
atomneb Documentation

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USER DOCUMENTATION

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- *User Documentation*
- *API Reference*

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 **CHIANTI 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), `AtomOmi j .fits` for *Collision Strengths* (Ω_{ij}), and `AtomAi j .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 I, He II, C I, C II, C III, C VI, N II, 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 (Kisieliuss 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 \(2017\)](#).

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

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 (E_j)*:

```
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** library:

```
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:*

```
atom_rc_file = os.path.join(base_dir, data_dir, 'rc_collection.fits')
atom='c'
ion='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])
```

which gives:

```

914.00000    0.69280000    0.021400000    -0.016300000    -0.
↪24310000    -0.88000000
962.00000    1.0998000    -0.004200000    -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:*

```

atom_rc_file = os.path.join(base_dir, data_dir, 'rc_PPb91.fits')
atom='c'
ion='iii'
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], cii_rc_
↪data.y[i])

```

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:*

```

atom_rc_file = os.path.join(base_dir, data_dir, 'rc_he_ii_PFSd12.fits')
atom='he'
ion='ii'
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,
↪ wavelength=True)
print(hei_rc_data.aeff[0])

```

which gives:

```

5000.0000      10.000000      -25.379540      -25.058970      -25.
↪948440      -24.651820      -25.637660
...

```

- *FSL13 N II data:*

```

atom_rc_file = os.path.join(base_dir, data_dir, 'rc_n_iii_FSL13.fits')
atom='n'
ion='iii'
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])
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])

```

which gives:

```

255.000      79.5000      47.3000      12.5000      6.20000      4.00000↪
↪      2.86000
258.000      54.4000      29.7000      7.92000      4.11000      2.72000↪
↪      2.00000
310.000      48.1000      23.7000      5.19000      2.55000      1.65000↪
↪      1.21000
434.000      50.3000      23.2000      4.71000      2.26000      1.45000↪
↪      1.05000

6413.23 6g - 4f2p6g G[9/2]o4 - 2p4f 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:

```

aatom_rc_file = os.path.join(base_dir, data_dir, 'rc_o_iii_SSB17_orl_case_
↪b.fits')
atom='o'
ion='iii'
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)

```

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```

oii_rc_data_wave = atomneb.read_aeff_o_ii_ssb17(atom_rc_file, atom, ion,
↪casel, wavelength_range, wavelength=True)
print(oii_rc_data.aeff[0])
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])

```

which gives:

```

1.64100e-30  1.60000e-30  1.56400e-30  1.54100e-30  1.52100e-30  1.
↪50900e-30
...

5327.17  2s22p2 (1S) 3p  2Po
5325.42  2s22p2 (1S) 3p  2Po
5327.18  2s22p2 (1D) 3d  2Ge
5326.84  2s22p2 (1D) 3d  2Ge
...

```


REFERENCES

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

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:

```
>> 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
>> citation = atomneb.get_aeff_collection_reference_citation(atom_rc_file,
↪ atom, ion)
>> print(citation)
Davey, A. R., Storey, P. J. and Kisielius, R., Astron.Astrophys.Suppl.,
↪ 142, 85, 2000
```

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_PFS12.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_he_ii_PFS12.fits')
>> atom='he'
>> ion='ii' # He I
>> reference='PFS13'
>> citation = atomneb.get_aeff_he_i_pfs12_reference_citation(atom_rc_
↪ file, atom, ion, reference=reference)
>> print(citation)
Porter, R. L., Ferland, G. J., Storey, P. J. and Detisch, M. J., MNRAS,
↪ 433L, 89, 2013
```

Returns This function returns the Citation.

Return type str

Parameters

- **atom_rc_file** (*str*) – the FITS data file name ('rc_he_ii_PFS12.fits')
- **atom** (*str*) – atom name e.g. 'he'
- **ion** (*str*) – ionic level e.g. 'ii'
- **reference** (*str, optional*) – set for the reference e.g. 'PFS13'

`atomneb.get_aeff_n_iii_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_iii_fsl13_reference_citation(atom_rc_file,
↪ atom, ion)
>> print(citation)
Fang X., Storey P.J., and Liu X.-W., R. 2011, Astron.Astrophys. 530,
↪ A18; 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:

```
>> 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_o_iii_SSB17_orl_
↪case_b.fits')
>> atom='o'
>> ion='iii' # O II
>> reference='SSB17'
>> citation = atomneb.get_aeff_o_ii_ssb17_reference_citation(atom_rc_file,
↪ atom, ion)
>> print(citation)
Storey, P.J., Sochi, T. and Bastin, R. 2017, MNRAS, 470, 379; VizieR,
↪ On-line Data Catalog: VI/150
```

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,
↪ atom, ion)
```

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```
>> print(citation)
    Pequignot, D., Petitjean, P. and Boisson, C. Astron.Astrophys., 251, ↵
↵ 680, 1991
```

Returns This function returns the Citation.

