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Angular and radial correlation in doubly excited systems when 1 <or=Z<or=4: the 2p^{2 3}P state

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Angular and radial correlation io doubly excited systems when $1 \leq Z \leq 4$: the $2p^2$ ³P state

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Abstract. The angular and radial components of electron correlation have each been examined in detail for the discrete $2p^2$ ³P states of H⁻, He, Li⁺ and Be²⁺. These doubly excited sytems were described by highly accurate explicitly correlated wavefunctions. The analysis involved the **use** *or* angular Coulomb holes, changes in the one- and two-particle radial density distributions and several angular and radial expectation values. Additionally, various statistical correlation coefficients were used which emphasized. in turn, angular and radial correlation properties in different regions of the two-particle density.

The angular holes and related properties showed a clearly defined inverse- Z effect for He and the positive ions. This trend was not repeated for the radial curves. **However,** the radial densities did reveal a distinct 'in-out' correlation effect-similar in character to the split-shell behaviour for the ground state. By comparison with the findings for $Z \ge 2$, the angular and radial correlation effects for H⁻ were always exceedingly large, thus setting it apart from the other systems. For He, the angular hole for the comparatively slow moving 2p2'P electmns was found to be *over 50%* deeper than that *for* the gmund state and about six times the depth of a $1s2p³P$ hole. The statistical correlation coefficients highlighted a steady growth, with *Z,* in the relative importance of angular correlation. Nevertheless, for each system, these coefficients indicated that the radial effect was the prevailing influence **on** the two-particle probability distribution.

1. Introduction

By comparison with the gound state, doubly excited states **(DES)** of simple atoms are occupied by relatively slow moving electrons and, consequently, they should be more responsive to the influence of Coulomb correlation. This was demonstrated to be the case for the $2p^2$ ³P states of H⁻, He, Li⁺ and Be²⁺ by Banyard *et al* (1992, hereafter referred to as BKD). In that work, we analysed the correlation-induced changes in the probability distribution for r_{12} , the interelectronic separation, by means of Coulomb holes, partial Coulomb holes and various $\langle r_1^2 \rangle$. Comparisons were made with the ground state and the singly-excited state $1s2p³P$. The latter comparison arose since, for low *Z*, an important decay mechanism for the $2p^2$ ³P state is by a radiative transition to the $1s2p^3P$ level. It is observed that, being discrete, the $2p^2^3P$ state exhibits no wavefunction mixing with open channels. Transitions involving this pes have been considered by, for example, Westerveld **ef** *ai* (1979), Auderbert *er* al **(1984)** and Karim and Bhalla **(1988).** See our previous report **(BKD)** for further references.

In the present article, we perform an in-depth appraisal of the separate angular and radial components of electron correlation associated with the $2p^2$ ³P state when $1 \leq Z \leq 4$. As before, the correlated descriptions are provided by the highly accurate

wavefunctions of Drake **(1986);** the non-correlated reference state is again represented by the restricted Hartree-Fock **(HF)** functions obtained from a program by Froese-Fischer **(1987).** in addition to reporting angular Coulomb holes and correlation-induced changes in the one- and two-particle radial densities, we also determine a series of expectation values and calculate various statistical correlation coefficients τ (see Kutzelnigg *et a/* **1968** and, more recently, Banyard and Mobbs **1981,** and Thakkar **1987).** Atomic units are used throughout this work.

Other studies of either angular or radial correlation, or both, for simple systems in excited states have been undertaken by many workers. Some typical references are Sinanoglu and Herrick **(1975),** Rehmus *et* **al(1978),** Ezra and Berry **(1983),** Nicolaides *el al* **(1987),** Ojha and Berry **(1987),** Dmitrieva and Plindov **(1988),** Rau and Molina **(1989),** lvanov **(1992)** and Chen **er** *at* **(1992).**

2. Calculations and **results**

Drake **(1986)** wrote the space part of the explicitly correlated wavefunction for the $He(2p² P)$ -like systems as

$$
\Psi_{\text{corr}}(\boldsymbol{r}_1, \boldsymbol{r}_2) = (1 - P_{12}) \sum_{i,j,k}^{N} A_{ijk} r_1^i r_2^j r_{12}^k \exp(-\alpha r_1 - \beta r_2) Y_{l_1=1, l_2=1, L=1}^{M=1} (\theta_1, \phi_1, \theta_2, \phi_2)
$$

where

$$
Y_{i_1=1,i_2=1,L=1}^{M=1}(\theta_1,\phi_1,\theta_2,\phi_2)=2^{-1/2}\left[Y_1^1(\theta_1,\phi_1)\,Y_1^0(\theta_2,\phi_2)-Y_1^0(\theta_1,\phi_1)\,Y_1^1(\theta_2,\phi_2)\right].\eqno(1)
$$

