-16)); # exp(x) transpl. to [-1,1]
190
                                            # Cheb coeffs of F
        c = np.real(np.fft.fft(F))/nf;
        index=reversed(np.arange(1,K+2,1))
        partofc=[]
        for i in index:
           partofc.append(c[i-1])
195
        #f = np.polyval(c(K+1:-1:1), w); # analytic part f of F
        f = np.polyval(partofc,w);  # analytic part f of F
        \#[U, S, V] = svd(hankel(c(2:K+1)));
                                            # SVD of Hankel matrix
        hankie=scil.hankel(c[1:K+1])
        U,S,V=np.linalg.svd(hankie,full_matrices=False)
        #s = S(n+1, n+1);
                                              # singular value
        s=S[n]
        \#u = U(K:-1:1,n+1); v = V(:,n+1); \# singular vector
205
        u=[]
        index=reversed(np.arange(0,K,1))
        for i in index:
           u.append(U[i,n])
        \#v=np.array(V[:,n].copy())
210
        v=np.array(V[n,:].copy())
        \#zz = zeros(1, nf-K);
                                             # zeros for padding
        zz=np.zeros([1,nf-K])
        \#b = fft([u zz])./fft([v zz]);
                                            # finite Blaschke product
        b=np.fft.fft(Append(u,zz))/np.fft.fft(Append(v,zz))
215
        \#rt = f-s*w.^K.*b;
                                              # extended function r-tilde
        rt=f-s*(w**K)*b;
        #rtc = real(fft(rt))/nf;
                                              # its Laurent coeffs
        rtc=np.real(np.fft.fft(rt))/nf;
        \#zr = roots(v); qk = zr(abs(zr)>1); \# poles
220
        zr=np.roots(v);qk=np.array(absG(zr));
        #qc = poly(qk);
                                              # coeffs of denominator
        qc=np.poly(qk);
        #pt = rt.*polyval(qc,w);
                                             # numerator
        pt=rt*np.polyval(qc,w);
                                             # coeffs of numerator
        #ptc = real(fft(pt)/nf);
        ptc=np.real(np.fft.fft(pt)/nf);
        \#ptc = ptc(n+1:-1:1); ck = 0*qk;
        index=reversed(np.arange(0,n+1,1))
        ptc2=[]
        for i in index: #Can I just reversed ptc?
           ptc2.append(ptc[i])
        ptc=ptc2.copy()
        ck=0*qk
        #N+1?
235
        #for k = 1:n
                                 # calculate residues
             q = qk(k); q2 = poly(qk(qk\sim=q));
             ck(k) = polyval(ptc,q)/polyval(q2,q);
        for k in range (0, n):
            if len(qk) == k:
240
                print("we are short a qk")
                continue
```

```
q=qk[k];
            q2 = [];
            for item in qk:
245
                if not q==item:
                    q2.append(item)
            q2=np.poly(q2);
            ck[k]=np.polyval(ptc,q)/np.polyval(q2,q)
        \#zk = scl*(qk-1).^2./(qk+1).^2;
                                             # poles in z-plane
250
        zk=scl*((qk-1)**2)/((qk+1)**2)
        \#ck = 4*ck.*zk./(qk.^2-1);
                                             # residues in z-plane
        ck=4*ck*zk/(qk**2-1)
        #Cut down ck and zk to half the original points
        ck2=[];zk2=[]
        for i in range(0,len(ck)):
            if i % 2 == 0:
                ck2=np.append(ck2,ck[i])
                zk2=np.append(zk2,zk[i])
        return (ck2, zk2)
    def RationalPrep(N,Phi):
        """Calculate constants for a rational approximation
265
        Inputs:
                         Number of Quadrature points
        N:
        Phi:
                         'Parabola',
                          'Cotangent', or
                          'Hyperbola' (shape of Phi)
270
        Outputs:
        ck:
                        First set of constants for approximation
        zk:
                        Second set of constants for approximation
        theta=np.pi*np.arange(1,N,2)/N
275
        if Phi=='Parabola':
            zk=N*(0.1309-0.1194*theta**2+0.2500j*theta)
            w=N*(-2*0.1194*theta+0.2500j)
        elif Phi=='Cotangent':
            cot=1/np.tan(0.6407*theta)
280
            ncsc=-0.6407/(np.sin(0.6407*theta)**2)
            zk=N*(0.5017*theta*cot-0.6122+0.2645j*theta)
            w=N*(0.2645j+0.5017*cot+0.5017*theta*ncsc)
        elif Phi=='Hyperbola':
            zk=2.246*N*(1-np.sin(1.1721-0.3443j*theta))
            w=2.246*N*(0.3443j*np.cos(1.1721-0.3443j*theta))
        elif Phi=='Best':
            ck, zk=DeterminePolesNResidues(N)
            return (ck, zk)
        else:
            print("Did not pick proper rational approximation dude")
            print("Quiting now")
            quit()
        ck=1.0j/N*np.exp(zk)*w
```

```
return (ck, zk)
    def RationalApprox(A,n0,t,N,ck,zk,tol=1e-12,maxits=2000):
        Calculate the rational approximation solution for n(t)
300
        Inputs:
                   Matrix with system to be solved
        A:
       n0:
                   initial conditions of the system
                   time at which solution is determined
        t.:
                   Number of quadrature points (should be less than 20)
        N:
305
        ck:
                   constants for quadrature solution
                   constants for quadrature solution
        zk:
                   Tolerence for convergence for GMRES
                   Maximium iterations for GMRES
       maxits:
       Outputs:
310
        nt:
                   Solution at time t
        .....
       nt=np.zeros(len(n0))
        for k in range (int (N/2)):
315
            if len(n0)>1:
                \#phi, code=spla.gmres(zk[k]*sparse.identity(len(n0))-A*t,n0,
                                     tol=tol, maxiter=maxits)
                phi=np.dot(np.linalg.inv(zk[k]*np.identity(len(n0))-A*t),
                           n0)
                #if (code):
                #
                  print(code)
            else:
                phi = (zk[k]-A*t)**(-1)*n0
            nt=nt-2*np.real(ck[k]*phi)
325
        return (nt)
    def MakeAb(hi_flux_frac = 0.5,phi = 1.0e14):
        """Interaction functions
        @ In, nuclides: dictionary with isotope keywords and
330
                        corresponding indices
        @ In, parent: parent nuclides undergoing a decay or interaction
                        new value in interaction matrix, either a half
        @Out, value:
                         life [secs] or 2.45 MeV and 14.1 MeV cross
                         sections [barns]
335
        def betanegdecay(nuclides, parent):
                parent == 'F20': return nuclides['Ne20'], 11.1 # s
            elif parent == '019': return nuclides['F19'], 26.9 # s
            elif parent == 'N16': return nuclides['016'], 7.13 # s
            elif parent == 'N17': return nuclides['017'], 4.174 # s
            elif parent == 'C14': return nuclides['N14'], 1.803517E11 # s
            elif parent == 'C15': return nuclides['N15'], 2.45 # s
            elif parent == 'B12': return nuclides['C12'], 0.0202 # s
            elif parent == 'Be10': return nuclides['B10'], 4.73364E13 # s
            elif parent == 'Bell': return nuclides['B11'], 13.8 # s
            elif parent == 'Li8': return nuclides['Be8'], 0.840 # s
```

```
elif parent == 'He6': return nuclides['Li6'], 0.807 # s
            elif parent == 'H3': return nuclides['He3'], 3.887896E8 # s
350
            else: return -1, 0.0
        def betaposdecay(nuclides, parent):
               parent == 'F18': return nuclides['018'], 6586.2 # s
            elif parent == 'N13': return nuclides['C13'], 598.2 # s
355
            else: return -1, 0.0
        def twoalphadecay(nuclides, parent):
            if parent == 'Be8': return nuclides['He4'], 7.0E-17 # s
            else: return -1, 0.0
360
        def n_gamma(nuclides, parent):
               parent == 'F19':
               return nuclides['F20'], 8.649107E-5, 3.495035E-5
            elif parent == '016':
365
               return nuclides['017'], 1.0E-4,
                                                       1.0E-4
            elif parent == '017':
               return nuclides['018'], 2.2675E-4,
                                                        2.087114E-4
            elif parent == 'N14':
               return nuclides['N15'], 2.397479E-5,
                                                        1.679535E-5
370
            elif parent == 'N15':
               return nuclides['N16'], 8.121795E-6,
                                                        8.56E-6
            elif parent == 'Be9':
               return nuclides['Be10'],1.943574E-6,
                                                       1.660517E-6
            elif parent == 'Li6':
               return nuclides['Li7'], 1.106851E-5,
                                                       1.017047E-5
            elif parent == 'Li7':
               return nuclides['Li8'], 4.677237E-6,
                                                       4.105546E-6
            elif parent == 'He3':
               return nuclides['He4'], 9.28775E-5,
                                                       3.4695E-5
            elif parent == 'H2':
               return nuclides['H3'], 8.413251E-6,
                                                       9.471512E-6
            else:
               return -1, 0.0, 0.0
385
        def n_2n(nuclides, parent):
           if parent == 'F19':
               return nuclides['F18'], 0.0,
                                                        0.04162
            elif parent == '017':
390
               return nuclides['016'], 0.0,
                                                        0.066113
            elif parent == 'N14':
               return nuclides['N13'], 0.0,
                                                        0.006496
            elif parent == 'N15':
               return nuclides['N14'], 0.0,
                                                        0.112284
395
            elif parent == 'B11':
               return nuclides['B10'], 0.0,
                                                        0.018805
            elif parent == 'Be9':
               return nuclides['Be8'], 0.0205,
                                                       0.484483
            elif parent == 'Li7':
400
               return nuclides['Li6'], 0.0,
                                                        0.031603
```

```
elif parent == 'H3':
                return nuclides['H2'],
                                         0.0,
                                                          0.0497
            elif parent == 'H2':
                return nuclides['H1'], 0.0,
                                                         0.166767
            else:
                return -1, 0.0, 0.0
        def n_alpha(nuclides, parent):
               parent == 'F19':
410
                return [nuclides['N16'], nuclides['He4']], 2.1667E-5, 0.028393
            elif parent == '016':
                return [nuclides['C13'], nuclides['He4']], 0.0, 0.144515
            elif parent == '017':
                return [nuclides['C14'], nuclides['He4']], 0.117316, 0.260809
415
            elif parent == 'N14':
                return [nuclides['B11'], nuclides['He4']], 0.104365, 0.080516
            elif parent == 'N15':
                return [nuclides['B12'], nuclides['He4']], 0.0,0.069240
            elif parent == 'B10':
420
                return [nuclides['Li7'], nuclides['He4']], 0.281082, 0.044480
            elif parent == 'B11':
                return [nuclides['Li8'], nuclides['He4']], 0.0,0.031853
            else:
                return [-1,-1], 0.0, 0.0
425
        def n_2alpha(nuclides, parent):
               parent == 'N14':
                return [nuclides['Li7'], nuclides['He4']], 0.0,0.031771
            elif parent == 'B10':
430
                return [nuclides['H3'], nuclides['He4']], 0.038439, 0.095487
                return [-1,-1], 0.0, 0.0
        def n_nalpha(nuclides, parent):
               parent == 'F19':
                return [nuclides['N15'], nuclides['He4']], 0.0,0.3818
            elif parent == '017':
                return [nuclides['C13'], nuclides['He4']], 0.0,0.043420
            elif parent == 'N15':
440
                return [nuclides['B11'], nuclides['He4']], 0.0,0.012646
            elif parent == 'B11':
                return [nuclides['Li7'], nuclides['He4']], 0.0,0.286932
            elif parent == 'Be9':
                return [nuclides['He6'], nuclides['He4']], 0.0825,0.0104
445
                return [-1,-1], 0.0, 0.0
        def n_2nalpha(nuclides, parent):
            if parent == 'Li6':
450
                return [nuclides['H1'], nuclides['He4']], 0.0,0.0783
            elif parent == 'Li7':
                return [nuclides['H2'], nuclides['He4']], 0.0,0.020195
            else:
```

