# Many-Electron Atoms

## Thornton and Rex, Ch. 8

In principle, can now solve Sch. Eqn for any atom.

In practice, -> Complicated!

#### Goal--

To explain properties of elements from principles of quantum theory (without exact solutions)

- Elements distinguished by nuclear charge Z (= number of electrons)
- To first approx., each electron moves in electric field of nucleus + remaining electrons:

electron state: 
$$(n, \ell, m_{\ell}, m_{s})$$

shell subshell

n labels energy, but no simple formula.
 $\ell$  subshells no longer completely degenerate.

- Principles for filling electron states:
  - 1) Always fill lowest energy state first.
  - 2) No two electrons can have same quantum numbers  $(n, \ell, m_{\ell}, m_{s})$ .

#### Pauli Exclusion Principle

No two electrons can occupy the same quantum state.

### Building up atomic structure of atoms

Hydrogen

+1/2

Helium

0 + 1/2

-1/2

Helium has a closed shell.

For Lithium, now add n=2 electron, but  $\ell = 0$  or  $\ell = 1$ ?

Smaller ℓ always has lower energy.

Lithium

+1/2

-1/2

+1/2

	<u>n</u>	<u>ℓ</u>	$\underline{m}_{\underline{\ell}}$	$\underline{m}_{\mathtt{s}}$
Hydrogen	1	0	0	+1/2
Helium	1	0	0	-1/2
Lithium	2	0	0	+1/2
Beryllium	2	0	0	-1/2
Boron	2	1	-1	+1/2
Carbon	2	1	0	+1/2
Nitrogen	2	1	+1	+1/2
Oxygen	2	1	-1	-1/2 <del>Y</del>
Flourine	2	1	0	-1/2
Neon	2	1	+1	+1/2 +1/2 -1/2 -1/2 -1/2 +1/2
Sodium	3	0	0	· •/ •
Magnesium	3	0	0	-1/2 tsp
Aluminum	3	1	-1	+1/2
Silicon	3	1	0	+1/2
Phosporus	3	1	+1	+1/2
Sulfur	3	1	-1	-1/2
Chlorine	3	1	0	-1/2
Argon	3	1	+1	-1/2
Potassium	4	0	0	+1/2

#### Chemical properties of elements

Electrons in outermost, largest n orbits are most weakly bound. They determine the chemical properties of the elements. Elements with similar electron structure have similar properties.

Inert or Noble Gases
 Closed p subshell (s for He).
 He (1s²), Ne (2s²2p6), Ar (3s²3p6)

#### Alkalis

Have single electron electron outside closed shell.

Li (2s1), Na (3s1), K (4s1)

#### · Halogens

Are one electron short of a closed shell. F  $(2s^22p^5)$ , Cl  $(3s^23p^5)$ 

## Total Angular Momentum

Consider a 1-electron atom (or with just 1 electron outside closed shell).

It has Orbital Angular momentum L and Spin Angular momentum 5.

These can be combined to give Total Angular momentum  $\vec{J} = \vec{L} + \vec{5}$ .

J is quantized with

$$J = \sqrt{j(j+1)}$$
 h

and

$$J_z = m_j \hbar$$

where  $j = \ell \pm s$ 

or 
$$j = \ell \pm 1/2$$
 (since  $s = 1/2$ )

j will be half-integral (1/2, 3/2, 5/2, ...) m<sub>j</sub> will also be half-integral, ranging from -j to j.

Example:  $\ell = 1$ , s = 1/2

$$m_{\ell} = (1,0,-1)$$
  $m_{s} = (-1/2,+1/2)$ 

3 2 = 6 states

Can combine into

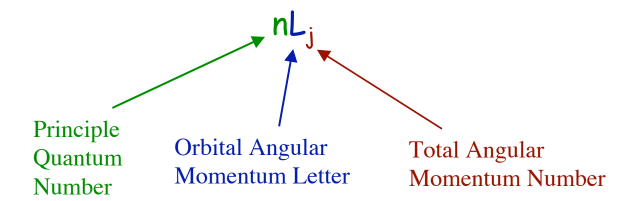
$$j = 3/2 = 1 + 1/2$$
  
 $m_j = (-3/2, -1/2, +1/2, +3/2)$  (4 states)

or

$$j = 1/2 = 1 - 1/2$$
  
 $m_i = (-1/2,+1/2)$  (2 states)

Total number of j-states is 6 = 4 + 2.