P12 is a permutation operator and, as shown, the angular term involves products **of** spherical harmonic functions. **A** summary of the remaining notation is given in BKD. Since *N* is the upper limit on the summation, it is used to designate a particular wavefunction. As previously, we examine two correlated wavefunctions for each *Z:* the energetically best function and, for comparison, the function with the fewest terms. For H⁻, the Drake wavefunction with least terms has $N = 20$ and recovers 99.72% of the correlation energy. All other Drake functions describing the $1 \le Z \le 4$ systems yielded percentages in *excess* of that obtained by the **D-20** (Drake: equation (1) with $N = 20$) wavefunction for H⁻. *N* values, total energies *E* and correlation percentages are quoted in table **1** of **BKD.** Each numerical **HF** wavefunction, **HFCNUM),** derived from the Froese-Fischer **(1987)** program is represented by the same linear combination of Slater-type orbitals **(STO)** as used before. These fitted **HF(STOI** functions are employed throughout this work but, for comparison, expectation values are reported for both $HF(NUM)$ and $HF(STO)$.

Following Youngman and Banyard (1987), the distribution function for $\gamma = \theta_{12}$, the angle between the position vectors $r₁$ and $r₂$ for the two electrons, is given in terms of a normalized space wavefunction $\Psi(\mathbf{r}_1, \mathbf{r}_2)$ by

$$
P(\gamma) = \int \Psi^*(\mathbf{r}_1, \mathbf{r}_2) \Psi(\mathbf{r}_1, \mathbf{r}_2) d\tau_1 d\tau_2/d\gamma
$$
 (2)

where, unlike an earlier definition by Banyard and Ellis **(1972).** we note that

$$
\int_0^{\pi} P(\gamma) d\gamma = 1.
$$
 (3)

The **origin** of coordinates is always located at the nucleus; see figure 1 of BKD **(1992)** where, in that instance, the inter-vector angle was labelled θ_{12} . The angular Coulomb hole is defined here as

$$
\Delta P(\gamma) = P_{\text{corr}}(\gamma) - P_{\text{HF}}(\gamma) \tag{4}
$$

where the integral of $\Delta P(\gamma)$ with respect to $0 \leq \gamma \leq \pi$ is zero. For the 2p²³P state

$$
P_{HF}(\gamma) = \frac{3}{4}\sin^3(\gamma). \tag{5}
$$

The plot of $P_{HF}(\gamma)$ is shown in figure 1A along with the $\Delta P(\gamma)$ curves for H⁻ derived from the **D-20** and **D-84** correlated wavefunctions. In figure lB, we show the angular holes associated with the energetically best Drake functions: D-84 for H⁻ and D-70 for He, Li⁺ and Be²⁺. For each $Z \ge 2$, the correlated wavefunction with the fewest terms, **D-13, produced a** $\Delta P(\gamma)$ **curve which was graphically indistinguishable from that for** the D-70 function. Figure 2 shows the curves for $1 \le Z \le 4$ when scaled to give $Z\Delta P(\gamma)$ against γ . In figure 3 the angular hole for He is compared with the corresponding

Figure 1. A: the normalized angular function $P_{HF}(\gamma)$ for all Hartree-Fock (HF) $2p^{2.3}P$ wavefunctions; *y* is the angle subtended by electrons I and 2 at *the* nuclear origin. *Also* shown are the angular Coulomb holes, $\Delta P(\gamma) = P_{\text{corr}}(\gamma) - P_{\text{HF}}(\gamma)$, for H⁻(2p²³P). The $P_{\text{corr}}(\gamma)$ functions are derived from the Drake wavefunction in equation (1) using, in turn, $N = 20$ and $N = 84$. The $\Delta P(\gamma)$ curves are denoted by their correlated descriptions $D-20$ and **D-84.**

B: the $\Delta P(\gamma)$ curves for the $2p^2$ ³P state when $1 \leq Z \leq 4$. For each system, $P_{\text{corr}}(\gamma)$ is obtained from the energetically best version or equation *(1):* the **D-84** wavefunction for H⁻ and the *D*-70 versions for He, Li⁺ and Be²⁺. When $Z \ge 2$, the *D*-13 functions yielded $\Delta P(\gamma)$ curves which were graphically indistinguishable from the corresponding D-70 curve.

Figure 2. The Z-scaled angular holes for the $2p^2$ ³P states of H⁻, He, Li⁺ and Be²⁺ using the energetically best correlated wavefunctions: **D-84** for **H-** and the **D-70** functions **when** *z>2*

Figure 3. Comparison of the angular holes, $\Delta P(\gamma)$ against γ , for the 2p²³P, 1s2p³P and 1s²¹S states of He. The full curve is for the 2p²³P state, the broken curve and the dotted curve represent the **ls2p** 'P and **Is'** *'S* states, respectively. **For** this order of these **He** stales, the correlated descriptions are provided by the energetically best wavefunctions of Drake **(1986),** Tweed (1973) and Weiss **(1961).**

results for the Is' **'S** and ls2p **3P** states taken, respectively, from Banyard and Ellis (1972) and (1975). Their angular Coulomb curves for both states have been revised **to** satisfy equations (2) and **(3).** Thc correlated description of the ground state was provided by a 35-term configuration-interaction **(CI)** wavefunction (Weiss 1961) which recovered 98.8% of the correlation energy whereas, for the singly excited state, they used an explicitly correlated wavefunction (Tweed 1973) which accounted for 78.6% of the correlation energy.

