
User's Manual

NTP-ERSN: A new package for solving the multigroup neutron transport equation in a slab geometry, Version 1.1

M. Lahdour, T. El Bardouni*

¹*Radiations and Nuclear Systems Laboratory, University Abdelmalek Essaadi,
Faculty of Sciences of Tetuan, Morocco
Corresponding Address:
mlahdour@uae.ac.ma

<https://github.com/mohamedlahdour/NTP-ERSN.git>

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Introduction

Welcome to the NTP-ERSN User's Guide! This tutorial will describe the essential aspects to perform multigroup neutron transport equation physics calculations by using NTP-ERSN code, available for download at <https://github.com/mohamedlahdour/NTP-ERSN.git>.

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I Installation and Quick start

I.I Obtaining the source

All NTP-ERSN source code with its GUI is hosted on GitHub and it can be downloaded for free. You can download the source code directly from GitHub or, if you have the Git version control software installed on your computer, you can use git to obtain the source code. The latter method has the benefit that it is easy to receive updates directly from the GitHub repository. GitHub has a good set of instructions for how to set up git to work with GitHub since this involves setting up ssh keys. With git installed and setup, the following command will download the full source code from the GitHub repository:

```
git clone https://github.com/mohamedlahdour/NTP-ERSN.git
```

or by downloading directly the archive of the actual release:

```
wget https://github.com/mohamedlahdour/NTP-ERSN/archive/v-1.1.tar.gz
```

I.2 Installing Prerequisites on Ubuntu

1- untar the archive NTP-ERSN.tar.gz

In the case where the source code was downlaod directly from GitHub, you can untar the archive and executed the installation script (NTP-ERSN-latest.sh). A second case, the source code can be downlaod by using git if you have the Git version control software installed on your computer.

NTP-ERSN_installer.sh

The script (NTP-ERSN-latest.sh) will install NTP-ERSN with its GUI and its dependencies on the system. Install script is available for *UBUNTU distribution of linux* Ubuntu operating system version 17.10 and all the latest versions.

2- In your system, run the commands below.

```
cd NTP-ERSN/script  
chmod +x NTP-ERSN_latest.sh  
sh NTP-ERSN_latest.sh
```

```
cd NTP-ERSN  
sh script/NTP-ERSN_installer.sh
```

This NTP-ERSN and its GUI has been released under the MIT GPL license. Since, the software is a Python-based application; it requires a Python Runtime Environment to work correctly.

3- You can started the software by typing the following command-line argument in a Linux terminal:

```
python interface.py
```

Then a Main window (GUI) of NTP-ERSN package on Ubuntu Linux machine ~~well~~ be displayed as in Figure I.

