# **Photoionization of endohedral atoms: Molecular and interchannel-coupling effects**

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Calculations of the photoionization cross section of the 2 $p$  and 3s subshells of free Ar and Ar@C<sub>60</sub> as examples have been performed using the molecular structure of the confined system and time-dependent density functional theory for the dynamical quantities. The results for Ar 2*p* in the combined system exhibit significant confinement resonances with the lower-energy ones being quite sharp, in contrast to the results of jellium-model calculations. In addition, calculations done with and without interchannel coupling between the photoionization channels of the 2*p* subshell of the Ar atom and the 1*s* subshell of the  $C_{60}$  shell show that, in this case, the coupling is of negligible importance, even though the C 1*s* cross section is more than an order of magnitude larger than that of Ar 2*p* in the 300 eV range. The Ar 3*s*, which is not hybridized, also exhibits confinement resonances, but is very strongly affected by interchannel coupling with photoionization channels from the  $C_{60}$  shell. The phenomenology of both 2*p* and 3*s* subshells is explained in terms of the interchannel-coupling matrix elements. These results should be applicable to inner-shell ionization of essentially any endohedral fullerene system.

DOI: [10.1103/PhysRevA.92.023405](http://dx.doi.org/10.1103/PhysRevA.92.023405) PACS number(s): 33*.*80*.*Eh*,* 33*.*90*.*+h

### **I. INTRODUCTION**

Over the past decade or so there have been a large number of studies of the photoionization of endohedral atoms—atoms confined in a fullerene or other type of cage  $[1-14]$ . These investigations were stimulated by the fact that endohedral atoms are of both basic and applied interest. On the basic side, they present an exceptionally clean and stable "laboratory" to study the effects of confinement upon the properties of the encapsulated atom; to understand how the confinement alters the static and dynamic properties of the atom. In addition, many possible applications of endohedral atoms include quantum computing [\[15\]](#page-5-0), drug delivery [\[16\]](#page-5-0), photovoltaics [\[17\]](#page-5-0), and hydrogen storage [\[18\]](#page-5-0), among others.

Of the many extant photoionization studies, the overwhelming majority are theoretical, owing to the difficulties in the fabrication of endohedral systems in large enough quantities to investigate experimentally, although some experimental photoionization work has been reported [\[10\]](#page-5-0). Furthermore, almost all of the theoretical investigations have treated the potential of the confining shell in a spherical jellium model of one sort or another [\[18,10–13,19\]](#page-5-0). The few studies considering the full molecular potential have been at the density functional theory (DFT) (static-mean-field) [\[20\]](#page-5-0) or static-exchange (Hartree-Fock) level [\[9\]](#page-5-0). Thus, there are no reports of calculations which include both molecular structure and correlation; such calculations are, however, important to provide some notion of the qualitative and quantitative accuracy of the previous calculations. This is needed owing to the relative scarcity of experimental results.

To ameliorate the situation, we have embarked upon a program to do just that, to perform calculations of the photoionization of confined atoms including both the molecular structure of the target and correlation. The correlation is included through the use of time-dependent density functional theory (TDDFT) [\[21,22\]](#page-5-0) which is an outgrowth of earlier work at just the DFT [\[20\]](#page-5-0) level. As a first step we have chosen Ar $@C_{60}$  which has been the subject of a number of earlier studies [\[1,2,4,9,11,23–28\]](#page-5-0). There are two questions which we are particularly interested in: how confinement resonances are affected by the inclusion and interaction of both the molecular structure and correlation; and how the inclusion of the C 1 $s$  channels of  $C_{60}$ , and the interchannel coupling with these channels, affects the inner-shell atomic photoionization channels. We consider the Ar 2*p* and 3*s* cross sections to explore these questions, but the results should be generally applicable to any endohedral fullerene. Ar 2*p* and 3*s* were chosen because they are inner shells (no hybridization and negligible interaction with the confining shell) [\[27,29\]](#page-5-0) so that we could focus on the effect(s) of the confining environment and correlation on the final-state continuum wave function which results in the changes to the cross section. These are important questions. Confinement resonances in any confined system give significant information about the physical dimensions and shape of the confining shell [\[2\]](#page-5-0). And an understanding of the effect(s) of the C 1*s* ionization channels of the  $C_{60}$  fullerene on the inner-shell ionization of the caged atom should be widely applicable to any atom trapped in any fullerene cage.

