Photoassociation of a cold-atom-molecule pair: Long-range quadrupole-quadrupole interactions

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I. INTRODUCTION

Since it was proposed by Thorsheim et al. [1] in 1987, and first observed for sodium [2] and for rubidium [3] atoms in 1993, the photoassociation (PA) of pairs of ultracold atoms has had a tremendous impact on research in atomic, molecular, and optical physics at low temperatures. There are several recent review articles devoted to the various aspects of PA [4–7]; therefore, we briefly recall in what follows some of the main features of the PA process, which gave rise to a new high-resolution spectroscopic technique, that is, the PA spectroscopy. Due to their extremely low relative kinetic energy, atoms from an ultracold gas can be associated via a quasiresonant free-bound dipolar transition to form an electronically excited molecule, which is often created in a highly excited rovibrational level. As the PA process is mainly controlled by the long-range electrostatic interactions between cold atoms, it has been used as a high-resolution spectroscopy technique for highly excited rovibrational levels. Such levels observed using PA correspond to vibrational motion of a molecule with much larger extension than the usual chemical bond [8–12]. Such molecules with a very large amplitude of vibration were predicted 15 years before the mentioned experiments [13,14]. The spectroscopy of the highly excited rovibrational levels of photoassociated dimers made it possible, in particular, to determine the most accurate values of the radiative lifetime of the first excited state of alkali-metal atoms (see, for instance, [15]). Another example of PA application is the formation of stable ultracold molecules, reported initially for Cs₂ [16] and later for many other homonuclear and heteronuclear alkali-metal diatomic molecules [17–24].

With the improvement of the experimental techniques at ultracold temperatures, the study of the quantum dynamics of few-body systems in the ultracold regime has become possible, as illustrated by the recent observations of cold collisions between atoms and molecules [25–29]. Such phenomena attract at present a lot of interest as they represent the first manifestation of a novel ultracold chemistry, which is controlled by the quantum nature of the colliding partners [30,31]. In particular, at certain conditions, the ultracold few-body dynamics exhibits universal (i.e., species-independent) properties for long-range bound states and resonances (see, for example, Refs. [32,33] and references therein), nowadays referred to as the Efimov physics [34,35]. The Efimov states have recently been observed experimentally [36–38].

All these developments concern atoms and molecules in their electronic ground state. The purpose of the present study, as the first of a series of articles, is to investigate the next step toward ultracold chemistry: the association of ultracold atoms and molecules with a laser field to create weakly bound trimers or tetramers in an excited electronic state. Just as for pairs of atoms, the PA probability is determined by the long-range interactions between the colliding partners. Here, we consider the long-range interaction between a ¹Σ₂⁺ molecule in a given rovibrational level (νd,j) with an atom in a P electronic level without fine structure. This situation will be illustrated with the interaction between a ground-state Cs₂ molecule and an excited Cs(6²P) atom, but can be easily generalized to other species. The leading term of this interaction at large interparticle distances $R$ is a quadrupole-quadrupole term varying as $R^{-5}$. The present work can also be viewed as a step beyond several related studies. The quadrupole-quadrupole interaction between two excited ²P atoms has been calculated for alkali-metal atom pairs [39] and for the LiB molecule [40]. In Refs. [41,42], the van der Waals interaction (varying as $R^{-6}$) between alkali-metal dimers in the $(ν_d = 0, j = 0)$ level of their lowest triplet state and a ground-state alkali-metal atom has been determined, while in Refs. [43,44] the interaction between a ²Π molecule and a ³P atom at fixed geometries is obtained as a sum of a dipole-quadrupole term (in $R^{-4}$) and a quadrupole-quadrupole term (in $R^{-5}$).

