

Advanced Usage

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Additional tools



Not just FISPACT-II, additional tools included:

- compress_xs_endf Turn nuclear data into binary format for quicker processing
- extract_xs_endf Get the energy dependent cross section per reaction
- listreactions Prints information about reaction data in ENDF-6 file
- makenuclideindex Create a new index file given nuclide names

Other tools online and open source:

- GitHub https://github.com/fispact/
 - Pypact python3 package to parse output file and other functions
 - SPECTRA-PKA primary knock-on atom spectra
 - FFM Fispact File Maker
- Docker premade container environments with nuclear data included
 - DockerHub https://hub.docker.com/u/fispact/





Extract_xs_endf



Previously we compared different incident particle spectra and how this caused different collapsed cross section values. We showed the energy dependent cross section for Al 24 (n,p), but I did not show how to get this data.

FISPACT-II has a tool – extract_xs_endf, which will allow you to extract this data.

- Needs a lightweight files file
- No input file, just command line args
- Does need a fluxes file even if you just want the XS

Try the exercise in 'extended' folder, to understand how to use it and plot cross sections.

Listreactions



FISPACT-II has a tool – listreactions, which will allow you to print available channels in ENDF-6 files.

- No input files, just two args:
 - 1. "n", "p", "d", "g", "t", "a"
 - 2. ENDF-6 file path

First argument is really redundant, since you can give it a proton file and input "g", but for correctness use the correct particle for that file.

Group Convert



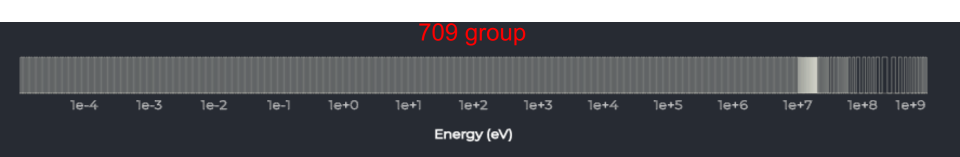
FISPACT-II uses binned (group-wise) data

A major problem with this:

- Group structure of F-II nuclear data is usually 709 for most libraries
- Whilst 709 is generally pretty good, your flux must match this structure
- If you have a spectra in another structure how do you use it with F-II?

Two solutions:

- Make your own nuclear data for that group use NJOY, PREPRO (not recommended)
- Use group convert F-II keyword (GRPCONVERT)



Group Convert



Input arbitrary group structure

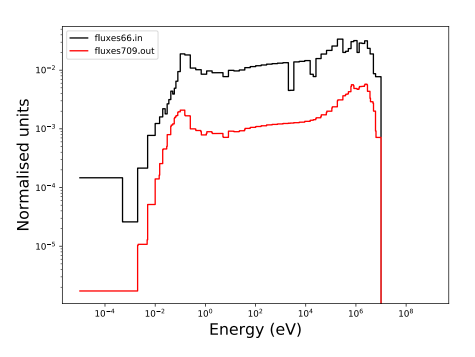
- Lethargy = $u = ln(E_0/E)$
- Output F-II known group structure
- Conversion can be done using equal lethargy or equal energy per bin method (CNVTYPE)
- Splits groups based on weighting each input and output energy group
- Not a separate tool use GRPCONVERT keyword in input file
- Requires a files file with:
 - ind_nuc needed to run but not used
 - arb flux input flux file
 - fluxes output flux file

Recommended to use groups "close" to each other. But is possible to do 66 -> 1102 if desired. Results depend on how groups "line" up.

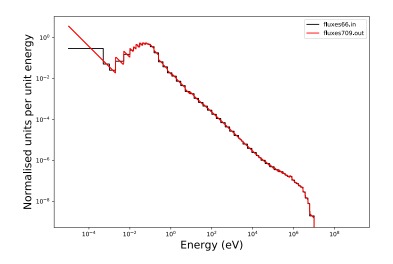
Group Convert

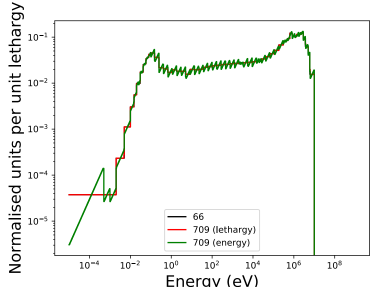


Exercise 'grpconvert' shows conversion from group 66 to 709. Must compare using bin width (per unit energy) or unit lethargy.



Try the exercise in 'extended' folder.