```
return [-1,-1], 0.0, 0.0
455
        def n_3nalpha(nuclides, parent):
            if parent == 'Li7':
                return [nuclides['H1'], nuclides['He4']], 0.0,6.556330E-5
            else:
460
                return [-1,-1], 0.0, 0.0
        def n_p(nuclides, parent):
                parent == 'F19':
                return [nuclides['019'], nuclides['H1']], 0.0,0.018438
            elif parent == '016':
                return [nuclides['N16'], nuclides['H1']], 0.0,0.042723
            elif parent == '017':
                return [nuclides['N17'], nuclides['H1']], 0.0, 0.041838
            elif parent == 'N14':
470
                return [nuclides['C14'], nuclides['H1']], 0.014102, 0.043891
            elif parent == 'N15':
                return [nuclides['C15'], nuclides['H1']], 0.0,0.019601
            elif parent == 'B10':
                return [nuclides['Be10'], nuclides['H1']], 0.018860, 0.034093
475
            elif parent == 'B11':
                return [nuclides['Bell'], nuclides['H1']], 0.0,0.005564
            elif parent == 'Li6':
                return [nuclides['He6'], nuclides['H1']], 0.0,0.00604
            elif parent == 'He3':
480
                return [nuclides['H3'], nuclides['H1']], 0.714941, 0.121
            else:
                return [-1,-1], 0.0, 0.0
        def n_np(nuclides, parent):
485
            if parent == 'F19':
                return [nuclides['018'], nuclides['H1']], 0.0, 0.061973
            elif parent == 'N15':
                return [nuclides['C14'], nuclides['H1']], 0.0, 0.044827
            elif parent == 'B11':
490
                return [nuclides['Be10'], nuclides['H1']], 0.0, 0.001016
            else:
                return [-1,-1], 0.0, 0.0
        def n_d(nuclides, parent):
                parent == 'F19':
                return [nuclides['018'], nuclides['H2']], 0.0, 0.022215
            elif parent == '016':
                return [nuclides['N15'], nuclides['H2']], 0.0,0.017623
            elif parent == '017':
500
                return [nuclides['N16'], nuclides['H2']],  0.0,0.020579
            elif parent == 'N14':
                return [nuclides['C13'], nuclides['H2']], 0.0, 0.042027
            elif parent == 'N15':
                return [nuclides['C14'], nuclides['H2']], 0.0,0.014926
505
            elif parent == 'B10':
                return [nuclides['Be9'], nuclides['H2']], 0.0, 0.031270
```

```
elif parent == 'Li7':
                return [nuclides['He6'], nuclides['H2']], 0.0, 0.010199
            elif parent == 'He3':
510
                return [nuclides['H2'], nuclides['H2']], 0.0,0.07609
            else:
                return [-1,-1], 0.0, 0.0
        def n_t(nuclides, parent):
515
               parent == 'F19':
                return [nuclides['017'], nuclides['H3']],  0.0,0.01303
            elif parent == 'N14':
                return [nuclides['C12'], nuclides['H3']],  0.0,0.028573
            elif parent == 'N15':
                return [nuclides['C13'], nuclides['H3']],  0.0,0.020163
            elif parent == 'B11':
                return [nuclides['Be9'], nuclides['H3']], 0.0,0.015172
            elif parent == 'Be9':
                return [nuclides['Li7'], nuclides['H3']], 0.0,0.020878
            elif parent == 'Li6':
                return [nuclides['He4'], nuclides['H3']], 0.206155,0.0258
            else:
                return [-1,-1], 0.0, 0.0
530
        # Create Activation and Decay Matrix and initial
        # nuclide quantity vector
        A = np.zeros((len(nuclides), len(nuclides)))
535
        lo_flux_frac = (1.0-hi_flux_frac)
        phi = phi * 60 * 60 * 24 #10^14 1/cm^2/s in 1/cm^2 /day
       phi_hi = hi_flux_frac*phi*1.0e-24
       phi_lo = lo_flux_frac*phi*1.0e-24
540
        for isotope in nuclides:
            row = nuclides[isotope]
            row_betanegdecay = betanegdecay(nuclides, isotope)
545
            row_betaposdecay = betaposdecay(nuclides, isotope)
            row_2alphadecay = twoalphadecay(nuclides, isotope)
            row_n_gamma =
                                n_gamma(nuclides, isotope)
                                n_2n(nuclides, isotope)
            row_n_2n =
            row_n_alpha =
                                n_alpha(nuclides, isotope)
                                n_2alpha(nuclides, isotope)
            row_n_2alpha =
                                n_nalpha(nuclides, isotope)
            row_n_nalpha =
            row_n_2nalpha =
                                n_2nalpha(nuclides, isotope)
            row_n_3nalpha =
                                n_3nalpha(nuclides, isotope)
            row_n_p =
                                n_p(nuclides, isotope)
                                n_np(nuclides, isotope)
            row_n_np =
            row_n_d =
                                n_d(nuclides, isotope)
            row_n_t =
                                n_t(nuclides, isotope)
            row_lo_act_sum = row_n_gamma[1] + row_n_2n[1] + 
                             row_n_alpha[1] + row_n_2alpha[1] +\
560
```

```
row_n_nalpha[1] + row_n_2nalpha[1] + \
                              row_n_3nalpha[1] + row_n_p[1] + 
                             row_n_p[1] + row_n_d[1] + 
                             row_n_t[1]
            row_hi_act_sum = row_n_gamma[2] + row_n_2n[2] +\
565
                             row_n_alpha[2] + row_n_2alpha[2] + \\
                             row_n_nalpha[2] + row_n_2nalpha[2] + \
                             row_n_3nalpha[2] + row_n_p[2] +\
                             row_n_np[2] + row_n_d[2] +row_n_t[2]
            # try:
570
                  if row_n_alpha[0] >= 0:
                      print(row_n_alpha)
                      donotuse=100
                  continue
            # except TypeError:
575
                  print(row_n_alpha)
                  print(row_n_alpha[0][0])
                  quit()
580
            if row_betanegdecay[0] >= 0:
                # [days^-1]
                row_lambda = np.log(2)*60*60*24/row_betanegdecay[1]
            elif row_betaposdecay[0] >= 0:
                # [davs^-1]
                row_lambda = np.log(2)*60*60*24/row_betaposdecay[1]
            elif row_2alphadecay[0] >= 0:
                # [days^-1]
                row_lambda = np.log(2)*60*60*24/row_2alphadecay[1]
            else:
590
                row_lambda = 0.0
            # Diagonal Assignment
            A[row,row] = -row_lambda - phi_lo*row_lo_act_sum -\
                         phi_hi*row_hi_act_sum
            # Off Diagonal Assignment
595
            if row_betanegdecay[0] >= 0:
                A[row\_betanegdecay[0], row] = np.log(2)*60*60*24/
                                              row_betanegdecay[1]
            if row_betaposdecay[0] >= 0:
                A[row\_betaposdecay[0], row] = np.log(2)*60*60*24/
600
                                              row_betaposdecay[1]
            if row_2alphadecay[0] >= 0:
                A[row_2alphadecay[0], row] = np.log(2)*60*60*24/
                                             row_2alphadecay[1]
            if row_n_qamma[0] >= 0:
                A[row_n_gamma[0], row] = phi_lo*row_n_gamma[1] +
                                         phi_hi*row_n_gamma[2]
            if row_n_2n[0] >= 0:
                A[row_n_2n[0], row] = phi_lo*row_n_2n[1] + 
                                      phi_hi*row_n_2n[2]
            if row_n_alpha[0][0] >= 0:
                for i in row_n_alpha[0]:
                    A[i,row] = phi_lo*row_n_alpha[1] +
```

```
phi_hi*row_n_alpha[2]
615
           if row_n_2alpha[0][0] >= 0:
               for i in row_n_2alpha[0]:
                   A[i,row] = phi_lo*row_n_2alpha[1] + 
                             phi_hi*row_n_2alpha[2]
           if row_n_nalpha[0][0] >= 0:
               for i in row_n_nalpha[0]:
620
                   A[i,row] = phi_lo*row_n_nalpha[1] + 
                             phi_hi*row_n_nalpha[2]
           if row_n_2nalpha[0][0] >= 0:
               for i in row_n_2nalpha[0]:
                  A[i,row] = phi_lo*row_n_2nalpha[1] +\
625
                             phi_hi*row_n_2nalpha[2]
           if row_n_3nalpha[0][0] >= 0:
               for i in row_n_3nalpha[0]:
                   A[i,row] = phi_lo*row_n_3nalpha[1] + 
                             phi_hi*row_n_3nalpha[2]
630
           if row_n_p[0][0] >= 0:
               for i in row_n_p[0]:
                  A[i,row] = phi_lo*row_n_p[1] + phi_hi*row_n_p[2]
           if row_n_p[0][0] >= 0:
               for i in row_n_np[0]:
635
                   A[i,row] = phi_lo*row_n_np[1] + phi_hi*row_n_np[2]
           if row_n_d[0][0] >= 0:
               for i in row_n_d[0]:
                  A[i,row] = phi_lo*row_n_d[1] + phi_hi*row_n_d[2]
           if row_n_t[0][0] >= 0:
               for i in row_n_t[0]:
                  A[i,row] = phi_lo*row_n_t[1] + phi_hi*row_n_t[2]
       b = np.zeros(len(nuclides))
645
       # N_0 expressed as kg nuclide per kg FLiBe
       #b[nuclides['F19']] = 0.7685
       #b[nuclides['Be9']] = 0.0911
       #b[nuclides['Li6']] = 0.01065636
650
       #b[nuclides['Li7']] = 0.12974364
       AtomsofFLiBe=6.0899894727155e24
       b[nuclides['F19']] = AtomsofFLiBe * 4
       b[nuclides['Be9']] = AtomsofFLiBe*1
       b[nuclides['Li6']] = AtomsofFLiBe*2*0.0759
655
       b[nuclides['Li7']] = AtomsofFLiBe*2*0.9241
       return (A,b)
   ################## Plotting Function #############################
   def reduceList(List, N):
       List2=[List[0]]
665
       Div=int(len(List)/N)
```

