Photoemission from hybrid states of Cl@C_{60} before and after a stabilizing charge transfer

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Abstract
Photoionization calculations of the endofullerene molecule Cl@C_{60} with an open-shell chlorine atom are performed in the linear response density functional theory based on a spherical jellium model of C_{60}. Cross sections for atom–fullerene hybrid photoemission studied show the effects of the hybridization symmetry, the giant plasmon and the molecular cavity. Comparisons with the results of Ar@C_{60} provide insights into the role of a shell-closing electron and its influence on the dynamics. The results for Cl@C_{60} are further compared with those of a more stable configuration that results after a C_{60} electron transfers to Cl forming Cl\textsuperscript{−}@C_{60}\textsuperscript{+}. This comparison reveals noticeable differences in the ionization properties of the antibonding hybrid state while the bonding hybrid remains nearly unaltered showing a magnification covering the entire giant plasmon energy range. These predictions should raise interest in probing the molecular configuration using photoelectron spectroscopy.

Keywords: photoionization, endofullerenes, hybridization

(Some figures may appear in colour only in the online journal)

1. Introduction
Significant success in the synthesis of endofullerene molecules—systems of an atom or a smaller molecule incarcerated within the fullerene cage [1]—has spawned a series of experiments [2–4] using merged beam techniques at the Berkeley Advanced Light Source. The experiments accessed photoionization properties of these materials in gas phase. In addition, endofullerenes, being natural entrapment of atoms, have led to a series of theoretical studies of the effects of ionizing radiation on these systems, i.e., photoionization [5, 6]. Such fundamental spectroscopic knowledge is particularly useful due to a broad horizon of applied importance of these materials in, namely, (i) solid state quantum computations [7, 8], (ii) improving the superconducting ability of materials [9], (iii) biomedical fields [10], (iv) contrast-enhancement research for magnetic resonance imaging (MRI), (v) improving organic photovoltaic devices [11], and even in (vi) astrophysics [12].

Endofullerenes confining open-shell atoms have potential applied interests of rather exotic nature [13]. For instance, N@C_{60} has attracted interest due to its uniquely long spin relaxation times driven by the confinement [14]. Electron paramagnetic resonance study [15] of P@C_{60} has shown enhancement in hyperfine coupling of the phosphorous’ unpaired electrons with its nucleus which is attributed to the admixture of
excited states acquiring angular momentum from the cage. In contrast, the hyperfine interaction between the positively-charged muon and the unpaired electrons in the muonium atom in C_{60}, relative to free muonium, is predicted to diminish from the confinement [16]. These discoveries render particular relevance to study open-shell atomic endofullerenes, including assessing the spectroscopy of their more stable configurations resulting from electron transfers to atomic vacancies.

One unique phenomenon that ubiquitously occurs across endofullerene systems is the emergence of atom–fullerene ground state orbital hybridization. This entails the formation of symmetric (bonding) and antisymmetric (antibonding) hybrid states as the eigenstates of the whole system from the mixing of an atomic and a fullerene orbital of identical angular momentum symmetry. A number of our previous studies have predicted such hybrid states in various endofullerenes and their broad spectrum of photoionization properties [17–21]. It is therefore of particular value to scrutinize the sensitivity of the hybridization and the photoemission dynamics of these hybrid states via gentle changes of configuration features by, for instance, comparing the effects of confinement upon successive atoms in the periodic table. It is important to note that while the simple static annular square well (ASW) model potential is often useful [5, 22], the model does not include C_{60} electrons; consequently, the possibility of hybridization is omitted in ASW calculations. Besides, the absence of electrons in the ASW model vastly limits both ground-state and dynamical electron correlations.

