THE AUFBAU PRINCIPAL, KRAMERS RELATION, SELECTION RULES, AND RYBERG ATOMS

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ABSTRACT. A properly prepared Rydberg atom can be used in experiments effectively as a two-state atomic system. In this compendium we briefly discuss the nature of Rydberg Atoms.

1. FROM THE HYDROGENIC: THE AUFBAU PRINICIPAL

Though we demonstrated the basic theory of the hydrogen atom in previous sections, we now wish to consider a simple model for more complex atoms. An atom with atomic number Z, i.e. Z protons, if it is neutral, will have Z matching electrons whereby we might write the Hamiltonian as,

(1.1)
$$H = \sum_{j=1}^{Z} \left(-\frac{-\hbar^2}{2m} \nabla_j^2 - \left(\frac{1}{4\pi\epsilon_0}\right) \frac{Ze^2}{r_j} \right) + \frac{1}{2} \left(\frac{1}{4\pi\epsilon_0}\right) \sum_{j\neq k}^{Z} \frac{e^2}{|r_j - r_k|}$$

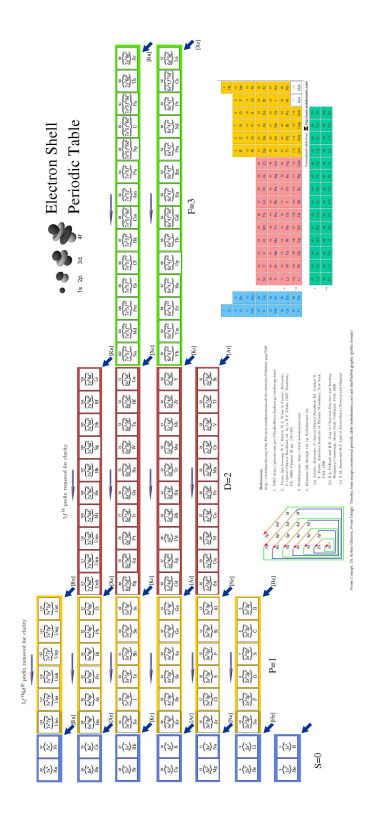
A simple model for hydrogenic atoms invokes the Aufbau principal, of building up atoms from the hydrogen case. We know that the electron requires anti-symetric wave functions, and a generally (though not completely) accurate set of rules called Hund's Rules give us a good guide for "building" up the other atoms from hydrogen, namely:

- States with larger S lie lower
- For an S state, the higher the L, the lower the energy
- For any L and S, if the shell is less than half-filled, the lower state has J = |L S|, else the lower state is J = L + S.

Gilmore and Jones have produced a periodic table that demonstrates this shell filling model (though it ignores the exceptional cases for simplicity) that is reproduced on in Figure 1. A completely filled shell is much less reactive (has larger ionization binding energy) than a nearly empty shell (the Alkali metals for example), and so the latter type are useful for many optical experiments.

We are particulary interested in atoms which have very hydrogenic properties and whose outer most electron (typically, an Alkali metal with only one valence electron) is in a state with an extremely highly principal quantum number $n \ge 50$. Rubidium is a common example. (In the experiments done on trapped ions at the National Institute of Standards and Technology, positively ionized Beryllium is used. The ionization is necessary for trapping; and the missing valance electron leaves the ion with only one valance. However, the ion need not be in the Rydberg state for these experiments).

Here we briefly survey the systematic way in which the periodic table is built up (Aufbau). We demonstrate a system this author uses to simplify the process of spectral identification of elements, where the spectral identification is recorded as ${}^{2S+1}L_J$. Here the capital letters are used to indicate we are dealing with *totals*.



From Figure 1 we have the following rules for energy level buildup (excluding the 1s starter). Here we start from the bottom right, work our way diagonally through the shells, and then loop back to the next diagonal circuit. We find:

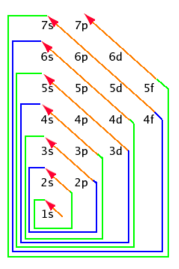


FIGURE 1.1. General order of energy levels in the Aufbau buildup process

- (1) Every n level will starts off exactly two loops.
- (2) Each l value starts off two loops as well, with a higher n for the second case.
- (3) Following the diagonal, the l value decreases as n increases so that their binary sum is exactly equal to the given loop.