The following section gives a brief presentation of the theoretical methodology. In Sec. [III,](#page-1-0) the results of the calculations are presented and discussed. The final section includes a summary, conclusions, and some prospectus as to where we go from here.

# **II. BRIEF DESCRIPTION OF THE THEORETICAL METHODOLOGY**

The computational approach used in our research has been described in detail in previous work [\[30–32\]](#page-5-0) and has been shown to produce results in excellent agreement with the experiment; here we limit the discussion to a summary of only

<span id="page-1-0"></span>the main steps. The method is based on accurate solution of the scattering problem in a density functional theory framework and utilizes a discretization of both bound and continuum wave functions in a multicenter basis of *B*-spline functions, *Bi*, times real spherical harmonics  $Y_{lm}(\theta, \varphi)$ :

$$
\chi_{ilm} = 1/r B_i(r) Y_{lm}(\theta, \varphi),
$$

which are further fully adapted to the molecular symmetry. The bound states are calculated by a conventional generalized diagonalization of the Kohn-Sham Hamiltonian matrix while the continuum states are obtained by a least-squares approach. The self-consistent-field initial electronic density of the ground state is first obtained by a standard linear combination of atomic orbitals (LCAO) approach utilizing the ADF program [\[33,34\]](#page-5-0). The exchange-correlation potential LB94 [\[35\]](#page-5-0) is employed, with a double-zeta with polarization (DZP) basis set for Ar and C taken from the optimized database included in the ADF package. From the ground-state density, a fixed Kohn-Sham Hamiltonian is obtained and it is further diagonalized in the *B*-spline basis for both the bound and continuum states (static-exchange DFT approach). The *B*-spline basis set comprises a long-range expansion around a common origin (one-center expansion), with large angular momenta, to describe the continuum wave functions up to the asymptotic Coulomb region, where they are fitted to analytical solutions, and a set of additional functions centered on the various nuclei, in the spirit of the LCAO approach, which take care of the Coulomb singularities at the nuclei and ensure rapid convergence of the expansion. In the present calculation an interval up to  $R_{\text{max}} = 25$  atomic units (a.u.) has been employed for the one-center expansion, with a step size of 0.2 a.u., and maximum angular momentum up to  $L_{\text{max}} = 25$ . For the expansions around the carbon atoms  $L_{\text{max}} = 2$  was employed. The radial expansions around the carbon atoms were  $R_{\text{max}} =$ 1*.*32 au. These choices ensured complete convergence of the calculated cross sections reported.

Cross sections and asymmetry parameters can finally be obtained by the calculation of transition dipole moments, through standard angular momentum analysis [\[36\]](#page-5-0). For the TDDFT calculations, we employed the noniterative algorithm previously developed [\[32\]](#page-5-0). The basic dynamical variable becomes the linear response potential  $\dot{V}^{SCF} = V^{ext} + \delta V$ . This, as well as the wave functions, is expanded in the full multicenter *B*-spline basis, and all equations are recast as matrix equations: both

$$
\delta \rho = \chi V^{\text{SCF}}
$$
 and 
$$
\delta V = K \delta \rho
$$

are linear, and define  $\chi$  and  $K$  matrices, the linear susceptibility *χ* and the kernel *K* that give the potential generated by a charge density change *δρ*. Combining, we arrive at the equation for  $V^{\text{SCF}}$ .