will

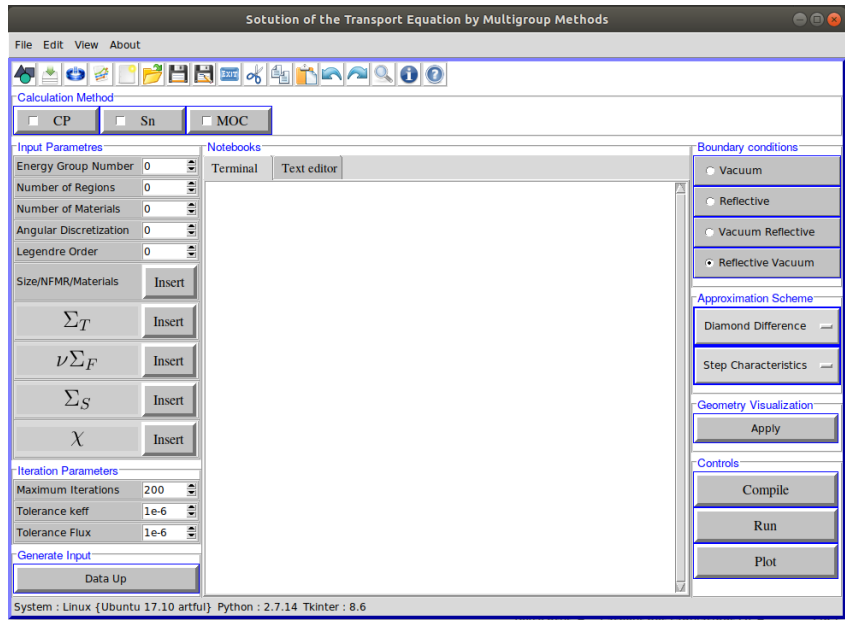


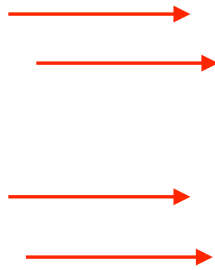
Figure 1: Main window of the GUI of the NTP-ERSN package

2 Examples and test^s suite

2.1 Writing JSON Input Files

The input data file must be in JSON format. The first method is to write it directly in **Text editor** as is shown in the figure 2 witch illustrates an example input file of a critical Benchmark taken from [Avneet Sood(2003)]. This benchmark contains one region and one energy groups.

2.2 Generating JSON Input Files



Another method is to use a set of buttons on the left side of the main window 3, these buttons allowing users to insert input data automatically without requiring an in-depth knowledge of JSON file syntax. Once the user click^s on **Data Up** button the input file will be automatically generated in the window **Text editor**. The figure 4 represents the generated input file named INPUT.JSON contains another example of a Criticality Verification Suite Problems taken from [Avneet Sood(2003)]. This benchmark contains three region^s and two energy groups with scattering source^s isotropic. The parameters of this benchmark are written in INPUT.JSON input file as follows: We define the region^s number, material^s number and the scattering order of the medium where the neutron is moving. Then the size of each region (coarse mesh), within which the number of fine meshes is defined. To each region, is assigned a material index. The fission spectrum and the total^s diffusion and fission cross sections have values for different materials. Each material can be placed in any region using