We also need use of binary addition, where the number five is written, for example, as:

$$\underbrace{1}_{2^2} \underbrace{1}_{2_1} \underbrace{0}_{2_0} = 1 \times 4 + 1 \times 2 + 0 \times 1 = 5$$

With these rules, as well as Hund's rules, we can identify the elements. The system is introduced by examples. Our first element is Hydrogen (Z=1), with a single (1s) shell filled. We thus write this as,

$$\overbrace{1}^{Z} - \overbrace{001}^{n=1} - \overbrace{000}^{l=0} - \overbrace{\bigcirc}^{\text{Shell half-filled}} \Rightarrow (1s) : {}^{2}S_{1/2}$$

Here, l = 0 so our main label is S. There is one unpaired electron, so that our spin is 1/2, whereby we have ²S. Finally, the shell is at least half filled, so our J is additive, and we have ²S_{1/2}.

Next we have Helium, and we finish filling the 1s shell:

$$\overbrace{2}^{Z} - \overbrace{001}^{n=1} - \overbrace{000}^{l=0} - \overbrace{\otimes}^{\text{Shell filled}} \Rightarrow (1s)^2 : {}^{1}S_0$$

In this case, the shell is completely filled (the ionization energy is at a maxium for this loop) so we have no unpaired electrons (the Pauli exclusion principal demands that a filled shell unit will have one spin up and one spin down electron) and total spin is zero.

Now that the (1s) shell is filled, and our loop takes us the n = 2 level with Lithium,

$$\overbrace{2}^{Z} - \overbrace{010}^{n=2} - \overbrace{000}^{l=0} - \overbrace{000}^{\text{Shell half-filled}} \otimes (He)(2s) : {}^{2}S_{1/2}$$

Once we get to the third loop, l = 1 and we have three shells to fill (corresponding to $m = 0, \pm 1$). For example,

$$\overbrace{2}^{Z} - \overbrace{010}^{n=2} - \overbrace{001}^{l=1} - \overbrace{001}^{\text{Shell}} < \frac{1/2 \text{ filled}}{\circ \circ \otimes} \Rightarrow He(2s)^2(2p) : {}^2P_{1/2}$$

Here the shell is less than half filled, so J = |L - S| = 1/2.

How do we account for the *total* angular momentum? Based on Hund's rule that higher l corresponds to lower energy, we consider an amendment to the system that takes account of the vectoral addition of l such that we match the convention. For any set of subshells, the center shell corresponds to zero, the rightward shells correspond to increasing levels of l, and the leftward decreasing levels from zero. Consider, for example, Sc, Z = 21, whose shells will fill as

$$\overbrace{\circ}^{-2} \xrightarrow{-1} \overbrace{\circ}^{0} \xrightarrow{1} \overbrace{\circ}^{2} \xrightarrow{2} \Rightarrow L = 2$$

Mn as another example gives,

$$\overbrace{\bigcirc}^{-2} \overbrace{\bigcirc}^{-1} \overbrace{\bigcirc}^{0} \overbrace{\bigcirc}^{1} \overbrace{\bigcirc}^{2} \Rightarrow L = 0$$

While Co has,

$$\underbrace{\bigcirc 2}_{2\times2+2\times1+1\times0+1\times(-1)+1\times(-2)=3}^{-2} \xrightarrow[]{0} 0 \xrightarrow[]{0} 1 \xrightarrow[]{0} 2 \xrightarrow[]{$$

Of course, there are exceptions such as Cr, where we would expect,

$$\overbrace{\circ}^{-2} \overbrace{\oslash}^{-1} \overbrace{\oslash}^{0} \overbrace{\oslash}^{1} \overbrace{\oslash}^{2} \Rightarrow L = 2$$

