# PyNeb\_manual\_2

June 2, 2020

## 0.1 Using Atom to compute populations and emissivities

The **Atom** class is equipped with method which give access to the populations, emissivities, and other atomic quantities:

```
[1]: import pyneb as pn
     03 = pn.Atom('0', 3)
     N2 = pn.Atom('N', 2)
     S2 = pn.Atom('S', 2)
[2]: 03.getEnergy(4, unit='eV')
[2]: 2.513565018800101
[3]: 03.getStatWeight(level=4)
[3]: 5.0
[4]: 03.getPopulations(tem=1.0e4, den=1e2)
[4]: array([7.81643685e-01, 1.93870586e-01, 2.44814197e-02, 4.30930821e-06,
            2.97166313e-10])
[5]:
     O3.getPopulations(tem=1.5e4, den=1e2)
[5]: array([8.02454816e-01, 1.75280839e-01, 2.22544005e-02, 9.94241529e-06,
            2.05075367e-09])
[6]: 03.getPopulations(tem=1e4, den=1e2).sum()
[6]: 1.0
[7]: print('Critical densities')
     print('N2' + ' '.join('{:8.2e}'.format(cd) for cd in N2.
     →getCritDensity(tem=12000)))
     print('03' + ' '.join('{:8.2e}'.format(cd) for cd in 03.

    getCritDensity(tem=12000)))
```

Critical densities N20.00e+00 4.06e+01 2.47e+02 9.55e+04 1.74e+07 9.83e+09 030.00e+00 5.31e+02 3.70e+03 7.31e+05 2.61e+07

- [8]: 03.getEmissivity(tem=1e4, den=1e2, wave=5007) # in erg.s-1.cm3
- [8]: array(3.49707371e-21)

Wavelengths should be entered with enough precision to avoid confusion with nearby lines. Use **printTransition** to check if the transition selected by the code actually correponds to the one you intended:

[9]: 03.printTransition(5007)

Input wave: 5007.0

Closest wave found: 5006.8 Relative error: 3E-05 Transition: 4 -> 3

**getTransition** is an abridged version which only returns the tuple of levels rather than an extended output, and is therefore apt to be used in scripts.

- [10]: 03.getTransition(5007)
- [10]: (4, 3)

Or you can explicitly use the levels corresponding to the transition:

- [11]: 03.getEmissivity(tem=1e4, den=1e2, lev\_i=4, lev\_j=3)
- [11]: array(3.49707371e-21)

In the case of **getEmissivity**, tem and den can be arrays. In such a case, if they have different dimensions N and M, the function will return an array of NxM emissivities corresponding to all tem-den combinations; if both arrays have the same dimension, you can obtain the emissivities of either the NxN array of tem-den combinations as in the previous case, or of the 1D, N-length array obtained pairing tem and den element by element. This is controlled by the "product" parameter, the default being True (results is NxN matrix):

```
[12]: 03.getEmissivity([10000, 12000], [100, 500], 4, 2, product=True)
```

- [12]: array([[1.17195376e-21, 1.18350278e-21], [1.79323749e-21, 1.80805851e-21]])
- [13]: 03.getEmissivity([10000, 12000], [100, 500], 4, 2, product=False)
- [13]: array([1.17195376e-21, 1.80805851e-21])

### 0.2 Physical conditions determined from line ratios

The **Atom** object also contains a method to compute the electron temperature or density given a line ratio:

```
[14]: 03.getTemDen(int_ratio=150., den=100., wave1=5007, wave2=4363)
```

[14]: 10070.1086130086

The keyword tem (or den) specifies the supplied value of the temperature (or density). Which quantity is computed (temperature or density) is determined by which quantity is provided to the method: if "den" is given, then "tem" is computed, and vice versa.

