atomneb Documentation

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USER DOCUMENTATION

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INTRODUCTION

atomneb is a library written in Python for reading atomic data from a database containing atomic data stored in the Flexible Image Transport System (FITS) file format for *collisionally excited lines* and *recombination lines* typically observed in spectra of ionized gaseous nebulae. The AtomNeb database were generated for use in pyEQUIB, proEQUIB, and other nebular spectral analysis tools.

1.1 Collisional Excitation Unit

AtomNeb for collisionally excited lines contains sets of atomic datasets, which include energy levels (E_j) , collision strengths (Ω_{ij}) , and transition probabilities (A_{ij}) of the most ions commonly observed in ionized nebulae.

The atomic datasets for collisionally excited lines are as follows:

- Collection from the National Institute of Standards and Technology (NIST) Atomic Spectra Database, the CHI-ANTI atomic database, and some improved atomic data from Cloudy v13.04 and pyNeb v1.0. This collection was compiled according to the atomic data used in pyNeb v1.0.
- Chianti52 from the CHIANTI atomic database version 5.2. This dataset was compiled according to the atomic data used in MOCASSIN.
- Chianti60 from the CHIANTI atomic database version 6.0. This dataset was compiled according to the atomic
 data used in MOCASSIN.
- Chianti70 from the CHIANTI atomic database version 7.0. This dataset was compiled according to the atomic
 data used in MOCASSIN.

Each dataset contains the following atomic data FITS files: AtomElj.fits for *Energy Levels* (E_j) , AtomOmij. fits for *Collision Strengths* (Ω_{ij}) , and AtomAij.fits for *Transition Probabilities* (A_{ij}) .

1.2 Recombination Unit

AtomNeb for recombination lines contains sets of effective recombination coefficients (α_{eff}) of recombination lines of H I, He II, C II, C III, C III, C VI, N III, N III, N IV, N V, N VI, N VII, O II, O III, O IV, O V, O VI, O VIII, and Ne II ions typically observed in ionized nebulae, as well as Branching ratios (Br) of O II and N II lines.

The atomic datasets for recombination lines are as follows:

- RC Collection, effective recombination coefficients for C II (Davey et al. 2000), N II (Escalante and Victor 1990), O II (Storey 1994; Liu et al. 1995), and Ne II ions (Kisielius et al. 1998), including Branching ratios (Br) for O II and N II ions. This collection was compiled according to the atomic data used in MOCASSIN.
- SH95 Collection, hydrogenic ions for Z=1 to 8, namely H I, He II, Li III, Be IV, B V, C VI, N VII, and O VIII ions from Storey and Hummer (1995).

- PPB91 Collection, effective recombination coefficients for H, He, C, N, O, Ne ions from Pequignot, Petitjean and Boisson (1991).
- PFSD12 He I data, effective He I recombination coefficients from Porter et al (2012) and (2013a).
- FSL13 N II data, effective N II recombination coefficients (corrigendum) from Fang, Storey and Liu (2011) and (2013b).
- SSB17 O II data, effective O II recombination coefficients of 8889 recombination lines for Cases A, B, and C, and 2433 optical (3500-9000Å) recombination lines for Case B from Storey, Sochi and Bastin (2017).

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INSTALLATION

To install the last version, all you should need to do is

```
$ python setup.py install
```

To install the stable version, you can use the preferred installer program (pip):

```
$ pip install atomneb
```

or you can install it from the cross-platform package manager conda:

```
$ conda install -c conda-forge atomneb
```

To get this package with all the FITS file, you can simply use git command as follows:

```
git clone https://github.com/atomneb/AtomNeb-py
```

If you plan to use the recent O II recombination coefficients (Storey, Sochi and Bastin 2017), you need to unpack them:

```
cd AtomNeb-py/atomic-data-rc/
tar -xvf *.fits.tar.gz
```

This package requires the following packages:

- NumPy
- pandas
- Astropy

THREE

USAGE

The Documentation of the functions provides in detail in the *API Documentation* (atomneb.github.io/AtomNeb-py/doc). There are two main categories: *collisionally excited lines (CEL)* and *recombination lines (RC)*.

3.1 Collisional Excitation Unit

The atomic data for **collisional excitation unit** (CEL) contain Energy Levels (E_j) , Collision Strengths (Ω_{ij}) , and Transition Probabilities (A_{ij}) . We have four atomic datasets for them: collection, chianti52, chianti60, and chianti70.

You need to load the **atomneb** library as follows:

```
import atomneb
```

Also:

```
import numpy as np
import os

atom_elj_file = os.path.join(base_dir,data_dir, 'AtomElj.fits')
atom_omij_file = os.path.join(base_dir,data_dir, 'AtomOmij.fits')
atom_aij_file = os.path.join(base_dir,data_dir, 'AtomAij.fits')
elj_data_list = atomneb.read_elj_list(atom_elj_file)
omij_data_list = atomneb.read_omij_list(atom_omij_file)
aij_data_list = atomneb.read_aij_list(atom_aij_file)
```

Now you have access to:

• Energy Levels (Ej):

```
atom='o'
ion='iii'
oiii_elj_data = atomneb.read_elj(atom_elj_file, atom, ion, level_num=6)
print(np.asarray(oiii_elj_data.j_v))
print(np.asarray(oiii_elj_data.ej))
```

which gives:

```
      0.00000
      1.00000
      2.00000
      2.00000
      0.00000
      2.00000

      0.00000
      113.200
      306.200
      20273.30
      43185.69
      60324.80
```