As usual, the normalized radial distribution is defined as

$$
D(r_1) = \int \Psi^*(\mathbf{r}_1, \mathbf{r}_2) \Psi(\mathbf{r}_1, \mathbf{r}_2) d\tau_1 d\tau_2 / d\mathbf{r}_1
$$
 (6)

and, hence, we formed a one-particle radial hole

$$
\Delta D(r_1) = D_{\text{corr}}(r_1) - D_{\text{HF}}(r_1). \tag{7}
$$

Similarly, a two-particle radial function can be expressed as

$$
D(r_1, r_2) = \int \Psi^*(r_1, r_2) \Psi(r_1, r_2) d\tau_1 d\tau_2 / dr_1 dr_2 \tag{8}
$$

and the corresponding correlation change is given by

$$
\Delta D(r_1, r_2) = D_{\text{corr}}(r_1, r_2) - D_{\text{HF}}(r_1, r_2). \tag{9}
$$

Obviously we have

$$
\int_0^\infty \int_0^\infty \Delta D(r_1, r_2) dr_1 dr_2 = \int_0^\infty \Delta D(r_1) dr_1 = 0.
$$
 (10)

When plotted against Z_{r_1} , figure 4A shows $Z^{-1}D_{HF}(r_1)$ for each Z and figure 4B shows $\Delta D(r_1)$ for $2 \le Z \le 4$ (the H⁻ curve is omitted because of its size, see figure 5). In each instance, the correlated description used in forming $\Delta D(r_1)$ is derived from the energetically best Drake wavefunction. Included in figure **4A,** as a broken curve, is the radial distribution generated from an independent-particle $2p^2$ ³P wavefunction

Figure 4. A: radial densities $D_{HF}(r_1)$ derived from the HF wavefunctions for the $2p^2$ ³P states when $1 \le Z \le 4$. The broken curve is $D(r_1)$ generated from an independent-particle **2p"P wavefunction comprising unoptimized hydrogenic orbitals.** *Both* **axes are scaled to preserve normalization.**

B: correlation-induced changes in the one-particle density, $\Delta D(r_1)$, for He, Li⁺ and Be^{2+} plotted against Zr_1 . For each system, a D-70 correlated wavefunction is used to determine $D_{\text{corr}}(r_1)$ in equation (7).

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Figure 5. Comparisons of $\Delta D(r_1)$ for the 2p²³P state of H⁻, He, Li⁺ and Be²⁺ using, in **each case, two energetically different correlated wavefunctions; see the diagrams A-D,** respectively. For $Z=1$, results for $\Delta D(r_1)$ are shown when $D_{corr}(r_1)$ in equation (7) is derived, in turn, from the correlated D-20 and D-84 wavefunctions. For each $Z \ge 2$, the $\Delta D(r_1)$ curves correspond to the appropriate D-13 and D-70 correlated functions.

based on unoptimized hydrogenic orbitals. The $\Delta D(r_1)$ curves arising from the two versions of equation (1) for each system are compared in figure **5,** as indicated. The $D_{HF}(r_1, r_2)$ and $\Delta D(r_1, r_2)$ functions are shown as surfaces in figure 6 for H⁻ and He; $D_{\text{corr}}(r_1, r_2)$ in equation (9) was obtained from **D-84** for H⁻ and **D-70** for He. Since the results for Li^+ and Be^{2+} differ, essentially, only in scale from those for He, they are not included in figure **6.**

The expectation values $\langle r_1^n r_2^n \cos \gamma \rangle$, $\langle r_1^n \rangle$ and $\langle r_1^n r_2^n \rangle$ for these $2p^2$ ³P states are given in tables 1-3, where *n* is an integer. $\langle \gamma \rangle$, obtained from $P(\gamma)$, is quoted in table 1 and the standard deviation σ for each $D(r_1)$ is given in table 2. For each *Z*, tables 2 and 3 contain values derived from HF(NUM) and its HF(STO) representation. The correlationinduced changes in $P(\gamma)$ and $D(r_1)$ can also be assessed by evaluating

$$
Y_{\gamma} = \frac{1}{2} \int_{0}^{\pi} |\Delta P(\gamma)| d\gamma \text{ and } Y_{r} = \frac{1}{2} \int_{0}^{\infty} |\Delta D(r_{1})| dr_{1}.
$$
 (11)

The results for Y , and Y _r, expressed as percentages of the integrated HF distributions, are given in tables 1 and 2. In addition, defining the change in a statistical correlation coefficient as $\Delta \tau = \tau$ (corr) - τ (HF), we determined $\Delta \tau_{1/r}$, $\Delta \tau_r$, $\Delta \tau_{\gamma}$, $\Delta \tau_{\gamma}$ and $\Delta \tau_{\gamma}$. Definitions of these radial and angular τ , involving $\langle r_1^n r_2^n \rangle$ and $\langle r_1^n r_2^n \cos \gamma \rangle$ respectively, have been quoted recently by Banyard **(1990),** see equations **(1)-(5). For** radial correlation, $\Delta \tau_{1/r}$ places the emphasis on the *inner* regions of the two-particle density in position space whereas $\Delta \tau_r$ locates the emphasis in the outer regions of the density. Similarly, when discussing angular correlation, we note that $\Delta \tau_{\rm v}$, $\Delta \tau_{\rm v}$ and $\Delta \tau_{\rm v}$ possess, in that order, an emphasis which moves progressively from the inner to the outer regions of the two-particle density. The values are presented in table 4.