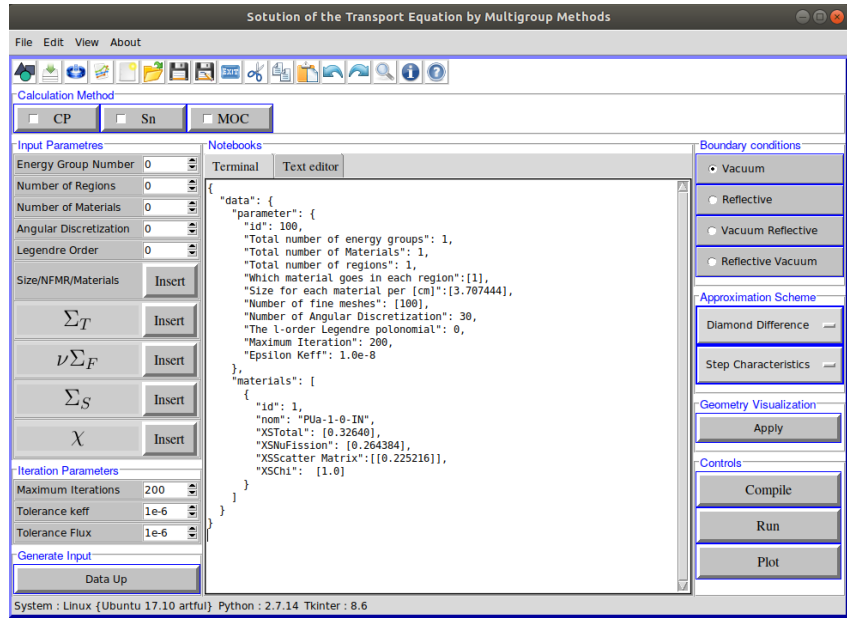


Figure 2: Setting Up Input File for slab geometry in one energy group

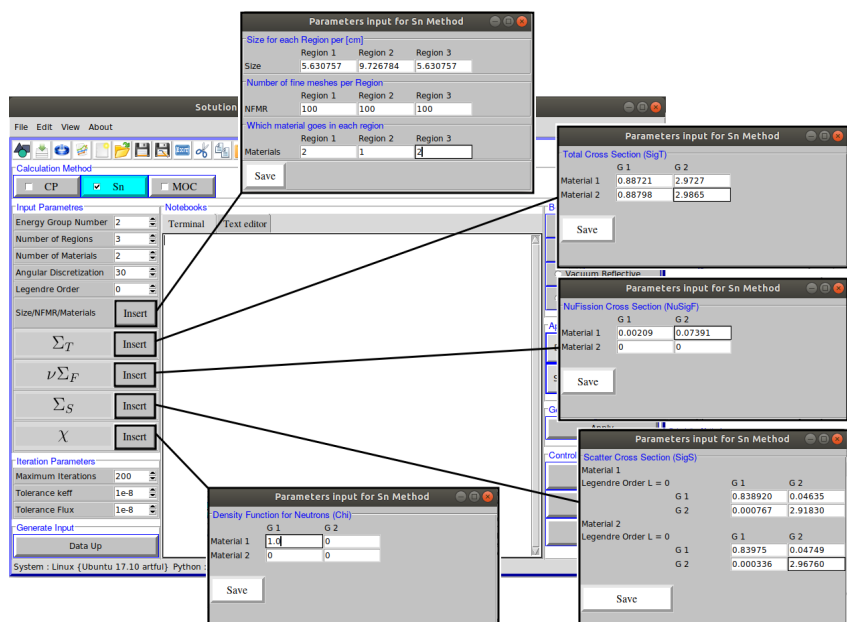


Figure 3: Example of inserting input data in GUI

the appropriate index. This input format allows one to investigate fairly easily a wide range of slab compositions.

```
{
  "data": {
    "parameter": {
      "id": 100,
      "Total number of energy groups": 2,
      "Total number of Materials": 2,
      "Total number of regions": 3,
      "Which material goes in each region": [2, 1, 2],
      "Size for each material for [cm]": [5.630757, 9.726784, 5.630757],
      "Number of fine meshes": [5, 5, 5],
      "Number of Angular Discretization": 8,
      "The l-order Legendre polynomial": 0,
      "Maximum Iteration": 200,
      "Epsilon_Keff": 1e-8
    },
    "materials": [
      {
        "name": "material 1",
        "id": 1,
        "XSTotal": [0.88721, 2.9727],
        "XSNUFission": [0.00209, 0.07391],
        "XSScatter Matrix": [[[0.838920, 0.04635], [0.000767, 2.91830]]],
        "XSChi": [1.0, 0.0]
      },
      {
        "name": "material 2",
        "id": 2,
        "XSTotal": [0.88798, 2.9865],
        "XSNUFission": [0.0, 0.0],
        "XSScatter Matrix": [[[0.83975, 0.04749], [0.000336, 2.96760]]],
        "XSChi": [0, 0]
      }
    ]
  }
}
```

*fills
Size of each region*

*Maximum number of Iterations
Criterion Keff convergence*

name

name

polynomial

Figure 4: INPUT.JSON input file

3 Running NTP-ERSN

3.1 Running NTP-ERSN under a GUI

Once you have created your input file (*.json), it is relatively straightforward to run the code. *running the code* the steps of *execution* are:

- • Set up input file as described in Section 2.1, 2.2 or import one of the existing input files into the `./tests` folder using the `Open data file` button on the menu bar. *You* will find in this folder a set of test cases named (*.json) taken from [Avneet Sood(2003)] (see figure 5).
- • Select one of the `CP`, `SN` or `MOC` methods in `Calculation Method`.
- • *Click* `One-click` on the button `Compile`, a Python C/API extension modules to call Fortran 90/95 modules subroutines will be created. *This task can be done only at the first time you use the software.*
- Check `Boundary Condition` radio button.
- Check `Approximation Scheme` radio button for the case where `SN` or `MOC` methods are selected.
- Click `Run` button for solve multigroup scheme.
- When completed with criticality calculation, the software will display the message "finished running case" (see figure 6).