A prototype case that we examine here is the hybrid level photoionization of Cl@C_{60} versus Ar@C_{60}; Cl has just one electron less than Ar in the outer 3p shell and an atomic number lower by one. Closed-shell Ar, being chemically inert, almost certainly locates at the center of the spherical C_{60}. We treat the barely open-shell Cl also within the spherical geometry so as to retain the same spherical calculation as was done for the Ar case. We then consider a system of Cl^{-} @C_{60}^{+} produced by the transfer of a C_{60} electron to fill in the Cl hole. This configuration likely attains more stability by forming closed-shell Cl^{-}. There has been experimental evidence, based on laser desorption mass spectroscopy, of C_{60} with a single Cl^{-} inside [23]. While it is expected that the polarization interaction of the ion can induce some offset in its position from the center of C_{60}, a density functional theory calculation with Born–Oppenheimer molecular dynamics indicates that this offset is quite small within neutral C_{60} [24]. Earlier studies showed only small effects of the cage polarization except very close to the ionization threshold [25]. Likewise, a relatively weak effect on the process from a small offset of the atomic location was predicted [26]. Therefore, we treat Cl^{-} @C_{60}^{+} assuming spherical geometry as we. Then compare the hybrid photoionization of this new configuration with Cl@C_{60}. Ultimately, the general comparison among these three endofullerene systems unfolds the delicate dependence of the hybridization and resulting photoionization cross sections on a shell-closing electron as well as on an electron transfer from the cage to the atom.

2. Brief description of theory

The details of the theory are described in reference [18]. Choosing the photon polarization along the z-axis, the photoionization dipole transition cross section in a linear response approximation of time-dependent density functional theory is given by

$$\sigma_{nl-kl'} \sim \left| \langle \psi_{nl} | z + \delta V | \phi_{kl'} \rangle \right|^2.$$  \hspace{1cm} (1)

Here k is the momentum of the continuum electron, z is the one-body dipole operator, $\phi_{kl'}$ is the single electron bound wavefunction of the target level, and $\psi_{nl}$ is the respective outgoing dipole-allowed continuum wavefunction, with $l' = l \pm 1$. $\delta V$ represents the complex induced potential that accounts for electron correlations within the linear response framework.

We model the bound and continuum states self-consistently using the independent particle local density approximation (LDA) modified by van Leeuwen–Baerends (LB) exchange–correlation functional [27]. The jellium potentials, $V_{jel}(r)$, representing 60 C^{+} ions for C_{60} is constructed by smearing the total positive charge over a spherical shell with known molecular radius $R = 3.54$ Å (6.70 a.u.) [28] and thickness $\Delta$. A constant pseudopotential $\pi$ is added to $V_{jel}$ for quantitative accuracy. The Kohn–Sham equations for the system of 240 electrons (four valence 2s^{2}2p^{2} electrons from each carbon atom), plus all electrons of the central atom with atomic number $\zeta$, are then solved in the LDA + LB potential

$$V_{LDA+LB}(r) = -\frac{\zeta}{r} + V_{jel}(r) + \int dr' \frac{\rho(r')}{|r-r'|} + V_{XC}[\rho(r)],$$  \hspace{1cm} (2)

to obtain the bound and continuum orbitals in equation (1) self-consistently as described in reference [18]. Besides the LDA term, equation (2) also includes the LB exchange–correlation functional in $V_{XC}$ involving the gradient of the electron density in the scheme described earlier [29]. This approach provides an accurate asymptotic description of the ground state potential. The values of $\Delta$ and $\pi$ are determined both by requiring charge neutrality and obtaining the experimental value [30] of the first ionization threshold of C_{60}. $\Delta = 1.30$ Å (2.46 a.u.) thus obtained closely agree with that extracted from measurements [28]. $\pi$ was found to be 12.1 eV. Obviously, omitting either $\zeta/r$ or $V_{jel}$ in equation (2) produces, respectively, the empty C_{60} and the free atomic results.

