



Review

# Momentum evaluations of the electron-nuclear shielding constant for many-electron systems

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Hartree Fock (HF) wave function in momentum space Nuclear magnetic shielding constant calculation are determined for the ground states of the following member of Li-isoelectric series: Li, Be<sup>+</sup>, B<sup>+2</sup>, C<sup>+3</sup>, N<sup>+4</sup>, O<sup>+5</sup>, Fe<sup>+6</sup>, and Ne<sup>+7</sup>. For each species, the density distribution function  $D(p_1)$ , the electron density at nucleus, moments of  $\langle p^n \rangle$  for  $n = -2, -1$  to 2, and diamagnetic susceptibility are reported using MathCad program.

**Keywords :** Momentum, Electron, Nuclear, Electron Systems

## INTRODUCTION

A useful working description of the electronic structure in atoms or ions is provided by the independent particle model. The analysis is applied here to the ground state of Li. By partitioning the second order density as proposed by (Banyard and Youngman, 1987) (Valley and Alley, 2002).

The prime example of this search is HF treatment. HF theory is the use of a single-determinant wave function  $\Psi$  which is formed from a set of occupied spin orbitals  $x_1, x_2, x_3, \dots, x_n$  (Thakkarjit and Vedene, 1978)

$$\Psi = A \det \{ x_1, x_2, x_3, \dots, x_n \} \quad (1)$$

Where  $n$  is the number of electrons.  $A$  is a normalization constant and the diagonal element product of determinant is satisfied explicitly in eq. 1.

$x_n$  is the doublet spin eigenfunction and the functional form employed is

$$x = \alpha(1)\beta(2)\alpha(3) - \beta(1)\alpha(2)\alpha(3) \quad (2)$$

## THEORY

A general analysis of the <sup>2</sup>S-state of the Li I isoelectronic series in the same spirit as King has recently been carried out (Banyard et al., 1988). The radial electronic density distribution functions  $D_o(p)$  is evaluated for N-electron systems using (Banyard et al., 1988):

$$D_o(p) = \int_0^\pi \int_0^{2\pi} p^2 \rho(p) d\Omega \quad (3)$$

$$\rho(p) = N \int \Psi^*(x_1, x_2, \dots, x_n) X \Psi(x_1, x_2, \dots, x_n) ds_1 ds_2 ds_3 \dots dx_n \quad (4)$$

Where standard notation  $d\Omega = \sin\theta d\theta d\phi$

And  $x_i$  denoted a combined spatial and spin coordinate and

$\Psi$  is normalized wave function and in present work  $N = 3 \cdot \Psi(x_1, x_2, \dots, x_n)$

**Table I.** Radial Density Distribution Fuunction , Electron Density At The Nucleus:

Species	shell	D <sub>o</sub> (r)x.	Location ρ(0)	p1
Li	KαKβ	0.544	0.127	0.3
Be+	KβKα	0.324	0.083	0.5
B <sup>+2</sup>	KαKα	0.228	0.015	0.6
C <sup>+3</sup>	KαKα	0.182	2.175-03	0.9
N <sup>+4</sup>	KαKα	0.150	5.545-03	1.1
O <sup>+5</sup>	KαKα	0.128	1.780-04	1.3
F <sup>+6</sup>	KαKα	0.111	7.595-05	1.4
Ne <sup>+7</sup>	KαKα	0.099	3.710-05	1.5

**Table II.** Expectation Values For The Intra Shell(K-Shell)For Li –Isoelectronic Series :

Species	n				
	+2	-2	-1	0	+1
Li	0.76422	0.657109	0.999999	2.246038	7.223937
Be+	0.396380	0.474361	0.999999	3.092023	13.57525
B <sup>+2</sup>	0.241883	0.371072	1.000000	3.938945	21.92341
C <sup>+3</sup>	0.162729	0.304664	1.000000	4.786131	32.26986
N <sup>+4</sup>	0.11683	0.258356	0.999999	5.63409	44.61596
O <sup>+5</sup>	0.087981	0.224287	0.999999	6.482112	58.96018
F <sup>+6</sup>	0.06859	0.198129	0.999999	7.330403	75.30776
Ne <sup>+7</sup>	0.054269	0.177433	0.999998	8.17861	93.653734