Return type str

Parameters

- **atom_rc_file**(*str*) – the FITS data file name ('rc_PP91.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*,
case1=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, 41S, 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', default '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')

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'
>> 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:

```
>> 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'
>> citation = atomneb.get_omij_reference_citation(atom_omij_file, atom,
↪ion, reference)
>> print(citation)
    Storey, P. J., Sochi, T., and Badnell, N. R. 2014, Astron.Astrophys.,
↪441, 3028
```

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:

```
>> 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_reference = atomneb.list_aeff_collection_references(atom_
↪rc_file, atom, ion)
>> print(list_cii_aeff_reference)
```

Returns This function returns the references.

Return type an array of strings

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)

`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_PFS12.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_he_ii_PFS12.fits')
>> atom='he'
>> ion='ii' # He I
>> list_hei_aeff_reference = atomneb.list_aeff_he_i_pfsd12_
↪references(atom_rc_file, atom, ion)
>> print(list_hei_aeff_reference)
PFS12 PFS13
```

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_PFS12.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_
↪rc_file, atom, ion)
>> print(list_nii_aeff_reference)
```

Returns This function returns the references.

Return type an array of strings

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’

`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:

```
>> 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_o_iii_SSB17_orl_
↳ case_b.fits')
>> atom='o'
>> ion='iii' # O II
>> list_oii_aeff_reference = atomneb.list_aeff_o_ii_ssb17_references(atom_
↳ rc_file, atom, ion)
>> print(list_oii_aeff_reference)
```

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

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.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:

```
>> 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
>> list_hi_aeff_reference = atomneb.list_aeff_sh95_references(atom_rc_
↪file, atom, ion)
>> print(list_hi_aeff_reference)
```

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:

```
>> 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'
>> list_oiii_aij_reference = atomneb.list_aij_references(atom_aij_file,
↪atom, ion)
>> print(list_oiii_aij_reference)
FFT04-SZ00 FFT04 GMZ97-WFD96 SZ00-WFD96
```

Returns This function returns the references.

Return type an array of data

Parameters

- **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 (Ω_{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_omij_reference = atomneb.list_omij_references(atom_omij_file,
→ atom, ion)
>> print(list_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 (A_{eff}) from the table extensions of the FITS data file (`‘rc_collection.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_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    0.02140000    -0.01630000    -0.
→24310000    -0.88000000
    962.00000    1.0998000    -0.004200000    -0.02790000    -0.
→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')

`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_PFS12.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_PFS12.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, wavelength=True)
```

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```
>> print(hei_rc_data.aeff[0])
5000.0000      10.000000      -25.379540      -25.058970      -25.
→948440      ...
>> n_line = len(hei_rc_data_wave.wavelength)
>> for i in range(0, n_line):
>>     print(hei_rc_data_wave.wavelength[i],
>>           hei_rc_data_wave.lowerterm[i], hei_rc_data_wave.upperterm[i])
2945.00005p{3}P2s{3}S
3188.00004p{3}P2s{3}S
3614.00005p{1}P2s{1}S
...
```

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_PFS12.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_PFS12.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_PFS12.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_PFS12.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_PFS12.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').

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_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])
    255.000      79.5000      47.3000      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', default '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

`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],
↵cii_rc_data.y[i])
C2+A      9903.4600      0.69700000      -0.78400000      ...
C2+A      4267.1500      1.0110000      -0.75400000      ...
...
```

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

`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', default '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  0.0000000  0.020290000  0.
↪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.
↪0000000  0.0000000
```

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

Return type an array of data

Parameters

- `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.
↪740 60324.790
```

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      2      2      2      3
↪ 3      4
>> print(np.asarray(oiii_omij_data.level2))
    0      2      3      4      5      3      4      5      4
↪ 5      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'

Parameters

- **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_PFS12.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_PFS12.fits')
>> atom='he'
>> ion='ii' # He I
>> list_hei_aeff_data = atomneb.search_aeff_he_i_pfs12(atom_rc_file,
↪ atom, ion)
>> print(list_hei_aeff_data)
he_ii_aeff_PFS12 he_ii_aeff_PFS13
```

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_PFS12.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.

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_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      850.000      151.000      74.4000      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_
↪ case_b.fits')
>> atom='o'
>> ion='iii' # O II
>> case1='B'
>> wavelength=5325.42
>> list_oi_aeff_data = atomneb.search_aeff_o_ii_ssb17(atom_rc_file, atom,
↪ ion, case1, wavelength)
>> print(np.asarray(list_oi_aeff_data.wavelength))
5325.42
>> print(np.asarray(list_oi_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', default '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.

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'
>> list_cii_aeff_data = atomneb.search_aeff_ppb91(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_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.

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'
>> list_oiii_aij_data = atomneb.search_aij(atom_aij_file, atom, ion)
>> print(list_oiii_aij_data)
    o_iii_aij_FFT04-SZ00 o_iii_aij_FFT04 o_iii_aij_GMZ97-WFD96 o_iii_aij_
↪SZ00-WFD96
```

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:

```
>> 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_data = atomneb.search_omij(atom_omij_file, atom, ion)
>> print(list_oiii_omij_data)
    o_iii_omij_AK99 o_iii_omij_LB94 o_iii_omij_Pal12-AK99 o_iii_omij_SSB14
```

Returns This function returns the Omij_Data.

Return type an array of data

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'