3. Discussion

3.1. Angular correlation

For a 2p²³P state, the $P_{HF}(\gamma)$ curve in figure 1A is applicable for any *Z*. Since the two electrons singly occupy, say, the $2p_0$ and $2p_1$ orbitals at the independent-particle level, the inter-electronic angular distribution is symmetric about its maximum. Hence in table 1, $\langle \gamma \rangle = 90^{\circ}$ for the HF descriptions. Naturally, the sin γ term arising from the Jacobian and contained, by implication, in the definition of $P(\gamma)$, see equation (2), renders the distribution function zero when $\gamma = 0^{\circ}$ or 180° for *any* state.

As we progress from the **D-20** to the **D-84** wavefunction for H-, the correlation energy is improved by 0.27%. Figure **1A** shows that this improvement is reflected in $\Delta P(\gamma)$. Although the crossover points are virtually coincident at about 90°, the energetically better curve (D-84) is seen to be not quite so deep in the $0 < \gamma < 90^{\circ}$ region and, consequently, it is less positive when $90^{\circ} < y < 180^{\circ}$. As found for $P_{HF}(y)$, each angular Coulomb hole in figure 1 is effectively zero at small γ ; a similar behaviour occurs as $\gamma \rightarrow 180^\circ$.

When $Z \ge 2$, the **p**-13 wavefunction recovers over 99.9% of the correlation energy and the use of the D -70 function improves the result by $\leq 0.08\%$. Therefore, for He and the positive ions, it is not surprising that the two $\Delta P(\gamma)$ curves are indistinguishable to within graphical accuracy: see figure 1B. For general comparison, figure 1B includes the $H^-(D-84)$ curve. Table 1 reveals that the *introduction* of correlation produces a

characteristics of the surfaces are similar to those for He.

Table 1. Expectation values $\langle r_1^n r_2^n \cos \gamma \rangle$ and $\langle \gamma \rangle$ for the He(2p²³P)-like systems when $1 \le Z \le 4$, where $n = -1$, 0, +1, and γ is the angle subtended by the position vectors r_1 and r_2 at the nucleus as origin. Y_{γ} is the percentage of the normalized probability function $P_{HF}(\gamma)$ redistributed due to electron correlation. Throughout this work, all distances are measured in atomic units.

System	Wavefunction ³	$\langle r_1^{-1}r_2^{-1}\cos\gamma\rangle$	$\langle \cos \gamma \rangle$	$\langle r_1^{+1} r_2^{+1} \cos \gamma \rangle$	$\langle \gamma \rangle$ (deg)	$Y_{\infty}(\%)$
$H^{-}(2p^{23}P)$	$D-20b$	-0.0019780	-0.097 966	-7.7127	96.200	9.18
	$D-84$	-0.0019176	-0.094344	-7.3833	95.971	8.84
$He(2p^2)^3P$	$D-13$	-0.010331	-0.071 449	$-0.744.38$	94.528	6.66
	$D - 70$	-0.010299	-0.071404	-0.74310	94.526	6.65
$Li^+(2p^2)^3P$	$D-13$	-0.017932	-0.048310	$-0.195.94$	93.065	4.49
	$D-70$	-0.017890	-0.048287	-0.19554	93.064	4.49
$Be^{2+}(2p^{2}P)$	$D - 13$	$-0.025,560$	-0.036378	-0.077893	92.310	3.37
	$D - 70$	-0.025494	-0.036363	-0.077804	92.308	3.38

^a Hartree-Fock (HF) values for $\langle r_1^n r_2^n \cos \gamma \rangle$ are zero in each instance. We also note that $\langle \gamma \rangle_{\text{HF}} = 90^\circ$.

 b_{D-20} , for example, refers to the Drake (1986) wavefunction with $N=20$ in the summation limit in equation (1).

Table 2. Values of $\langle r_1^n \rangle$ for the 2p² ³P states when $1 \le Z \le 4$ and $-2 \le n \le +2$. Also quoted is σ , the radial standard deviation, and Y_r, the percentage of the normalized density $D_{\text{HF}}(r_1)$ which is redistributed due to electron correlation.