The wave function employed in this study are Hartree-Fock type (Landau and Lishitz, 1974).

$$\Psi(x_1, x_2, x_3) = A \sum_{\mu}^N C_{\mu} \phi_{\mu} \tag{6}$$

Where A is the antisymmetrizer , are the variation ally determined expansion coefficients ,N is the number of basis function .

**COMPACT FORMULA OF**

$$D_0(p)$$

The one particle radial Expectation values can be obtain [5]

$$\langle p_i^n \rangle = \sum_{-1}^3 \langle \Psi | p_i^n | \Psi \rangle$$

$$\langle p_i^n \rangle = \int_0^{(\infty)} D_o(p) p^n dp$$

The required formula of the one particle radial density distribution function is:

$$D_o(p) = C_{\mu} C_N \left[ \frac{(2 \xi_{\mu})^{n_{\mu} + 0.5}}{\sqrt{(2 n_{\mu})}} \frac{(2 \xi_N)^{n_N + 0.5}}{\sqrt{(2 n_N)}} \right]$$

$$p^{n_{\mu} - 1} p^{n_N - 1} e^{-(\xi_{\mu} + \xi_N) p}$$

The one particle expectation values is:

**Table Iii:** Expectation Values For The Intra Shell(L-Shell) For Li –Isoelectronic Series :

Species	n				
	-2	-1	0	+1	+2
Li	5.046617	3.872695	0.999999	0.413496	0.417639
Be+	8.224346	2.205598	0.999999	0.766091	1.404733
B <sup>+2</sup>	4.141884	2.371072	1.000000	1.112376	2.906504
C <sup>+3</sup>	2.501313	1.207308	0.99989	1.456210	4.913969
N <sup>+4</sup>	1.674565	0.985904	1.000000	1.798962	7.424515
O <sup>+5</sup>	1.200417	0.833497	0.999999	2.231450	10.435781
F <sup>+6</sup>	0.903436	0.722215	0.999999	2.481965	13.949432
Ne <sup>+7</sup>	0.721308	0.654871	0.999998	2.862836	18.043641

Table VI : Expectation Values For The Inter Shell(Kl-Shell) For Li –Isoelectronic Series :

Species	n				
	-2	-1	0	+1	+2
Li	12.905423	2.264902	1.000000	1.329767	3.820788
Be+	4.310366	1.339979	0.999999	1.929057	7.489979
B <sup>+2</sup>	2.191884	0.964646	0.999999	2.525660	12.414957
C <sup>+3</sup>	1.332021	0.755976	1.000000	3.121264	18.59191
N <sup>+4</sup>	0.895698	0.622130	0.999999	3.716531	26.02024
O <sup>+5</sup>	0.644199	0.528892	0.999999	4.311427	34.69798
F <sup>+6</sup>	0.486013	0.460172	1.000000	4.966185	44.628599
Ne <sup>+7</sup>	0.388139	0.4161521	0.999999	5.520758	55.848681

$$\langle p_i^n \rangle = \langle R_{1s} | p^{n+2} | R_{1s} \rangle \tag{10}$$

$$\langle p_i^n \rangle = \langle \sum_{\mu} S_{\mu\ell} C_{\mu\ell} | p^{n+1} | \sum_N S_{N\ell} C_{N\ell} \rangle \tag{11}$$

$$\langle p^n \rangle = \langle S_{1\ell} C_{1\ell n} | p^{n+2} | S_{1\ell} C_{1\ell n'} \rangle \tag{12}$$

$$\langle p_i^n \rangle = \langle \sum_{\mu} N_{\mu\ell} p^{(n\mu\ell-1)} e^{(-\xi\mu\ell)p} C_{\mu\ell n} | p^{n+2} | \sum_N N_{N\ell} p^{(nN\ell-1)} e^{(-\xi\mu\ell)p} C_{N\ell n} \rangle$$