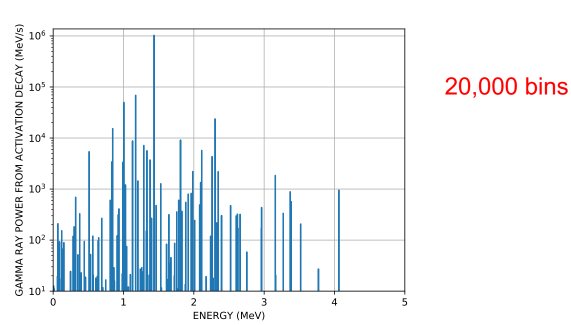


Gamma spectrum



FISPACT-II also outputs the gamma spectrum rate (power) at each timestep.

- If you use ATOMS or SPECTRUM the data will be in the output file
- Default gamma group is <u>24-energy group structure</u>
- Use GROUP 1 to use <u>22-group 'Steiner' structure</u>
- If this is too coarse you can input custom bin boundaries with READGG



Try the exercise in 'advanced' folder, to understand how to use it and plot cross sections.

FISPACT-II Output Format



- Fairly easy to look inside and read a value
- Hard to script and parse values
- Output files can be quite large (often > 5MB)
- Grep will only take you so far

Solutions:

- Write your own parser (not recommended)
- Use TAB files for easier parsing (limited output, limited precision)
- Use JSON output (standardised format, high precision, since v4 only)
- Use pypact (python3 package designed for parsing output)

Exercise on each in advanced section

FISPACT-II JSON



FISPACT-II 4.0 now has a JSON output

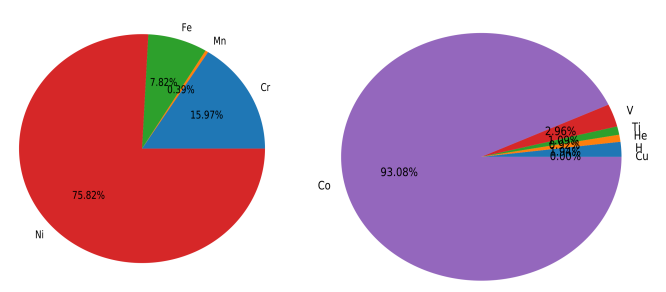
```
{
    "name" : "json",
    "description" : "I am JSON",
    "id" : 4,
    "value" : 4.3e+9
}
```

Simply use JSON keyword in control section
We did this in FNS Inconel example

```
<< ----set initial switches and get nuclear data--
--- >>
CLOBBER
JSON
GETXS 0
GETDECAY 0
FISPACT
* FNS 5 Minutes Inconel-600
```

Exercise JSON

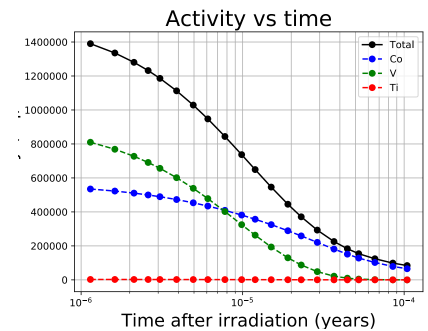




Left: initial

Right: final (excluding initial)

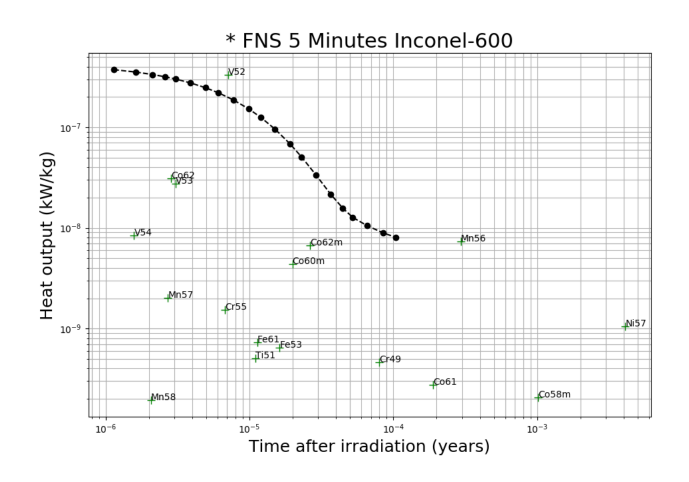
Exercise JSON (advanced) shows how to do this