• Collision Strengths (Ω_{ij}) :

```
atom='o'
ion='iii'
oiii_omij_data = atomneb.read_omij(atom_omij_file, atom, ion)
print(np.asarray(oiii_omij_data.level1))
print(np.asarray(oiii_omij_data.level2))
print(np.asarray(oiii_omij_data.strength)[0])
```

which gives:

```
0 1 1 1 1 ...
0 2 3 4 5 ...
100.0 158.50 251.20 398.10 631.0 ...
```

• Transition Probabilities (Aij):

```
atom='o'
ion='iii'
oiii_aij_data = atomneb.read_aij(atom_aij_file, atom, ion)
pprint(oiii_aij_data.aij)
```

which gives:

```
0.0000 2.5969e-05 0.0000 2.3220e-06 ...
```

3.2 Recombination Unit

The atomic data for **recombination unit (RC)** contain effective recombination coefficients (α_{eff}) of emission lines from different collections: RC Collection, SH95 Collection, PPB91 Collection, PFSD12 He I data, FSL13 N II data, and SSB17 O II data.

You need to load the atomneb libary:

```
import atomneb
```

Also:

```
import numpy as np import os
```

Now you have access to effective recombination coefficients (α_{eff}) of the following collections:

• RC Collection:

which gives:

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```
914.00000
             0.69280000
                           0.021400000
                                         -0.016300000

→24310000

             -0.88000000
962.00000
             1.0998000 -0.0042000000
                                         -0.027900000
                                                         -0.
→22940000
             -0.96560000
997.00000
            0.78210000
                          -0.36840000 0.00030000000
                                                         -0.
→12170000
             -0.78740000
. . .
```

• SH95 Collection:

```
atom_rc_file = os.path.join(base_dir,data_dir, 'rc_SH95.fits')
atom='h'
ion='ii'
hi_rc_data = atomneb.read_aeff_sh95(atom_rc_file, atom, ion)
print(hi_rc_data.aeff[0])
```

which gives:

```
100.00000 500.00000 0.0000000 4.2140000e-27 1.7560000e-

→27 1.0350000e-27 ...
```

• PPB91 Collection:

which gives:

```
C2+A 9903.4600 0.69700000 −0.78400000 4.2050000 

→ 0.72000000 1.0000000 1.6210000 

C2+A 4267.1500 1.0110000 −0.75400000 2.5870000 

→ 0.71900000 0.95000000 2.7950000 ...
```

• PFSD12 He I data:

which gives:

```
5000.0000 10.000000 −25.379540 −25.058970 −25.

→948440 −24.651820 −25.637660

...
```

• FSL13 N II data:

which gives:

```
255.000
             79.5000
                           47.3000
                                        12,5000
                                                      6.20000
                                                                    4.00000
      2.86000
258.000
                           29.7000
             54.4000
                                        7.92000
                                                      4.11000
                                                                    2.72000_
      2.00000
310.000
            48.1000
                           23.7000
                                        5.19000
                                                      2.55000
                                                                    1.65000_
      1.21000
                           23.2000
434.000 50.3000
                                        4.71000
                                                      2.26000
                                                                    1.45000
     1.05000
6413.23 \text{ } 6g - 4f2p6g \text{ } G[9/2]o4 - 2p4f \text{ } F[7/2]e3
6556.32 6g - 4f2p6g G[9/2]o5 - 2p4f G[7/2]e4
6456.97 6g - 4f2p6g G[9/2]o5 - 2p4f F[7/2]e4
6446.53 6g - 4f2p6g F[7/2]o3 - 2p4f D[5/2]e2
6445.34 6g - 4f2p6g F[7/2]o4 - 2p4f D[5/2]e3
```

• SSB17 O II data: You first need to unpack rc_o_iii_SSB17_orl_case_b.fits.tar.gz, e.g.:

```
tar -xvf rc_o_iii_SSB17_orl_case_b.fits.tar.gz
```

If you need to have access to the full dataset (entire wavelengths, case A and B):

```
tar -xvf rc_o_iii_SSB17.fits.tar.gz
```

Please note that using the entire atomic data will make your program very slow and you may need to have a higher memory on your system. Without the above comment, as default, the cose uses rc_o_iii_SSB17_orl_case_b.fits:

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which gives:

```
1.64100e-30 1.60000e-30 1.56400e-30 1.54100e-30 1.52100e-30 1.

$\infty$50900e-30 \\
\dots$

5327.17 2s22p2(1S) 3p 2Po \\
5325.42 2s22p2(1S) 3p 2Po \\
5327.18 2s22p2(1D) 3d 2Ge \\
5326.84 2s22p2(1D) 3d 2Ge \\
\dots$
```

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REFERENCES

• Danehkar, A. (2019). AtomNeb: IDL Library for Atomic Data of Ionized Nebulae. *J. Open Source Softw.*, **4**, 898. doi:10.21105/joss.00898

FIVE

ATOMNEB.MAIN PACKAGE

5.1 atomneb main module

This module contains functions for Atomic Data of Ionized Nebulae

```
atomneb.get_aeff_collection_reference_citation (atom\_rc\_file, atom, ion, br=None, reference=None)
```

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_collection.fits').

For example:

Returns This function returns the Citation.

Return type str

Parameters

- atom_rc_file (str) the FITS data file name ('rc_collection.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'
- **br** (boolean, optional) set for the branching ratios (Br)
- reference (str, optional) set for the reference