$$
(K\chi - 1)V^{\text{SCF}} = V^{\text{ext}}.
$$

The computationally most demanding part is the evaluation of the *χ* matrix, which is energy dependent and has to be recomputed at each selected energy. It is obtained by firstorder time-dependent perturbation theory. Finally the TDDFT transition matrix elements are obtained by substituting *V* SCF in place of the dipole operator  $V^{\text{ext}}$  in the transition dipole

matrix elements,

$$
\mathbf{d}_{Elm,\gamma,i}^{(-)} = \langle \varphi_{Elm}^{(-)} | V_{\gamma}^{\text{SCF}} | \varphi_i \rangle,
$$

where  $\varphi_i$  and  $\varphi_{Elm}^{(-)}$  are the initial and final (incoming-wave boundary conditions) wave functions of the system, and *γ* represents the photon polarization.

## **III. RESULTS AND DISCUSSION**

The results of our calculations for the 2*p* subshell of free Ar and confined  $Ar@C_{60}$  are given in Fig. 1; in the confined case, in molecular notation, Ar 2*p* becomes the 3*T* 1*u* state of the combined  $Ar@C_{60}$  system. For the free case, the DFT cross section exhibits a small rise from threshold resulting from the shape resonance in the *d*-wave continuum of the  $2p \rightarrow 3d$  transition. Including correlation with the TDDFT result, the shape resonances are still there, although somewhat altered in shape from DFT, and a significant difference in magnitude is seen at threshold, a difference which decreases with increasing energy, eventually disappearing completely. This difference arises from both initial-state correlation (configuration interaction with double excitations), and final-state correlation (in the form of interchannel coupling among photoionization channels) [\[37\]](#page-5-0). It is of importance to point out that the present TDDFT results are in excellent agreement with those of experiment and previous sophisticated calculations (not shown) [\[38\]](#page-5-0). This agreement is of importance because it shows that the present TDDFT calculation includes all of the important atomic effects for this particular case. It is also to be noted that the cross sections, except for the near-threshold shape resonance, are quite smooth as a function of energy.

Looking now at the confined case, a rather different picture emerges. The atomic cross section is very substantially modulated; these modulations are known as confinement resonances [\[39\]](#page-5-0) and have been seen experimentally for the case of 4*d* photoionization in the  $Xe@C_{60}$  endohedral fullerene [\[10\]](#page-5-0). Basically these are caused by interference between the photoelectron wave emerging directly and waves that are scattered from the inner and outer walls of the confining



FIG. 1. (Color online) Photoionization cross section for the free and confined Ar 2*p* subshells each calculated using DFT and TDDFT methods without coupling to the C 1s channels of the  $C_{60}$  shell.

 $C_{60}$  shell [\[12,39\]](#page-5-0). It is important to note that these confinement resonances are almost exactly the same in DFT and TDDFT calculations, thereby confirming that confinement geometry is the dominant determinant of this phenomenon; correlation, although it is seen to have some effect, does not play a large role in their behavior, i.e., the confinement resonances are determined almost solely by confinement geometry.

Note also that the three lowest resonances closest to threshold are quite sharp, but the higher-energy resonances are much smoother. The sharpness of the resonances near threshold is evidently due to the more granular nature of the confinement potential when the molecular structure is considered; the slower photoelectrons have more time to "experience" the different confinement potential. This is rather different from the predictions of a spherical jellium model of the confining potential, which predicts all confinement resonances to be rather smooth [\[11\]](#page-5-0). However, it does agree with a previous study at the Hartree-Fock level which also predicted sharp near-threshold confinement resonances [\[9\]](#page-5-0). The confinement resonances are also stronger than those predicted by a spherical jellium model of the confining potential  $[11]$ ; even 150 eV above threshold, the amplitude is about 10% of the cross section. The amplitude of the confinement resonances diminishes much more rapidly with energy in a jellium model. This difference too is evidently the effect of using the full molecular, as opposed to jellium, confining potential in the calculations. In general, it is expected that deficiencies of the jellium model stems from the excessive delocalization of the valence electron charge (typically 240 electrons) compared with a full molecular calculation. In the latter, electrons are more tightly bound close to the carbon atoms, and to the C-C bonds. So scattering of the photoelectrons from the hard atomic cores is significantly enhanced; and, on the other hand, response effects are diminished, due to the lower mobility of the electron cloud.