Exercise Pypact



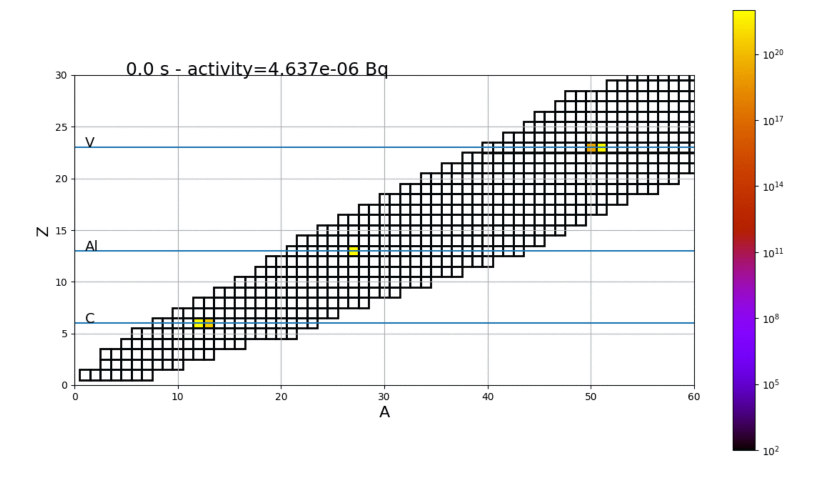
Use pypact to make simple plots vs time



Exercise Pypact



Chart of nuclides animation – less than 70 lines of code!



Multiflux



- In most practical scenarios, energy spectra changes with time and space
- Important for burn-up calculations
- Need to allow for different spectra in inventory calculations, but...
 - One simulation/run = one collapse
- ⇒ When the spectra is changing, we must re-collapse
- Chain input1, output1 => input2, output2 =>...
- Can be difficult to parse output file
 - As mentioned before, a few options TAB, JSON, PYPACT
- We will use the TAB1 file to help automate this, since the input just needs to know the number of atoms

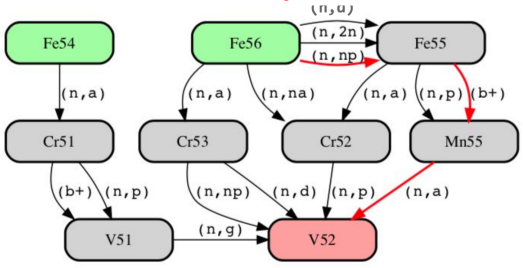
Exercise multiflux (advanced) shows how to do this

Pathways



- Nuclear data represents a major contributor to uncertainties in nuclear simulations
- There are known knowns, known unknowns, and unknown unknowns
- We can estimate the known unknowns via pathways analysis built into FISPACT-II
- Pathways also provides information on the path from parent to child
 - How much of X comes from Y?
 - What if we isotopically tailored X?

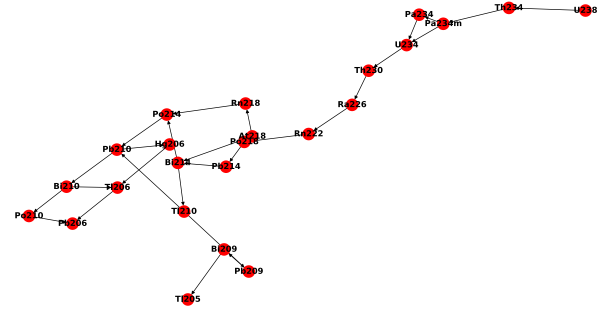
Where did my 52V come from?



Pathways



- Use the PATH keyword to specify a path of interest
- Use ROUTES keyword to specify a parent and child to find all routes
- Other keywords to related to pathways analysis
 - PATHRESET
 - LOOKAHEAD
 - UNCERTAINTY
 - ZERO
 - FISCHOSE
 - GENERIC



Exercise pathways (extended) shows how to use it

Summary and more



- FISPACT-II is an advanced multigroup Multiphysics code capable of:
 - Providing radio properties per nuclide at times from 1e-20 to 1e20 seconds!
 - Associated gamma spectrum at time interval
 - Extract + plot XS data
 - Uncertainty analysis
 - Pathways analysis
- Open source tools in active development
- Many ways to analyse FISPACT-II output
 - GNU plotting built in
 - TAB files can make life easier but difficult hard to script
 - JSON output has many parsers ready C++, Fortran, Python, js, ...
 - Pypact
- Docker
 - Many images ready made with nuclear data inside
 - Self contained environment to run fispact easily
 - Nuclear data in binary format performance improvement

Coming soon



There has been recent development into FISPACT-II to bring you:

A kernel with APIs

- No files
- C, C++, Fortran, and Python bindings
- Significant performance boost
- Makes some analysis trivial and fast

A web application

- No installation required
- No command line
- No output file
- Dashboard approach
- Select and click!

I will give a demo of the API later....