If the intensity ratio is a simple ratio of two transitions, you can:

• either give the wavelengths of the two transitions involved:

```
[15]: 03.getTemDen(0.02, den=1.e4, wave1=4363, wave2=5007)
```

[15]: 14923.313659752961

• or give the four levels that define the two transitions, in the following order: (upper level of numerator) (lower level of numerator) (upper level of denominator) (lower level of denominator); e. g.:

```
[16]: 03.getTemDen(0.02, den=1.e4, lev_i1=5, lev_j1=4, lev_i2=4, lev_j2=3)
```

[16]: 14923.313659752961

In the general case of an intensity ratio formed by any number of transitions, an algebraic expression must be supplied as the argument of the keyword **to eval**:

```
[17]: 03.getTemDen(0.02, den=1.e4, to_eval="I(5, 4) / (I(4, 3) + I(4, 2))")
```

[17]: 17201.456195718863

```
[18]: N2.getTemDen(150., den=100., to_eval = "(L(6584) + L(6548)) / L(5755)")
```

[18]: 8303.6441184966

The **to\_eval** argument accepts either I(i, j) or L(wavelength) to identify the transitions involved in the diagnostic. Both can be mixed in the same string. If you do not know what transition corresponds to a given wavelength, use **printTransition** to find it.

The parameters **tem** and **den**, as well as the line ratio, may be arrays (1D or 2D, as in the case of observations obtained from IFUs), in which case the result will have the same shape. Some restrictions can be set to the domain explored by the method when looking for the solution; see the method's documentation for further details.

```
[19]: 03.getTemDen([0.015, 0.019], den=[1.e4, 1.1e4], to_eval="I(5, 4) / (I(4, 3) + \cup + \cup I(4, 2))")
```

```
[19]: array([14936.84544261, 16724.87927208])
```

```
[20]: 03.getTemDen([0.015, 0.019], den=1.e4, to_eval="I(5, 4) / (I(4, 3) + I(4, 2))")
```

```
[20]: array([14936.84544261, 16755.22374476])
```

Notice that if you want to simultaneously determine both temperature and density combining two diagnostics (from two different atoms), you need to use the **getCrossTemDen** method of the **Diagnostic** class

Example of use in the case of density determination:

```
[21]: S2.getTemDen(1.1, tem = 1e4, wave1=6730, wave2 = 6716)
```

[21]: 710.3124452343512

#### 0.3 Ionic abundance determination

The ionic abundance is obtained from the intensity of a line normalized to Hbeta=100.

```
[22]: 03.getIonAbundance(int_ratio=127, tem=1.5e4, den=100., wave=5007)
```

[22]: 1.3536707442595606e-05

```
[23]: S2.getIonAbundance(int_ratio=72, tem=1.5e4, den=100., to_eval='L(6716)+L(6731)')
```

[23]: 7.570636624108791e-07

The ionic abundance from recombination lines is treated below

## 0.4 Creating a dictionary of Atom objects

You can define all the atoms at once and put them in a dictionary by creating each atom at a time through the commands:

```
[24]: 03 = pn.Atom('0', '3')
02 = pn.Atom('0', '2')
N2 = pn.Atom('N', '2')
```