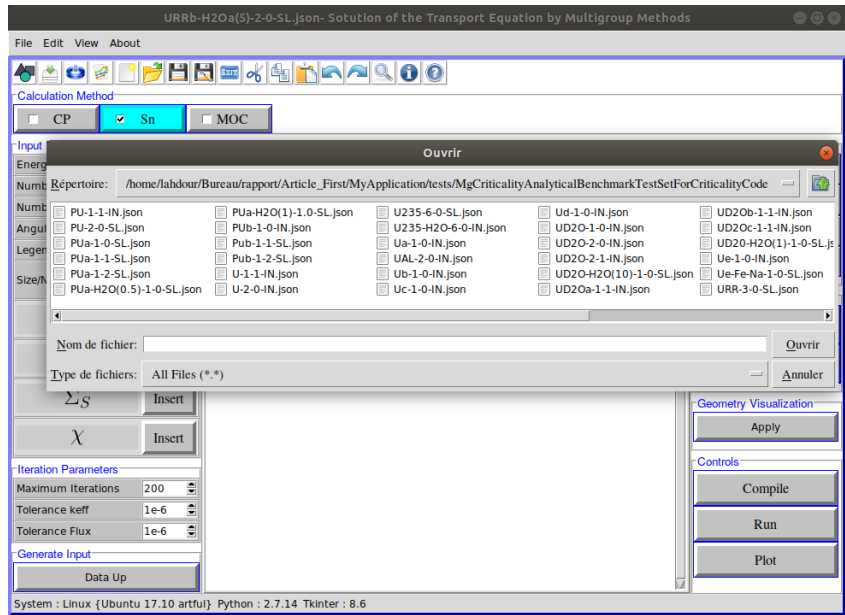
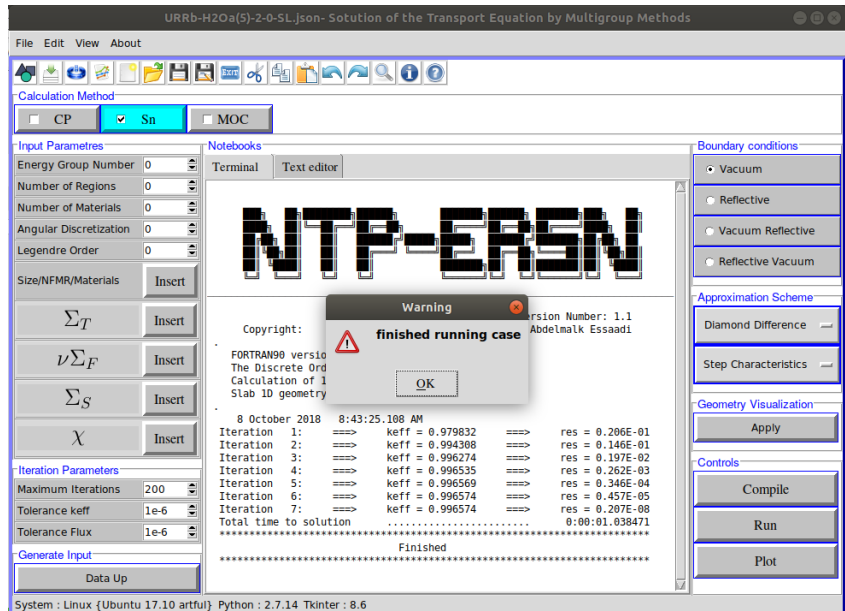


Figure 5: Import input file



When
Figure 6: Finished running case *is finished*

3.2 Flux Visualization

The **Plot** button refers to a set of routines to plot the scalar flux in space and in each energy group. The figure 7 shows the flux for the example input file of figure 4 with three regions and two materials after clicking on the **Plot** button.