System	Wavefunction	$\langle r_1^{-2} \rangle$	$\langle r_1^{-1} \rangle$	$\langle r_1^{+1} \rangle$	(r_1^2)	σ	Y_r (%)
$H^{-}(2p^{2}P)$	HF(NUM)	0.041 392	0.16731	8.3220	92.467	4.8177	
	HF(STO)	0.041 388	0.167 29	8.3243	92.530	4.8203	
	$D-20$	0.043 008	0.161 54	10.304	199.22	9.0830	14.2
	$D - 84$	0.042999	0.16067	11.481	251.16	10925	15.3
$He(2p^{23}P)$	HF (NUM)	0.24082	0.41790	3.0814	11.691	1.4817	
	HF (STO)	0.240 82	0.41790	3.0814	11.691	1.4817	
	$D-13$	0.24170	0.418 10	3.0895	11.782	1.4958	0.404
	$D-70$	0.24172	0.418 10	3.0899	11.791	1.4979	0.378
$Li^{+}(2p^{2}^{3}P)$	HF (NUM)	0.60697	0.66795	1.9043	4.4165	0.88892	
	HF (STO)	0.60697	0.66795	1.9043	4.4166	0.88893	
	$D-13$	0.607 65	0.668 05	1.9057	4.4258	0.89118	0.130
	$D-70$	0.607 68	0.668 05	1.9058	4.4271	0.89174	0.114
$Be^{2+}(2p^2)^3P$	HF (NUM)	1.1398	0.91796	1.3786	2.3045	0.635 60	
	HF (STO)	1.1398	0.91796	1.3786	2.3045	0.635 59	
	$D-13$	1.1403	0.91802	1.3790	2.3066	0.636 27	0.0691
	D-70	1.1403	0.91802	1.3791	2.3071	0.636 54	0.0530

noticeable increase in *(y)* **for** each system, particularly for H-. However, the *improvement* in the correlated wavefunctions discussed above is seen to cause a reduction in *(y)* for H- of about *0.23"* whereas, **for** He, **Li'** and **Be2',** the decrease is **only** marginal. We also note that, for each system, the quantum mechanical results for $\langle \gamma \rangle$ and the angular probability distributions $P_{HF}(\gamma)$ and $\Delta P(\gamma)$ lend support to the quasiclassical picture **of** a 2p2 **'P** state as being a floppy linear 'triatomic molecule', see Krause *et al* **(1987).** This molecular model **of** electron correlation, initially proposed by Herrick and Kellman **(1980),** assumes that the **two** electrons are localized diametrically at equal

System	Wavefunction	$(r^{-2}r_2^{-2})$	$\langle r_1^{-1}r_2^{-1} \rangle$	$\langle r_1^{+1}, r_2^{+1} \rangle$	$\langle r_1^{-2}r_2^{+2} \rangle$
$H^-(2p^2)^3P$	HF(NUM)	0.001 7133	0.027991	69.256	8549.5
	HF (STO)	0.0017129	0.027 987	69.294	8561.8
	$D-20$	0.000 786 24	0.020 255	85.210	11596
	D-84	0.000 766 24	0.019757	91.951	14 727
He(2p ²³ P)	HF (NUM)	0.057994	0.174 64	9.4952	136.67
	$HF(STO)$.	0.057994	0.174 64	9.4952	136.67
	$D-13$	0.050 515	0.16882	9.2444	122.83
	$D-70$	0.050 595	0.16883	9.2471	123.08
$Li^+(2p^2)^3P$	HF (NUM)	0.36841	0.446 15	3.6264	19.506
	HF (STO)	0.36841	0.446 15	3.6264	19.506
	$D-13$	0.33971	0.437 58	3,5676	18.303
	$D-70$	0.340 02	0.437 60	3.5680	18.323
$Be^{2+}(2p^2)^3P$	HF(NUM)	1.2991	0.842 64	1.9006	5,3109
	HF(STO)	1.2991	0.842 64	1,9006	5.3109
	$D-13$	1.2272	0.831 25	1,8785	5.0789
	D-70	1.2273	0.831 26	1.8786	5.0800

Table 3. Values of $\langle r_1^n r_2^n \rangle$ for the $2p^2 {}^3P$ states when $1 \le Z \le 4$ and $-2 \le n \le +2$.

Table 4. Changes due to correlation in various statistical correlation coefficients τ for the $2p^2$ ³P states when $1 \le Z \le 4$. $\Delta \tau_{1/2}$ and $\Delta \tau_1$ assess the radial component of correlation and $\Delta \tau_{\gamma}$, $\Delta \tau_{\gamma}$ and $\Delta \tau_{\gamma}$ assess the angular component. For each component, the chosen $\Delta \tau$ represent **a** progressive shift of emphasis from the inner to the outer regions of the two-particle probability distribution. **We** note that, since **T(HF)** is zero far these systems, each $\Delta \tau = \tau$ (corr).