or rather use one of the following shortcuts:

```
[25]: atoms = pn.getAtomDict() # a method always requires parenthesis, even without

→ argument
```

```
warng _ManageAtomicData: rec data not available for A12 warng _ManageAtomicData: rec data not available for Ar2 warng _ManageAtomicData: rec data not available for Ar3 warng _ManageAtomicData: rec data not available for Ar4 warng _ManageAtomicData: rec data not available for Ar5 warng _ManageAtomicData: rec data not available for Ba2
```

```
warng _ManageAtomicData: rec data not available for Ba4
ERROR None: No data for this case B
ERROR None: No data for this case B
warng ManageAtomicData: rec data not available for Ca5
warng ManageAtomicData: rec data not available for C12
warng ManageAtomicData: rec data not available for Cl3
warng ManageAtomicData: rec data not available for Cl4
warng _ManageAtomicData: rec data not available for Fe3
warng ManageAtomicData: rec data not available for K4
warng _ManageAtomicData: rec data not available for K5
warng _ManageAtomicData: rec data not available for Mg5
warng _ManageAtomicData: rec data not available for Mg7
ERROR None: No data for this case B
warng _ManageAtomicData: rec data not available for Na4
warng _ManageAtomicData: rec data not available for Na6
warng _ManageAtomicData: rec data not available for Ne3
warng _ManageAtomicData: rec data not available for Ne4
warng _ManageAtomicData: rec data not available for Ne5
warng _ManageAtomicData: rec data not available for Ne6
warng ManageAtomicData: rec data not available for Ni3
ERROR None: No data for this case B
warng ManageAtomicData: rec data not available for S2
warng _ManageAtomicData: rec data not available for S3
warng _ManageAtomicData: rec data not available for S4
warng _ManageAtomicData: rec data not available for Si2
warng _ManageAtomicData: rec data not available for Si3
warng _ManageAtomicData: rec data not available for Xe3
warng _ManageAtomicData: rec data not available for Xe4
warng _ManageAtomicData: rec data not available for Xe6
warng _ManageAtomicData: rec data not available for Kr3
warng _ManageAtomicData: rec data not available for Kr4
warng _ManageAtomicData: rec data not available for Kr5
warng _ManageAtomicData: rec data not available for Se3
warng _ManageAtomicData: rec data not available for Se4
warng ManageAtomicData: rec data not available for Br3
warng ManageAtomicData: rec data not available for Br4
warng ManageAtomicData: rec data not available for Rb4
warng _ManageAtomicData: rec data not available for Rb5
warng _ManageAtomicData: rec data not available for Rb6
warng _ManageAtomicData: rec data not available for Fe4
warng _ManageAtomicData: rec data not available for Fe5
warng _ManageAtomicData: rec data not available for Fe6
warng _ManageAtomicData: rec data not available for Fe7
warng _ManageAtomicData: rec data not available for Fe2
warng _ManageAtomicData: rec data not available for P2
warng _ManageAtomicData: rec data not available for Te3
```