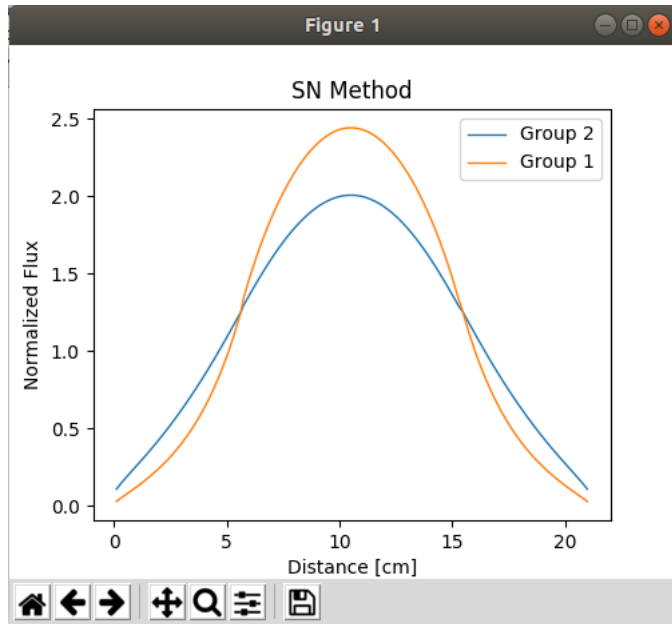


Figure 7: The fast and thermal fluxes in the URRb-H₂Oa(5)-2-o-SL benchmark problem.

3.3 Geometry Visualization

To plot the geometry, click the **Apply** button.
A depiction of the geometry for the example input file of figure 4 with three regions and two materials is illustrated in Figure 8.

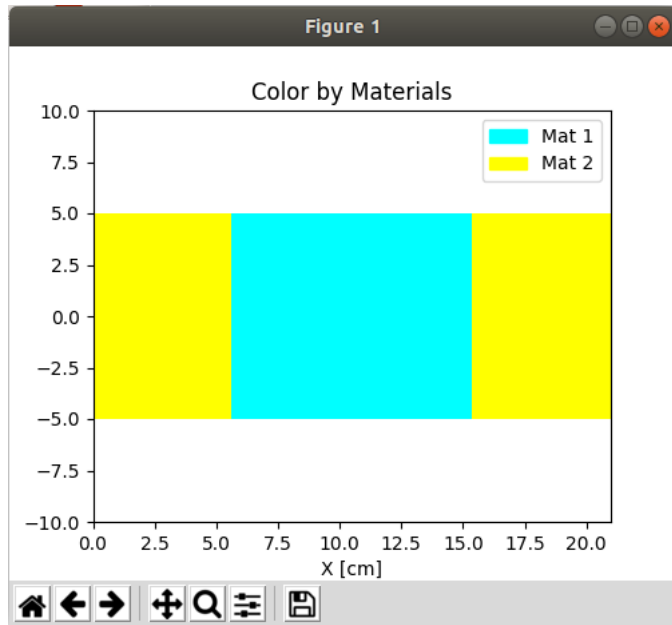


Figure 8: A slab geometry colored by material.

3.4 Simple Output

The following is the corresponding output to the above case. The contents of the output file are briefly described: version and run time information, input parameter – values from input, calculation run-time parametres method, scalar flux solution and output parameter_s – solution to transport eqaution.