In Sec. II, we briefly review the main ingredients of the perturbative approach based on the multipolar expansion of the long-range interaction between the two fragments. Section III is devoted to the calculation of $C_j$ coefficients of the long-range behavior of molecular potentials. We consider the general case of an arbitrary rotational state $J$ of the dimer as well as give an analytical solution for the particular case of $J = 1$. Atomic units (a.u.) for distances (1 a.u. = 0.052 917 7 nm) and for energies (1 a.u. = 219 474.631 37 cm⁻¹) are used throughout the article, unless otherwise stated.

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II. INTERACTION POTENTIAL AND PERTURBATION THEORY

We start the description of the present theory from the general case, as for instance in Refs. [43,45,46]. We consider two charge distributions, A and B, far from each other such that they do not overlap with each other. A criterion for such a condition is given by the so-called Le Roy radius [47,48] defined as \( R_{LR} = 2\sqrt{\langle r_A^2 \rangle + \langle r_B^2 \rangle} \), where \( \langle r_A^2 \rangle \) and \( \langle r_B^2 \rangle \) are the averaged squared distances of the outermost electron from the origin of each charge distribution A and B, respectively. The electrostatic potential energy of interaction between A and B can be written as an expansion over products of multipole moments of A and B located at a distance \( R \) from each other:

\[
\hat{V}_{AB}(R) = \sum_{L_A,L_B=0}^{\infty} \sum_{M=-L}^{L} \frac{1}{R^{1+L_A+L_B}} \langle L_{x}\rangle \hat{Q}_{L_{x}}^{M}(\hat{r}_A) \hat{Q}_{L_{x}}^{-M}(\hat{r}_B),
\]

where \( L_\text{min} = \text{min}(L_A, L_B) \). The operator \( \hat{Q}_{L_{x}}^{M}(\hat{r}_X) \) is associated with the \( 2L_{x} \)-pole of the charge distribution \( X \) (\( X = A \) or \( B \)), expressed in the body-fixed coordinate system (CS) with the origin at the center of mass of \( X \),

\[
\hat{Q}_{L_{x}}^{M}(\hat{r}_X) = \sqrt{\frac{4\pi}{2L+1}} \sum_{i \in X} q_i \hat{f}_{i}^{L_{x}} Y_{L_{x}}^{M}(\hat{\theta}_i, \hat{\phi}_i),
\]

where \( q_i \) is the value of each charge \( i \) composing the distribution \( X \). The two coordinate systems (centered at A and B) are assumed to have parallel axes with the Z axis that goes from the center of mass of \( A \) toward \( B \) (see Fig. 1). This choice of Z implies in Eqs. (1) and (2) that \( M_A = -M_B = M \), where \( M_A \) and \( M_B \) are the projections of \( L_A \) and \( L_B \), so that the factor \( \hat{f}_{L_{x}L_{x}M} \) is equal to

\[
\hat{f}_{L_{x}L_{x}M} = \frac{(-1)^{L_{x}}(L_A + L_B)!}{\sqrt{(L_A + M)!(L_A - M)!(L_B + M)!(L_B - M)!}}.
\]

The energy of interaction between the two charge distributions is calculated using perturbation theory. To the lowest (zeroth) order of perturbation theory, the two systems are independent and the total energy is the sum of the individual energies,

\[
E_{0} = E_{0A} + E_{0B},
\]

and the total wave function is the product of individual wave functions,

\[
|\Psi_{0}\rangle = |\Psi_{A0}\rangle |\Psi_{B0}\rangle.
\]

In Eqs. (4) and (5) and in what follows, the superscript labels the perturbation order and the subscript labels the unperturbed states.