## [26]: atoms # All the available atoms [26]: {'Al2': Atom Al2 from al\_ii\_atom\_JSP86-HK87-VVF96-KS86.dat and al\_ii\_coll\_KHAF92-TBK85-TBK84.dat, 'Ar2': Atom Ar2 from ar\_ii\_atom\_Bal06.dat and ar\_ii\_coll\_PB95.dat, 'Ar3': Atom Ar3 from ar iii atom MB09.dat and ar iii coll MB09.dat, 'Ar4': Atom Ar4 from ar\_iv\_atom\_MZ82.dat and ar\_iv\_coll\_RB97.dat, 'Ar5': Atom Ar5 from ar\_v\_atom\_LL93-MZ82-KS86.dat and ar\_v\_coll\_GMZ95.dat, 'Ba2': Atom Ba2 from ba\_ii\_atom\_CO4.dat and ba\_ii\_coll\_SB98.dat, 'Ba4': Atom Ba4 from ba\_iv\_atom\_BHQZ95.dat and ba\_iv\_coll\_SB98.dat, 'C1': Atom C1 from c i atom FFS85.dat and c i coll JBK87-PA76.dat, 'C1r': Atom C1 from c\_i\_rec\_P91.func, 'C2': Atom C2 from c ii atom GMZ98.dat and c ii coll BP92.dat, 'C2r': Atom C2 from c\_ii\_rec\_P91.func, 'C3': Atom C3 from c\_iii\_atom\_G83-NS78-WFD96.dat and c\_iii\_coll\_Bal85.dat, 'C4': Atom C4 from c\_iv\_atom\_WFD96.dat and c\_iv\_coll\_AKO4.dat, 'Ca5': Atom Ca5 from ca\_v\_atom\_M83-KS86.dat and ca\_v\_coll\_GMZ95.dat, 'Cl2': Atom Cl2 from cl\_ii\_atom\_MZ83.dat and cl\_ii\_coll\_T04.dat, 'Cl3': Atom Cl3 from cl\_iii\_atom\_M83-KS86.dat and cl\_iii\_coll\_BZ89.dat, 'C14': Atom C14 from cl\_iv\_atom\_KS86-MZ82-EM84.dat and cl\_iv\_coll\_GMZ95.dat, 'Fe3': Atom Fe3 from fe\_iii\_atom\_Q96\_J00.dat and fe\_iii\_coll\_Z96.dat, 'K4': Atom K4 from k\_iv\_atom\_M83-KS86.dat and k\_iv\_coll\_GMZ95.dat, 'K5': Atom K5 from k\_v\_atom\_M83-KS86.dat and k\_v\_coll\_BZL88.dat, 'Mg5': Atom Mg5 from mg\_v\_atom\_GMZ97.dat and mg\_v\_coll BZ94.dat, 'Mg7': Atom Mg7 from mg\_vii\_atom\_GMZ97.dat and mg\_vii\_coll\_LB94-U.dat, 'N1': Atom N1 from n i atom KS86-WFD96.dat and n i coll PA76-DMR76.dat, 'N1r': Atom N1 from n\_i\_rec\_P91.func, 'N2': Atom N2 from n\_ii\_atom\_FFTO4.dat and n\_ii\_coll\_T11.dat, 'N2r': Atom N2 from n\_ii\_rec\_FSL11.func, 'N3': Atom N3 from n\_iii\_atom\_GMZ98.dat and n\_iii\_coll\_BP92.dat, 'N3r': Atom N3 from n\_iii\_rec\_P91.func, 'N4': Atom N4 from n\_iv\_atom\_WFD96.dat and n\_iv\_coll\_RBHB94.dat, 'Na4': Atom Na4 from na\_iv\_atom\_GMZ97.dat and na\_iv\_coll\_BZ94.dat, 'Na6': Atom Na6 from na\_vi\_atom\_GMZ97.dat and na\_vi\_coll\_LB94.dat, 'Ne2': Atom Ne2 from ne\_ii\_atom\_Bal06.dat and ne\_ii\_coll\_GMB01.dat, 'Ne2r': Atom Ne2 from ne\_ii\_rec\_KSDN98.func, 'Ne3': Atom Ne3 from ne\_iii\_atom\_GMZ97.dat and ne\_iii\_coll\_McLB00.dat, 'Ne4': Atom Ne4 from ne\_iv\_atom\_BBZ89-BK88.dat and ne\_iv\_coll\_G81.dat, 'Ne5': Atom Ne5 from ne\_v\_atom\_GMZ97-U-BD93.dat and ne\_v\_coll\_DPNP13.dat, 'Ne6': Atom Ne6 from ne\_vi\_atom\_GMZ98.dat and ne\_vi\_coll\_ZGP94.dat, 'Ni3': Atom Ni3 from ni\_iii\_atom\_B01.dat and ni\_iii\_coll\_B01.dat, 'O1': Atom O1 from o\_i\_atom\_WFD96.dat and o\_i\_coll\_BK95.dat, 'O1r': Atom O1 from o\_i\_rec\_P91.func, 'O2': Atom O2 from o\_ii\_atom\_FFTO4.dat and o\_ii\_coll\_KalO9.dat, 'O2r': Atom O2 from o\_ii\_rec\_SSB17-B-opt.hdf5,