The sharpness of the lower-energy confinement resonances would probably not be seen in a room-temperature experiment owing to the vibrations which would move some of the confined atoms off center, thereby smearing out the confinement resonances. Such a mechanism was suggested earlier as an explanation as to why confinement resonances had not been seen; they have since been observed [\[40\]](#page-5-0). This suggests that, to compare with experiment, a broadening factor would have to be applied to account for this vibrational excitation. This also suggests that experimental work on cold target systems would change the observed shape of the low-energy near-threshold confinement resonances.

Now, in the results presented in Fig. [1,](#page-1-0) the Ar 2*p* photoionization channels for the confined system were not coupled to the C 1 $s$  photoionization channels arising for the C<sub>60</sub> cage, which were kept frozen. This could be of some importance since the C 1*s* threshold is at about 300 eV, so that above 300 eV the Ar 2*p* and C 1*s* photoionization channels are both open; furthermore, the threshold C<sub>60</sub> 1s photoionization cross section is about 100 Mb, which is estimated by normalizing experimental results at higher energy as 60 times the cross section of the free carbon atom  $[41]$ , and this is more than an order of magnitude larger than the Ar 2*p* cross section at that energy, as seen in Fig. [1.](#page-1-0) Thus, with a small mixing of the  $C_{60}$  1s cross section, the Ar 2p cross section could be considerably altered. To explore this possibility, the



FIG. 2. (Color online) Calculated photoionization cross section for confined Ar 2*p* at the TDDFT level with and without coupling with the C 1s channels of the  $C_{60}$  cage. The fact that only a single curve is seen shows that the coupling has negligible effect.

calculation for the confined system has been redone using TDDFT to include interchannel coupling with C 1*s*; the Ar 2*p* photoionization cross sections are shown in Fig. 2, with and without the interchannel coupling between Ar 2*p* and C 1*s*. The outstanding feature of this comparison is that the curves are almost exactly the same; thus, despite the C 1*s* cross section being about a factor of 10 larger than the Ar 2*p* cross section around 300 eV, the interchannel coupling is negligible. The same can be said about the absence of autoionization resonances due to the huge C  $1s \rightarrow C 2p\pi^*$  below-threshold excitation. In fact similar coupling of core excitations on one site with another site's ionization was expected and first reported [so called MultiAtom Resonant PhotoEmission (MARPE) effect] in solid state photoemission [\[42\]](#page-6-0), but it proved elusive afterwards, and has been barely detected in isolated molecules [\[43\]](#page-6-0).

To understand this result, it is important to note that two conditions must be fulfilled for interchannel coupling to alter a cross section of a particular photoionization channel. First, the cross section for that channel must be degenerate with a channel with a significantly larger cross section. And, second, there must be a non-negligible interchannel coupling matrix element connecting the two channels [\[37\]](#page-5-0). The first condition is clearly satisfied for Ar  $2p$  with  $C_{60}$  1*s*, as discussed above. Thus, given the results depicted in Fig. 2, it must be the case that the interchannel coupling matrix elements are extremely small. The direct part of the matrix element for this case, the Coulomb interaction between the  $2p^5 \varepsilon l$ 1*s*<sup>60</sup> final state and the  $2p^6 1s^{59} \varepsilon' l'$  final state (2p is Ar 2p and 1s is C<sub>60</sub> 1s), can be written as  $\langle 2p\varepsilon l | \frac{1}{r_{12}} | \varepsilon' l' 1s \rangle$ , and is included in the calculation. This coupling matrix element is, however, negligible because each of the discrete orbitals in the direct matrix element overlaps with the continuum orbital of the other channel. Even at the C 1*s* threshold, where the exchange matrix element should maximize, while the  $\varepsilon'$ *l'* continuum orbital (arising from C 1*s* ionization) is slowly varying, the *εl* orbital, arising from Ar 2*p* ionization, is moderately energetic and oscillates rather rapidly. Thus, while the 2*p* orbital overlaps somewhat