```
atomneb.get_aeff_he_i_pfsd12_reference_citation(atom_rc_file, atom, ion, reference=None)
```

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_he_ii_PFSD12.fits').

For example:

Returns This function returns the Citation.

Return type str

Parameters

- atom_rc_file (str) the FITS data file name ('rc_he_ii_PFSD12.fits')
- atom (str) atom name e.g. 'he'
- ion (str) ionic level e.g 'ii'
- reference (str, optional) set for the reference e.g. 'PFSD13'

```
atomneb.get_aeff_n_ii_fsl13_reference_citation(atom_rc_file, atom, ion, reference=None)
```

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_n_iii_FSL13.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = ['atomic-data-rc']
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_n_iii_FSL13.fits')
>> atom='n'
>> ion='iii' # N II
>> reference='FSL13'
>> citation = atomneb.get_aeff_n_ii_fsl13_reference_citation(atom_rc_file, atom, ion)
>> print(citation)
Fang X., Storey P.J., and Liu X.-W., R. 2011, Astron.Astrophys. 530, Ale; 2013, Astron.Astrophys. 550, C2
```

Returns This function returns the Citation.

Return type str

Parameters

• atom_rc_file (str) - the FITS data file name ('rc_n_iii_FSL13.fits')

```
• atom (str) - atom name e.g. 'n'
```

- ion (str) ionic level e.g 'iii'
- reference (str, optional) set for the reference e.g. 'FSL13

atomneb.get_aeff_o_ii_ssb17_reference_citation(atom_rc_file, atom, ion, reference=None)

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_o_iii_SSB17.fits').

For example:

Returns This function returns the Citation.

Return type str

Parameters

- atom_rc_file (str) the FITS data file name ('rc_o_iii_SSB17.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- reference (str, optional) set for the reference e.g. 'SSB17

atomneb.get_aeff_ppb91_reference_citation(atom_rc_file, atom, ion, reference=None)

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_PPB91.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = ['atomic-data-rc']
>> atom_rc_file = os.path.join(base_dir,data_dir,'rc_PPB91.fits')
>> atom='c'
>> ion='iii'
>> citation = atomneb.get_aeff_ppb91_reference_citation(atom_rc_file,u_datom, ion)
```

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```
>> print(citation)
Pequignot, D., Petitjean, P. and Boisson, C. Astron.Astrophys., 251, 4680, 1991
```

Returns This function returns the Citation.

Return type str

Parameters

- atom_rc_file (str) the FITS data file name ('rc_PPB91.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'
- reference (str, optional) set for the reference

atomneb.get_aeff_sh95_reference_citation(atom_rc_file, atom, ion, reference=None, case I=None)

This function returns the reference citation for a recombination coefficient (Aeff) from the 2nd binary table extension of the FITS data file ('rc_SH95.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = ['atomic-data-rc']
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_SH95.fits')
>> atom='h'
>> ion='ii' # H I
>> citation = atomneb.get_aeff_sh95_reference_citation(atom_rc_file, atom, ion)
>> print(citation)
Storey, P. J. and Hummer, D. G., MNRAS, 272, 415, 1995
```

Returns This function returns the Citation.

Return type str

Parameters

- atom_rc_file (str) the FITS data file name ('rc_SH95.fits')
- atom (str) atom name e.g. 'h'
- ion (str) ionic level e.g 'ii'
- reference (str, optional) set for the reference
- case1 (str, optional) set for the case 'a' or 'b', defualt 'b'

atomneb.get_aij_reference_citation(atom_aij_file, atom, ion, reference)

This function returns the reference citation for a transition probability (Aij) from the 2nd binary table extension of the FITS data file ('AtoAij.fits')

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data', 'collection')
>> atom_aij_file = os.path.join(base_dir, data_dir, 'AtomAij.fits')
>> atom='o'
>> ion='iii'
>> reference='FFT04'
>> citation = atomneb.get_aij_reference_citation(atom_aij_file, atom, ion, reference)
>> print(citation)
Froese Fischer et al 2004, ADNDT 87, 1
```

Returns This function returns the Citation.

Return type str

Parameters

- **Atom_Aij_file** (str) the FITS data file name ('AtoAij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- **reference** (*str*) set for the reference e.g. 'FFT04'

```
atomneb.get_elj_reference_citation(atom_elj_file, reference)
```

This function returns the reference citation for energy levels (Ej) from the 2nd binary table extension of the FITS data file ('AtomElj.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..','atomic-data', 'collection')
>> atom_elj_file = os.path.join(base_dir,data_dir, 'AtomElj.fits')
>> reference='L7288'
>> citation=atomneb.get_elj_reference_citation(atom_elj_file, reference)
>> print(citation)
    C. E. Moore, in CRC Series in Evaluated Data in Atomic Physics, 339 pp.