We remark that treating open-shell, thus non-spherical, 3p orbital density of CI in a spherical frame may produce inaccuracies in Hartree–Fock type methods that treat the non-local exchange exactly [31]. In the density functional model as in the current study, on the other hand, one may adjust the parameters of the functional, or use a different functional, to force accurate ground state properties [32]. However, we could not find a unique set of parameter values of our LB functional to work for both free Cl and empty C_{60}. Therefore, we have used the set, that is successful for C_{60}, for the endohedral composite-systems as well. But these values overestimate CI ionization potential of NIST database [33] by about 7%. They further produce an electron affinity of −4.15 eV as opposed to the measured value of −3.6 eV [34], even though CI^{-} has a closed-shell spherical 3p shell. But these
small inaccuracies should not take away much from the main results of this study which explores the dominant effects of C_{60}.

The time-dependent (TD) LDA + LB-derived $\varepsilon + \delta V(r)$ in equation (1) is proportional to the induced changes in the electron density as a function of the photon frequency $\omega$ [35]. This change is

$$
\delta \rho(r'; \omega) = \int \chi(r, r'; \omega) \delta \rho(r, \omega)
$$

where the full susceptibility, $\chi$, builds the dynamical correlation from the independent-particle LDA + LB susceptibilities

$$
\chi^0(r, r'; \omega) = \sum_{nl} \phi^0_n(r) \phi^0_n(r') \tilde{G}(r, r'; \epsilon_{nl} + \omega)
+ \sum_{nl} \phi_{nl}(r) \phi^*_{nl}(r') \tilde{G}^*(r, r'; \epsilon_{nl} - \omega)
$$

through the matrix equation

$$
\chi = \chi^0[1 - (\partial V / \partial \rho)]^{-1}
$$

involving the variation of the ground-state potential $V$ with respect to the ground-state density $\rho$. The radial components of the full Green’s functions in equation (4) are constructed with the regular ($f_L$) and irregular ($g_L$) solutions of the homogeneous radial equation

$$
\left( \frac{1}{r^2} \frac{\partial}{\partial r} r^2 \frac{\partial}{\partial r} - \frac{L(L+1)}{r^2} - V_{\text{LDA+LB}} + E \right) f_L(r; r; E) = 0
$$

as

$$
G_L(r, r'; E) \equiv \frac{2 f_L(r; E) h_L(r; E)}{W[f_L, h_L]}
$$

where $W$ represents the Wronskian and $h_L = g_L + if_L$ and $E$ is the photoelectron energy. Obviously, TD(LDA + LB) thus includes the dynamical many-electron correlation by improving upon the mean-field LDA + LB description.

3. Results and discussion

3.1. Cl@C_{60} versus Ar@C_{60}

3.1.1. Ground state atom–C_{60} hybridization: In an endo-fullerene system, an eigenstate of the free atomic Hamiltonian can admix with an eigenstate of the empty C_{60} Hamiltonian of the same angular momentum symmetry to produce hybrid states which are eigenstates of the combined system. Thus, ground state LDA + LB results for Cl@C_{60} reveal hybridization between valence 3pCl and a deeper 2pC_{60} state (2p state of a jellium C_{60}), both of the same angular character, which jointly produce symmetrically and antisymmetrically combined states of Cl@C_{60} that can be written as:

$$
|\text{Cl} + \text{C}_{60}\rangle = \sqrt{\alpha} |\phi_{3p\text{Cl}}\rangle + \sqrt{1 - \alpha} |\phi_{2p\text{C}_{60}}\rangle
$$

(7a)

$$
|\text{Cl} - \text{C}_{60}\rangle = \sqrt{\alpha} |\phi_{3p\text{Cl}}\rangle - \sqrt{1 - \alpha} |\phi_{2p\text{C}_{60}}\rangle
$$