<span id="page-3-0"></span>with the threshold  $\varepsilon'$ *l'* function, the overlap of the 1*s* with the *εl* orbital is negligible, owing to the oscillation in the latter. And this matrix element clearly decreases rapidly with increasing energy because the continuum wave functions oscillate more rapidly in space as the energy increases so that the overlap with the discrete wave functions decreases. The exchange part of the interchannel coupling matrix element,  $\langle 2p \varepsilon l | \frac{1}{r_{12}} | 1 s \varepsilon' l' \rangle$ , is treated only approximately in the local density approximation (LDA), substituting the derivative of the exchange-correlation potential  $V_{\text{XC}}$  in place of the Coulomb operator  $1/r_{12}$ , so that the exchange matrix element is calculated as  $\langle 2p \varepsilon l | V_{XC}/r | 1s \varepsilon' l' \rangle$ . This matrix element too is quite small in the present case since the overlap of the discrete Ar 2*p* and C 1*s* wave functions is so small because these two wave functions are quite compact (∼0*.*2 atomic units in radius) and centered about 6 a.u. apart. Since this exchange matrix element is negligible, in the present case, it is evident that the approximation here causes no inaccuracies. Furthermore, it is indeed a general feature that core ionizations arising from different inequivalent sites of the same atomic species, e.g., C 1*s* ionization from inequivalent molecular sites, are very weakly coupled, and that is generally borne out in TDDFT calculations. On the other hand, the inadequacy of the TDDFT exchange matrix elements in describing charge transfer excitation between spatially separated sites, due to the exponential compared to Coulombic decay of the exchange matrix element, has been documented in the literature [\[44,45\]](#page-6-0)

This understanding suggests that it might be possible for interchannel coupling to be important if the confined atom or molecule has an inner subshell whose ionization energy is very close to the C 1*s* ionization energy, in which case, interchannel coupling in the threshold region could be significant, owing to the exchange part of the interchannel coupling matrix element. A molecule containing a carbon atom is a good candidate for this since the 1*s* electrons of the carbon in the confined molecule will have a threshold energy very close to that of the 1 $s$  electrons of the carbon atoms of the  $C_{60}$  shell.

Next consider the Ar 3*s* subshell, which has the same symmetry as the  $2Ag$  level of free  $C_{60}$  so they are considered together. The same problem hase been recently considered by Jose and Lucchese [\[9\]](#page-5-0), at the static-exchange level. In Fig. 3, the DFT and TDDFT results for the photoionization cross sections of free Ar 3*s* and the 2*Ag* state of free  $C_{60}$ are shown. The Ar 3*s* DFT cross section is quite smooth and similar to the Hartree-Fock (HF) one [\[9\]](#page-5-0), but the addition of interchannel coupling through the TDDFT calculation induces a Cooper minimum and a completely different spectral shape in the threshold region, in excellent agreement with experiment and previous theoretical work [\[46\]](#page-6-0). Note that, above about 70 eV, where interchannel coupling with the Ar 3*p* ionization channels becomes unimportant, the DFT and the TDDFT results are essentially the same. For the 2*Ag* state of free  $C_{60}$ , the DFT result is seen to exhibit confinement resonances and is quite similar to the Hartree-Fock result [\[9\]](#page-5-0). The introduction of correlation through the TDDFT calculation induces some changes in the 2*Ag* cross section over most of the energy range; the confinement resonance at about 40 eV is increased by about 50%. The region very close to threshold is changed markedly, however, owing to the autoionizing resonances.



FIG. 3. (Color online) Calculated photoionization cross sections for the free Ar 3s and the free  $C_{60}$  2Ag states at both DFT and TDDFT levels of approximation.