It so happens, though, that Cr works out to have L = 0. This methodology has the benefit of being systematic enough to lend to easy comprehension, once the user gets used to the rules. As well, once could quite easily program these rules into a computer to output a generalized table. We present a partial list of the periodic table (the first three loops) as a final example.

| Loop | Ζ | *n* - *l* - *shell* | Shell | Code |
|------|----|--|--------------------|-----------------|
| 1 | 1 | $001 - 000 - \oslash$ | (1s) | $^{2}S_{1/2}$ |
| | 2 | $001 - 000 - \otimes$ | $(1s)^2$ | $^{1}S_{0}$ |
| 2 | 3 | $010 - 000 - \oslash$ | (He)(2s) | ${}^{2}S_{1/2}$ |
| | 4 | $010 - 000 - \otimes$ | $(He)(2s)^2$ | ${}^{1}S_{0}$ |
| 3 | 5 | $010 - 001 - \circ \circ \oslash$ | $(He)(2s)^2(2p)$ | $^{2}P_{1/2}$ |
| | 6 | $010 - 001 - \circ \oslash \oslash$ | $(He)(2s)^2(2p)^2$ | ${}^{3}P_{0}$ |
| | 7 | $010 - 001 - \oslash \oslash$ | $(He)(2s)^2(2p)^3$ | ${}^{4}S_{3/2}$ |
| | 8 | $010 - 001 - \oslash \oslash$ | $(He)(2s)^2(2p)^4$ | ${}^{3}P_{2}$ |
| | 9 | $010 - 001 - \oslash \otimes \bigotimes$ | $(He)(2s)^2(2p)^5$ | $^{2}P_{3/2}$ |
| | 10 | $010 - 001 - \otimes \otimes \otimes$ | $(He)(2s)^2(2p)^6$ | ${}^{1}S_{0}$ |
| | 11 | $011 - 000 - \oslash$ | Ne(3s) | $^{2}S_{1/2}$ |
| | 12 | $011 - 000 - \otimes$ | $Ne(3s)^2$ | ${}^{1}S_{0}$ |

2. KRAMER'S RELATION AND THE CIRCULAR STATE

An atom which has a valance electron in an extremely high principal quantum number state is in a Rydberg state. These states are called circular states for the following reason. (We use the simpler hydrogen case due to the fact that it is not a terrible approximation to hydrogenic atoms.)

The radial equation (see the compendium on the hydrogen atom) was found to be simplifiable to,

(2.1)
$$-\frac{\hbar^2}{2m}u'' + \left(-\frac{e^2}{4\pi\epsilon_0}\frac{1}{r} + \frac{\hbar^2}{2m}\frac{l(l+1)}{r^2}\right)u = Eu$$

With,

(2.2)
$$E = -\frac{m}{2\hbar^2} \left(\frac{e^2}{4\pi\epsilon_0}\right)^2 \frac{1}{n^2}$$

and $a = 4\pi\epsilon_0 \hbar^{@}/me^2$, we can rewrite this equation as,

(2.3)
$$u'' = \left(\frac{l(l+1)}{r^2} - \frac{2}{ar} + \frac{1}{n^2 a^2}\right)u$$

We then consider that (using integration by parts and knowing that the infinite limit of the radial component must be zero)

$$\begin{aligned} \int ur^{s}u''dr &= \int ur^{s} \left(\frac{l(l+1)}{r^{2}} - \frac{2}{ar} + \frac{1}{n^{2}a^{2}}\right)u \, dr \\ &= \langle r^{s-2} \rangle l(l+1) - \frac{2}{a} \langle r^{s-1} \rangle + \frac{1}{n^{2}a^{2}} \langle r^{s} \rangle \\ &= -\int (ur^{s})'u'dr \\ &= -\int (ur^{s})'u'dr - s \int ur^{s-1}u'dr \\ &= -\left(\frac{2}{s+1} \int u''r^{s+1}u'dr\right) - s\left(-\frac{s-1}{2} \int ur^{s-2}u \, dr\right) \\ &= -\left(\frac{2}{s+1} \int \left(\frac{l(l+1)}{r^{2}} - \frac{2}{ar} + \frac{1}{n^{2}a^{2}}\right)ur^{s+1}u'dr\right) + \frac{s(s-1)}{2} \langle r^{s-2} \rangle \\ (2.4) &= -\frac{l(l+1)(s-1)}{s+1} \langle r^{s-2} \rangle + \frac{2s}{a(s+1)} \langle r^{s-1} \rangle - \frac{1}{n^{2}a^{2}} \langle r^{s} \rangle + \frac{s(s-1)}{2} \langle r^{s-2} \rangle \end{aligned}$$