In the present study, the system A is the alkali-metal dimer and B is the alkali-metal atom. We consider the dimer in a vibrational level \( v_d \) of its ground electronic state \( |X \Sigma_{+}^{e}\rangle \), and in an arbitrary rotational state \( |j,m_{j}\rangle \). In order to investigate a realistic approach for atom-molecule PA, we consider the atom B with a single outer electron being excited to the \( p \) state \( |n,\ell = 1,\lambda\rangle \). However, we ignore in the following the fine structure of the excited atom for clarity, as discussed later in the text. The projections \( m_{j} \) and \( \lambda \) are defined with respect to the Z axis. The energy origin corresponds to an infinite separation between the atom and the dimer. Thus, the unperturbed energy reduces to

\[
E_{0} = B_{0j} (j + 1),
\]

where \( B_{0j} \) is the rotational constant of the dimer in its vibrational level \( v_d \). The atomic state of B is expressed in the \( LS \) coupling case, because the operators in the interaction potential of Eq. (1) act only on the coordinate part of wave functions. The first-order correction \( E_{1} \) to the energy is due to the permanent multipoles of A and B. In our case, both distributions exhibit a permanent quadrupole moment in their body-fixed frame so that the most important contribution comes from the quadrupole-quadrupole interaction with an asymptotic coefficient \( C_5 \),

\[
E_{1} = \frac{C_5}{R^6}.
\]

III. CALCULATION OF THE \( C_5 \) COEFFICIENT

The \( C_5 \) coefficient is calculated for arbitrary values of \( m_j \) and \( \lambda \), using the degenerate perturbation theory. We define two body-fixed CSs (Fig. 1). The first CS (which we call the dimer CS, or D-CS) with axes \( X_A, Y_A, \) and \( Z_A \) has as the origin the center of mass of the dimer. The \( Z_A \) axis is the dimer axis and the \( Y_A \) axis is orthogonal to the plane of the trimer. The second CS (trimer CS, or T-CS) with axes \( XYZ \) is such that the \( X, Z \) axes are also (as \( X_A \) and \( Z_A \))
in the plane of the trimer, while Z is oriented from the center of the dimer toward the atom B; the \( Y_A \) and \( Y \) axes are identical. The T-CS is deduced from the D-CS by a rotation with an angle \( \delta \) around the \( Y \) axis.

The perturbation Hamiltonian \( V_{AB}^{qq}(R) \) for the quadrupole-quadrupole interaction is given by setting \( L_A = L_B = 2 \) in Eq. (1):

\[
\hat{V}_{AB}^{qq}(R) = \frac{24}{R^5} \sum_{M=-2}^{2} \hat{Q}_2^M(\hat{r}_A)\hat{Q}_2^{-M}(\hat{r}_B) (2 + M)! (2 - M)!. \tag{8}
\]

The Hamiltonian has the form of a sum of tensor products, composed of operators acting in subspaces of the unperturbed eigenstates of A and B. Usually, the \( 2^L \)-pole tensor components \( \hat{q}_L^M \) of a charge distribution are defined in its proper CS, that is, \( X_A Y_A Z_A \) for the dimer A. Therefore, the \( \hat{Q}_L^M \) tensor components in the T-CS are written as

\[
\hat{Q}_L^M = \sum_{M' = -L}^{L} d_{LM'}(\delta) \hat{q}_{L}^{M'}, \tag{9}
\]

where \( d_{LM'}(\delta) \) are the reduced Wigner matrix elements. In the case of the alkali-metal dimer in the \( 1 \Sigma^+ \) state, the only non-zero component of the quadrupole moment is \( \hat{q}_2^0 \) and Eq. (9) reduces to

\[
\hat{Q}_2^M = d_{20}^M(\delta) \hat{q}_2^0. \tag{10}
\]

The component \( \hat{q}_2^0 \) is just a scalar parameter, which in what follows is referred to as \( q_2^0 \).