We note that $\tau_{\mathbf{v}} = \langle \cos \gamma \rangle$ **. The definition of each** τ **used here has been quoted recently by Banyard** (1990), equations **(1-5).**

distances from the nucleus: the system then experiences quantized collective rotations and large bending vibrations. Ezra and Berry **(1983)** have used this model to examine, in detail, a 'molecular' interpretation of the dynamics of correlation in various **DES** for a series of He-like ions. Of particular interest is their comment that the $2p^2$ ³P state can be regarded as a floppy molecule with one quantum of bending vibration. Figure 7 of Ezra and Berry (1983) shows that their conditional probability density for the $2p^{2}$ ³P state has a maximum around $\gamma = 90^{\circ}$ and, because of symmetry constraints, possesses nodes at $\gamma = 0^{\circ}$ and 180°. More importantly, however, they observe that, in the region when r_1 equals the most probable value for r_2 , angular correlation shifts

the maximum in the density to values of γ which are slightly greater than 90°. As Z was increased, Ezra and Berry (1983) found that the conditional probability density tended to become more symmetric about $\gamma = 90^{\circ}$.

The simple Z-scaling employed here in figure 2 for the angular holes produces a near-coincidence over the whole range, except of course for H-. Clearly, even though $\Delta P(\gamma)$ for **H**⁻ has the largest magnitude in figure 1B, the angular hole is too small to fit in with the inverse-Z behaviour observed here. When $Z \ge 2$, the inverse-Z effect is also shown in the Y_{γ} values and in the *excess* of the correlated $\langle \gamma \rangle$ values over the HF result of go", see table 1 **in** each case. For He, the comparison in figure 3 of the angular hole for $2p^2$ ³P with those for the 1s2p³P and 1s²¹S states reveals large differences in magnitude, the curves being ordered as *DES>* ground state > singly excited state. Indeed, the **DES** hole **is** over 50% deeper than the 1s' **IS** hole and several times deeper than that for 1s2p³P. Nevertheless, we note that the value for γ which *locates* the minimum in each curve increases according *to* the occupancy of a p orbital. Such an ordering is not applicable to the crossover or nodal point where, as figure 3 shows, the 1s2p³P curve has the largest γ value. However, in a study of angular holes for the ground state (Banyard and Ellis 1972), it was found that a wavefunction which recovered about 80% of the correlation energy produced a nodal point with a γ value larger than that obtained when the description was over 95% correlated. Thus, for the $1s2p³P$ state, the use of a correlated wavefunction which is less accurate than those for the other states may have yielded a nodal *y* which is, in reality, too large. Obviously, for He, the differences between these curves are sufficiently large to conclude that angular correlation has a much greater effect on the slow moving **DES** electrons than on the **ls2 'S** or ls2p **'P** electrons, in spite of the diffuse nature of the **DES** charge cloud.

3.2. Radial correlation

When examining the influence of correlation on the radial distributions $D(r_1)$ and $D(r_1, r_2)$, defined in equations (6) and (8), respectively, it is to be expected that the one-particle function $D(r_1)$ will be less sensitive than $D(r_1, r_2)$. In figure 4A, the $D(r_1)$ curves have been Z-scaled in such a way that normalization is preserved. The curves for $1 \le Z \le 4$, derived from the *HF(STO)* results, are observed to converge towards $D(r_1)$ generated from an independent-particle $2p²$ **P** wavefunction using unoptimized hydrogenic orbitals. As in BKD, this confirms the *decreasing* importance of the interelectronic repulsion term as Z increases. In figure 4B, the $\Delta D(r_1)$ against Zr_1 curves obtained from the energetically best correlated wavefunctions show a roughly common *form* when $Z \ge 2$. The curve for H⁻ is not included due to its extensive r_1 range *and* the comparatively massive magnitude for $\Delta D(r_1)$, see figure 5. The 'double-occupancy' of the 2p orbital, with its non-zero angular momentum, is seen to produce curves for $D_{HF}(r_1)$ and $\Delta D(r_1)$ which have negligible values at small r_1 . By contrast, the HF curves for **1s' 'S** and ls2p **'P,** each involving a 1s orbital, are known to rise rapidly from the origin. *An* attempt was made to reduce the ranges of the cross over regions in figure 4B by using a $(Z - k)$ scaling factor. Unfortunately, the screening parameter $k = 0.28$ required for a minimized spread of inner nodes was quite different from that of 0.49 required for the outer nodes.

For each system, the radial 'in-out' effect of correlation is immediately apparent. Like the 'split-shell' effect for the 1s² ¹S state, a $\Delta D(r_1)$ curve is seen to be most negative at a Z_r value slightly in excess of that for the maximum in the corresponding $D_{HF}(r_1)$: we also note that each set of Zr_1 values approaches a limit as Z increases. Obviously,

as *Z* becomes larger, $\Delta D(r_1)$ decreases in magnitude and relative importance. This is supported by the **Y,** values in table *2* where the *decrease* with *Z* is considerably greater than that observed for **Y,** in table I.