-- (CRC Press, Boca Raton, FL, 1993)
```

Returns This function returns the Citation.

Return type str

Parameters

- atom_elj_file (str) the FITS data file name ('AtomElj.fits')
- **reference** (str) set for the reference e.g. 'L7288'

```
atomneb.get_omij_reference_citation(atom_omij_file, atom, ion, reference)
```

This function returns the reference citation for collision strengths (Omega_ij) from the 2nd binary table extension of the FITS data file ('AtomOmij.fits').

For example:

Returns This function returns the Citation.

Return type str

Parameters

- atom_omij_file (str) the FITS data file name ('AtomOmij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- **reference** (*str*) set for the reference e.g. 'SSB14'

```
atomneb.list aeff collection references (atom rc file, atom, ion, br=None)
```

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc_collection.fits').

For example:

Returns This function returns the references.

Return type an array of strings

- atom_rc_file (str) the FITS data file name ('rc_collection.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'

• **br** (boolean, optional) – set for the branching ratios (Br)

atomneb.list_aeff_he_i_pfsd12_references(atom_rc_file, atom, ion)

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc_he_ii_PFSD12.fits').

For example:

Returns This function returns the references.

Return type an array of strings

Parameters

- atom_rc_file (str) the FITS data file name ('rc_he_ii_PFSD12.fits')
- atom (str) atom name e.g. 'he'
- ion (str) ionic level e.g 'ii'

atomneb.list_aeff_n_ii_fsl13_references(atom_rc_file, atom, ion)

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc_n_iii_FSL13.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = ['atomic-data-rc']
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_n_iii_FSL13.fits')
>> atom='n'
>> ion='iii' # N II
>> list_nii_aeff_reference = atomneb.list_aeff_n_ii_fsl13_references(atom_irc_file, atom, ion)
>> print(list_nii_aeff_reference)
```

Returns This function returns the references.

Return type an array of strings

- atom_rc_file (str) the FITS data file name ('rc_n_iii_FSL13.fits')
- atom (str) atom name e.g. 'n'

• ion (str) – ionic level e.g 'iii'

```
atomneb.list_aeff_o_ii_ssb17_references(atom_rc_file, atom, ion)
```

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc o iii SSB17.fits').

For example:

Returns This function returns the references.

Return type an array of strings

Parameters

- atom_rc_file (str) the FITS data file name ('rc_o_iii_SSB17.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'

atomneb.list_aeff_ppb91_references(atom_rc_file, atom, ion)

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc_PPB91.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = ['atomic-data-rc']
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_PPB91.fits')
>> atom='c'
>> ion='iii'
>> list_cii_aeff_reference = atomneb.list_aeff_ppb91_references(atom_rc_
--file, atom, ion)
>> print(list_cii_aeff_reference)
```

Returns This function returns the references.

Return type an array of strings

- atom_rc_file (str) the FITS data file name ('rc_PPB91.fits')
- atom (str) atom name e.g. 'c'

• ion (str) – ionic level e.g 'iii'

atomneb.list_aeff_sh95_references(atom_rc_file, atom, ion)

This function returns a list for all references of recombination coefficients (Aeff) for given element and ionic level from the FITS data file ('rc_SH95.fits').

For example:

Returns This function returns the references.

Return type an array of strings

Parameters

- atom rc file (str) the FITS data file name ('rc SH95.fits')
- atom (str) atom name e.g. 'h'
- ion (str) ionic level e.g 'ii'

atomneb.list aij references (atom aij file, atom, ion)

This function returns a list for all references of transition probabilities (Aij) for given element and ionic level from the FITS data file ('AtoAij.fits').

For example:

Returns This function returns the references.

Return type an array of data

- atom_rc_file (str) the FITS data file name ('AtoAij.fits')
- atom (str) atom name e.g. 'o'

• ion (str) – ionic level e.g 'iii'

atomneb.list_omij_references(atom_omij_file, atom, ion)

This function returns a list for all references of collision strengths (Omega_ij) for given element and ionic level from the FITS data file ('AtomOmij.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data', 'collection')
>> atom_omij_file = os.path.join(base_dir, data_dir, 'AtomOmij.fits')
>> atom='o'
>> ion='iii'
>> list_oiii_omij_reference = atomneb.list_omij_references(atom_omij_file, atom, ion)
>> print(list_oiii_omij_reference)
AK99 LB94 Pal12-AK99 SSB14
```

Returns This function returns the references.

Return type str

Parameters

- atom_omij_file (str) the FITS data file name ('AtomOmij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'

atomneb.read_aeff_collection(atom_rc_file, atom, ion, br=None, reference=None)

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc_collection.fits').

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_collection.fits')
>> atom='c'
>> ion='iii' # C III
>> cii_rc_data = atomneb.read_aeff_collection(atom_rc_file, atom, ion)
>> n_line = len(cii_rc_data.wavelength)
>> for i in range(0, n_line):
>>
       print(cii_rc_data.wavelength[i], cii_rc_data.a[i],
            cii_rc_data.b[i], cii_rc_data.c[i],
>>
              cii_rc_data.d[i], cii_rc_data.f[i])
  914.00000 0.69280000
                               0.021400000 -0.016300000
                                                                -0.
             -0.88000000

→24310000

  962.00000
               1.0998000 -0.0042000000
                                               -0.027900000
                                                                -0.
\hookrightarrow 22940000
              -0.96560000
```

Returns This function returns the effective recombination coefficients. aeff_data (c_iii_aeff) { Wavelength:0.0,a: 0.0, b: 0.0, c: 0.0, d: 0.0, f: 0.0}, aeff_data (n_iii_aeff) { a: 0.0, b: 0.0, c: 0.0}, aeff_data (n_iii_br) {Wavelength: 0.0, BR: 0.0, g1:0, g2:0, Mult1:", LowerTerm:", UpperTerm:" }, aeff_data (o_iii_aeff) {Term: ", Case1: ", a2: 0.0, a4: 0.0, a5: 0.0, a6: 0.0, b: 0.0, c: 0.0, d: 0.0}, aeff_data (o_iii_br) {Wavelength:double(0.0), Br_A: 0.0, Br_B: 0.0, Br_C: 0.0, g1: 0, g2: 0, Mult1: ", LowerTerm: ", UpperTerm: "}, aeff_data (ne_iii_aeff) {Wavelength:0.0, a: 0.0, b: 0.0, c: 0.0, d: 0.0, f: 0.0, br: 0.0},

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_collection.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'
- **br** (boolean, optional) set for the branching ratios (Br)
- reference (str, optional) set for the reference