Where  
 $S_{\mu\ell}$  and  $S_{N\ell}$  takenas Slatertypeorbitals  
 $S_{\mu\ell}$  and  $S_{N\ell}$  is the normalization factor  
 $n_{\mu\ell}$  and  $n_{N\ell}$  is the principle quantum number  
 $\xi_{\mu\ell}$  and  $\xi_{N\ell}$  is the orbital exponent

$\ell$  is a azmeithal quantum number

$$N_{\mu\ell} = \frac{(2\xi_{\mu\ell})^{n_{\mu\ell}+0.5}}{\sqrt{(2n_{\mu\ell})}} \quad \text{and} \quad N_{N\ell} = \frac{(2\xi_{N\ell})^{n_{N\ell}+0.5}}{\sqrt{(2n_{N\ell})}} \tag{14}$$

$$\langle p^n \rangle = \langle S_{1\ell} C_{1\ell n} | p^{n+2} | S_{1\ell} C_{1\ell n'} \rangle \tag{15}$$

$$\langle p^n \rangle = \langle S_{1\ell} C_{1\ell n} | p^{n+2} | S_{1\ell} C_{1\ell n'} \rangle \langle S_{2\ell} C_{2\ell n} | p^{n+2} | S_{2\ell} C_{2\ell n'} \rangle \tag{13}$$

$$\langle S_{3\ell} C_{3\ell n} | p^{n+2} | S_{3\ell} C_{3\ell n'} \rangle \tag{16}$$

$$\rho(0) = \left\{ \frac{D_o(p)}{4\pi p^2} \right\}_{p \rightarrow 0} \tag{17}$$

$$\rho(0) = \langle \delta(p_i) \rangle \tag{18}$$

**Table IV :** Nuclear Magnetic Shielding Constant And Diamagnetic Suscubtability For Li –Isoelectronic Series :

Species	shell	$\sigma$	$\chi$
Li	K $\alpha$ K $\beta$	1.166-05	-6.411-05
	L $\alpha$ L $\beta$	6.874-05	-3.707-06
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	4.021-05	-3.391-05
Be+	K $\alpha$ K $\beta$	8.420-06	-1.205-04
	L $\alpha$ L $\beta$	3.915-05	-1.247-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	2.379-05	-6.648-04
B <sup>+</sup>	K $\alpha$ K $\beta$	6.586-06	-1.946-04
	L $\alpha$ L $\beta$	2.766-05	-2.580-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	1.712-01	-1.102-04
C <sup>+3</sup>	K $\alpha$ K $\beta$	5.408-06	-2.864-04
	L $\alpha$ L $\beta$	2.143-05	-4.361-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	1.342-05	-1.650-04
N <sup>+4</sup>	K $\alpha$ K $\beta$	4.586-06	-3.960 -04
	L $\alpha$ L $\beta$	1.750-05	-6.589-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	1.104-05	-2.309-04
O <sup>+5</sup>	K $\alpha$ K $\beta$	3.981-06	-5.233-04
	L $\alpha$ L $\beta$	1.479-05	-9.262-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	9.388-6	-3.081-04
F <sup>+6</sup>	K $\alpha$ K $\beta$	23.51-06	-6.684-04
	L $\alpha$ L $\beta$	1.282-05	-8.238-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	8.168-05	-3.961-04
Ne <sup>+7</sup>	K $\alpha$ K $\beta$	3.151-06	-8.312-04
	L $\alpha$ L $\beta$	1.162-05	-1.601-05
	K $\alpha$ L $\beta$ = K $\beta$ L $\alpha$	7.387-05	-4.957-04