In the T-CS, the wave function of the rotational state of the dimer \( |jm_j\rangle \) is written as \( \sqrt{(2j + 1)/2} d_{j0,0}(\delta) \), depending only on the internal angle \( \delta \). The normalization constant is such that the integral over angle \( \delta \) is unity. We obtain the following expression for matrix elements of the operator \( \hat{Q}_2^M \):

\[
\langle jm'_j|\hat{Q}_2^M|jm_j\rangle = \frac{2j + 1}{2} q_2^0 \int_0^{\pi} \sin \delta d\delta d_{jm'_j,0}(\delta) d_{j0,0}(\delta)
\]

\[
= C_{jm_j}^{00} \right\}
\]

where the Clebsch-Gordan coefficients \( C_{jm}^{00} \) appear after integrating the product of three \( d_{jm}^{LM} \) functions [49]. The zeroth-order energy \( E_0 \) depends on \( j \) and is degenerate for all values of \( m_j \). Thus, the perturbation Hamiltonian Eq. (8) has to be evaluated with the degenerate perturbation theory. Indeed, Eq. (11) shows that the quadrupole moment has matrix elements for different values of \( m_j \) because of the \( m'_j = m_j + M \) selection rule. The degeneracy between different \( m_j \) values will be removed, leading to different values of \( C_5 \).

Assuming the alkali-metal atom are in the state labeled \( |n\ell\lambda\rangle \) (the spin is neglected here), we calculate the matrix elements of the atomic quadrupole moment operator for a given \( \ell \) between two different Zeeman sublevels \( \lambda \) and \( \lambda' \) following the same treatment as previously. We obtain

\[
\langle n\ell\lambda'\rangle \hat{Q}_2^M |n\ell\lambda\rangle = -\frac{4\pi}{5} \left( q_2^0 \right)^2 \int_0^{2\pi} \sin \theta d\phi \int_0^{\pi} d\theta Y_2^0 Y_2^{\lambda'} Y_2^{\lambda}. \tag{12}
\]

where the negative sign comes from the electron charge. The mean squared position \( \langle \hat{r}_A^2 \rangle \) of the valence electron is independent of \( \lambda \). Using the properties of spherical harmonics, we rewrite Eq. (12) as

\[
\langle n\ell\lambda'\rangle |\hat{Q}_2^M|n\ell\lambda\rangle = -\frac{C_{j0}^{00}}{200} C_{2M\lambda}^{00} (r_2^0)^2. \tag{13}
\]

The situation is analogous to the molecular case: If \( M \neq 0 \), the operator \( \hat{Q}_2^M \) couples \( \lambda \) to \( \lambda' = \lambda + M \), and the perturbation Hamiltonian lifts the degeneracy with respect to \( \lambda \) also.

Summarizing the preceding results, the perturbation operator of Eq. (8) couples the \( (2j + 1) \) rotational states of the molecule with a given value of \( j \) and the \( (2\ell + 1) \) Zeeman states of the atom with a given value of \( \ell \). The \( C_5 \) coefficients are then given by \( (2j + 1) \times (2\ell + 1) \) eigenvalues of the operator \( \hat{V}_{AB}^{qq} \). Using Eqs. (11) and (13), the matrix elements of \( \hat{V}_{AB}^{qq} \) are written

\[
\langle jm'_j,\ell^\prime\lambda'\rangle |\hat{V}_{AB}^{qq}|jm_j,\ell\lambda\rangle = -\frac{C_{j0}^{00}}{200} C_{2M}^{00} (r_2^0)^2 \frac{C_{j0}^{00}}{R^5} \times \sum_{M=-2}^{2} \frac{C_{j0}^{00}}{2Mj0} C_{2M}^{00}. \tag{14}
\]

From the integration over Euler angles and the properties of Clebsch-Gordan coefficients, the following selection rules for \( \hat{V}_{AB}^{qq} \) are derived: (1) The projection \( m_J \) of the total orbital momentum \( \tilde{J} = j + \ell \) on the laboratory \( z \) axis is conserved. (2) The projection \( m_J = m_j + \ell \) of the total orbital momentum \( \tilde{J} \) on the \( z \) axis of T-CS is conserved. This rule can also be deduced by the combination of Eqs. (11) and (13).