```
for i in range(1,len(List)-1):
            if i % Div == 0:
                List2.append(List[i])
        List2.append(List[-1])
670
        return (List2)
    def PlotPoints(Ntot, Nplot):
   def loop_values(list1,index):
        This function will loop through values in list even if
        outside range (in the positive sense not negative)
        while True:
680
            try:
                list1[index]
                break
            except IndexError:
                index=index-len(list1)
685
        return(list1[index])
    def Legend(ax):
        handles, labels=ax.get_legend_handles_labels()
        ax.legend(handles, labels, loc='best',
690
                  fontsize=LegendFontSize,prop=font)
        return (ax)
    # def Legend(ax):
              handles, labels = ax.get_legend_handles_labels()
695
              box=ax.get_position()
              ax.set_position([box.x0, box.y0, box.width*SquishGraph,
                                box.height])
              ax.legend(handles, labels, loc='center',
700
                        bbox_to_anchor=(BBOXX, BBOXY),
                        fontsize=LegendFontSize,prop=font,
                        ncol=NumberOfLegendColumns)
              return(ax)
   def InList(item2,List):
     TF=False
      for item1 in List:
        if item1 == item2:
          TF=True
      if not TF:
710
        print("Invalid selection for plotting")
        print("Shuting down")
        quit()
   def plot(df,Plotting,Name,NumOfPoints):
        #Plot In grams
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
```

```
List=list(df.columns.values)
720
        x=df[List[0]].values[2:-1]
        Check=0
        for Item in Plotting:
          InList(Item, List) #Check if we have the isotope
725
          y=((df[Item].values[2:-1])/Na)*df[Item].values[0]
          if len(x)>NumOfPoints:
            x=reduceList(x,NumOfPoints)
            y=reduceList(y,NumOfPoints)
          ax.plot(x,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
735
                  label=Item)
          Check=Check+1
        #Log or linear scale?
740
        ax.set_xscale(XScale)
        ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
745
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                      fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        ax.set_ylabel(Ylabel,
750
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
755
        Legend (ax)
        plt.savefig("Plots/"+Name+'_grams.pdf')
        #Plot in Bq ##################################
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
        List=list(df.columns.values)
        x=df[List[0]].values[2:-1]
765
        Check=0; Sum=np.zeros(len(x))
        for Item in Plotting:
          InList(Item,List) #Check if we have the isotope
          y=((df[Item].values[2:-1]))*df[Item].values[1]
          Sum=Sum+y
770
          if len(x)>NumOfPoints:
            xP=reduceList(x, NumOfPoints)
```

```
y=reduceList(y,NumOfPoints)
          ax.plot(xP,y,
                  linestyle=loop_values(LineStyles, Check),
775
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item)
780
          Check=Check+1
        if len(x) > NumOfPoints:
          Sum=reduceList (Sum, NumOfPoints)
        ax.plot(xP,Sum,
                linestyle=loop_values(LineStyles, Check),
785
                marker=loop_values (MarkerType, Check),
                color=loop_values(Colors, Check),
                markersize=loop_values(MarkSize,Check),
                alpha=loop_values(Alpha_Value, Check),
                label="Sum")
        #Log or linear scale?
        ax.set_xscale(XScale)
        if sum(Sum) == 0:
          ax.set_yscale('linear')
795
        else:
          ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                      fontweight=TitleFontWeight, fontdict=font, ha='center')
800
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        YlabelBq="Activity $\\left[\\frac{Bq}{kg \\text{ FLiBe}}\\right]$"
805
        ax.set_ylabel(YlabelBq,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
810
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Bq.pdf')
    def plots2(df, df2, Plotting, Name, NumOfPoints, Method1, Method2):
        #Plot in grams
815
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
        List=list(df.columns.values)
        x=df[List[0]].values[2:-1]
        Check=0
        for Item in Plotting:
          InList(Item, List) #Check if we have the isotope
          y=((df[Item].values[2:-1])/Na)*df[Item].values[0]
825
```

```
y2=((df2[Item].values[2:-1])/Na)*df2[Item].values[0]
          if len(x)>NumOfPoints:
            x=reduceList(x,NumOfPoints)
            y=reduceList(y,NumOfPoints)
            y2=reduceList(y2,NumOfPoints)
830
          ax.plot(x,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values (MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check) *1.5,
835
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method1)
          Check=Check+1
          ax.plot(x,y2,
                  linestyle=loop_values(LineStyles, Check),
840
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method2)
          Check=Check+1
        #Log or linear scale?
        ax.set_xscale(XScale)
        ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
855
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                      fontsize=XFontSize, fontweight=XFontWeight,
                      fontdict=font)
        ax.set_ylabel(Ylabel,
                      fontsize=YFontSize,
860
                      fontweight=YFontWeight,
                      fontdict=font)
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Grams.pdf')
865
        #Plot in Bq
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
        List=list(df.columns.values)
        x=df[List[0]].values[2:-1]
        Check=0; Sum=np.zeros(len(x)); Sum2=np.zeros(len(x))
        for Item in Plotting:
          InList(Item, List) #Check if we have the isotope
          y=((df[Item].values[2:-1]))*df[Item].values[1]
```

```
y2=((df2[Item].values[2:-1]))*df2[Item].values[1]
          Sum=Sum+y
          Sum2=Sum2+y2
          if len(x) > NumOfPoints:
            xP=reduceList(x, NumOfPoints)
            y=reduceList(y,NumOfPoints)
            y2=reduceList(y2,NumOfPoints)
          ax.plot(xP,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values (MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check) *1.5,
890
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method1)
          Check=Check+1
          ax.plot(xP,y2,
                  linestyle=loop_values(LineStyles, Check),
895
                  marker=loop_values (MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method2)
900
          Check=Check+1
        if len(x) > NumOfPoints:
          Sum=reduceList(Sum, NumOfPoints)
          Sum2=reduceList(Sum2,NumOfPoints)
        ax.plot(xP,Sum,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check) *1.5,
                  alpha=loop_values(Alpha_Value, Check),
910
                  label="Sum "+Method1)
        Check=Check+1
        ax.plot(xP,Sum2,
                linestyle=loop_values(LineStyles, Check),
                marker=loop_values(MarkerType, Check),
915
                color=loop_values (Colors, Check),
                markersize=loop_values(MarkSize,Check),
                alpha=loop_values(Alpha_Value, Check),
                label="Sum "+Method2)
920
        #Log or linear scale?
        ax.set_xscale(XScale)
        if sum(Sum) == 0:
          ax.set_yscale('linear')
        else:
925
          ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                      fontweight=TitleFontWeight, fontdict=font, ha='center')
930
        #Set X and y labels
```

```
ax.set_xlabel(Xlabel,
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        YlabelBq="Activity $\\left[\\frac{Bq}{kg \\text{ FLiBe}}\\right]$"
935
        ax.set_ylabel(YlabelBq,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
940
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Bq.pdf')
    def ListToStr(List):
      Str=''
945
      for i in range(0,len(List)):
        if not i==len(List)-1:
          Str=Str+str(List[i])+","
        else:
          Str=Str+str(List[i])+"\n"
950
      return (Str)
    def PrepFile(Name, n0):
      File=open(Name,'w')
      File.write("Mass then Time (d),"+','.join(nuclide_names)+'\n')
      File.write("Masses,"+ListToStr(atom_mass)) #New line already included
      File.write("DecayConts,"+ListToStr(decay_consts))
      File.write("0,"+ListToStr(n0))
      return (File)
960
    def Print (Method, nuclide, Results, Time):
      Index=nuclides[nuclide]
      MassConversion=atom_mass[Index]/Na
      string="Isotope "+nuclide_names[Index]+", Mass (g) = "
      Mass=Results[Index] *MassConversion
      Mass="%.4e" % Mass
      print (Method+" :", string, Mass, "Time=%.2f" % Time)
    def Years (Method, nuclide, Results):
     Index=nuclides[nuclide]
      LambdaY=decay_consts[Index] *60 *60 *24 *365.25
      Lambdas=decay_consts[Index]
      string="Isotope "+nuclide_names[Index]+", Years to 444 Bq = "
      Years=(-1/LambdaY)*np.log(444/(Results[Index]*Lambdas))
      print (Method+" :", string, "%.3e" % Years)
975
```

Project

For the project I wanted to solve a depletion problem using the algorithms developed in the previous homework. The bateman equations will be written in a different form to align more with how the system was built in homework 5, and for the project.

The production of an isotope is dictated by production and loss

$$\frac{dn_i}{dt} = -\lambda_i^{eff} n_i + \sum_{j=1}^N b_{j \to i}^{eff} n_j$$

Where N is the number of nuclides and,

$$\lambda_i^{eff} = \lambda_i + \phi \sum_{i=1}^N \sigma_{i \to j}$$

and

$$b_{j \to i}^{eff} = b_{j \to i} \lambda_j + \sigma_{j \to i} \phi + \gamma_{j \to i} \sigma_{j,f} \phi$$

where $\gamma_{j\to i}$ is the fission yield for isotope i from fission of isotope j. $b_{j\to i}$ is the fraction of radioactive disintegration by nuclide j, which leads to nuclide i. In the case of spontaneous fission $b_{j\to i}$ is the product of spontenous fission fraction for nuclide j and the yield for fission from nuclide j producing nuclide i.

For a system of isotopes, the system can be reduced to:

$$\frac{d\vec{n}}{dt} = \mathbf{A}\vec{n}(t)$$

Where \boldsymbol{A} is a matrix whose diagonal elements are $[-\lambda_1^{eff}, -\lambda_2^{eff}, ..., -\lambda_N^{eff}]$, all off diagonal elements are $b_{j \to i}^{eff}$ (i for the row, and j is for the column) and $\vec{n}(t) = [n_1, n_2, ..., n_N]$.

The solution to this system is so obvious, I won't even write it down.

For the current problem 1 metric ton of PWR fuel will be irradiated for 400 days at a constant flux of 2.9896 $\cdot 10^{14}$ n/cm²·s (37.5 W/g initially) and some of the fission product masses will be determined as a function of time. If we ignore the oxygen, the initial amount of atoms for each of the heavy isotopes is

$$N_{^{234}U} = 270 \text{ g} \frac{6.022E23 \text{ atoms}}{234.0409523 \text{ g}} = 6.94741E23 \frac{\text{atoms of }^{234}\text{U}}{\text{tHM}}$$

 $N_{^{235}U} = 30000 \text{ g} \frac{6.022E23 \text{ atoms}}{235.0439301 \text{ g}} = 7.6864E25 \frac{\text{atoms of }^{235}\text{U}}{\text{tHM}}$
 $N_{^{238}U} = 969730 \text{ g} \frac{6.022E23 \text{ atoms}}{238.0507884 \text{ g}} = 2.4532E27 \frac{\text{atoms of }^{238}\text{U}}{\text{tHM}}$

The results will be compared with ORIGEN2 results for a similar problem that will irradiate the same fuel with a constant power of 37.5 W/g (compare with my results from UQ).

Project Results

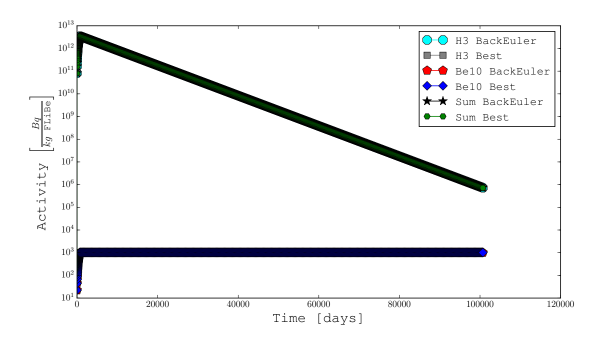
The results for the code are shown below. The hardest part of this problem was building the A matrix. In order to determine if the system was solved correctly the information fed into ORIGEN2 from its TAPE9.inp file was used so that results from the two codes could be compared. I wanted to be able to recreate what ORIGEN2 does with my own code so I could complete the project for UQ in the future after I graduate. The only part I am not sure about on how to do is how to sample the cross section spectrum.

Comparison of results from the two codes are shown in the table below, and it can be seen that they are fairly close. The main difference is from the constant power (ORIGEN2) constant flux (MINE) different assumption.

Table 1: Comparison between ORIGEN2 and my code results in grams

Nuclide	ORIGEN2	My Code
^{137}Cs	553	544
^{137}Ba	7.10	7.13
^{136}Xe	1.04E3	1.02E3
^{134}Ba	3.70	3.60
^{133}Cs	549	538
^{106}Ru	74.3	73.2
^{105}Pd	136	134
^{99}Tc	374	371
^{95}Zr	83.1	79.9
^{235}U	1.629E4	1.65 E4
^{238}U	9.597 E5	9.6E5
^{239}Pu	3.994E3	3.85E3
^{241}Pu	5.279E2	5.838E2

Below are some graphs of the production and loss of nuclides to show nuclide production trends.



Project Code

Listing 7: Main Code

