

*AHELP for CIAO 3.4*

# ionbal

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## Synopsis

Calculate collisional ionization equilibrium ion balance.

## Syntax

```
result = ionbal(z, ion, temperature)
```

## Description

'ionbal' calculates the fractional abundance of an ion in collisional ionization equilibrium at a particular temperature or temperatures. By default, the ionization balance of Mazzotta et al. (1998, A&AS, v133, p403) is used.

ionbal is a GUIDE routine, which must be initialized using the require("guide") command in chips or sherpa. GUIDE uses the optional ATOMDB database, and this command will fail if the ATOMDB is not available on your system.

## Example 1

```
sherpa> require("guide")
GUIDE Initialized using ATOMDB v1.3.0
sherpa> o7 = ionbal(8,7,1.e6)
sherpa> print(o7)
0.986722
sherpa> o8 = ionbal(8,8,1.e6)
sherpa> print(o8)
0.00876597
```

Puts the fractional abundance of O VII in collisional equilibrium at T = 1.e6 K into the S-lang variable o7, and the fractional abundance of O VIII at the same temperature into the variable o8. The variable name is arbitrary. This demonstrates that observing an O VII line in a 1.e6 K plasma is not unusual, but a strong O VIII would be unexpected.

## Example 2

```
chips> require("guide")
GUIDE Initialized using ATOMDB v1.3.0
chips> T = [1.e5:1.e7:1.e4]
chips> fe17 = ionbal(26, 17, T)
chips> curve(T,fe17)
```

This shows how to set up a vector of temperatures T (from 1.e5 to 1.e7 K, in steps of 1.e4 K) and then calculate the fractional abundance of Fe XVII at those temperatures, putting the result into the S-lang variable fe17. Alternatively, a logarithmically spaced temperature vector could be created with  $T = 10^{[5:7:0.1]}$ . The ionization balance is then plotted using the command "curve( T, fe17 )".

## Parameters

name	type	ftype	min	max	units	reqd
Z	integer	input	1	28		yes
Ion	integer	input	1	28		yes
T	float	input	1.E4	1.E9	K	yes

## Detailed Parameter Descriptions

**Parameter=Z (integer required filetype=input min=1 max=28)**

*The atomic number for the element of interest.*

**Parameter=Ion (integer required filetype=input min=1 max=28)**

*The ion number (starting at 1 for the neutral ion) for the ion of interest.*

**Parameter=T (float required filetype=input min=1.E4 max=1.E9 units=K)**

*The temperature(s) (in K or keV) of interest. Values below 100 are assumed to be in keV; values above this are in K. Data are only available between  $10^4 - 10^9$  K.*

## Bugs

See the [Sherpa bug pages](#) online for an up-to-date listing of known bugs.

## See Also

*guide*

[describe](#), [identify](#), [mdl2latex](#), [strong](#)

## Ahelp: ionbal – CIAO 3.4

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URL:  
<http://cxc.harvard.edu/ciao3.4/ionbal.html>  
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