'03': Atom 03 from o\_iii\_atom\_FFT04-SZ00.dat and o\_iii\_coll\_SSB14.dat,

'O3r': Atom O3 from o\_iii\_rec\_P91.func,

```
'O4r': Atom O4 from o iv rec P91.func,
       '05': Atom 05 from o_v_atom_H80-NS79.dat and o_v_coll_BBDK85.dat,
       'S2': Atom S2 from s_ii_atom_PKW09.dat and s_ii_coll_TZ10.dat,
       'S3': Atom S3 from s_iii_atom_PKW09.dat and s_iii_coll_TG99.dat,
       'S4': Atom S4 from s_iv_atom_JKD86-DHKD82.dat and s_iv_coll_DHKD82.dat,
       'Si2': Atom Si2 from si_ii_atom_BL93-CSB93-N77.dat and si_ii_coll_DK91.dat,
       'Si3': Atom Si3 from si_iii_atom_M83-OKH88-FW90-KS86.dat and
      si iii coll DK94.dat,
       'Xe3': Atom Xe3 from xe_iii_atom_BHQZ95.dat and xe_iii_coll_SB98.dat,
       'Xe4': Atom Xe4 from xe_iv_atom_BHQZ95.dat and xe_iv_coll_SB98.dat,
       'Xe6': Atom Xe6 from xe_vi_atom_BHQZ95.dat and xe_vi_coll_SB98.dat,
       'Kr3': Atom Kr3 from kr_iii_atom_BH86.dat and kr_iii_coll_S97.dat,
       'Kr4': Atom Kr4 from kr_iv_atom_BH86.dat and kr_iv_coll_S97.dat,
       'Kr5': Atom Kr5 from kr_v_atom_BH86.dat and kr_v_coll_S97.dat,
       'Se3': Atom Se3 from se_iii_atom_BH86.dat and se_iii_coll_S97.dat,
       'Se4': Atom Se4 from se_iv_atom_B05.dat and se_iv_coll_B05.dat,
       'Br3': Atom Br3 from br_iii_atom_BH86.dat and br_iii_coll_S97.dat,
       'Br4': Atom Br4 from br_iv_atom_BH86.dat and br_iv_coll_S97.dat,
       'Rb4': Atom Rb4 from rb_iv_atom_BH86.dat and rb_iv_coll_S97.dat,
       'Rb5': Atom Rb5 from rb_v_atom_BH86.dat and rb_v_coll_S97.dat,
       'Rb6': Atom Rb6 from rb_vi_atom_BH86.dat and rb_vi_coll_S97.dat,
       'Fe4': Atom Fe4 from fe_iv_atom_FFRRO8.dat and fe_iv_coll_ZP97.dat,
       'Fe5': Atom Fe5 from fe v atom Nal00.dat and fe v coll BGMcL07.dat,
       'Fe6': Atom Fe6 from fe_vi_atom_CP00.dat and fe_vi_coll_CP99.dat,
       'Fe7': Atom Fe7 from fe vii atom WB08.dat and fe vii coll WB08.dat,
       '3He2': Atom 3He2 from 3he_ii_atom_cloudy.dat and 3he_ii_coll_cloudy.dat,
       'Fe2': Atom Fe2 from fe_ii_atom_B15.dat and fe_ii_coll_B15.dat,
       'P2': Atom P2 from p_ii_atom_MZ82.dat and p_ii_coll_T04.dat,
       'Te3': Atom Te3 from te_iii_atom_M18.dat and te_iii_coll_M18.dat}
     It is also possible to select only a subset of the elements or ions available by specifying the arguments
     elem list or atom list:
[27]: print(len(atoms))
     73
[28]: atoms = pn.getAtomDict(elem_list=['C', 'N', 'O']) # all the ions with spectral
       → from 1 to 6 are created
     ERROR None: No data for this case B
     ERROR None: No data for this case B
     warng _ManageAtomicData: data for C5 not available
     warng _ManageAtomicData: data for C5 not available
     warng _ManageAtomicData: data for C5 not available
     warng _ManageAtomicData: data for C6 not available
```