```
#!/usr/bin/env python3
#Means for improving speed. Make dir and save all variables
#made for making a, so that no parsing is needed,
#Just multiplying by phi
import time
start_time = time.time()
import Functions as f
NumOfPoints=30
                  # Max Number of points for plots
PlottingG1=["922350"] #List elements you want to Plot
#CompareG1=["922350","922380","942390"]
CompareG1=["942390"]
PlottingG2=["922350"] #List elements you want to Plot
#CompareG2=["551370","631540","601480"]
CompareG2=["551370"]
#List elements you want to compare between methods
########## Initialize Matrix #########################
phi=1.0e14; Power = 37.5 \# MW (bc 1 ton HM power den is W/q)
Fissile_Isotopes=["922350","922380","942390","942410"]
FissionXSections=f.FindFissionXSection(Fissile_Isotopes)
MeVperFission=f.CalMevPerFiss(Fissile_Isotopes)
ToAdd=[]
Nuclides, Nuclide_Names=f.Isotopes(ToAdd)
n0=f.Makeb(Nuclides)
phi=f.Calculatephi(FissionXSections, MeVperFission, n0, Power, Fissile_Isotopes, Nuclides)
#This variable takes a while to make, so if its already made
#Just load it
#f.os.system("rm Saved_Variables/Masses.npy")  #To start over and remake
if not f.os.path.isfile('Saved_Variables/Masses.npy'):
   #Grab atomic weight information
   df=f.pd.read_csv('Data/AtomicWeights.csv')
   Atom_Mass=f.GatherMasses(df, Nuclides)
   f.np.save('Saved_Variables/Masses', Atom_Mass)
else:
  Atom_Mass=f.np.load('Saved_Variables/Masses.npy')
```

```
#Load up all the decay constants
   #f.os.system("rm Saved_Variables/Decay_Consts.npy")  #To start over and remake
   if not f.os.path.isfile('Saved_Variables/Decay_Consts.npy'):
      Decay_Conts=f.GatherDecay(Nuclide_Names)
      f.np.save('Saved_Variables/Decay_Consts', Decay_Conts)
   else:
      Decay_Conts=f.np.load('Saved_Variables/Decay_Consts.npy')
  #load up A and b with phi
   #f.os.system("rm Saved_Variables/Amatrix.npy")  #To start over
   #f.os.system("rm Saved_Variables/n0vector.npy") #to start over
   if not f.os.path.isfile('Saved_Variables/Amatrix.npy'):
      #A is in terms of years
      A, n0=f.MakeAb (phi, Nuclides, Nuclide_Names, Decay_Conts)
      f.np.save('Saved_Variables/Amatrix',A)
      f.np.save('Saved_Variables/n0vector',n0)
   else:
      A=f.np.load('Saved_Variables/Amatrix.npy')
      n0=f.np.load('Saved_Variables/n0vector.npy')
  phi=0
   #load up A and b without phi (decay)
   #f.os.system("rm Saved_Variables/Amatrixdecay.npy") #To start over
  #f.os.system("rm Saved_Variables/n0vectordecay.npy") #to start over
   if not f.os.path.isfile('Saved_Variables/Amatrixdecay.npy'):
      #A is in terms of years
      Adecay, n0=f.MakeAb (phi, Nuclides, Nuclide_Names, Decay_Conts)
      f.np.save('Saved_Variables/Amatrixdecay', A)
      f.np.save('Saved_Variables/n0vectordecay', n0)
80
      Adecay=f.np.load('Saved_Variables/Amatrixdecay.npy')
      n0decay=f.np.load('Saved_Variables/n0vectordecay.npy')
  if not A.shape[0] == A.shape[1] or not A.shape[0] == len(n0):
      print("A is not a square matrix")
      quit()
  print("A is complete")
  ########### Initialize Time ############################
   t=1.09514; #years
  Nt=100;
          #Number of Time Steps
  dt=t/Nt;
  Time=f.np.linspace(dt,t,Nt) #Time steps
  tDecay=0; #No time of decay
  NtDecay=1; #Number of Decay Time Steps
  dtDecay=tDecay/NtDecay
  TimeDecay=f.np.linspace(t+dtDecay,t+tDecay,NtDecay)
```

```
############## And Time How long it takes ######################
  ############## And Plot at Each Time Step ##################
  110
  #Matrix Exp - Unstable
  #Current_Time=time.time()
  #maxits=20
  # nt_Mat=f.MatExp(A, n0, t, maxits)
# Mat_Time=time.time()-Current_Time
  #nt_Mat=n0.copy();TIMEOLD=0
  #for TIME in Time[1:len(Time)]: #Unstable
     #nt_Mat=f.MatExp(A, n0, TIME, maxits) No Converge
     #Step through
     #nt_Mat=f.MatExp(A, nt_Mat, TIME-TIMEOLD, maxits)
120
     #TIMEOLD=TIME.copy()
  #nt_Mat=f.RationalApprox(A, n0, t, maxits) #one Step
  #Mat_Time=time.time()-Current_Time
  Current_Time=time.time()
  File=f.PrepFile('Data/BackEuler.csv',n0,Nuclide_Names,Atom_Mass,Decay_Conts) #Prep File
  nt_Back=n0.copy();
  #Irradiation Time
  for TIME in Time:
     nt_Back=f.BackEuler(A, nt_Back, dt)
    File.write(str(TIME)+","+f.ListToStr(nt_Back))
  #Decay Time
  #for TIME in TimeDecay:
  # nt_Back=f.BackEuler(Adecay, nt_Back, dtDecay)
    File.write(str(TIME)+","+f.ListToStr(nt_Back))
140
  File.close()
  Back_Time=time.time()-Current_Time
  #Irradiation Time
150 | Current_Time=time.time()
  N=10;
  Method="Best" #Parabola, Cotangent, Hyperbola, Best
  File=f.PrepFile("Data/"+Method+".csv", n0, Nuclide_Names, Atom_Mass, Decay_Conts)
  ck, zk=f.RationalPrep(N,Method); nt_Rational=n0.copy(); TIMEOLD=0
  #for TIME in Time:
  # nt_Rational=f.RationalApprox(A,n0,TIME,N,ck,zk)
```

```
File.write(str(TIME) +", "+f.ListToStr(nt_Rational))
      #Step through
      #nt_Rational=f.RationalApprox(A, nt_Rational, TIME-TIMEOLD, N, ck, zk)
      #TIMEOLD=TIME.copy()
   nt_Rational=f.RationalApprox(A,n0,t,N,ck,zk) #one Step
   File.write(str(TIME) +", "+f.ListToStr(nt_Rational))
   #Non Irradiation Time
  n0decay=nt_Rational
   #for TIME in TimeDecay:
      nt_Rational=f.RationalApprox(Adecay, n0decay, TIME-t, N, ck, zk)
       File.write(str(TIME) +", "+f.ListToStr(nt_Rational))
170
   File.close()
   Rational_Time=time.time()-Current_Time
   ################# Plot Solution ##########################
   dfBack = f.pd.read_csv('Data/BackEuler.csv',index_col=False)
   dfRational = f.pd.read_csv("Data/"+Method+".csv",index_col=False)
   # #Plot group 1 dudes Back Euler method
# f.plot(dfBack, PlottingG1, 'BackEulerG1', NumOfPoints)
   # f.plot(dfRational,PlottingG1,Method+'G1',NumOfPoints)
   # #Plot group 2 dudes Rational method
   # f.plot(dfBack,PlottingG2,'BackEulerG2',NumOfPoints)
  # f.plot(dfRational, PlottingG2, Method+'G2', NumOfPoints)
   #Plot group 1 dudes, compare both methods
   #Name='BackEuler_'+Method+"_G1"
   #f.plots2(dfBack, dfRational, CompareG1, Name,
          NumOfPoints, 'BackEuler', Method)
   #Plot group 2 dudes, compare both methods
   #Name='BackEuler_'+Method+"_G2"
   #f.plots2(dfBack, dfRational, CompareG2, Name,
          NumOfPoints, 'BackEuler', Method)
200
   #f.Print("Matrix Exp", "H3", nt_Mat, Mat_Time)
   f.Print("Backward Euler", "541360", nt_Back, Back_Time, Nuclides, Atom_Mass, Nuclide_Names)
   f.Print("Rational Approx ","541360",nt_Rational,Rational_Time,Nuclides,Atom_Mass,Nuclide_Names)
```

Listing 8: Functions holder

```
#!/usr/bin/env python3
  ################### Import packages ##############################
  import sys
  import numpy as np
  import scipy.sparse as sparse
 import scipy.sparse.linalg as spla
  import scipy.linalg as scil
  import scipy.special as sps
  import matplotlib.pyplot as plt
  plt.rcParams["font.family"] = "monospace"
15 import matplotlib
  matplotlib.rc('text',usetex=True)
  matplotlib.rcParams['text.latex.preamble']=[r"\usepackage{amsmath}"]
  import random as rn
  import matplotlib.mlab as mlab
 import copy
  import os
  import pandas as pd
  import os.path
  ####################### Variables ##########################
  # Basic information
30 FigureSize = (11, 6)
  FigureSize = (11, 6) # Dimensions of the figure
TypeOfFamily='monospace' # This sets the type of font for text
  font = {'family' : TypeOfFamily} # This sets the type of font for text
  LegendFontSize = 12
  Lfont = {'family' : TypeOfFamily} # This sets up legend font
  Lfont['size']=LegendFontSize
  Title = ''
  TitleFontSize = 22
  TitleFontWeight = "bold" # "bold" or "normal"
  #Xlabel='E (eV)' # X label
  XFontSize=18 # X label font size
  XFontWeight="normal" # "bold" or "normal"
  XScale="linear" # 'linear' or 'log'
  YFontSize=18
                            # Y label font size
  YFontWeight="normal" # "bold" or "normal"
```