Equation (14) demonstrates the equivalence between the atomic orbital momentum \( \ell \) and the dimer rotation \( \tilde{J} \) in the formalism, which describes long-range interaction between two charge distributions with defined angular momenta irrespective to their internal structure. If one of the two angular momenta is zero, the corresponding quadrupole moment vanishes, and the \( C_5 \) coefficient as well. Therefore, the interaction will be the usual \( C_6/R^6 \) van der Waals potential. If neither of the two angular momenta \( j \) and \( \ell \) is zero, the long-range interaction varies as \( C_5/R^5 \) and, therefore, the potential has a larger density of vibrational states close to the dissociation limit than the lowest electronic state of the system when the atom in its ground \( S \) state. Such a situation is favorable for the PA of atom-molecule pairs into excited trimers, just like for the PA of identical atom pairs (see, for example, experimental work of Ref. [50]).

IV. RESULTS AND DISCUSSION

To illustrate the previous formalism, we first consider analytically the simplest case \( j = \ell = 1 \). The perturbation Hamiltonian \( V_{AB}^{qq} \) reduces to a \( 9 \times 9 \) matrix with elements calculated from tensor products of the atomic and dimer states. For simplicity, we omit the \( Z_A \) and \( Z \) labels in the following, and the quantum states of the atom-molecule pair are denoted by projections \( |m_j,\lambda\rangle \) only. All such states form the basis of the representation. If we sort the states by values of the conserved projection of the angular momentum \( m_J = m_j + \lambda \), we obtain the matrix of \( \hat{V}_{AB}^{qq} \) in a block-diagonal form.
The two blocks defined by $|m_j, \lambda\rangle = | -1, -1\rangle$ and $| +1, +1\rangle$ ($m_j = \pm 2$) reduce to a single element with a negative value of the corresponding coefficient

$$C_5 = -\frac{6q_5^0 (r_{2m=1}^2)}{25}.$$  (15)

It produces an attractive interaction. Two other 2 \times 2 blocks (with $m_j = \pm 1$) are defined by the two subspaces $([-1, 0]; [0, -1])$ and $([0, 1]; [1, 0])$. The corresponding C5 coefficients are $\frac{3q_5^0 (r_{2m=0}^2)}{25}$ (positive value) and zero. Finally, the last 3 \times 3 block comes from the subspace $([-1, 1]; [0, 0]; [1, -1])$ ($m_j = 0$). Two of the corresponding C5 coefficients are zero, and the third one is

$$C_5 = -\frac{36q_5^0 (r_{2m=1}^2)}{25},$$  (16)

with the eigenvector $\frac{1}{\sqrt{2}}((-1, 1) + 2(0, 0) + (1, -1))$. The coefficient in Eq. (16) is negative with the largest magnitude out of all C5 coefficients obtained in the case of $j = \ell = 1$. It corresponds to the most attractive configuration between the atom and the dimer and is expected to be the most favorable for the PA.

The results of the calculation for the case $j = \ell = 1$ are summarized in Table I. The second column of the table gives the eigenvectors $|\Phi_0\rangle$ of the Hamiltonian of Eq. (8) in the $j = 1$ subspace. The third column gives the so-called reduced values of C5 in units of $q_5^0 (r_{2m=1}^2)$, which stresses the general character of our treatment: It can be applied to all alkali-metal trimers, but it can also be compared with the existing results for the PA.

The fourth column in Table I displays estimates for the C5 coefficients for Cs2 + Cs. To the best of our knowledge, there are no available values for the quadrupole moment of Cs2 in the literature. Therefore, we calculated it for the electronic ground state with the Gaussian quantum chemistry package [51] using the second-order Møller-Plesset method with the second definition of the triple-zeta-valence basis with polarization functions (so-called Def2-TZVPP), defined in Ref. [52]. To check the accuracy of such an estimation, we first calculated the quadrupole moment of K2 and compared it to available accurate ab initio calculations [53]. We obtained 12.258 a.u., which differs by a factor 1.28 from the value 15.689 a.u. of Ref. [53]. For cesium, the Def2-TZVPP basis [52] contains also effective core potentials (ECPs) standing for the 46 inner electrons of the core. We obtained for Cs2 the value of 14.51 a.u. that we multiplied by the same factor to estimate the Cs2 quadrupole moment to $q_5^0 = 18.58$ a.u. The mean squared radius of the $6P$ orbital of cesium, which is 62.65 a.u., is calculated using a Dirac-Fock method [54]. It is worth mentioning that the values of C5 shown in the table are of the same order of magnitude as the values for Cs(6P) + Cs(6P) [39].