### Spectroscopic notation



Examples:  $2S_{1/2}$   $3P_{3/2}$  etc.

## Spin-Orbit Coupling

- Recall, coupling of spin to a magnetic field shifts the energy  $(V_B = -\vec{\mu}_s \cdot \vec{B})$ .
- Motion of electron produces an "internal" magnetic field.

So there is an additional contribution to the energy:

$$V_{SL} = -\mu_{s} \cdot \overrightarrow{B}_{int}$$
Proportional to -S

Proportional to L

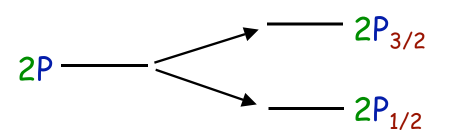
$$V_{SL} \propto \vec{S} \cdot \vec{L}$$

#### This is the Spin-Orbit Coupling:

$$V_{SL} \propto \vec{S} \cdot \vec{L}$$

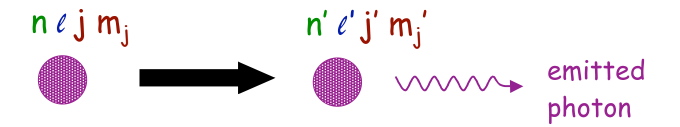
Now states with definite energy do not have unique L and S quantum numbers  $(m_e, m_s)$ . We must use J quantum numbers  $(j, m_i)$ .

States with  $j = \ell - 1/2$  have slightly less energy than states with  $j = \ell + 1/2$ .



(States with different m<sub>j</sub> are still degenerate for each j.)

## Selection Rules



#### Allowed transitions:

• lifetimes  $\tau \sim 10^{-9}$  sec

$$\Delta n = anything$$
,  $\Delta \ell = \pm 1$ ,

$$\Delta j = 0, \pm 1, \qquad \Delta m_j = 0, \pm 1$$

#### Forbidden transitions:

lifetimes much longer

Ex. 
$$2s \rightarrow 1s$$
,  $\tau \sim 1/7$  sec

## Many-Electron Atoms

A careful analysis involving  $\vec{L}$  and  $\vec{S}$  in multi-electron atoms is very complicated.

#### Hund's Rules

(Empirical rules for filling a subshell, while minimizing the energy)

- 1) The total Spin should be maximized (without violating Pauli Exclusion Principle).
- 2) Without violating Rule 1, the Orbital Angular momentum should also be maximized.

#### Handwaving explanation:

Electrons repel each other, so we want them as far from each other as possible.

- 1) If spins of two electrons are aligned (for maximum  $\vec{s}$ ), then Pauli Exclusion Principle says they must have different  $\vec{L}$  orbits. They will tend to be farther apart.
- 2) If the L orbits are aligned (although with different magnitudes), then the electrons will travel around the nucleus in the same direction, so they don't pass each other as often.

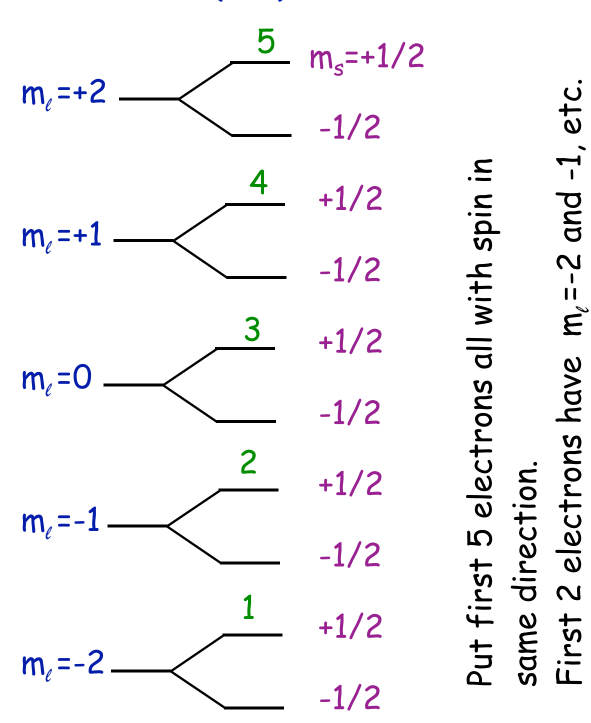
#### Example:

A d subshell ( $\ell = 2$ ) can contain 10 electrons.

$$m_{\ell}=+2$$
 $m_{s}=+1/2$ 
 $-1/2$ 
 $m_{\ell}=+1$ 
 $-1/2$ 
 $m_{\ell}=0$ 
 $-1/2$ 
 $m_{\ell}=-1$ 
 $-1/2$ 
 $m_{\ell}=-1$ 
 $-1/2$ 
 $m_{\ell}=-2$ 
 $-1/2$ 

#### Example:

A d subshell ( $\ell = 2$ ) can contain 10 electrons.