In figure 5, the $\Delta D(r_1)$ curves for H⁻ show a distinction between the **D-20** and **D-84** correlated wavefunctions and, **for** He and the positive ions, differences exist between results derived from the **D-13** and **D-70** functions. **As** expected, for these pairs of highly correlated wavefunctions the differences, although real, are indeed very small when compared with $D_{HF}(r_1)$. Nevertheless, figure 5 shows that the improvement in the correlated wavefunction for H⁻ deepens the radial hole and makes $\Delta D(r_1)$ more extensive. For $Z \ge 2$, improvement again extends each curve but the hole is *less* deep than that for **p**-13.

Examination of the two-particle radial densities in figure 6 reveals that the relatively massive correlation effect for H⁻ extends well beyond the region of the $D_{HF}(r_1, r_2)$ surface. For He, $\Delta D(r_1, r_2)$ is almost confined within the (r_1, r_2) range for the HF surface, a feature which was clearly apparent when comparing the surfaces for Li⁺ and for Be²⁺ (the pairs of surfaces for $Z = 3$ and 4 are not shown for reasons of space). The 'in-out' effect is highlighted in $\Delta D(r_1, r_2)$ by a maximum being located at small *r,* and large *r,,* and vice versa, coupled with the occurrence of a reduction in probability whenever $r_1 = r_2$.

3.3. Expectation values and AT

Table 1 shows that, in keeping with $\langle \gamma \rangle_{\text{corr}}$ being greater than $\langle \gamma \rangle_{\text{HF}} = 90^{\circ}$, the introduction of correlation produces a negative value for the angular-related properties $\langle r_1^n r_2^n \cos \gamma \rangle$. For each system, the magnitude of the value when $n = +1$ is several times larger than the result when $n = 0$ or -1 : in fact, for H^- , we note that in relative terms the size of $\langle r_1^{+1} r_2^{+1} \cos \gamma \rangle$ is extremely large. When *Z* is increased, $\langle r_1^{+1} r_2^{+1} \cos \gamma \rangle$ decreases rapidly in magnitude but, as mentioned, it remains the dominant value-in spite of $\langle r_1^{-1}r_2^{-1} \cos \gamma \rangle$ becoming steadily more negative. Improvement in the correlated wavefunctions is seen to decrease the magnitude of each expectation value for all *2,* thus indicating a reduction in angular correlation, as observed earlier.

In table 2, correlation increases $\langle r_1^n \rangle$ and σ for He and the positive ions. For H⁻, a noticeable reduction occurs for the energy-related property $\langle r_1^{-1} \rangle$, but all other quantities are increased. Indeed, the change in $\langle r_1^{+2} \rangle$ for H⁻ is over twofold and reflects the sizeable 'in-out' radial effect discussed above. When $Z \ge 2$, going from $D-13$ to **D-70** is seen to yield a marginal or zero increase in $\langle r_1^n \rangle$ and a slight increase in σ . However, improving the correlated description for **H'** causes a *signifcant* increase in $\langle r_1^{+1} \rangle$, $\langle r_1^{+2} \rangle$ and σ .

The $\langle r_1^n r_2^n \rangle$ values in table 3 show a marked sensitivity to correlation effects. Except when $n = +1$ and $+2$ for H⁻, all quantities are reduced in value by the use of the explicitly correlated wavefunctions: **as** might be anticipated, the reduction decreases with increasing *Z*. When $Z \ge 2$, these changes relative to the HF values range from a reduction of almost 13% down to about $4\frac{1}{2}$ %. However, when $Z = 1$, correlation produces a massive 55% reduction for $\langle r_1^{-2}r_2^{-2} \rangle$ and an *increase* of around 36% for $\langle r_1^{+2}r_2^{+2} \rangle$. Excluding H^- , the use of the energetically better correlated functions produces either a small or quite marginal increase in $\langle r_1^n r_2^n \rangle$. For H⁻, the use of the **D-84** description, instead of D-20, causes a decrease when $n \le -1$ and an increase when $n \ge +1$, the result for $\langle r_1^2 r_2^2 \rangle$ being changed by 27%. In passing, we note from tables 2 and 3 the high degree of similarity between the **HF(NUM)** values and those derived from its **HF(STO)** representation. Not surpisingly, the slightly reduced quality of the fitted function implied by the results for H⁻ is a reflection of the very diffuse nature of its $2p^2$ ³P wavefunction.

AU statistical correlation coefficients considered in table **4** are identically zero when derived from an **HF** wavefunction as, indeed, they are for *any* independent-particle representation of $2p^2$ ³P. Hence, $\Delta \tau = \tau$ (corr). Since, by definition, every τ is bounded between **-1** and +1, these different measures of angular and radial correlation are therefore capable of intercomparison: we recall that -1 and $+1$ correspond to perfect negative and positive correlation, respectively. **When** *2* **2,** table **4** reveals that the angular quantities $\Delta \tau_{\rm v}$ and $\Delta \tau_{\rm v}$ (emphasizing, in turn, the interemdiate and outer regions of the density) are always of larger magnitude than $\Delta \tau_{\gamma'}$ (emphasizing the inner regions of density). The largest value for H⁻ is $\Delta \tau_{\gamma}$. For each system, the radial quantities $\Delta\tau_{1/r}$ and $\Delta\tau_r$, are seen to be greater than the angular $\Delta\tau$; this is especially so **for H-** where both sets of radial values are greater by at least a factor of **3.5.** Once more, this highlights the distinctive importance of radial correlation in the description of $H^{-}(2p^{2}^{3}P)$.