For $\ell = 1$ and arbitrary $j$, the perturbation Hamiltonian is a $3(2j + 1) \times 3(2j + 1)$ matrix, which can be diagonalized numerically. The eigenvalues obtained numerically for $j = 2$–4 are given in Table II. The C5 coefficients are of the same order of magnitude as for $j = 1$, but on average they become smaller in magnitude as $j$ increases, due to smaller Clebsch-Gordan coefficients.

| Symmetry | $j$ | C5 (a.u.) | Symmetry | $j$ | C5 (a.u.) |
|----------|----|----------|----------|----|----------|
| $\Sigma^+$ | 2 | -913 | $\Delta$ | 2 | -140 |
|           | 2 | 116   |           | 2 | 1136    |
|           | 3 | -796  |           | 3 | 835     |
|           | 4 | 145   |           | 4 | 63      |
| $\Sigma^-$ | 4 | 157    |           | 4 | 721     |
|           | 3 | 465   |           | 4 | 623     |
|           | 4 | 489   | $\Phi$   | 2 | -399    |
| $\Pi$    | 2 | 964   | 3         | 245    |
|           | 2 | -19   |           | 3 | 1175    |
|           | 2 | 584   | 4         | -783   |
|           | 3 | -783  | 4         | -161   |
|           | 3 | 64    | 4         | 835    |
|           | 3 | 532   | $\Gamma$ | 3 | -465    |
|           | 4 | -739  | 4         | -320   |
|           | 4 | 108   | 4         | 1208   |
|           | 4 | 522   | $\Pi$    | 4 | -507    |

The values for $(r_{2m}^2)$ and $q_5^0$ are the same as in Table I.
coefficients. The $C_5$ coefficients are sorted by values of $|m_J|$, which, in analogy to diatomic molecules, are labeled $\Sigma$, $\Pi$, $\Delta$, $\Phi$, $\Gamma$, and $H$ for $|m_J| = 0$ to 5, respectively. For $\Sigma$ states, the reflection symmetry through the $Z$ axis is also considered, giving the usual $+$ (−) superscripts. For states other than $\Sigma$ the sign $+$ (−) is not specified because such states are degenerate (in the present approximation) with respect to the reflection.

We use the same symmetry notations in Figs. 2, 3, and 4, where we display the long-range potential energy curves $C_5/R^5$ for the Cs$_2 + Cs(6^2P)$ system calculated for the first five rotational levels $j$ of Cs$_2$ as a function of the atom-dimer distance $R$. The energies of dissociation are given by Cs$_2 + Cs(6^2P)$ system calculated for the first five rotational levels $j$ of Cs$_2$ as a function of the atom-dimer distance $R$. The energies of dissociation are given by Cs$_2$ rotational energies, $B_0j(j+1)$, $j = 0, \ldots, 4$. The rotational constant for ground vibrational level of Cs$_2$ is $B_0 = 1.17314 \times 10^{-2}$ cm$^{-1}$ [55].

The potential energy curves are shown up to $R = 500$ a.u. Beyond this limit, the distance between Cs$_2$ and Cs becomes comparable to the wavelengths of relevant atomic and molecular transitions, which are in the optical frequency domain. Therefore, in that region, electrodynamics effects, for example, due to retardation, should be taken into account [56].