(7b)

where the fraction $\alpha$ is the mixing parameter that renders the states orthonormal. Note that, we use the hydrogenic Coulomb notation of labeling for the atom and the standard harmonic oscillator notion for C_{60}. In figure 1, the radial components of these wavefunctions are shown and compared with the corresponding hybrid wavefunctions of Ar@C_{60}. Due to opposite mixing signs in equations (7a) versus (7b), the symmetric hybrid shows a radial oscillation at the shell region which is in-phase with that of 2pC_{60}, while it is out-of-phase for the antisymmetric hybrid. From a perturbation theory viewpoint, the strength of this mixing is proportional directly to the overlap of the participating (free) orbitals and inversely to the separation of their binding energies. As figure 1 indicates, the energy of 3pAr is extremely close to that of 2pC_{60}, while 3pCl moves a bit higher, leading to a stronger mixing with a value of $\alpha$ close to about 0.5 (equal share of atom–fullerene character) for Ar@C_{60}. This result is different from a weaker hybridization found in earlier LDA calculations of Ar@C_{60} [36, 37] which did not utilize the gradient corrected LB exchange–correlation functional of the current study that goes beyond LDA.

A somewhat reduced hybridization in Cl@C_{60} with a greater value of $\alpha$ thus implies enhanced Cl and enhanced C_{60} characters, respectively, for the symmetric and antisymmetric states (figure 1). This occurs in spite of a slightly increased wavefunction overlap due to a small displacement of 3pCl wavefunction toward the shell from that of 3pAr. Also note that the resulting symmetric hybrids of the systems are more separated energetically than the antisymmetric hybrids.

3.1.2. Photoionization of hybrid levels. Cross sections calculated at the correlated TD(LDA + LB) level for the hybrid
states of Cl@C₆₀ and Ar@C₆₀ are presented in figure 2 as a function of the photon energy. (Several narrow resonances are due to the many autoionization resonance channels that exist in the endfullerenes due to levels in the fullerene molecule that are not there in free atom.) Comparing these with 3p of free Cl and Ar indicates plasmon driven enhancements at low energies [20, 36]. This enhancement is significantly stronger for symmetric photoemission than the antisymmetric one. In the framework of interchannel coupling due to Fano, the correlation-modified [TD(LDA + LB)] matrix element of the photoionization of X ± C₆₀, X being Cl or Ar, can be written as [20],

\[
\mathcal{M}_±(E) = \mathcal{D}_±(E) + \sum_{nl} \int dE' \left| \psi_{nl}(E') \right|^2 \left| \phi_± \right|^2 \mathcal{D}_{nl}(E')
\]

(8)

in which the single electron (LDA + LB) matrix element is

\[
\mathcal{D}_±(E) = \langle \psi_{nl}(E) | \alpha \psi_{nl}(E) \rangle \langle \psi_{nl}(E) | \alpha \psi_{nl}(E) \rangle
\]

(9)

and \( |\psi_{nl} \rangle \) in the interchannel coupling integral is the (continuum) wavefunction of the \( nl \rightarrow kl' \) channel. Taking the hybridization into account, the channel wavefunctions in equation (8) become

\[
|\psi_+ \rangle = \sqrt{\alpha} |\psi_{3pX} \rangle + \sqrt{1 - \alpha} |\psi_{2pC_{60}} \rangle
\]

(10a)

\[
|\psi_- \rangle = \sqrt{1 - \alpha} |\psi_{3pX} \rangle - \sqrt{\alpha} |\psi_{2pC_{60}} \rangle
\]

(10b)

Substituting equations (7a) and (7b), but for a general X, and equations (10a) and (10b) in equation (8), and noting that the overlap between a pure X and a pure C₆₀ bound state is negligible, we separate the atomic and fullerene contributions to the integral to get the TD(LDA + LB) matrix element for \( X \pm C_{60} \) levels as,

\[
\mathcal{M}_+(E) = \sqrt{\alpha} \mathcal{M}_{3pX}(E) + \sqrt{1 - \alpha} \mathcal{M}_{2pC_{60}}(E)
\]

(11a)

\[
\mathcal{M}_-(E) = \sqrt{1 - \alpha} \mathcal{M}_{3pX}(E) - \sqrt{\alpha} \mathcal{M}_{2pC_{60}}(E),
\]