ERSN, UNIVERSITY ABDELMALEK ESSAADI FACULTY OF SCIENCES – TETOUAN, MOROCCO						
CODE DEVELOPED BY MOHAMED LAHDOUR, PHD STUDENT						
NTP-ERSN: SN DISCRETE ORDINATES METHOD						
VERSION NUMBER: 1.1						
VERSION DATE: 8 OCTOBER 2018						
RAN ON: 2018-10-07 13:44:25.78 (H/M/S)						

<hr/>						
INPUT PARAMETER – VALUES FROM INPUT						
<hr/>						
ENERGY GROUP NUMBER: 2						
REGIONS NUMBER: 3						
MATERIALS NUMBER: 2						
SIZE FOR EACH MATERIAL PER [CM]: 5.63076 9.72678 5.63076						
DISCRETIZATIONS ANGULAR: 8						
ORDER LEGENDRE POLONOMIAL: 0						
TOTAL NUMBER OF FINE MESHES: 15						
TOLERANCE KEFF AND FLUX: 1.0E-08						
<hr/>						
CALCULATION RUN-TIME PARAMETERS SN						
<hr/>						
GAUSS LEGENDRE QUADRATURE POINTS AND WEIGHTS:						
<hr/>						
N. GAUSS POINTS WEIGHTS						
<hr/>						
1	-9.60290E-01	1.01229E-01				
2	-7.96666E-01	2.22381E-01				
3	-5.25532E-01	3.13707E-01				
4	-1.83435E-01	3.62684E-01				
5	1.83435E-01	3.62684E-01				
6	5.25532E-01	3.13707E-01				
7	7.96666E-01	2.22381E-01				
8	9.60290E-01	1.01229E-01				
<hr/>						
PSEUDO CROSS SECTIONS DATA:						
<hr/>						
MATERIAL : 1						
<hr/>						
GROUP TOTAL ABSORPTION NU+FISSION SCATTERING						
<hr/>						
1	8.87210E-01	4.82900E-02	2.09000E-03	8.38920E-01		
2	2.97270E+00	5.44000E-02	7.39100E-02	2.91830E+00		
<hr/>						
MATERIAL : 2						
<hr/>						
GROUP TOTAL ABSORPTION NU+FISSION SCATTERING						
<hr/>						
1	8.87980E-01	4.82300E-02	0.00000E+00	8.39750E-01		
2	2.98650E+00	1.89000E-02	0.00000E+00	2.96760E+00		
<hr/>						
SCALAR FLUX SOLUTION						
<hr/>						
FLUXES PER MESH PER ENERGY GROUP:						
<hr/>						
MESH GROUP 1 GROUP 2						
<hr/>						
1	1.95059E-01	7.97230E-02				
2	3.90662E-01	2.10169E-01				
3	6.03844E-01	3.85998E-01				
4	8.53613E-01	6.45059E-01				
5	1.13254E+00	1.05059E+00				
6	1.50142E+00	1.69279E+00				
7	1.85024E+00	2.24946E+00				
8	1.97499E+00	2.41389E+00				
9	1.84576E+00	2.23982E+00				
10	1.49487E+00	1.68037E+00				
11	1.12032E+00	1.04146E+00				
12	8.48570E-01	6.40041E-01				
13	6.00118E-01	3.82054E-01				
14	3.88189E-01	2.08346E-01				
15	1.93810E-01	7.90349E-02				
<hr/>						
OUTPUT PARAMETER – SOLUTION TO TRANSPORT EQUATION						

K-EFF	=	0.996574
N. OUTER ITERATIONS	=	7
TOTAL INNER ITERATIONS	=	60050
TOTAL EXECUTION TIME	=	0:00:01.05 (H/M/S)

Figure 9: OUTPUT.TXT output file

References

- [Avneet Sood(2003)] D. Kent Parsons Avneet Sood, R.Arthur Forster. Analytical benchmark test set for criticality code verification. *Progress in Nuclear Energy*, 42(1):55 – 106, 2003. ISSN 0149-1970. doi: [https://doi.org/10.1016/S0149-1970\(02\)00098-7](https://doi.org/10.1016/S0149-1970(02)00098-7). URL <http://www.sciencedirect.com/science/article/pii/S0149197002000987>.