```
YScale="linear"
                        # 'linear' or 'log'
Check=0
Colors=["aqua", "gray", "red", "blue", "black",
            "green", "magenta", "indigo", "lime", "peru", "steelblue",
            "darkorange", "salmon", "yellow", "lime", "black"]
# If you want to highlight a specific item
# set its alpha value =1 and all others to 0.4
# You can also change the MarkSize (or just use the highlight option below)
# Can change all these to "." or "" for nothing "x" isn't that good
MarkerType=["8", "s", "p", "D", "*", "H", "h", "d", "^", ">"]
# LineStyles=["solid", "dashed", "dash_dot", "dotted", "."]
LineStyles=["solid"]
SquishGraph = 0.75
BBOXX = 1.24
BBOXY = 0.5
             # Set legend on right side of graph
NumberOfLegendColumns=1
#Xlabel='Time [days]'
Xlabel='Time [years]'
Ylabel="Mass $\\left[\\frac{g}{\\text{tHM}}\\right]$"
Na=6.0221409E23
############## Lists
                                ######################
def Returnfloat(string):
   11 11 11
   string has format 238.023249814(23)
         or format [15.99903-15.99977]
         or format 235.04+/-0.0000019
   Returns just the number, no uncertainties
   if "(" in string:
      Number=str(string.split('(')[0])
      LastErrorNumber=str(string.split("(")[1].replace(")",""))
```

```
NumberOfZeros=len(Number.split(".")[1])-len(LastErrorNumber)
            Error="0."
            for i in range(0, NumberOfZeros):
                Error=Error+"0"
            Error=Error+LastErrorNumber
105
        elif "[" in string:
            FirstNum=float(string.split('-')[0].replace("[",''))
            SecondNum=float(string.split('-')[1].replace(']',''))
            Number=str((FirstNum+SecondNum)/2)
            Error=str(float(Number)-FirstNum)
110
        elif "+/-" in string:
            Number=string.split("+/-")[0]
            Error=string.split("+/-")[1]
115
        return (float (Number))
    def Isotopes(ToAdd):
      This function will create a dictionary 'Nuclides'
120
      with nuclides found in tape9.inp, excluding activation isotopes
     Nuclides={}
     Nuclide_Names=()
125
     with open ('Data/tape9.inp') as f:
       TAPE9Content=f.readlines()
      count=0
      for line in TAPE9Content:
130
       hold=line.split()
        #No activation products or the -1 between libraries
        if not '1' in hold[0] and not '601' in hold[0] and "-" not in hold[0]:
          #No repeats
                                       #No decimals
                                                                  #No text
135
          if hold[1] not in Nuclides and "." not in hold[1] and hold[1].isdigit():
            #Filter out lower mass isotopes
            if len(hold[1]) == 6:
              Nuclides[hold[1]]=count
              Nuclide_Names=Nuclide_Names+(hold[1],)
140
              count=count+1
      for isotope in ToAdd:
          Nuclides[isotope]=count
          Nuclide_Names=Nuclide_Names+(isotope,)
          count=count+1
      return (Nuclides, Nuclide_Names)
   def GatherDecay(Nuclide_Names):
      11 11 11
      This function will create an array 'Decay_Consts'
      that has all the half-life information for our system.
```

```
.....
155
      Decay_Consts = np.zeros(len(Nuclide_Names))
      with open ('Data/tape9.inp') as f:
        TAPE9Content=f.readlines()
160
      for i in range(0,len(Nuclide_Names)):
          Nuclide=Nuclide Names[i]
          for line in TAPE9Content:
              hold=line.split()
165
              #Look for half life information, and decay type information
              #No activation products or the -1 between libraries
              if hold[0] == '2' or hold[0] == '3':
                   if hold[1] == Nuclide:
170
                       Thalf=float (hold[3])
                       if hold[2] == '1': #seconds
                           const=np.log(2)/Thalf
                       elif hold[2] == '2': #minutes
                           const=np.log(2)/(Thalf*60)
175
                       elif hold[2] == '3': #hours
                           const=np.log(2) / (Thalf * 60 * 60)
                       elif hold[2]=='4': #days
                           const=np.log(2)/(Thalf*60*60*24)
                       elif hold[2] == '5': #years
                           const=np.log(2)/(Thalf*60*60*24*365.25)
                       elif hold[2] == '6': #Stable
                           const=-1
                       elif hold[2]=='7':
                           const=np.log(2) / (Thalf*60*60*24*365.25*10**3)
                       elif hold[2] == '8':
                           const=np.log(2)/(Thalf*60*60*24*365.25*10**6)
                       elif hold[2] == '9':
                           const=np.log(2)/(Thalf*60*60*24*365.25*10**9)
                       else:
190
                           print("could not find a proper halflife")
                           print(line)
                           quit()
                       Decay_Consts[i]=const
195
      return (Decay_Consts)
   def FindAtomicMass(df,proton,Isotope):
200
        This function will take in a dataset 'df' look through the
        'df.Protons' column and find the column that matches with
        'proton'. If the row that contains 'proton' also contains
        'Isotope' in the 'df. Isotope' column, then the value stored
205
        in 'df.Relative_Atomic_Mass' is reported for that row.
```

```
Because the proton numbering scheme can have a format
        '10' for hydrogen and '10' for neon (following MCNP ZAID
        naming conventions) if we don't find a value with the whole
        string of 'proton' then the program looks through the first
210
        element of string and tries to match that 'proton[0]'
        If no matches are found, and error is thrown out.
        df = dataset with columns 'Protons' 'Isotopes' and
        'Relative_Atomic_Mass'. Dataset created with pandas
215
        proton = string with proton number (follow MCNP zaid format)
        Isotope = string with isotope number (just put the atomic mass
        do not follow MCNP format - different for few cases)
220
        #print (df)
        for i in range(0,len(df.Protons)):
            dfPro=str(df.Protons[i])
            if proton==dfPro:
225
                dfIso=str(df.Isotope[i])
                if Isotope==dfIso:
                    Mass=df.Relative_Atomic_Mass[i]
                    break
        try:
230
            Mass
        except NameError:
            for i in range(0,len(df.Protons)):
                dfPro=str(df.Protons[i])
                if proton[0] == dfPro:
                    dfIso=str(df.Isotope[i])
                     if Isotope==dfIso:
                        Mass=df.Relative_Atomic_Mass[i]
                         break
        try:
            Mass
        except NameError:
            print("Could not find atomic mass for proton = "\
                  +proton+" and for Isotope = "+Isotope)
            Mass='10000.09(23)'
245
        Mass=Returnfloat (Mass)
        return (Mass)
    def GatherMasses(df, Nuclides):
      11 11 11
250
     Make numpy array of masses which correspond to isotopes in Nuclides
      Nuclides is a dictionary. Where each key is a zaid number
     Atom_Mass=np.zeros(len(Nuclides))
255
      for key, value in Nuclides.items():
        if not len(key) == 6:
          print("Did not filter out the lower mass isotopes, quitting")
          quit()
```

```
proton=key[0:2]
260
       if key[2] == "0":
           Isotope=key[3:5]
       elif key[4] == "0" and float (proton) \star 3<100:
           Isotope=key[2:4]
       elif key[2]!="0":
265
           Isotope=key[2:5]
       Mass=FindAtomicMass(df,proton,Isotope)
       if Mass>9000:
           print("Mass is over 9,000!!!")
           print (key)
           quit()
       Atom_Mass[value] = Mass
     return (Atom_Mass)
275
    ####### Functions for solving the system Ax=b (kind of) #######
    def MatExp(A, n0, t, maxits, tolerance=1e-12, LOUD=False):
     converged = False
     sum_old=n0.copy()*0
285
     while not (converged):
       if m==0:
           APowerm=np.identity(A.shape[0])
           Factorial=1
290
       else:
           APowerm=np.dot(APowerm, A)
           Factorial=Factorial*m
       Sum=sum_old+(1/Factorial)*np.dot((APowerm)*(t**m),n0)
295
       #Avoid dividing by zero
       if sum(Sum) ==0: m+=1; sum_old=Sum.copy(); continue
       change = np.linalg.norm(Sum-sum_old)/np.linalg.norm(Sum)
       converged = (change < tolerance) or (m > maxits)
       if (LOUD>0) or (converged and LOUD<0):</pre>
         print("Iteration", m, ": Relative Change =", change)
       if (m > maxits):
305
         print("Warning: Source Iteration did not converge: "+\
               " m : "+str(m)+", Diff : %.2e" % change)
       #Prepare for next iteration
       m += 1
       sum_old = Sum.copy()
310
     return (Sum)
```

```
def BackEuler(A, no, dt):
        I=np.identity(A.shape[0])
        return(np.dot(np.linalq.inv(I-A*dt),no))
315
    def DeterminePolesNResidues(n):
        This program takes the algorithm from the reference
        and converts to a python script... I know its janky
320
        but it works
        11 11 11
        def Append(List1, List2):
            for item in List2:
                for item2 in item:
                    List1=np.append(List1, item2)
            return (List1)
        def absG(List):
            List2=[]
            for item in List:
                if abs(item)>1:
                    List2.append(item)
            return(List2)
        #function [zk, ck] = cf(n);
                                             # no of Cheb coeffs
        K = 75;
335
                                             # no of pts for FFT
        nf = 1024;
        #Roots correct?
        roots=np.arange(0,nf,1)/nf
        #w = np.exp(2i*pi*(0:nf-1)/nf);
                                             # roots of unity
340
        w=np.exp(2j*np.pi*roots)
        t = np.real(w);
                                             # Cheb pts (twice over)
        scl = 9;
                                             # scale factor for stability
        \#F = np.exp(scl*(t-1)./(t+1+1e-16)); \# exp(x) transpl. to [-1,1]
        F = np.exp(scl*(t-1)/(t+1+1e-16)); # exp(x) transpl. to [-1,1]
345
        c = np.real(np.fft.fft(F))/nf;
                                             # Cheb coeffs of F
        index=reversed(np.arange(1,K+2,1))
        partofc=[]
        for i in index:
            partofc.append(c[i-1])
        #f = np.polyval(c(K+1:-1:1), w);
                                            # analytic part f of F
        f = np.polyval(partofc, w);  # analytic part f of F
        \#[U,S,V] = svd(hankel(c(2:K+1))); \#SVD of Hankel matrix
        hankie=scil.hankel(c[1:K+1])
        U,S,V=np.linalg.svd(hankie,full_matrices=False)
        #s = S(n+1, n+1);
                                              # singular value
        s=S[n]
        \#u = U(K:-1:1,n+1); v = V(:,n+1); \# singular vector
        u = []
        index=reversed(np.arange(0,K,1))
        for i in index:
            u.append(U[i,n])
        \#v=np.array(V[:,n].copy())
```