**DISCUSSION**

**A- CALCULATION OF ELECTRON DENSITY AT THE NUCLUES**

$$\rho(0)$$

From equation (3), the electron density at the nucleus can be evaluated using (King, 1988):

$$(17) \rho(0) = \left\{ \frac{D_0(p)}{4\pi p^2} \right\}_{p \rightarrow 0}$$

Because of the simple analytic form obtained for one particle radial density in the integral (eq.8) is trivial evaluate the electron density at the nucleus

$$\rho(0) = \langle \delta(p_i) \rangle \tag{18}$$

In table I, the electron density at the nucleus  $\rho(0)$  depends on the one particle radial density distribution  $D(p_i)$  and the results of  $D(p_i)$  is decreases as atomic

number increases the electron nuclear cusp condition are infairly reasonable agreement, which improves with increasing charge.

**B- ONE PARTICLE RADIAL DENSITY DISTRIBUTION FUNCTION  $\langle p_i^n \rangle$ :**

From the radial expectation values given in tables II,III,VI respectively we notice that the effect of electron correlation is to reduce  $\langle p_i^n \rangle$  for each  $n$ , indicating that the magnitude of the on the one particle radial density distribution has been decreases for all  $p_i$ . For each Z the one particle radial expectation values decreases when  $n$  goes from -1 to -2 ,and increases when  $n$  goes from 1 to 2 , that means the expectation value weight different regions of space (i.e the probability of finding the electrons in the region near the nucleus or farther away from it).when  $n=1$ ,and 2, the one particle expectation value  $\langle p_i^n \rangle$  increases by increasing the atomic number Z, this is due to the Coulomb attraction force of the nucleus to the charge which leads to increase the probability of finding the electron near the nucleus .For negative values

of  $n$ , the expectation values decreases by increasing the atomic number  $Z$  due to the weakly Coulomb attraction force between the nucleus and the electrons in the outer shells.

Correlation also produces a shift in the towards higher momentum since we observe an increase in both  $\langle p \rangle$  and  $\langle p_i^2 \rangle$ . The normalized condition can be obtained from the calculation of one particle expectation value  $\langle p_i^n \rangle$  at ( $n=0$ ) and this state may be applied for all shells. By examining all expectation tables for the Li-atom, Li-like ions we found that the value of  $L\alpha L\beta$  is greater than those for  $K\alpha K\beta$  because it is the outer most shell (i.e in momentum view the inner most shell  $L\alpha L\beta$  to the outer most shell  $K\alpha K\beta$ ), the two state decreases when then atom number ( $Z$ ) increase too.

### C-NUCLER MAGNETIC SHEILDING CONSTANT $\sigma$ AND DIAMAGNETIC SUSCUBTABILITY

Table IV show the results of the nuclear magnetic shielding constant  $\sigma$   $K\alpha K\beta$  is greater than that found for  $L\alpha L\beta$  due to the diamagnetic shielding factor ( ) for  $L\alpha L\beta$ , and this is not found for  $K\alpha K\beta$  because the charge for the two electron in the  $K\alpha K\beta$  will reduce the nucleus shielding to the  $L\alpha L\beta$ . As  $Z$  increases the nuclear magnetic shielding constant  $\sigma$  decreases in momentum conclusions.

The diamagnetic susceptibility for Li-isoelectronic series for  $K\alpha K\beta$  is less than that for  $L\alpha L\beta$ ,  $K\beta L\alpha$ ,  $K\alpha L\beta$  because the diamagnetic susceptibility depends on the radius of 1s and 2s respectively.

Also this can be confirmed by comparing the moment  $\langle p_i^n \rangle$  at  $n = 2$  From comparison between  $\chi$  for Li-atom

and  $\chi$  for Li-like ions, it is observed that the diamagnetic susceptibility decreases as  $Z$  (atomic number) increases, because the radius of 1s and 2s for positive ions is smaller than that for Li-atom due to the attraction force between the electron and the proton. This conclude agree with the relation between  $\sigma$  and  $\langle p_i^{-1} \rangle$  from equation (8)

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