Algebraic exercise yields Kramer's relation,

(2.5)
$$\frac{s+1}{n^2} \langle r^s \rangle - (2s+1)a \langle r^{s-1} \rangle + \frac{s}{4} \left((2l+1)^2 - s^2 \right) a^2 \langle r^{s-2} \rangle = 0$$

When we set s = 0 we find,

(2.6)
$$\frac{1}{n^2}\langle 1\rangle - a\langle \frac{1}{r}\rangle = 0 \quad \ni \quad \langle \frac{1}{r}\rangle = \frac{1}{n^2 a}$$

Thus setting s = 1 yields,

(2.7)
$$\frac{2}{n^2} \langle r \rangle - 3a \langle 1 \rangle + \frac{1}{4} \left((2l+1)^2 - 1 \right) a^2 \langle \frac{1}{r} \rangle = 0 \quad \ni \quad \langle r \rangle = \frac{a}{2} \left(3n^2 - l(l+1) \right)$$

When we use the states such that l = n - 1 and |m| = n - 1, i.e. the maximum allowed, we have

(2.8)
$$\langle r \rangle = an^2 + \frac{an}{2}$$

For very large n, this approximates the classical result for circular orbits, $\langle r \rangle = n^2 a$, thus the name circular Rydberg states.

3. Selection Rules and Available Transitions

A change in energy levels for a Rydberg atom in a circular state must obey selection rules so that $\Delta m = \pm 1$ or 0, and $\Delta l = \pm 1$. Thus an atom in a Rydberg state under guarded environmental conditions can only transition as

$$n \rightarrow n-1, l \rightarrow l-1, |m| \rightarrow |m|-1$$

Thus a Rydberg atom approximates a two-level system.

We demonstrate the theory behind elementary selection rules with a simple example. A two-state system in the presence of puturbing Hamiltonian can be described as,

(3.1)
$$\Psi(t) = c_a(t)\psi_a e^{-iE_at/\hbar} + c_b(t)\psi_b e^{-iE_bt/\hbar}$$

Its evolution is described by

$$(H + H'(t))\Psi = i\hbar \frac{\partial \Psi}{\partial t}$$

from which we have,

$$(3.2) \quad c_a(t)\langle\psi_a|H'|\psi_a\rangle e^{-iE_at/\hbar} + c_b(t)\langle\psi_a|H'|\psi_b\rangle e^{-iE_bt/\hbar} = i\hbar c_a(t)e^{-iE_at/\hbar bar}$$

Rejoining convention and writing $H'_{ij} = \langle \psi_i | H' | \psi_j \rangle$, $\omega_0 = (E_b - E_a)/\hbar$, and assuming (as is warented in the experiments we discuss) that the diagonal of the perturbing part of the Hamiltonian is zero, we can obtain a set of equations for the prefactors,