(11b)

where the second terms on the right-hand side are responsible for the plasmonic enhancements at the lower energies, as seen in figure 2. Obviously, \( |\mathcal{M}_{2pC_{60}}|² \) will produce the 2p cross section of empty C₆₀ which we also include in figure 2 for comparison. As seen, while the cross sections for the symmetric hybrid of both endohedral systems enhance far more than that of 2pC₆₀ at plasmon energies, they are comparable for the antisymmetric hybrids. Writing the atomic and C₆₀ contributions in equations (11a) and (11b) respectively as complex quantities \( X_i + iX_r \) and \( C_i + iC_r \) (subscript r and i for the real and imaginary part), and recalling that the cross section \( \sigma \) is proportional to the square modulus of the matrix element, we can express the cross sections as,

\[
\sigma_+(E) \sim (\sqrt{\alpha} X_i + \sqrt{1 - \alpha} C_i)^² + (\sqrt{\alpha} X_r + \sqrt{1 - \alpha} C_r)^²
\]

(12a)

\[
\sigma_-(E) \sim (\sqrt{1 - \alpha} X_i - \sqrt{\alpha} C_i)^² + (\sqrt{1 - \alpha} X_r - \sqrt{\alpha} C_r)^².
\]

(12b)

Evidently, the enhancement of the symmetric state cross section, equation (12a), involving the sum of real and imaginary components, will be universally larger than that of the antisymmetric state which include their differences. Indeed, this is seen for both Cl@C₆₀ and Ar@C₆₀ in figure 2 and a direct consequence of the in-phase versus out-of-phase radial oscillation of the hybrid wavefunctions at C₆₀ shell (figure 1). However, the detailed similarities and differences of enhanced cross section between two systems must depend on their respective values of \( \alpha \) in a rather complicated way that involves the interferences between the atomic and the C₆₀ components of the matrix elements in equations (12a) and (12b). We note that while this enhancement is of a comparable size for the symmetric states of both systems in figure 2(a), it is significantly weaker for antisymmetric Cl–C₆₀ compared to Ar–C₆₀ (figure 2(b)). The situation further complicates from the effect of lowering binding energies of Cl@C₆₀ hybrid levels as is discussed in section 3.2 below.

As the plasmonic effect weakens with increasing energy, the cross sections largely follow their free atom curves, as seen in figure 2. Cooper minimum-like structures develop [38] around 50 eV on both the symmetric curves with the minimum of Cl@C₆₀ being deeper due to its higher atomic character. Note that the Cooper minima in 3p cross sections for free atoms are clearly visible in figure 2. In any case, such minima also show up around 60 eV for antisymmetric emissions where the structure for Cl@C₆₀ is very weak because of its weaker atomic character.

Above these energies the cross sections oscillate as a consequence of a well-known multipath interference mechanism [39] due to the cavity structure of C₆₀ which was modeled.

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Figure 2. TD(LDA + LB) photoionization cross sections of symmetric (a) and antisymmetric (b) levels of Cl@C₆₀ compared to Ar@C₆₀. Cross sections for 3p of free Cl and free Ar, and for 2p of empty C₆₀ are also plotted for comparisons.
earlier in detail in reference [40]. At such high energies the interchannel coupling in equation (8) vanishes to simplify equations (11a) and (11b) to,

$$D_+(E) = \sqrt{\alpha D_{3p} x(E)} + \sqrt{1 - \alpha D_{2pC_{60}}(E)} \quad \text{(13a)}$$

$$D_-(E) = \sqrt{1 - \alpha D_{3p} x(E)} - \sqrt{\alpha D_{2pC_{60}}(E)}. \quad \text{(13b)}$$