Going now to  $Ar@C_{60}$ , the situation changes dramatically. The cross sections for the  $4Ag$  state in Ar $@C_{60}$ , essentially the Ar 3*s* (which is not hybridized), and the 5*Ag* is what the 2*Ag* of free  $C_{60}$  becomes in the combined system, as shown in Fig. 4. To begin with, note that the reverse ordering is obtained at the HF level [\[9\]](#page-5-0), with the two levels almost degenerate. It is hard to speculate on the exact ordering in the absence of more accurate calculations, and it is, in any case, not very relevant for the present discussion, except to note that both calculations agree in finding no hybridization between Ar  $3s$  and  $C_{60}$  orbitals, despite the closeness in energy of the two levels. Most important, however, is that the 4*Ag* (3*s*) cross section is virtually unrecognizable compared to the free case. The DFT result shows a cross section which displays a number of confinement resonances owing to the confinement; this result is qualitative



FIG. 4. (Color online) Calculated photoionization cross sections of the 4*Ag* and 5*Ag* subshells of Ar@C<sub>60</sub> at DFT and TDDFT levels of approximation. The 4*Ag* state is essentially Ar 3*s* (which is not hybridized) and the  $5Ag$  is almost exactly the  $2Ag$  state of free  $C_{60}$ .

similar to the earlier HF result [\[9\]](#page-5-0). This occurs even though the initial state, the Ar 3*s*, remains virtually unaltered by the confinement since it is not hybridized; the final continuum state is significantly altered by the molecular potential of the  $C_{60}$  cage. The confinement resonances are part and parcel of the effect of the  $C_{60}$  on the continuum wave function.

However, looking at the TDDFT cross section, it is clear that interchannel coupling engenders significant changes in the 4*Ag* (3*s*) cross section, particular at the lower energies; the prominent confinement resonance at about 45 eV is reduced in magnitude by a factor of 5 or so by the coupling. This could also be tied to the Cooper minimum in free Ar, caused by interchannel coupling and properly included at the TDDFT level. At the higher energies, however, the effects of the coupling die out and the DFT and the TDDFT results are virtually identical. To understand this phenomenology, note that the interchannel coupling matrix element where the Ar 3*s* wave function overlaps with a discrete wave function of the  $C_{60}$  cage is negligible because the overlap is negligible. On the other hand, since the binding energy of Ar 3*s* is quite close to the binding energy of a number of  $C_{60}$  shell orbitals, the interchannel coupling matrix elements where each discrete orbital overlaps with a continuum orbital are not small because the continuum orbitals, both being near threshold, oscillate only very slowly with *r*. With increasing energy, however, they oscillate more rapidly, thereby "killing" the interchannel coupling matrix element. This explains why there is significant interchannel coupling near threshold, but essentially none above about 70 eV, as seen in Fig. [4.](#page-3-0) These results are qualitatively similar to what was found in calculations using a jellium model to approximate the molecular potential of the fullerene cage [\[11,23\]](#page-5-0).

The  $5Ag$  cross section, essentially a pure  $C_{60}$  state, behaves somewhat differently. The DFT cross section is quite similar to the cross section for the  $2Ag$  of free C<sub>60</sub>, although there are some differences which reflect the fact that the molecular field changes somewhat owing to the introduction of the endohedral Ar atom. At the TDDFT level, significant alteration is evident at the lower energies, particularly around 45 eV, for the same reasons as discussed above in connection with the 4*Ag* (3*s*) cross section. At the higher energies, however, there remain effects of interchannel coupling, like the roughly 30% increase in the confinement resonance at 70 eV. This is because the 5*Ag* orbital, being a  $C_{60}$  orbital, overlaps reasonably well with other  $C_{60}$  orbitals so that the interchannel coupling matrix elements, where the discrete orbital overlap is not negligible as it was in the case of the 4*Ag* (3*s*), overlap. Thus, while interchannel coupling between the 4*Ag* (3*s*) photoionization channel and the  $C_{60}$  channels is essentially gone at 70 eV, this is not the case for  $5Ag$ , since it is a  $C_{60}$  orbital itself.