Further interest in table 4 arises from the *ratios* of various $\Delta\tau$ and, in particular, their Z-dependent trends. For example, the ratio $\Delta \tau_r / \Delta \tau_{1/r}$ is 0.95, 1.50, 1.49 and 1.49 for H⁻, He, Li⁺ and Be²⁺, respectively, showing that the balance of *radial* correlation at different distances from the nucleus is similar in the neutral and positive systems, but is again distinct for H⁻. When ordered as before, the angular ratio of the outer to the inner effect, $\Delta \tau_v / \Delta \tau_v$, yields 0.66, 1.48, 1.50 and 1.51. Thus, when $Z \ge 2$, the relative strengths of *angular* correlation in different radial regions of the density are seen to be almost constant. Moreover, from the quite remarkable similarity between these two sets of ratios, we conclude that, for each system except **H-,** the relative measure of the angular effects is virtually *equal* to that for the radial effects. By contrast, the ratios for H^- are seen to be significantly different. It is also informative to examine the ratio of a radial to an angular $\Delta \tau$. For increasing Z, $\Delta \tau_{1/r} / \Delta \tau_{\gamma'}$ is 7.90, 2.10, 1.83 and 1.73 and $\Delta\tau_r/\Delta\tau_r$ is 11.4, 2.12, 1.82 and 1.71. Hence, we note that, in both the inner and outer regions of space, angular correlation becomes relatively more important as *Z* increases. The change in going from H⁻ to He is seen to be quite dramatic in both regions. Nevertheless, for each *2,* these ratios suggest that radial correlation remains, overall, the prevailing influence on the two-particle probability density.

Generally, the improvement in these correlated wavefunctions causes a decrease in magnitude for the listed $\Delta \tau$, the exceptions being $\Delta \tau_{1/r}$ for **H**⁻ and $\Delta \tau_{1/r}$ and $\Delta \tau_r$ for Be²⁺ which, as seen from table 4, become more negative.

4. Summary

Highly accurate explicitly correlated wavefunctions, of benchmark quality, have been used to examine angular and radial correlation effects in the doubly excited state $2p^2$ ³P when $1 \leq Z \leq 4$. Following our recent article on the total correlation effect, these separate components of electron correlation are assessed here in terms of an angular Coulomb hole, a radial hole, several expectation values and various angular and radial statistical correlation coefficients.

For He and the positive ions, a clearly defined inverse-2 relationship exists for the angular holes $\Delta P(\gamma)$ against γ , where γ is the inter-electronic angle subtended at the nucleus. Naturally, this behaviour also occurs in Y_{γ} : the overall change caused by

correlation in the normalized angular distribution $P(\gamma)$. In addition, the Z-effect is reflected in the difference between the correlated and Hartree-Fock **(HF)** expectation values for γ . Although H⁻ possesses much the largest angular hole in absolute terms, the curve is still too small to form part of the inverse-Z pattern; a similar conclusion holds for Y_y and the correlation-induced excess for $\langle y \rangle$. For He, the curve for the doubly excited state was compared with the angular holes for the 1s' **'S** and ls2p **'P** states. The $2p^2$ ³P state, with its diffuse probability density and relatively slow moving electrons, was found to have an angular hole which is noticeably *grealer* than that for each energetically lower state.

The influence of correlation on the one-particle radial density $D(r_1)$ reveals a marked 'in-out' effect which parallels the correlated 'split-shell' behaviour ofthe doubly occupied ground state. By analogy with the angular behaviour, the size of the radial hole $\Delta D(r_1)$ for H⁻ is much larger than that obtained for each of the other systems. However, irrespective of the H^- result, no simple Z-scaling effect could bring the remaining $\Delta D(r_1)$ curves into any overall agreement. For the two-particle radial function $D(r_1, r_2)$, the outward redistribution caused by correlation for H⁻ extends well beyond the (r_1, r_2) -range observed for $D_{HF}(r_1, r_2)$. For each system, the maxima in $\Delta D(r_1, r_2)$ at large r_1 and small r_2 , and vice versa, emphasized most forcefully the 'in-out' feature found in $\Delta D(r_1)$.

Correlation produces changes in the angular and radial expectation properties for **H**^{$-$} which are massive by comparison with the findings for $Z \ge 2$. Ratios of statistical correlation coefficients, each formulated from specific expectation values, proved to be highly informative. For every system except H^- , the ratio of the outer to the inner effect for both the radial and angular measures of correlation are virtually identical. Further, although statistical coefficients indicate that angular correlation increases in *relafive* importance as *Z* increases, they also suggest that the major influence on the two-particle distribution arises from radial correlation.

Finally, the dramatic changes created by each component of correlation for **H-** set it apart from He and the positive ions.

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