```
v=np.array(V[n,:].copy())
        \#zz = zeros(1, nf-K);
                                              # zeros for padding
        zz=np.zeros([1,nf-K])
        \#b = fft([u zz])./fft([v zz]);
                                              # finite Blaschke product
        b=np.fft.fft(Append(u,zz))/np.fft.fft(Append(v,zz))
370
        \#rt = f-s*w.^K.*b;
                                              # extended function r-tilde
        rt=f-s*(w**K)*b;
        #rtc = real(fft(rt))/nf;
                                              # its Laurent coeffs
        rtc=np.real(np.fft.fft(rt))/nf;
        \#zr = roots(v); qk = zr(abs(zr)>1); \# poles
375
        zr=np.roots(v); qk=np.array(absG(zr));
        #qc = poly(qk);
                                              # coeffs of denominator
        qc=np.poly(qk);
                                              # numerator
        #pt = rt.*polyval(qc,w);
        pt=rt*np.polyval(qc,w);
380
        #ptc = real(fft(pt)/nf);
                                              # coeffs of numerator
        ptc=np.real(np.fft.fft(pt)/nf);
        \#ptc = ptc(n+1:-1:1); ck = 0*qk;
        index=reversed(np.arange(0,n+1,1))
        ptc2=[]
        for i in index: #Can I just reversed ptc?
            ptc2.append(ptc[i])
        ptc=ptc2.copy()
        ck=0*ak
        \#N+1?
        #for k = 1:n
                                  # calculate residues
             q = qk(k); q2 = poly(qk(qk\sim=q));
             ck(k) = polyval(ptc,q)/polyval(q2,q);
        for k in range (0, n):
395
            if len(qk) == k:
                print("we are short a qk")
                continue
            q=qk[k];
            q2 = [];
            for item in qk:
400
                if not q==item:
                    q2.append(item)
            q2=np.poly(q2);
            ck[k]=np.polyval(ptc,q)/np.polyval(q2,q)
        \#zk = scl*(qk-1).^2./(qk+1).^2; # poles in z-plane
405
        zk=scl*((qk-1)**2)/((qk+1)**2)
        \#ck = 4*ck.*zk./(qk.^2-1);
                                             # residues in z-plane
        ck=4*ck*zk/(qk**2-1)
        #Cut down ck and zk to half the original points
        ck2=[];zk2=[]
        for i in range(0,len(ck)):
            if i % 2 == 0:
                ck2=np.append(ck2,ck[i])
                zk2=np.append(zk2,zk[i])
        return (ck2, zk2)
    def RationalPrep(N, Phi):
```

```
"""Calculate constants for a rational approximation
420
        Inputs:
        N:
                         Number of Quadrature points
        Phi:
                         'Parabola',
                         'Cotangent', or
                         'Hyperbola' (shape of Phi)
425
        Outputs:
                        First set of constants for approximation
        ck:
        zk:
                        Second set of constants for approximation
        theta=np.pi*np.arange(1,N,2)/N
430
        if Phi=='Parabola':
            zk=N*(0.1309-0.1194*theta**2+0.2500j*theta)
            w=N*(-2*0.1194*theta+0.2500j)
        elif Phi=='Cotangent':
            cot=1/np.tan(0.6407*theta)
435
            ncsc=-0.6407/(np.sin(0.6407*theta)**2)
            zk=N*(0.5017*theta*cot-0.6122+0.2645j*theta)
            w=N*(0.2645j+0.5017*cot+0.5017*theta*ncsc)
        elif Phi=='Hyperbola':
            zk=2.246*N*(1-np.sin(1.1721-0.3443j*theta))
440
            w=2.246*N*(0.3443j*np.cos(1.1721-0.3443j*theta))
        elif Phi=='Best':
            ck, zk=DeterminePolesNResidues(N)
            return (ck, zk)
        else:
            print("Did not pick proper rational approximation dude")
            print("Quiting now")
            quit()
        ck=1.0j/N*np.exp(zk)*w
        return (ck, zk)
    def RationalApprox(A, n0, t, N, ck, zk, tol=1e-12, maxits=2000):
        Calculate the rational approximation solution for n(t)
455
        Inputs:
       A:
                   Matrix with system to be solved
       n0:
                   initial conditions of the system
        t:
                    time at which solution is determined
                   Number of quadrature points (should be less than 20)
       N:
460
        ck:
                   constants for quadrature solution
                    constants for quadrature solution
        zk:
                    Tolerence for convergence for GMRES
        tol:
       maxits:
                   Maximium iterations for GMRES
       Outputs:
465
        nt:
                    Solution at time t
       nt=np.zeros(len(n0))
        for k in range(int(N/2)):
            if len(n0)>1:
470
                \#phi, code=spla.gmres(zk[k]*sparse.identity(len(n0))-A*t,n0,
```

```
tol=tol, maxiter=maxits)
              phi=np.dot(np.linalg.inv(zk[k]*np.identity(len(n0))-A*t),
                        n0)
              #if (code):
475
                  print(code)
          else:
              phi = (zk[k]-A*t)**(-1)*n0
          nt=nt-2*np.real(ck[k]*phi)
       return (nt)
480
   485
   def reduceList(List, N):
       List2=[List[0]]
       Div=int(len(List)/N)
       for i in range(1,len(List)-1):
          if i % Div == 0:
490
              List2.append(List[i])
       List2.append(List[-1])
       return (List2)
   def PlotPoints(Ntot, Nplot):
       t=1
   def loop_values(list1, index):
       This function will loop through values in list even if
       outside range (in the positive sense not negative)
500
       11 11 11
       while True:
          try:
              list1[index]
              break
          except IndexError:
              index=index-len(list1)
       return(list1[index])
   def Legend(ax):
      handles, labels = ax.get_legend_handles_labels()
       ax.legend(handles, labels, loc='best',
                fontsize=LegendFontSize,prop=font)
       return (ax)
515
   # def Legend(ax):
            handles, labels=ax.get_legend_handles_labels()
   #
            box=ax.get_position()
            ax.set_position([box.x0, box.y0, box.width*SquishGraph,
                           box.height])
520
            ax.legend(handles, labels, loc='center',
   #
                     bbox_to_anchor=(BBOXX,BBOXY),
   #
                     fontsize=LegendFontSize,prop=font,
                     ncol=NumberOfLegendColumns)
```

```
return(ax)
525
    def InList(item2,List):
      TF=False
      for item1 in List:
        if item1 == item2:
530
          TF=True
      if not TF:
        print("Invalid selection for plotting")
        print("Shuting down")
        quit()
    def plot(df,Plotting,Name,NumOfPoints):
        #Plot In grams
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
540
        List=list(df.columns.values)
        x=df[List[0]].values[2:]
        Check=0
545
        for Item in Plotting:
          InList(Item,List) #Check if we have the isotope
          y=((df[Item].values[2:])/Na)*df[Item].values[0]
          if len(x)>NumOfPoints:
            x=reduceList(x,NumOfPoints)
550
            y=reduceList(y,NumOfPoints)
          ax.plot(x,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item)
          Check=Check+1
560
        #Log or linear scale?
        ax.set_xscale(XScale)
        ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
        #Set X and y labels
        ax.set_xlabel(Xlabel,
570
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        ax.set_ylabel(Ylabel,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
575
        Legend (ax)
```

```
plt.savefig("Plots/"+Name+'_grams.pdf')
        #Plot in Bq #################################
580
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
        List=list(df.columns.values)
585
        x=df[List[0]].values[2:]
        Check=0; Sum=np.zeros(len(x))
        for Item in Plotting:
          InList(Item, List) #Check if we have the isotope
          y=((df[Item].values[2:]))*df[Item].values[1]
          Sum=Sum+y
          if len(x)>NumOfPoints:
            xP=reduceList(x, NumOfPoints)
            y=reduceList(y,NumOfPoints)
          ax.plot(xP,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
600
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item)
          Check=Check+1
        if len(x) > NumOfPoints:
          Sum=reduceList(Sum, NumOfPoints)
605
        ax.plot(xP,Sum,
                linestyle=loop_values(LineStyles, Check),
                marker=loop_values (MarkerType, Check),
                color=loop_values(Colors, Check),
                markersize=loop_values(MarkSize,Check),
610
                alpha=loop_values(Alpha_Value, Check),
                label="Sum")
        #Log or linear scale?
        ax.set_xscale(XScale)
615
        if sum(Sum) == 0:
          ax.set_yscale('linear')
        else:
          ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        YlabelBq="Activity $\\left[\\frac{Bq}{\\text{tHM}}}\\right]$"
        ax.set_ylabel(YlabelBq,
                       fontsize=YFontSize,
630
                       fontweight=YFontWeight,
```

```
fontdict=font)
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Bq.pdf')
635
    def plots2(df, df2, Plotting, Name, NumOfPoints, Method1, Method2):
        #Plot in grams
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
640
        List=list(df.columns.values)
        x=df[List[0]].values[2:]
        x2=df2[List[0]].values[2:]
        Check=0
645
        for Item in Plotting:
          InList(Item,List) #Check if we have the isotope
          y=((df[Item].values[2:])/Na)*df[Item].values[0]
          y2=((df2[Item].values[2:])/Na)*df2[Item].values[0]
          if len(x) > NumOfPoints:
            x=reduceList(x,NumOfPoints)
            y=reduceList(y,NumOfPoints)
          if len(x2)>NumOfPoints:
            x2=reduceList(x2, NumOfPoints)
            y2=reduceList(y2, NumOfPoints)
          ax.plot(x,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
660
                  markersize=loop_values(MarkSize,Check) *1.5,
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method1)
          Check=Check+1
          ax.plot(x2,y2,
                  linestyle=loop_values(LineStyles, Check),
665
                  marker=loop_values (MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method2)
670
          Check=Check+1
        #Log or linear scale?
        ax.set_xscale(XScale)
        ax.set_yscale(YScale)
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
        #Set X and y labels
        ax.set_xlabel(Xlabel,
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
```

```
ax.set_ylabel(Ylabel,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Grams.pdf')
690
        #Plot in Ba
        fig=plt.figure(figsize=FigureSize)
        ax=fig.add_subplot(111)
695
        List=list(df.columns.values)
        x=df[List[0]].values[2:]
        x2=df2[List[0]].values[2:]
700
        Check=0; Sum=np.zeros(len(x)); Sum2=np.zeros(len(x2))
        for Item in Plotting:
          InList(Item, List) #Check if we have the isotope
          y=((df[Item].values[2:]))*df[Item].values[1]
          y2=((df2[Item].values[2:]))*df2[Item].values[1]
705
          Sum=Sum+y
          Sum2=Sum2+v2
          if len(x) > NumOfPoints:
            xP=reduceList(x,NumOfPoints)
            y=reduceList(y,NumOfPoints)
          else:
            xP=x.copy()
          if len(x2) > NumOfPoints:
            xP2=redcueList(x2, NumOfPoints)
            y2=reduceList(y2, NumOfPoints)
715
          else:
            xP2=x2.copy()
          ax.plot(xP,y,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
720
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check) *1.5,
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method1)
          Check=Check+1
725
          ax.plot(xP2,y2,
                  linestyle=loop_values(LineStyles, Check),
                  marker=loop_values(MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check),
730
                  alpha=loop_values(Alpha_Value, Check),
                  label=Item+" "+Method2)
          Check=Check+1
        if len(x) > NumOfPoints:
          Sum=reduceList(Sum, NumOfPoints)
735
        if len(x2)>NumOfPoints:
```