#### **IV. SUMMARY AND CONCLUSIONS**

A calculation of the photoionization of an atom confined in a fullerene cage taking both correlation and molecular structure has been performed. The Ar $@C_{60}$  system was investigated and the near-threshold confinement resonances were found to be quite sharp, unlike the findings of calculations using spherical jellium models of the confining potential [\[1–8\]](#page-5-0). In addition, as a result of the inclusion of molecular structure, the confinement resonances were found to extend to much higher energy as compared to jellium calculations  $[1-8]$ . Furthermore, it was found that correlation has only very minor effects on the confinement resonances, which means that they are determined largely by the detailed structure of the confining system, a very general result. This also means that experimental observation of confinement resonances provides information on confinement geometry, including where the trapped atom or molecule resides in the confinement cavity.

It was also seen that interchannel coupling of the small Ar 2*p* photoionization cross section with the (degenerate) much larger  $C_{60}$  1s channels was not found, and the finding explained. As a result of the understanding of why interchannel coupling was negligible in this case, it was also suggested under what conditions interchannel coupling between the C60 1*s* and the photoionization channels of a confined atom or molecule might be found. However, it is evident that this is a general result; interchannel coupling in the photoionization of essentially any inner subshell of any atom confined in a fullerene will be negligible. This understanding is of importance in that theoretical studies of inner-shell ionization have been avoided in the past because the conventional wisdom was that, in the energy neighborhood of the very large C 1*s* cross section of the fullerene shell, the atomic cross section would be greatly affected by interchannel coupling. Thus, the present results open the door to a number of additional studies that were hitherto avoided. It should also be mentioned that inner-shell ionization investigations of confined atoms are particularly useful in that the initial states, being so compact, are essentially entirely atomic, so that all the changes from the free-atom cross sections result from confinement effects on the final continuum state wave functions, i.e., inner-shell ionization studies of confined atoms amount to spectroscopy of the continuum wave functions.

In addition, our study of the photoionization of Ar 3*s*, and the related  $C_{60}$  2*Ag*, in the Ar $@C_{60}$  system has revealed both significant effects of the molecular potential along with significant interchannel coupling. Even though Ar 3*s* is still an inner shell and not hybridized, still interchannel coupling has a large effect on the cross section in the threshold region, an effect that gradually disappears with increasing energy; this behavior was explained in terms of the details of the interchannel coupling matrix element. Investigation of the 5*Ag* cross section (which is essentially the  $C_{60}$  2Ag in the combined system) revealed a somewhat different phenomenology. This cross section is perturbed only a small amount from the free  $C_{60}$  case owing to the introduction of the Ar atom. In the threshold region, here too extensive alteration of the cross section was found, and these alterations, while they decreased with increasing energy, decreased much less rapidly than in the 3*s* case for reasons explained.

As far as future work is concerned, we have begun investigating the photoionization of all of the subshells of the encapsulated Ar atom with an eye to understanding where the simpler jellium results are correct and where they need correction. Based upon that investigation, other systems will be scrutinized. In addition, the influence of the vibrational motion on the damping of the resonances will be investigated.

Finally, by producing cross sections of significant accuracy, it is our hope that these and future results will <span id="page-5-0"></span>stimulate laboratory investigations of the photoionization of endohedral fullerenes. And with the increasing capabilities of fabricating significant quantities of samples, we urge that photoelectron spectroscopic studies be performed to separate the cross section by subshell and test theory stringently.

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## **ACKNOWLEDGMENTS**

P.D. gratefully acknowledges support from the European COST Action No. CM1204 XLIC. S.T.M. is grateful for support from the U.S. Department of Energy, Office of Chemical Sciences under Grant No. DE-FG02-03ER15428.

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