The multipath interference model is based on the following mechanism. The matrix element $D$ can generally be isolated in to two components, one arising from the atomic region and other from the $C_{60}$ shell region, and can be written down respectively as follows [40]

$$D_x \sim D_{\text{atom}}(k) + A_{\text{refl}}(k) \left[ e^{-ikR_i} e^{-ikR_0} e^{-ikD} - e^{-ikD} \right] \quad \text{(14a)}$$

$$D_C \sim A_{\text{shell}}(k) e^{-ikD} \left[ a_i e^{-ikR_i} - a_o e^{-ikR_o} \right]. \quad \text{(14b)}$$

where the photoelectron momentum $k = \sqrt{2(E - \varepsilon_\pm)}$ in atomic units, $a_i$ and $a_o$ are the values of $\phi_\pm$ at the inner and outer radii $R_i$ and $R_o$ of $C_{60}$, and $v_0$ is the average depth of the shell potential. In equation (14a), $D_{\text{atom}}$ represents the direct ionization amplitude from the atomic region. The second term in this equation embodies the reflection of this outgoing photoelectron wave from both inner and outer surfaces of the shell. Quantitatively, this induces oscillations as a function of the photoelectron momentum with amplitude $A_{\text{refl}}$ and frequencies related to $D_i$ and $D_o$, the inner and outer diameters of the shell. The direct and the reflected parts coherently interfere in the cross section. Since, obviously, $A_{\text{refl}}$ is proportional to $D_{\text{atom}}$, the larger the atomic component of a hybrid wave-function, the stronger is the reflection and the higher is the chances that the oscillations occur about the free atom result. This is exactly what is seen for the high energy cross section of Cl + $C_{60}$ in figure 2(a). On the other hand, equation (14b), the portion of the overlap integral from the shell region, produces two localized emissions from shell edges, where the maximum ionizing forces become available due to rapid variations of the shell potential there. Such a diffraction-type effect translates into another oscillation in frequencies related to $D_i$ and $D_o$. This part will also add to coherent interference in the cross section and will dominate if a hybrid level has a stronger $C_{60}$ character, like for Cl–$C_{60}$. Indeed, the oscillations in the cross section of Cl–$C_{60}$ intensify at higher energies while the average value fall significantly lower than free 3pCl (figure 2(b)). For the Ar@$C_{60}$ hybrids, however, due to their almost equal share of atom–$C_{60}$ character (figure 1), the strength of high-energy cross sections are comparable, somewhat below 3pAr, but the differences in the details of their shapes again owe to the interference between reflective and diffusive emissions described above. We further note in figure 2 that for the pure 2p$C_{60}$ shell, which predominantly emits by the diffraction mechanism of equation (14b) at high energies, the result shows sharper oscillations with very rapid fall-off due to the absence of any atomic type steady emission [41].

3.2. Cl@$C_{60}$ versus Cl’@$C_{60}$

A reactive Cl atom is very likely to capture an electron from $C_{60}$ which will likely bring the compound to a more stable configuration Cl’@$C_{60}$+. If an electron transitions from the highest occupied (HOMO) level of $C_{60}$ of binding energy $-7.52$ eV to $3pCl$ level of electron affinity $-4.15$ eV, then the resulting configuration with the hole in HOMO will have the minimum total energy and, therefore, will be the ground state configuration for Cl’@$C_{60}$+. This does not even take into account the extra binding associated with the electrostatic Coulomb attraction between Cl’ and $C_{60}$+. Despite the electron affinity of Cl being less than the ionization potential of $C_{60}$ (see above), this extra binding is what that will enable the ionic compound to bind. Therefore, this ground state configuration of Cl’@$C_{60}$+ should be stable and abundantly formed.

We further note that if Cl@$C_{60}$ is already at the lowest energy, then a $C_{60}$ electron relocating to Cl will raise the energy of the molecule; the latter being stable will, thus, form a metastable configuration. On the other hand, the realistic ground state of the compound may as well be a mixture of Cl@$C_{60}$ and Cl’@$C_{60}$+ with more of the latter’s character. However, the comparison of the photoionization properties between these two diabatically unique configurations, as carried out in the present study, is of interest to understand the interplay between the two modes. Moreover, the similarities and variations of these properties based on two distinct configurations suggest the experimental need to access the real system via photoelectron spectroscopy [28], especially of the photoelectrons emitted from the atom–fullerene hybrid levels.