```
Sum2=reduceList(Sum2,NumOfPoints)
        ax.plot(xP,Sum,
                  linestyle=loop_values(LineStyles, Check),
740
                  marker=loop_values (MarkerType, Check),
                  color=loop_values(Colors, Check),
                  markersize=loop_values(MarkSize,Check) *1.5,
                  alpha=loop_values(Alpha_Value, Check),
                  label="Sum "+Method1)
        Check=Check+1
745
        ax.plot(xP2,Sum2,
                linestyle=loop_values(LineStyles, Check),
                marker=loop_values(MarkerType, Check),
                color=loop_values(Colors, Check),
                markersize=loop_values(MarkSize,Check),
750
                alpha=loop_values(Alpha_Value, Check),
                label="Sum "+Method2)
        #Log or linear scale?
        ax.set_xscale(XScale)
755
        if sum(Sum) == 0:
          ax.set_yscale('linear')
          ax.set_yscale(YScale)
760
        #Set Title
        fig.suptitle(Title, fontsize=TitleFontSize,
                     fontweight=TitleFontWeight, fontdict=font, ha='center')
        #Set X and y labels
        ax.set_xlabel(Xlabel,
765
                       fontsize=XFontSize, fontweight=XFontWeight,
                       fontdict=font)
        YlabelBq="Activity $\\left[\\frac{Bq}{\\text{tHM}}\\right]$"
        ax.set_ylabel(YlabelBq,
                       fontsize=YFontSize,
                       fontweight=YFontWeight,
                       fontdict=font)
        Legend (ax)
        plt.savefig("Plots/"+Name+'_Bq.pdf')
775
   def ListToStr(List):
      Str=''
      for i in range(0,len(List)):
        if not i==len(List)-1:
780
          Str=Str+str(List[i])+","
        else:
          Str=Str+str(List[i])+"\n"
      return (Str)
785
   def PrepFile(Name, n0, nuclide_names, atom_mass, decay_consts):
     File=open(Name,'w')
      File.write("Mass then Time (d),"+','.join(nuclide_names)+'\n')
     File.write("Masses,"+ListToStr(atom_mass)) #New line already included
```

```
File.write("DecayConts,"+ListToStr(decay_consts))
     File.write("0,"+ListToStr(n0))
     return (File)
   def Print(Method, nuclide, Results, Time, nuclides, atom_mass, nuclide_names):
     Index=nuclides[nuclide]
795
     MassConversion=atom_mass[Index]/Na
     string="Isotope "+nuclide_names[Index]+", Mass (g) = "
     Mass=Results[Index] *MassConversion
     Mass="%.4e" % Mass
     print (Method+" :", string, Mass, "Time=%.2f" % Time)
   def FindFissionXSection(Fissile_Isotopes):
      FissionXSections=np.zeros(len(Fissile_Isotopes))
810
       with open ('Data/tape9.inp') as f: #Save all X-section data to variable
          TAPE9Content=f.readlines()
       for i in range(0,len(Fissile_Isotopes)): #Loop through fissile isos
          parent=Fissile_Isotopes[i]
815
          for line in TAPE9Content: #Loop through x-section data
              hold=line.split()
              if '602' == hold[0] and hold[1] == parent: #Find x-section
820
                  FissionXSections[i]=hold[5]
                  break
       return (FissionXSections)
   def CalMevPerFiss(Fissile_Isotopes):
825
       11 11 11
       Given a list of fissile isotopes
       return a list of MeV/fission numbers
       calculated from an equation
       MeVperFission=np.zeros(len(Fissile_Isotopes))
       for i in range(0,len(Fissile_Isotopes)):
          isotope=Fissile_Isotopes[i]
          proton=isotope[0:2]
          if isotope[2] == "0":
835
              Anum=isotope[3:5]
           elif isotope[4] == "0" and float(proton) *3<100:</pre>
              Anum=isotope[2:4]
          elif isotope[2]!="0":
              Anum=isotope[2:5]
840
              print("Missed logic in finding A number")
```

```
quit()
          if int(Anum) < int(proton):</pre>
              print("Something is wrong, more protons than neutrons")
845
              print("Proton: ",proton, "A: ",Anum, "ZAID : ",isotope)
          MeVperFission[i]=1.29927*(10**-3)*(float(proton)**2)*(float(Anum)**0.5)+33.12
       return (MeVperFission)
850
   def Calculatephi (FissionXSections, MeVperFission, n0, Power, Fissile_Isotopes, Nuclides)
       Sim=0
       for i in range(0,len(FissionXSections)):
          sigma=FissionXSections[i] *10 **-24
855
          E=MeVperFission[i]
          N=n0[Nuclides[Fissile_Isotopes[i]]]
          Sum=Sum+sigma*E*N
       phi=(6.2414959617521E18*Power)/Sum
       return (phi)
   def Makeb(Nuclides):
       b = np.zeros(len(Nuclides))
       b[Nuclides['922340']] = 6.94741E23
865
       b[Nuclides['922350']] = 7.6864E25
       b[Nuclides['922380']] = 2.4532E27
       return (b)
870
   875
   class DecayClass:
       def __init__(self):
                   = 0. # 'The fraction of negatron beta decay transitions that results in
                           # in the daughter nuclide being in a relatively long-lived state'
                           # I think this should read the fraction of all decay events which are.
880
          self.IDFBX = '' # ZAID for daughter for FBX
          self.FPEC = 0. # Fraction of all decay events which are positron or EC
          self.IDFPEC = '' # ZAID for daughter for FPEC
          self.FPECX = 0. # Fraction of all EC or positron decays which result in excited state
          self.IDFPECX= '' # ZAID for daughter for FPECX
                    = 0. # Fraction of all decay events which are alpha
          self.FA
                     = ''  # ZAID for daughter for alpha
          self.IDFA
          self.FIT = 0. # 'fraction of all the decay events of an excited nuclear state
                           # which result in the production of the ground state of the same nucli
          self.IDFIT = '' # ZAID for daughter of FIT
          self.FSF
                    = 0. # Fraction that decay events that are spontaneous fission
          # No daughter listed, will loop through all elements for yields
                     = 0. # Fraction of all decay events that are beta + neutron decays
          self.FN
                     = '' # ZAID for daughter for FN
          self.IDFN
          #Note: Negatron beta decay = 1 - FBX - FPEC - FA - FIT - FSF - FN
```

```
= 0. # Fraction of all decay events which are beta
                        = ''
            self.IDFB
    def FindPotentialMatch(List, protons, A, Fraction, Excited, LOUD=False):
        if (Fraction<0 or Fraction>1):
            print("Fraction of decays is too low or high: ",Fraction)
            print("Inquire further")
            quit()
        if Fraction>0:
905
            for item in List:
                if protons in item[0:2] and A in item[:-1] and item[-1]==Excited:
                    Toreturn=item
        else:
            Toreturn=''
910
        try: #To make sure its defined
            Toreturn
        except NameError:
            if LOUD:
915
                print ("Could not find daughter in list of isotopes when expecting one")
                print("Looking for Protons : ",protons," Total Nucleons : ",A," Fraction)
                #print("Close items are")
                #for item in List:
                     if protons in item[:-1] and A in item[:-1]:
                         print(item)
            if (Fraction<2):</pre>
                if LOUD:
                    print("Assuming it doesn't matter, will let slide")
                Toreturn=''
925
            else:
                quit()
        return (Toreturn)
930
    def DecayInfo(Nuclide_Names, parent, Lambda, proton, A):
        This function will store and return decay information from
        the tape9.inp file
        11 11 11
935
        Info=DecayClass()
        with open ('Data/tape9.inp') as f:
            TAPE9Content=f.readlines()
940
        Found=False
        for line in TAPE9Content:
            hold=line.split()
            #Looking at second line in each library (put this if statement above below)
            if Found:
                #Spontaneous fission
```

```
Info.FSF=float(hold[1])
                 #Beta plus neutron
                 Info.FN=float(hold[2])
                 Info.IDFN=FindPotentialMatch(Nuclide_Names, str(int(proton)+1), str(int(A)-1), Info.FN,
                 break
             #Looking for fission product and actinide decay information
955
             #The libraries we are looking through for this information are '2' and '3'
             if ('2' == hold[0] \text{ or } '3' == hold[0]) and hold[1] == parent:
                 Found=True
                 #Beta minus to excited
960
                 Info.FBX=float(hold[4])
                 Info.IDFBX=FindPotentialMatch(Nuclide_Names, str(int(proton)+1), A, Info.FBX, '1')
                 #positron or EC
                 Info.FPEC=float(hold[5]) #Total positron or EC
                 Info.FPECX=float(hold[6]) #percent of above to excited
965
                 if Info.FPECX < 1:</pre>
                     Info.IDFPEC=FindPotentialMatch(Nuclide_Names, str(int(proton)-1), A, Info.FPEC, '0')
                 Info.IDFPECX=FindPotentialMatch(Nuclide_Names, str(int(proton)-1), A, Info.FPECX, '1')
                 Info.FA=float(hold[7])
                                            #Fraction of events that are alpha
                 Info.IDFA=FindPotentialMatch(Nuclide_Names, str(int(proton)-2), str(int(A)-4), Info.FA,
970
                 #Excited state to ground state
                 Info.FIT=float(hold[8])
                 Info.IDFIT=FindPotentialMatch(Nuclide_Names,proton,A,Info.FIT,'0')
         #Calculate the beta
        Info.FB=1-Info.FBX-Info.FPEC-Info.FA-Info.FIT-Info.FSF-Info.FN
        if (Info.FB<0 or Info.FB>1):
             #print("Fraction of beta decays is too low or high: ", Info.FB)
             if abs(Info.FB) < 7e-4 or abs(Info.FB-1) < 7e-4:
                 #print("But I'll let it slide and set to zero")
                 Info.FB=0
             else:
                 print("I can't let this slide, not small enough")
                 print("Beta excited", Info.FBX)
985
                 print("EC", Info.FPEC)
                 print("Alpha decay ground", Info.FA)
                 print("Excited to ground", Info.FIT)
                 print("Spontaneous fission", Info.FSF)
                 print("Beta Plus neutron", Info.FN)
990
                 quit()
        if Lambda>0:
             Info.IDFB=FindPotentialMatch(Nuclide_Names, str(int(proton)+1), A, Info.FB, '0'|)
995
        #Make sure all decay fractions add to one (for some reason the EC to excited is
        # probability given a EC
        FPEC = (1 - Info.FPECX) * Info.FPEC
        FPECX=Info.FPECX*Info.FPEC
        Info.FPEC=FPEC
1000
        Info.FPECX=FPECX
```