## Many-Electron Atoms

For many-electron atoms there is now orbit-orbit and spin-spin interactions, in addition to spin-orbit interactions.

Consider simplest case of 2 electrons with  $\vec{L}_1$ ,  $\vec{S}_1$  and  $\vec{L}_2$ ,  $\vec{S}_2$ .

Only "good" quantum number is associated with total angular momentum

$$\vec{J} = \vec{L}_1 + \vec{L}_2 + \vec{S}_1 + \vec{S}_2$$
.

(By "good", I mean states with definite energy have definite j and  $m_{j}$ .)

How can we describe atom to best understand energy levels?

### LS, or Russell-Saunders, Coupling

For most atoms the <u>spin-orbit</u> coupling is relatively weak. Then it makes sense to add the angular momentum in steps:

First, 
$$\vec{L} = \vec{L}_1 + \vec{L}_2$$
  
 $\vec{S} = \vec{S}_1 + \vec{S}_2$ 

Then 
$$\vec{J} = \vec{L} + \vec{S}$$

For 2 electrons the Total Spin Quantum Number S is = 0 (spins anti-parallel) or = 1 (spins parallel).

The Total Orbital Angular Momentum Quantum Number L is an integer in the range between  $|\ell_1 - \ell_2|$  and  $|\ell_1 + \ell_2|$ .

The Total Angular Momentum Quantum Number J is an integer in the range between |L-S| and |L+S|.

Note that for S=0, there is  $\underline{1}$  value of J, given by J=L. This state is called a <u>Singlet</u>.

For S=1, there are 3 values of J, given by J=L-1, J=L, J=L+1. These states are called a Triplet.

In general, the <u>multiplicity</u> of the states is given by (25+1).

The Spectroscopic notation is

$$n^{(25+1)}L_{J}$$

#### Example:

2 electrons, one in 4p, other in 4d.

I.e., n=4, 
$$\ell_1 = 1$$
,  $s_1 = 1/2$   
 $\ell_2 = 2$ ,  $s_2 = 1/2$ 

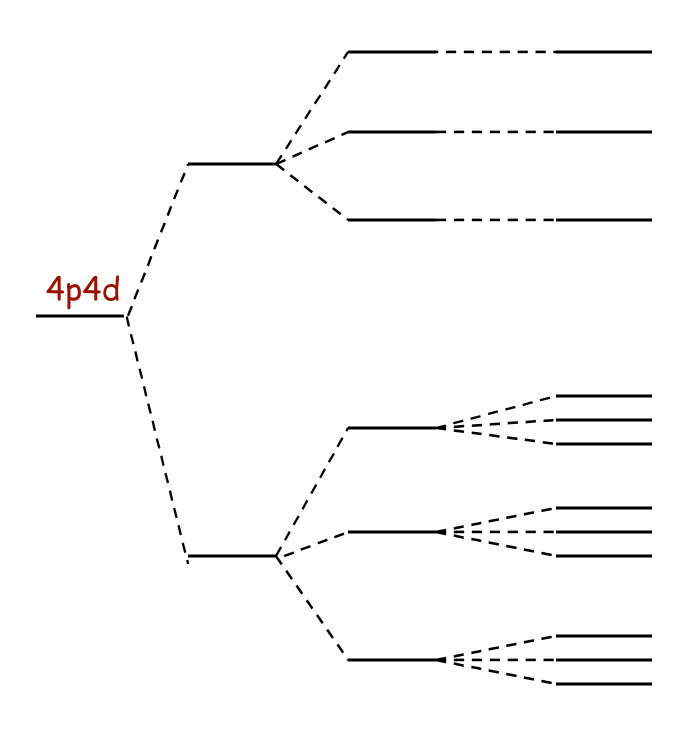
Possible values of 5:

Possible values of L:

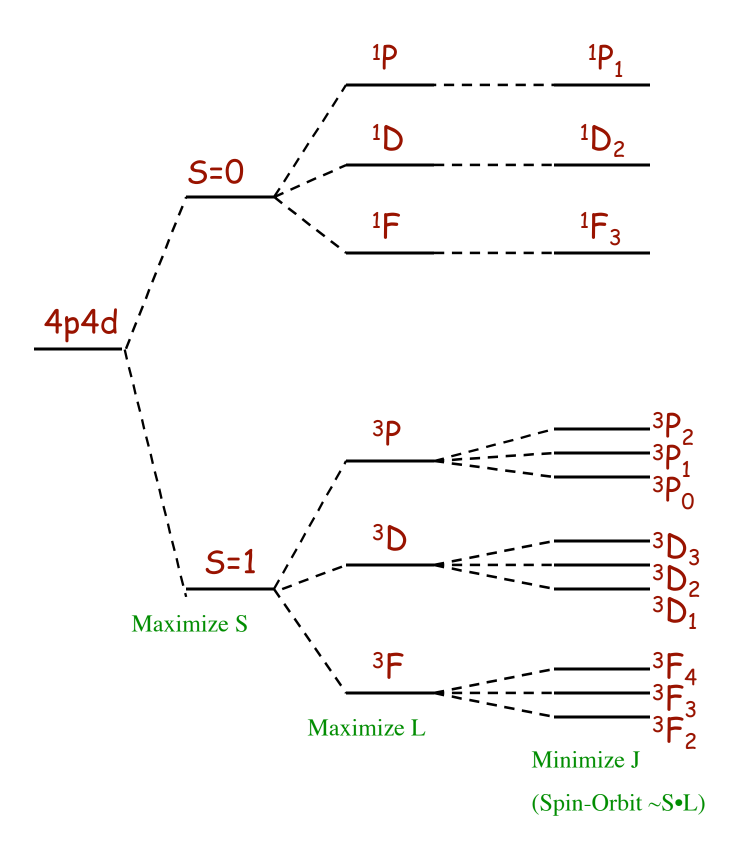
$$L=1, 2, or 3$$

Possible values of J:

Use Hund's rules to order the energies.



Use Hund's rules to order the energies.



#### Example:

Helium 152

$$\ell_1 = 0$$
,  $s_1 = 1/2$   
 $\ell_2 = 0$ ,  $s_2 = 1/2$ 

Possible values of S=0,1

Possible values of L=0

Possible values of J=0,1

States: <sup>1</sup>S<sub>0</sub>, (<sup>3</sup>S<sub>1</sub>)

not allowed by Pauli Exclusion (requires both electrons all same QN's)

If one electron is excited to 2s, so the state is 1s2s, then

both  ${}^{1}S_{0}$ ,  ${}^{3}S_{1}$  are allowed.

### jj Coupling

For high-Z elements the <u>spin-orbit</u> coupling is large for each electron. Now add the angular momentum:

First, 
$$\vec{J}_1 = \vec{L}_1 + \vec{S}_1$$
  
 $\vec{J}_2 = \vec{L}_2 + \vec{S}_2$ 

Then 
$$\vec{J} = \vec{J}_1 + \vec{J}_2$$

#### Anomalous Zeeman Effect

Recall, energy shift in external magnetic field:

$$V_B = -\overrightarrow{\mu} \cdot \overrightarrow{B}$$

The magnetic moment gets both orbital and spin contributions:

$$\vec{\mu} = \vec{\mu}_L + \vec{\mu}_S = \frac{-e}{2m} \left[ \vec{L} + 2 \vec{S} \right]$$

If S=0, this is simple. It is just the Normal Zeeman effect. Energy levels split according to  $m_{\ell}$  values:

$$V_B = m_\ell \mu_B B$$

But..... most atoms are not "Normal".

If both S and L are nonzero, the spin-orbit coupling requires us to use J-states. Projecting  $\vec{\mu}$  onto  $\vec{J}$  gives

$$V_{B} = \frac{e}{2m} g \vec{J} \cdot \vec{B}$$
$$= \mu_{B} g m_{J} B$$

where the projection factor (called the Landé g factor) is

$$g = 1 + \frac{J(J+1) + S(S+1) - L(L+1)}{2J(J+1)}$$

This is the Anomalous Zeeman Effect.