The excited state configurations of Cl’@$C_{60}$+ will be formed going higher in total energy as the transferred electron originates from gradually deeper states of $C_{60}$, ignoring the relaxation of the hole. For an empty $C_{60}$+, our LDA + LB ground state structure is insensitive to the location of the hole among the molecular levels which is not too surprising for a cloud of 240 delocalized electrons. Likewise, for the complex Cl’@$C_{60}$+ the LDA + LB energies and wavefunctions of all pure $C_{60}$ and hybrid states are found independent of which $C_{60}$ orbital the hole is situated at. This fact makes our current result rather robust. Furthermore, the energies and wavefunctions for pure $C_{60}$ levels of the compound remain practically identical to those of empty $C_{60}$, and therefore of little interest. However, the hybridization between 3pCl’ and 2p$C_{60}$+ is somewhat modified from that in Cl@$C_{60}$+, or, in other words, between after and before the electron makes a transition, as we describe below. Yet it is found that the hybridization in Cl’@$C_{60}$+ is still insensitive to the $C_{60}$ level the electron transitions from, that is the location of the hole. Therefore, our results presented here are robust being free of a specific choice of the hole level. On the other hand, we calculate Cl levels in Cl’@$C_{60}$+ to have higher binding energies than the Cl levels of Cl@$C_{60}$ to match the trend that it takes about 4.15 eV to remove a 3p electron from Cl’ to produce Cl.

The hybridization in Cl’@$C_{60}$+ is compared to that in Cl@$C_{60}$ in figure 3. Note that for Cl’@$C_{60}$+ the
participating free levels 3pCl$^-$ and 2pC$_{60}^+$ are significantly apart from each other (figure 3), the former becoming quite shallower and the latter becoming deeper compared to their counterparts in Cl@C$_{60}$ (figure 1). This strongly disfavors the hybridization. Conversely, as also seen in figure 3, the radial wavefunction of 3pCl$^-$ extends more radially outward than 3pCl to increase its overlap with 2pC$_{60}^+$ which favors the hybridization. In the tug-of-war between these two effects, the former wins resulting in some modification of hybridization for Cl$^-$ @ C$_{60}^+$ configuration as displayed in figure 3. It is seen that the atomic character stays unchanged, but the C$_{60}$ character weakens at the shell for the symmetric hybrid, while the reverse is true for the antisymmetric hybrid. Comparing the energy of the hybrid levels, the symmetric hybrid moves energetically higher (figure 3(a)) while the antisymmetric hybrids barely separate (figure 3(b)) as a result of the charge transfer. And this modified hybridization affects the resulting photoionization cross sections.

Figure 4 compares the TD(LDA + LB) cross sections for photoionization from hybrid levels of Cl$^-$ @ C$_{60}^+$ with Cl@C$_{60}$. Note that even though there is a slight modification in hybridization (figure 3) post electron transfer, the cross section of the symmetric level (figure 4(a)) is hardly modified, except that Cl$^-$ + C$_{60}^+$ starts at a lower photon energy due to the reduction of its binding energy. To be more precise, some reduction in the amplitude of the structure of Cl$^-$ + C$_{60}^+$ wavefunction (figure 3(a)) in the shell region results in little effect on the cross section. In the energy region of the plasmon, this can be understood in general from the interchannel coupling contribution of the matrix element in equation (8). Specifically, this term in part embodies the overlaps of the ‘fractional’ ionization channel emanating from the shell region of the symmetric hybrid with all C$_{60}$ channels which are accounted for in $\mathcal{M}_{2pC_{60}}$ in equation (12a). The channel overlap includes both the overlap between the bound and the continuum wavefunctions. While the reduction of the structure in Cl$^-$ + C$_{60}^+$ radial wave noted above lowers the aggregated strength of bound overlaps, the continuum overlaps will be slightly favored due to the following reason. A continuum overlap is more efficient at higher photoelectron momenta $k$, since differences between the momenta due to different level energies reduce enabling the continuum waves to oscillate progressively in phase with each other. Therefore, the Cl$^-$ + C$_{60}^+$ state opening at a lower photon energy benefits continuum overlaps to cause its net increase. This gain must be compensating the loss due to weakening hybridization to affect practically no change in the symmetric hybrid level cross section (figure 4) after the electron transfers. Note that the term $\mathcal{M}_{\text{f}(X)}$ in equation (12a) has no effect here due to about the same atomic character of hybrid wavefunctions before and after the electron switches locations. For the higher energy emission of this hybrid (figure 4(a)), a rather miniscule weakening of the multipath oscillations following the electron transfer traces to the reduction of its C$_{60}$ character that slightly reduces the collat-