```
return (Info)
1005
    def FindPotentialMatchX(List,protons,A,XSection,Excited,Lambda,LOUD=False):
        if (XSection<0):</pre>
            print("Fraction of decays is too low", XSection)
            print("Inquire further")
            quit()
1010
        if XSection>0:
            for item in List:
                if protons in item[0:2] and A in item[:-1] and item[-1]==Excited:
                    Toreturn=item
        else:
1015
            Toreturn=''
        try: #To make sure its defined
            Toreturn
        except NameError:
            if LOUD:
1020
                print ("Could not find daughter in list of isotopes when expecting one")
                print ("Looking for Protons : ",protons, " Total Nucleons : ",A, " XSection ", XSection)
                print ("Lambda = ", Lambda)
                #print("Close items are")
                #for item in List:
1025
                   if protons in item[:-1] and A in item[:-1]:
                         print(item)
            if (XSection<100000 or Lambda>0.1):
                if LOUD:
                    print("Assuming it doesn't matter, will let slide")
1030
                Toreturn=''
                XSection=0
            else:
                quit()
1035
        return (Toreturn, XSection)
    class XSectionClass:
        def __init__(self):
                              # the effective, one grounp (n,y) x-section leadin to ground state
            self.SNG = 0
1040
            self.IDSNG = ''
                               # ZAID ID for the above
            self.SN2N = 0. # the effective, one grounp (n,2n) x-section leading to ground state
            self.IDSN2N = ''
                              # ZAID ID for the above
            self.SN3N = 0 # effective to ground
            self.IDSN3N = '' # ZAID for daughter for above
1045
            self.SNA = 0.
                               # effective to ground n, alpha
            self.IDSNA = ''
                                # ZAID for daughter for above
            self.SNF = 0.
                         = 0.#
            self.SNP
            self.IDSNP = ''
                               # ZAID for daughter for above
1050
            self.SNGX
                         = 0.#
            self.IDSNGX = '' # ZAID for daughter of above
                          = 0.#
            self.SN2NX
            self.IDSN2NX = '' # ZAID for daughter for above
```

```
1055
    def XSectionInfo(Nuclide_Names, parent, L, proton, A):
         This function will store and return decay information from
         the tape9.inp file
1060
        Info=XSectionClass()
1065
        with open ('Data/tape9.inp') as f:
             TAPE9Content=f.readlines()
        Found=False
         for line in TAPE9Content:
1070
             hold=line.split()
             #Looking for fission product and actinide cross section information
             #The libraries we are looking through for this information are '602' and '603'
             #set a cross section to zero if daughter is not found (second return of FindPotentialMat
             if ('602' == hold[0] \text{ or } '603' == hold[0]) and hold[1] == parent:
1075
                 #(n,gamma)
                 Info.SNG=float(hold[2])
                 Info.IDSNG, Info.SNG=FindPotentialMatchX(Nuclide_Names, proton, str(int(A) +1), Info.SNG,
                 #n,2n
1080
                 Info.SN2N=float(hold[3])
                 Info.IDSN2N, Info.SN2N=FindPotentialMatchX(Nuclide_Names,
                                                              proton, str(int(A)-1), Info.SN2N, '0', L)
                 if '602' == hold[0]: #Actinides
                      #n,3n
1085
                     Info.SN3N=float(hold[4])
                     Info.IDSN3N, Info.SN3N=FindPotentialMatchX(Nuclide_Names,
                                                                  proton, str(int(A)-2), Info. SN3N, '0', L)
                      #n, f
                     Info.SNF=float(hold[5])
1090
                 if '603' == hold[0]: #Fission products
                      #n,alpha
                     Info.SNA=float(hold[4])
                     Info.IDSNA, Info.SNA=FindPotentialMatchX(Nuclide_Names, str(int(proton)-2), str(int
                                                      Info.SNA,'0',L)
                      #n,p
                     Info.SNP=float(hold[5])
                     Info.IDSNP,Info.SNP=FindPotentialMatchX(Nuclide_Names,
                                                                str(int(proton)-1), A, Info. SNP, '0', L)
                 # (n, gamma) excited
1100
                 Info.SNGX=float(hold[6])
                 Info.IDSNGX, Info.SNGX=FindPotentialMatchX (Nuclide_Names, proton,
                                                              str(int(A)+1), Info.SNGX, '1', L)
                 \#(n,2n) excited
                 Info.SN2NX=float(hold[7])
1105
                 Info.IDSN2NX, Info.SN2NX=FindPotentialMatchX (Nuclide_Names, proton,
                                                                str(int(A)-1), Info.SN2NX, '1', L)
```

```
return (Info)
1110
    def YieldInfo(yieldiso, holdIndex, LOUD=False):
        with open ('Data/tape9.inp') as f:
             TAPE9Content=f.readlines()
        Found=False; Yield=False
1115
         for line in TAPE9Content:
             hold=line.split()
             if Found:
                 if Yield:
                     returnyield=float(hold[holdIndex])
                     returnyield=0
             if '603' == hold[0] and hold[1]==yieldiso:
1125
                 Found=True
                 if float(hold[8])>0:
                     Yield=True
                 else:
                     Yield=False
1130
         if not Found:
             if LOUD:
                 print("Did not find a yield for ", yieldiso)
             returnyield=0
1135
        return(returnyield)
    def AddFission(A, Nuclides, isotope, c1, c2, row, LOUD=False):
         if isotope=="922320": #Th232
             #print("Th232")
1140
             holdIndex=1
             element="Th232"
         elif isotope=="922330": #U233
             #print("U233")
             holdIndex=2
1145
             element="U233"
         elif isotope=="922350": #U235
             #print ("U235")
             holdIndex=3
             element="U235"
1150
         elif isotope=="922380": #U238
             #print("U238")
             holdIndex=4
             element="U238"
         elif isotope=="942390": #Pu239
             #print("Pu239")
            holdIndex=5
             element="Pu239"
         elif isotope=="942410": #Pu241
             #print("Pu241")
1160
```

```
holdIndex=6
             element="Pu241"
         elif isotope=="962450": #Cm245
             #print("Cm245")
             holdIndex=7
1165
             element="Cm245"
         elif isotope=="982490": #Cf249
             #print("Cf249")
             holdIndex=8
             element="Cf249"
1170
         else:
             if LOUD:
                 print("Did not find yields for ",isotope," because not provided")
             holdIndex=100
1175
         if holdIndex<40:</pre>
             YieldSum=0 #Check what the yields sum up to
             for yieldiso in Nuclides:
                 actualrow=Nuclides[yieldiso]
1180
                 Yield=YieldInfo(yieldiso, holdIndex, LOUD=False)
                 A[actualrow,row]=A[actualrow,row]+c1*c2*Yield/100
                 YieldSum=YieldSum+Yield
             if LOUD:
                 print("Yield Sum for ", element," = ", YieldSum)
1185
         return (A)
    def MakeAb(phi, Nuclides, Nuclide_Names, Decay_Conts):
         # Create Activation and Decay Matrix and initial
1190
         # nuclide quantity vector
        A = np.zeros((len(Nuclides), len(Nuclides)))
         #10^14 1/cm^2/s in 1/cm^2 /year #Bars included
        phi = phi * 60 * 60 * 24 * 365.25 * (10**(-24))
1195
         for isotope in Nuclides:
             row = Nuclides[isotope]
             proton=isotope[0:2]
             if isotope[2]=="0":
1200
                 Anum=isotope[3:5]
             elif isotope[4] == "0" and float(proton) *3<100:</pre>
                 Anum=isotope[2:4]
             elif isotope[2]!="0":
                 Anum=isotope[2:5]
1205
             else:
                 print("Missed logic in finding A number")
                 quit()
             if int(Anum) < int(proton):</pre>
                 print("Something is wrong, more protons than neutrons")
                 print("Proton: ",proton, "A: ",Anum, "ZAID : ",isotope)
                 quit()
```

```
#print(row,isotope,Decay_Conts[row])
1215
             Lambda=Decay_Conts[row]
             #Store all decay and x section information for row (I think I mean column)
             row_decay = DecayInfo(Nuclide_Names,isotope,Lambda,proton,Anum)
             row_XSection=XSectionInfo(Nuclide_Names,isotope,Lambda,proton,Anum)
             #Convert LAmbda to years^-1 (If radioactive)
1220
             if Lambda>0:
                 Lambday=Lambda\star60\star60\star24\star365.25
             else:
                 Lambday=0
1225
             XList=["SN2N", "SN2NX", "SN3N", "SNA", "SNG", "SNGX", "SNF", "SNF"]
             DList=["FA", "FB", "FBX", "FIT", "FN", "FPEC", "FPECX", "FSF"]
             #USeful to see things
             #for a in dir(row_XSection):
1230
                  if not a.startswith('__'):
                      print(a)
                      print(getattr(row_XSection,a))
             #quit()
             #for a in dir(row_decay):
1235
                  if not a.startswith('__'):
                      print(a)
                      print(getattr(row_decay, a))
             #quit()
1240
             #Sum up all the sigma abs
             sigma_sum=0
             for xsec in XList:
                 sigma_sum=sigma_sum+getattr(row_XSection,xsec)
1245
             # Diagonal Assignment
             A[row,row] = -Lambday - phi*sigma_sum
             ##does a single isotope produce another isotope through more than one path
             ##like EC and NP
1250
             #for xsec in XList:
                  for decay in DList:
                      if not decay[-1] == 'F' and not "F" in xsec and len(getattr(row_decay, "ID"+decay)
                           if getattr(row_XSection, "ID"+xsec) == getattr(row_decay, "ID"+decay):
                               print("Isotope ",getattr(row_XSection,"ID"+xsec),"Produced from ",
1255
                                     isotope, " with RNS ", decay, xsec)
                               print("Both will be used")
             \#Off diagonal assignment adding x-sec productions except from fission
             for xsec in XList:
1260
                 if not "F" in xsec: #Don't do fission yet
                     Product=getattr(row_XSection,"ID"+xsec)
                     if len(Product)>1:
                         actualrow=Nuclides[Product]
                         A[actualrow, row] = A[actualrow, row] + phi * getattr (row_XSection, xsed)
1265
             #Off diagonal assignment adding all decay except from spontaneous fission
```

```
for decay in DList:
                 if not decay[-1]=='F':
                     Product=getattr(row_decay,"ID"+decay)
                     if len(Product)>1:
1270
                         actualrow=Nuclides[Product]
                         A[actualrow,row] = A[actualrow,row] + Lambday * getattr(row_decay,decay)
             #Now for xfission (these next two if statements take the longest)
             if row_XSection.SNF>0:
1275
                 A=AddFission(A, Nuclides, isotope, row_XSection.SNF, phi, row, LOUD=False)
             #Now for spontaneous fission
             if row_decay.FSF>0:
                 A=AddFission(A, Nuclides, isotope, row_decay.FSF, Lambday, row, LOUD=False)
1280
        b = np.zeros(len(Nuclides))
        b[Nuclides['922340']] = 6.94741E23
        b[Nuclides['922350']] = 7.6864E25
        b[Nuclides['922380']] = 2.4532E27
1285
        return (A,b)
```