```
atomneb.read_aeff_collection_list(atom_rc_file)
```

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc_collection.fits')

Returns This function returns the aeff data list: { Aeff Data:", Extension:0.0}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_collection.fits')

```
atomneb.read_aeff_collection_references(atom_rc_file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_collection.fits').

Returns This function returns the aeff_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_collection.fits')

 $\verb|atomneb.read_aeff_he_i_pfsd12| (atom_rc_file, atom, ion, wavelength=None, reference=None)|$

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc he ii PFSD12.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir, data_dir, 'rc_he_ii_PFSD12.fits')
>> atom='he'
>> ion='ii' # He I
>> hei_rc_data = atomneb.read_aeff_he_i_pfsd12(atom_rc_file, atom, ion)
>> hei_rc_data_wave = atomneb.read_aeff_he_i_pfsd12(atom_rc_file, atom, ion)
atom, wavelength=True)
```

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Returns This function returns the effective recombination coefficients.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_he_ii_PFSD12.fits')
- atom (str) atom name e.g. 'he'
- ion (str) ionic level e.g 'ii'
- wavelength (boolean, optional) set for returning the wavelengths
- reference (str, optional) set for the reference

```
atomneb.read_aeff_he_i_pfsd12_list(atom_rc_file)
```

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc he ii PFSD12.fits')

Returns This function returns the aeff_data_list: { Aeff_Data:", Extension:0.0}

Return type an array of data

Parameters atom_rc_file (str) - the FITS data file name ('rc_he_ii_PFSD12.fits')

```
atomneb.read_aeff_he_i_pfsd12_references(atom_rc_file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_he_ii_PFSD12.fits').

Returns This function returns the aeff data reference: {Reference: ', Citation:'}

Return type an array of data

Parameters atom rc file (str) - the FITS data file name ('rc he ii PFSD12.fits')

atomneb.read_aeff_n_ii_fsl13 (atom_rc_file, atom, ion, wavelength_range, wavelength=None, reference=None)

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc_n_iii_FSL13.fits').

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_n_iii_FSL13.fits')
>> atom='n'
>> ion='iii' # N II
>> wavelength_range=[4400.0, 7100.0]
>> nii_rc_data = atomneb.read_aeff_n_ii_fsl13(atom_rc_file, atom, ion,...
→wavelength_range)
>> nii_rc_data_wave = atomneb.read_aeff_n_ii_fsl13(atom_rc_file, atom,_
→ion, wavelength_range, wavelength=True)
>> print(nii_rc_data.aeff[0])
           79.5000 47.3000
   255.000
                                      12.5000
>> n_line = len(nii_rc_data_wave.wavelength)
>> for i in range(0, n_line):
      print(nii_rc_data_wave.wavelength[i], nii_rc_data_wave.tr[i], nii_
→rc_data_wave.trans[i])
  6413.236g - 4f2p6g G[9/2]o4 - 2p4f F[7/2]e3
   6556.326g - 4f2p6g G[9/2]o5 - 2p4f G[7/2]e4
   6456.976g - 4f2p6g G[9/2]o5 - 2p4f F[7/2]e4
```

Returns This function returns the effective recombination coefficients.

Return type an array of data

Parameters

- atom rc file (str) the FITS data file name ('rc n iii FSL13.fits')
- atom (str) atom name e.g. 'n'
- ion (str) ionic level e.g 'iii'
- wavelength (boolean, optional) set for returning the wavelengths
- reference (str, optional) set for the reference

```
atomneb.read_aeff_n_ii_fsl13_list(atom_rc_file)
```

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc_n_iii_FSL13.fits')

Returns This function returns the aeff_data_list: {Aeff_Data:", Extension:0, IND:long(0), Wavelength: float(0.0), Tr:", Trans: ", T_X: "}

Return type an array of data

Parameters atom rc file (str) - the FITS data file name ('rc n iii FSL13.fits')

```
atomneb.read_aeff_n_ii_fsl13_references(atom_rc_file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_n_iii_FSL13.fits').

Returns This function returns the aeff_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_rc_file (str) - the FITS data file name ('rc_n_iii_FSL13.fits')

atomneb.read_aeff_o_ii_ssb17 (atom_rc_file, atom, ion, case1, wavelength_range, wavelength=None, reference=None)