Differences between the cross sections of antisymmetric hybrid states of Cl$^-$ @ C$_{60}^+$ and Cl@C$_{60}$, shown in figure 4(b), are rather strong. The differences at plasmonic energies up to 40 eV are due to the reduction of the atomic character (figure 3(b)), particularly via the interference effects illustrated in equation (12b). Note that since the binding energies of this hybrid state suffer a very little change upon the charge transfer, the continuum overlap effect described above does not apply.
However, at higher energies, 80 eV and above, the difference in cross sections grows progressively stronger. This must be due to a cumulative effect of the reduction of the atomic component of direct ionization, as a result of the reduced atomic character in Cl$^{-}$ → C$_{60}^+$, the subsequent reduction of reflective amplitude in equations (14a) and (14b), and the interference between them. To this end, even after the molecule evolves to a stable configuration Cl$^{-}$ @C$_{60}^+$, the strong plasmonic magnification of the emission from the symmetric hybrid remains, but the high energy response of the antisymmetric state substantially changes. More generally, comparing results of figure 2 with 4, we note a slow trend of modification for the symmetric hybrid cross sections from Ar@C$_{60}$ to Cl@C$_{60}$ to Cl$^{-}$ @C$_{60}^+$, while the change is rather dramatic along this sequence for the antisymmetric level. Finally, we note from equations (14a) and (14b) that for a possible small offset of Cl or Cl$^{-}$ in the cage, the multipath interference oscillations will somewhat weaken from a dephasing effect, since the distance, the effective radius (R), of the atom to the cage will slightly vary from different directions.

4. Conclusions

Using a fairly successful methodology of the time-dependent local density approximation based on a spherical jellium modeling of C$_{60}$’s ion core, we compute the photoemission cross sections of the atom–fullerene hybrid levels of Cl@C$_{60}$. A comparison of the results with those of Ar@C$_{60}$ probes the modification effects of a shell-closing electron on the properties of these hybrid photoemissions. However, Cl@C$_{60}$ must be an unstable system and will likely induce an electron transfer from C$_{60}$ to Cl to reach a more stable configuration. We compared the results between these two configurations of Cl endofullerenes as well to assess the effects of this electron transition on the hybrid photodynamics. Thus, in general, tuning the configuration gently along the sequence of Ar@C$_{60}$ to Cl@C$_{60}$ to Cl$^{-}$ @C$_{60}^+$, a systematic evolution of the ionizing response of hybrid states is uncovered. A strong magnification of the low energy emission of the symmetric hybrid over the entire giant plasmon resonance region and an enhanced multipath interference effects for the high energy antisymmetric hybrid are the most robust features in our study. In specific, however, noticeable differences in the properties of the anti-bonding hybrid state ionization of Cl@C$_{60}$ versus Cl$^{-}$ @C$_{60}^+$ are predicted, while the bonding hybrid remains nearly unaltered. This is the first study of the photoionization properties of a halogen endofullerene to the best of our knowledge. Like for any level of a complex target, the emission cross sections of the hybrid levels reported in this study can be experimentally accessed by conventional photoelectron spectroscopy techniques. This may open a direction to probe the ground and metastable excited configurations of halogen endofullerenes where an internal electron transition is possible.

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