As already mentioned, the lower limit of the region where the present approximation is applicable can be estimated by the Le Roy radius $R_{LR} = 2(\langle r_0^2(Cs) \rangle + \sqrt{\langle r_0^2(Cs) \rangle^2})$, where $\langle r_0^2 \rangle$ and $\langle r_0^2 \rangle$ are related to the extension of the dimer and atomic electronic clouds, respectively. For the atom, one has $\langle r_0^2 \rangle = 62.65$ a.u., which is given in Table I. As for the dimer, $\langle r_0^2 \rangle$ is calculated from the elements of the quadrupole tensor. All its nondiagonal elements $Q_{\alpha\beta}$ are zero for $1\Sigma^+_g$ states in D-CS. Its diagonal elements $Q_{\alpha\alpha}$ ($\alpha = X_A, Y_A$, or $Z_A$) in D-CS are estimated using the GAUSSIAN package and can be formally written as a sum over all charges:

$$Q_{\alpha\alpha} = \sum_i q_i \langle \alpha_i^2 \rangle. \quad (17)$$
As the two nuclei of Cs$_2$ are along the $Z_A$ axis and it is a $^1\Sigma^+_g$ molecular state, $Q_{X_iX_k}$ is equal to $Q_{Y_iY_k}$ and both are functions of the coordinates of two valence electrons only ($i = 1, 2$):

$$Q_{X_iX_k} = -e^2 \sum_{i=1}^{2} \langle X_i^2 \rangle_i.$$  \hspace{1cm} (18)

Now considering for simplicity that the cores are fixed at the position $Z_A = z r_e / 2$ (the rigid rotor approximation, valid for $v_d = 0$), for $Q_{Z_iZ_k}$, we obtain

$$Q_{Z_iZ_k} = e^2 r_e^2 / 2 - e^2 \sum_{i=1}^{2} \langle Z_i^2 \rangle_i.$$  \hspace{1cm} (19)

Now, setting

$$\langle r_0^2 \rangle = \sum_{i=1}^{2} \sum_{a=(X_iX_k)Z_\alpha} \langle a^2 \rangle,$$  \hspace{1cm} (20)

we reach the final expression for $\langle r_0^2 \rangle$,

$$\langle r_0^2 \rangle = r_e^2 - Q_{Z_iZ_k} - 2Q_{X_iX_k},$$  \hspace{1cm} (21)

where $e = 1$ in atomic units. The $Q_{Q_{Q_{Q_{Q_{Q_{0}_{0}_{0}_{0}}}}}}$ matrix elements are calculated with the same method as $q_2^0$ (using the ratio $1.27$ to the $K_2$ value), which yields $Q_{X_iX_k} = -69$ a.u., $Q_{Z_iZ_k} = -41$ a.u., and $r_e = 8.7$ a.u. Therefore, we obtain $\langle r_0^2 \rangle = 216$ a.u. and the Le Roy radius $R_{LR} = 45$ a.u.

As we can see from Figs. 2, 3, and 4, the Le Roy radius is smaller than the distance at which the curves start to cross each other. This is the second main result of the article. Unlike the case of two atoms, the rotational structure of the dimer is small enough to compete with the quadrupole-quadrupole interaction. The lower limit of $R_m$ for the applicability of the present perturbation approach is thus fixed by the crossing of the potential energy curves. In order to estimate $R_m$, we note that the first crossing occurs between the curves dissociating toward to $j = 0$ limit and from the most attractive curve corresponding to $j = 1$. Putting $2B_0 \equiv C_5 / R_m^2$, yields a general estimate for $R_m$ [see Eq. (16)]:

$$R_m \sim \left( C_5 / B_0 \right)^{1/5} \left( 18q_0^2 / 25B_0 \right)^{1/5},$$  \hspace{1cm} (22)