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc o iii SSB17.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_o_iii_SSB17_orl_
⇔case_b.fits')
>> atom='o'
>> ion='iii' # O II
>> case1='B'
>> wavelength_range=[5320.0, 5330.0]
>> oii_rc_data = atomneb.read_aeff_o_ii_ssb17(atom_rc_file, atom, ion,_
→case1, wavelength_range)
>> oii_rc_data_wave = atomneb.read_aeff_o_ii_ssb17(atom_rc_file, atom,_
⇔ion,
>>
                                                   case1, wavelength_
→range, wavelength=True)
>> print(oii_rc_data.aeff[0])
  1.64100e-30 1.60000e-30 1.56400e-30 1.54100e-30 ...
>> n_line = len(oii_rc_data_wave.wavelength)
>> for i in range(0, n_line):
       print(oii_rc_data_wave.wavelength[i], oii_rc_data_wave.lower_
→term[i], oii_rc_data_wave.upper_term[i])
   5327.172s22p2(1S)3p 2Po
   5325.422s22p2(1S)3p 2Po
   5327.182s22p2(1D)3d 2Ge
```

Returns This function returns the effective recombination coefficients.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_o_iii_SSB17.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- case1 (str) set for the case 'a' or 'b', defualt 'b'
- wavelength_range (array) wavelength range e.g. [5320.0, 5330.0]
- wavelength (boolean, optional) set for returning the wavelengths
- reference (string, optional) set for the reference

```
\verb|atomneb.read_aeff_o_ii_ssb17_list| (atom\_rc\_file) \\
```

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc_o_iii_SSB17.fits')

Returns This function returns the aeff_data_list: {Aeff_Data:", Extension:0, IND:long(0), Wavelength: float(0.0), Case1:", lower_term: '', upper_term: ''}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_o_iii_SSB17.fits')

```
atomneb.read aeff o ii ssb17 references (atom rc file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_o_iii_SSB17.fits').

Returns This function returns the aeff_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_rc_file (str) - the FITS data file name ('rc_o_iii_SSB17.fits')

```
atomneb.read_aeff_ppb91 (atom_rc_file, atom, ion, reference=None)
```

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc_PPB91.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_PPB91.fits')
>> atom='c'
>> ion='iii' # C II
>> cii_rc_data = atomneb.read_aeff_ppb91(atom_rc_file, atom, ion)
>> n_line = len(cii_rc_data.wavelength)
>> for i in range(0, n_line):
            print(cii_rc_data.ion[i], cii_rc_data.case1[i], cii_rc_data.
→wavelength[i],
                  cii_rc_data.a[i], cii_rc_data.b[i], cii_rc_data.c[i],
                  cii_rc_data.d[i], cii_rc_data.br[i], cii_rc_data.q[i],
0.69700000
                                           -0.78400000
  C2+A 9903.4600
             4267.1500
                            1.0110000
                                           -0.75400000
   C2+A
                                                            . . .
```

Returns This function returns the effective recombination coefficients: {Ion: ', Case1:", Wavelength:0.0, a: 0.0, b: 0.0, c: 0.0, d: 0.0, br: 0.0, y: 0.0}

Return type an array of data

Parameters

- atom rc file (str) the FITS data file name ('rc PPB91.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'
- reference (str, optional) set for the reference

 $\verb|atomneb.read_aeff_ppb91_list| (atom_rc_file)$

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc_PPB91.fits')

Returns This function returns the aeff_data_list: { Aeff_Data:", Extension:0.0}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_PPB91.fits')

```
atomneb.read_aeff_ppb91_references(atom_rc_file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_PPB91.fits').

Returns This function returns the aeff_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_PPB91.fits')

```
atomneb.read aeff sh95 (atom rc file, atom, ion, reference=None, case1=None)
```

This function returns the effective recombination coefficients (Aeff) from the table extensions of the FITS data file ('rc_SH95.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_SH95.fits')
>> atom='h'
>> ion='ii' # H I
>> hi_rc_data = atomneb.read_aeff_sh95(atom_rc_file, atom, ion)
>> print(hi_rc_data.aeff[0])
   100.00000
               500.00000
                                  0.0000000 4.2140000e-27
                                                                1.
→7560000e-27 ...
   . . .
```

Returns This function returns the effective recombination coefficients.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_SH95.fits')
- atom (str) atom name e.g. 'h'
- ion (str) ionic level e.g 'ii'
- reference (str, optional) set for the reference
- case1 (boolean, optional) set for the case 'a' or 'b', defualt 'b'

```
atomneb.read_aeff_sh95_list(atom_rc_file)
```

This function returns the list of effective recombination coefficients (Aeff) from the 1st binary table extension of the FITS data file ('rc_SH95.fits')

Returns This function returns the aeff_data_list: { Aeff_Data:", Extension:0.0}

Return type an array of data

Parameters atom_rc_file (str) - the FITS data file name ('rc_SH95.fits')

```
atomneb.read_aeff_sh95_references(atom_rc_file)
```

This function returns the reference list of recombination coefficients (Aeff) from the 2nd binary table extension of the FITS data file ('rc_SH95.fits').

Returns This function returns the aeff_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_rc_file (str) – the FITS data file name ('rc_SH95.fits')

```
atomneb.read_aij (atom_aij_file, atom, ion, reference=None, level_num=None)
```

This function returns the transition probabilities (Aij) from the table extensions of the FITS data file ('AtomAij.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data', 'collection')
>> atom_aij_file = os.path.join(base_dir,data_dir, 'AtomAij.fits')
>> atom='o'
>> ion='iii'
>> reference='FFT04'
>> oiii_aij_data = atomneb.read_aij(atom_aij_file, atom, ion, reference)
>> print(oiii_aij_data.aij)
  0.0000000 2.5960000e-05
                            3.0300000e-11 2.3220000e-06
                                                                 0.
→0000000 0.0021910000
  0.0000000
                 0.0000000 9.6320000e-05
                                              0.0069510000
                                                                0.
→22550000
               230.80000
  0.0000000
                                 0.0000000
                                               0.020290000
                0.0000000
→00069980000
                  576.50000
                 0.0000000
  0.0000000
                                 0.0000000
                                                 0.0000000
                                                                 1.

→6850000 0.0057770000

  0.0000000
                 0.0000000
                                 0.0000000
                                                 0.0000000
                                                                 0.
→0000000 3.7600000e-11
  0.0000000
              0.0000000
                                 0.0000000
                                                 0.0000000
                                                                 0.
               0.0000000
→0000000
```

Returns This function returns the aij_data: { Aij:dblarr(n_level,n_level) }.