'04': Atom 04 from o\_iv\_atom\_GMZ98.dat and o\_iv\_coll\_BP92.dat,

warng \_ManageAtomicData: data for C6 not available

```
warng _ManageAtomicData: data for C6 not available
     warng _ManageAtomicData: data for C7 not available
     warng _ManageAtomicData: data for C7 not available
     warng _ManageAtomicData: data for C7 not available
     ERROR None: No data for this case B
     warng _ManageAtomicData: atom data not available for N5
     warng ManageAtomicData: coll data not available for N5
     ERROR None: No data for this case B
     warng ManageAtomicData: data for N6 not available
     warng _ManageAtomicData: data for N6 not available
     warng _ManageAtomicData: data for N6 not available
     warng _ManageAtomicData: data for N7 not available
     warng _ManageAtomicData: data for N7 not available
     warng _ManageAtomicData: data for N7 not available
     ERROR None: No data for this case B
     warng _ManageAtomicData: atom data not available for O6
     warng _ManageAtomicData: coll data not available for 06
     ERROR None: No data for this case B
     warng _ManageAtomicData: data for O7 not available
     warng ManageAtomicData: data for 07 not available
     warng _ManageAtomicData: data for O7 not available
[29]: atoms # All the CNO available atoms
[29]: {'C1': Atom C1 from c_i_atom_FFS85.dat and c_i_coll_JBK87-PA76.dat,
       'C1r': Atom C1 from c_i_rec_P91.func,
       'C2': Atom C2 from c_ii_atom_GMZ98.dat and c_ii_coll_BP92.dat,
       'C2r': Atom C2 from c_ii_rec_P91.func,
       'C3': Atom C3 from c_iii_atom_G83-NS78-WFD96.dat and c_iii_coll_Bal85.dat,
       'C4': Atom C4 from c_iv_atom_WFD96.dat and c_iv_coll_AKO4.dat,
       'N1': Atom N1 from n_i_atom_KS86-WFD96.dat and n_i_coll_PA76-DMR76.dat,
       'N1r': Atom N1 from n_i_rec_P91.func,
       'N2': Atom N2 from n_ii_atom_FFTO4.dat and n_ii_coll_T11.dat,
       'N2r': Atom N2 from n_ii_rec_FSL11.func,
       'N3': Atom N3 from n_iii_atom_GMZ98.dat and n_iii_coll_BP92.dat,
       'N3r': Atom N3 from n_iii_rec_P91.func,
       'N4': Atom N4 from n_iv_atom_WFD96.dat and n_iv_coll_RBHB94.dat,
       'O1': Atom O1 from o_i_atom_WFD96.dat and o_i_coll_BK95.dat,
       'O1r': Atom O1 from o_i_rec_P91.func,
       'O2': Atom O2 from o_ii_atom_FFTO4.dat and o_ii_coll_KalO9.dat,
       'O2r': Atom O2 from o_ii_rec_SSB17-B-opt.hdf5,
       '03': Atom 03 from o_iii_atom_FFT04-SZ00.dat and o_iii_coll_SSB14.dat,
       'O3r': Atom O3 from o_iii_rec_P91.func,
       'O4': Atom O4 from o_iv_atom_GMZ98.dat and o_iv_coll_BP92.dat,
       'O4r': Atom O4 from o_iv_rec_P91.func,
       'O5': Atom O5 from o_v_atom_H80-NS79.dat and o_v_coll_BBDK85.dat}
```

```
[30]: atoms = pn.getAtomDict(atom_list=['02', '03', 'Ar3', 'N2'])
     warng _ManageAtomicData: rec data not available for Ar3
[31]:
     atoms
[31]: {'O2': Atom O2 from o_ii_atom_FFTO4.dat and o_ii_coll_KalO9.dat,
       'O2r': Atom O2 from o_ii_rec_SSB17-B-opt.hdf5,
       '03': Atom 03 from o_iii_atom_FFT04-SZ00.dat and o_iii_coll_SSB14.dat,
       'O3r': Atom O3 from o_iii_rec_P91.func,
       'Ar3': Atom Ar3 from ar iii atom MB09.dat and ar iii coll MB09.dat,
       'N2': Atom N2 from n_ii_atom_FFTO4.dat and n_ii_coll_T11.dat,
       'N2r': Atom N2 from n_ii_rec_FSL11.func}
     In all these cases, a dictionary is created whose keys are the conventional atom names
     <element><spectrum>, and the corresponding entries the atoms themselves; e. g.:
[32]: atoms['N2']
[32]: Atom N2 from n_ii_atom_FFT04.dat and n_ii_coll_T11.dat
[33]: atoms['N2'].getEmissivity(tem=1e4, den=1e2, wave=6584)
                                                                   # example of use
[33]: array(5.9490128e-21)
     This can be useful if you need to loop on a list of atoms, to plot atomic data for example. To see
     what atoms have been created (which is limited by the data included in the selected atomic data
     set), enter:
[34]: atoms.keys()
[34]: dict_keys(['02', '02r', '03', '03r', 'Ar3', 'N2', 'N2r'])
     If you want to be able to access them directly rather than through a dictionary, input from the
     command line:
[35]: for key in atoms.keys():
          vars()[key] = atoms[key]
     and then you will be able to do the following:
[36]: Ar3.NLevels
```

[36]: 5