(3.3)
$$\dot{c_a} = -\frac{i}{\hbar} H'_{ab} e^{-i\omega_0 t} c_b$$

(3.4)
$$\dot{c}_b = -\frac{i}{\hbar} H'_{ba} e^{i\omega_0 t} c_a$$

Following [4] we consider a basic hydrogen atom in the n = 2 state in an electric field so that $H' = -eEr \cos \theta$. As we showed in a previous compendium, the wave functions for hydrogen, n = 1, n = 2, are

$$\psi_{100} = \frac{1}{\sqrt{\pi a^3}} e^{-r/a}$$

$$\psi_{200} = \frac{1}{\sqrt{8\pi a^3}} \left(1 - \frac{r}{2a}\right) e^{-r/2a}$$

$$\psi_{210} = \frac{1}{\sqrt{32\pi a^3}} \frac{r}{a} e^{r/a} \cos\theta$$

$$\psi_{211} = -\frac{1}{\sqrt{64\pi a^3}} \frac{r}{a} e^{-r/2a} \sin\theta e^{i\phi}$$

$$\psi_{21-1} = \frac{1}{\sqrt{64\pi a^3}} \frac{r}{a} e^{-r/2a} \sin\theta e^{-i\phi}$$

The perturbation matrix is simple in that all but one of these will be even in $r \cos \theta$, so the perturbation matrix $(\langle n'l'm'|H'|nlm \rangle)$ will have all zero elements except

(3.5)
$$\langle 100|H'|210\rangle = -\frac{eE}{4\sqrt{2}\pi a^4} \int e^{-3r/21} r^2 \cos^2\theta r^2 \sin\theta dr d\theta \ d\phi = -\frac{2^8}{\sqrt{2}3^5} eEa$$

| | ψ_{100} | ψ_{200} | ψ_{210} | ψ_{211} | ψ_{21-1} |
|---------------|-----------------------------------|--------------|-----------------------------------|--------------|---------------|
| ψ_{100} | 0 | 0 | $-\frac{2^{8}}{\sqrt{23^{5}}}eEa$ | 0 | 0 |
| ψ_{200} | 0 | 0 | 0 | 0 | 0 |
| ψ_{210} | $-\frac{2^{8}}{\sqrt{2}3^{5}}eEa$ | 0 | 0 | 0 | 0 |
| ψ_{211} | 0 | 0 | 0 | 0 | 0 |
| ψ_{21-1} | 0 | 0 | 0 | 0 | 0 |

Thus we see that under this perturbation, the $100 \leftrightarrow 210$ levels are "selected' and we have an approximate two-state system under the right conditions (ideal).

In general, we can derive selection rules for m and l transitions. In the case of m, we consider that $[L_z, z] = 0$ so that

(3.6)
$$0 = \langle n'l'm' | [L_z, z] | nlm \rangle = (m' - m) \langle n'l'm' | z | nlm \rangle$$

That $[L_z, x] = i\hbar y$ gives,

(3.7) $(m'-m)\langle n'l'm'|x|nlm\rangle = i\langle n'l'm'|y|nlm\rangle$

Finally, $[L_z, y] = -i\hbar x$ gives,

(3.8) $(m'-m)\langle n'l'm'|y|nlm\rangle = -i\langle n'l'm'|x|nlm\rangle$

Thus $(m'-m)^2 = 1$ or $\langle |x,y| \rangle = 0$. For the l, l' case, it can be shown that

$$L^{2}, [L^{2}, r]] = 2\hbar^{2}(rL^{2} + L^{2}r)$$

From this we can show that

(3.9) $2\hbar^4(l(l+1)+l'(l'+1))\langle n'l'm'|r|nlm\rangle = \hbar^4(l'(l_1')-l(l+1))^2)\langle n'l'm'|r|nlm\rangle$ Rewriting

l'(l'+1)-l(l+1) = (l'+l+1)(l'-l), and $2(l(l+1)+l'(l'+1)) = (l'+l+1)^2 + (l'-l)^2 - 1$

We thus conclude that $l' = l \pm 1$ and $m' = m \pm 1$ or m

4. The Preperation of Rydberg Atoms

References

- [1] G. C. Christopher, P. L. Knight, Introductory Quantum Optics, Cambridge University Press, 2005
- [2] S. Gasiorowicz, Quantum Physics, 3rd Ed., John Wiley & Sons, 2003
- [3] R. G. Hulet & D. Kleppner, *Physical Review Letters*, Vol. 51, Num 16, 1430
 [4] D. Griffiths, *Introduction to Quantum Mechanics*, Prentice Hall, 1995