Return type an array of data

- atom_rc_file (str) the FITS data file name ('AtoAij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- reference (str, optional) set for the reference
- level_num(str, optional) set for the maximum level number

```
atomneb.read_aij_list(atom_aij_file)
```

This function returns the list of transition probabilities (Aij) from the 1st binary table extension of the FITS data file ('AtomAij.fits').

Returns This function returns the aij_data_list: { Aij_Data:", Extension:0.0}

Return type an array of data

Parameters Atom Aij file (str) – the FITS data file name ('AtomAij.fits')

```
atomneb.read_aij_references(atom_aij_file)
```

This function returns the reference list of transition probabilities (Aij) from the 1nd binary table extension of the FITS data file ('AtomAij.fits').

Returns This function returns the aij_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters Atom_Aij_file (str) – the FITS data file name ('AtomAij.fits')

```
atomneb.read_elj_(atom_elj_file, atom, ion, level_num=None)
```

This function returns the energy levels (Ej) from the table extensions of the FITS data file ('AtomElj.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data', 'collection')
>> atom_elj_file = os.path.join(base_dir,data_dir, 'AtomElj.fits')
>> atom='o'
>> ion='iii'
>> oiii_elj_data=atomneb.read_elj(atom_elj_file, atom, ion, level_num=6)
>> print(np.asarray(oiii_elj_data.j_v))
   0.00000
               1.00000
                           2.00000
                                          2.00000
                                                       0.00000
                                                                    2.
→00000
>> print(np.asarray(oiii_elj_data.ej))
   0.0000000
              113.17800
                                   306.17400
                                                   20273.270
                                                                   43185.
            60324.790
→740
```

Returns This function returns the elj_data:{ Configuration:", Term:", J:", J_v:0.0, Ej:0.0, Reference:"}.

Return type an array of data

Parameters

- atom_elj_file(str) the FITS data file name ('AtomElj.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- **level_num**(*int*, *optional*) set for the maximum level number.

atomneb.read_elj_list(atom_elj_file)

This function returns the list of energy levels (Ej) from the 1st binary table extension of the FITS data file ('AtomElj.fits')

Returns This function returns the elj_data_list: { Elj_Data:", Extension:0.0}

Return type an array of data

Parameters atom elj file (str) – the FITS data file name ('AtomElj.fits')

```
atomneb.read_elj_references(atom_elj_file)
```

This function returns the reference list of energy levels (Ej) from the 2nd binary table extension of the FITS data file ('AtomElj.fits').

Returns This function returns the aij_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_elj_file (str) - the FITS data file name ('AtomElj.fits')

```
atomneb.read omij (atom omij file, atom, ion, reference=None, level num=None)
```

This function returns the collision strengths (omega_ij) from the table extensions of the FITS data file ('AtomOmij.fits').

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data', 'collection')
>> atom_omij_file = os.path.join(base_dir,data_dir, 'AtomOmij.fits')
>> atom='o'
>> ion='iii'
>> reference='SSB14'
>> oiii_omij_data=atomneb.read_omij(atom_omij_file, atom, ion,,,
→reference=reference)
>> print(np.asarray(oiii_omij_data.level1))
  0 1 1
                        1
                              1
        4
>> print (np.asarray(oiii_omij_data.level2))
        2
                3
                       4
                               5
>> print(np.asarray(oiii_omij_data.strength)[0])
   100.00000
                 125.89254
                            158.48932
                                                199.52623
                                                               251.
→18864
```

Returns This function returns the omij_data: { level1:0, level2:0, strength:array(temp_steps)}.

Return type an array of data

Params

```
atom_omij_file [in, required, type=string] the FITS data file name ('AtomOmij.fits')atom [in, required, type=string] atom name e.g. 'o'ion [in, required, type=string] ionic level e.g 'iii'
```

- reference (str, optional) set for the reference e.g. 'SSB14'
- level_num(int, optional) set for the maximum level number.

```
atomneb.read_omij_list(atom_omij_file)
```

This function returns the list of collision strengths (omega_ij) from the 1st binary table extension of the FITS data file ('AtomOmij.fits').

Returns This function returns the omij_data_list: { Omij_Data:", Extension:0.0}

Return type an array of data

Parameters atom_omij_file (str) - the FITS data file name ('AtomOmij.fits')

```
atomneb.read_omij_references(atom_omij_file)
```

This function returns the reference list of collision strengths (omega_ij) from the 2nd binary table extension of the FITS data file ('AtomOmij.fits').

Returns his function returns the aij_data_reference: { Reference:", Citation:"}

Return type an array of data

Parameters atom_omij_file (str) - the FITS data file name ('AtomOmij.fits')

```
atomneb.search_aeff_collection(atom_rc_file, atom, ion, br=None)
```

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rc collection.fits'), and returns the data entry.

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir, data_dir, 'rc_collection.fits')
>> atom='c'
>> ion='iii' # C III
>> list_cii_aeff_data = atomneb.search_aeff_collection(atom_rc_file, atom, ion)
>> print(list_cii_aeff_data)
c_iii_aeff
```

Returns This function returns the Aeff_Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_collection.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'
- **br** (boolean, optional) set for the branching ratios (Br), may not necessary

atomneb.search_aeff_he_i_pfsd12(atom_rc_file, atom, ion)

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rec_he_ii_PFSD12.fits'), and returns the data entry.

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir, data_dir, 'rc_he_ii_PFSD12.fits')
>> atom='he'
>> ion='ii' # He I
>> list_hei_aeff_data = atomneb.search_aeff_he_i_pfsd12(atom_rc_file, ueatom, ion)
>> print(list_hei_aeff_data)
he_ii_aeff_PFSD12 he_ii_aeff_PFSD13
```

Returns This function returns the Aeff_Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_he_ii_PFSD12.fits')
- atom (str) atom name e.g. 'he'
- ion (str) ionic level e.g 'ii'

atomneb.search_aeff_n_ii_fsl13 (atom_rc_file, atom, ion, wavelength)

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rc_n_iii_FSL13.fits'), and returns the data entry.

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_n_iii_FSL13.fits')
>> atom='n'
>> ion='iii' # N II
>> wavelength=5679.56
>> list_nii_aeff_data = atomneb.search_aeff_n_ii_fsl13(atom_rc_file, atom,
→ ion, wavelength)
>> print(np.asarray(list_nii_aeff_data.wavelength))
  5679.56
>> print(np.asarray(list_nii_aeff_data.aeff))
  7810.00 1780.00
                                                  74.4000
                         850.000
                                    151.000
                                                               53.
→1000 47.4000
  7370.00 1700.00 886.000
                                      206.000
                                                  110.000
                                                              80.
→1000 70.8000
  7730.00 1680.00 900.000
                                       239.000
                                                  138.000
                                                               103.
→000 92.9000
  8520.00 1710.00
                          905.000
                                       244.000
                                                   142.000
                                                               107.
→000 97.0000
```

Returns This function returns the Aeff Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_n_iii_FSL13.fits')
- atom (str) atom name e.g. 'n'
- ion (str) ionic level e.g 'iii'
- wavelength (int) set the wavelengths
- reference (str, optional) set for the reference

```
atomneb.search_aeff_o_ii_ssb17 (atom_rc_file, atom, ion, case1, wavelength)
```

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rc_o_iii_SSB17.fits'), and returns the data entry.

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('...', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir,data_dir, 'rc_o_iii_SSB17_orl_
>> atom='o'
>> ion='iii' # 0 II
>> case1='B'
>> wavelength=5325.42
>> list_oii_aeff_data = atomneb.search_aeff_o_ii_ssb17(atom_rc_file, atom,
→ ion, case1, wavelength)
>> print(np.asarray(list_oii_aeff_data.wavelength))
   5325.42
>> print(np.asarray(list_oii_aeff_data.aeff))
   3.41800e-32 3.33300e-32 3.25700e-32 3.20900e-32 3.16800e-32 ...
```

Returns This function returns the Aeff Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_o_iii_SSB17.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'
- case1 (str) set for the case 'a' or 'b', defualt 'b'
- wavelength (float) set the wavelengths

atomneb.search_aeff_ppb91(atom_rc_file, atom, ion)

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rec_PPB91.fits'), and returns the data entry.

Returns This function returns the Aeff Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_PPB91.fits')
- atom (str) atom name e.g. 'c'
- ion (str) ionic level e.g 'iii'

```
atomneb.search_aeff_sh95 (atom_rc_file, atom, ion)
```

This function searches effective recombination coefficients (Aeff) for given element and ionic levels in the FITS data file ('rec_SH95.fits'), and returns the data entry.

For example:

```
>> import atomneb
>> import numpy as np
>> import os
>> base_dir = os.getcwd()
>> data_dir = os.path.join('..', 'atomic-data-rc')
>> atom_rc_file = os.path.join(base_dir, data_dir, 'rc_SH95.fits')
>> atom='h'
>> ion='ii' # H I
>> list_hi_aeff_data = atomneb.search_aeff_sh95(atom_rc_file, atom, ion)
>> print(list_hi_aeff_data)
h_ii_aeff_a h_ii_aeff_b
```

Returns This function returns the Aeff_Data.

Return type an array of data

Parameters

- atom_rc_file (str) the FITS data file name ('rc_SH95.fits')
- atom (str) atom name e.g. 'h'
- ion (str) ionic level e.g 'ii'

```
atomneb.search_aij (atom_aij_file, atom, ion)
```

This function searches transition probabilities (Aij) for given element and ionic levels in the FITS data file ('AtomAij.fits'), and returns the data entry.

Returns This function returns the Aij_Data.

Return type an array of data

Parameters

- Atom_Aij_file (str) the FITS data file name ('AtomAij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii

```
atomneb.search_omij(atom_omij_file, atom, ion)
```

This function searches collision strengths (omega_ij) for given element and ionic levels in the FITS data file ('AtomOmij.fits'), and returns the data entry.

For example:

Returns This function returns the Omij_Data.

Return type an array of data

- atom_omij_file (str) the FITS data file name ('AtomOmij.fits')
- atom (str) atom name e.g. 'o'
- ion (str) ionic level e.g 'iii'