For cesium, Eq. (22) yields $R \approx 102$ a.u. In the rigid rotor approximation with $B_0 = 1/(2\mu r_e^2)$ and where $\mu$ is the reduced mass of the dimer, Eq. (22) shows that the value of $R_m$ is smaller for lighter atoms. For example, replacing in our treatment Cs by $^6$Li with the atomic parameters $\langle r_2^2 \rangle = 32.5$ a.u. $[57]$, $r_e = 5.05$ a.u., and $q_0^2 = 10.7$ a.u $[53]$, we obtain $R_m = 43$ a.u. This value of $R_m$ is larger than the Le

Roy radius for lithium, for which we obtained 26 a.u. using Eq. (21).

For distances such that $R < R_m$, the long-range potential (1) is still valid, but not the perturbative approach. The nonadiabatic interaction at the curve crossings (for a given symmetry) is expected to be strong. In particular, the interaction between permanent quadrupoles would couple the dimer rotational level $j$ with $j' = j \pm 2$, $j \pm 4$ near the crossings. Higher-order contributions in $1/R$ should also be considered.

The number $N$ of partial waves involved in the atom-dimer collisions depends on the temperature in the actual experiment. In order to give an upper bound for $N$, we consider a potential curve with the most attractive $C_5$, given by Eq. (16) and with the added centrifugal term. It is straightforward to show that the height of the potential barrier $E_N^{\text{max}}$ for a given $N$ is

$$E_N^{\text{max}} = \frac{9}{20} \left( \frac{5}{24} \right)^{2/3} \left( \frac{N(N+1)}{m} \right)^{5/3} \left( q_0^2 / r_c^2 \right)^{-2/3},$$  \hspace{1cm} (23)

where $m$ is the mass of a single atom. Converted to the temperature, $E_N^{\text{max}}$ is approximately $1$ $\mu$K for cesium. If we take typical temperatures 10–100 $\mu$K for which PA experiments are achieved, only a few partial waves ($6 \sim 7$ for the present case) will play a significant role in collisions. This contrasts with the PA of identical atom pairs, interacting with a long-range $R^{-3}$ potential which allow much more partial waves than here.

V. CONCLUSIONS AND PERSPECTIVES

In this article, we used the multipolar expansion to calculate the long-range interaction energy of a diatomic molecule in its electronic ground state and in an arbitrary rovibrational level and an excited atom. We applied our treatment to the case of a ground-state Cs$_2$ molecule and an excited Cs($6^2P$) atom, as a prospect for cold-atom–molecule PA. The dimer and the atom interact through their permanent quadrupole moment. In contrast with previous works, the anisotropic interaction is computed for arbitrary geometries of the atom-molecule pair and depends on their internal quantum numbers. We showed that the interaction lifts the degeneracy of their respective magnetic sublevels. Using the degenerate perturbation theory, we calculated the $C_5$ coefficients characterizing the quadrupole-quadrupole interaction for the five lowest rotational levels of the ground-state dimer. The PA of a ground-state $X^1\Sigma^+_g$ alkali-metal dimer molecule with a ground state $n^2S$ alkali-metal atom is found possible by exciting the dimer-atom system with a laser frequency red detuned from the $n^2S \rightarrow n^2P$ atomic transitions.

We demonstrated that the small-$R$ limit of applicability of our treatment is not due to the overlap of the electronic clouds of the partners, as in the atom-atom case, but to the competition between the rotational energy of the dimer and the long-range quadrupole-quadrupole interaction. This induces crossings between potential energy curves corresponding to different rotational levels. In the region $50 \leq R \leq 100$ a.u.,
the multipolar expansion is still valid but not the perturbation approach. The inclusion of nonadiabatic couplings is required in this region for an appropriate description of the long-range behavior, as well as higher-order effects in $1/R$. This will be discussed in detail in a forthcoming presentation.

It is important to stress that the preceding treatment has been developed in the framework of the $LS$ coupling case in order to keep our description simple. The next step is to account for the fine structure of the excited atom. The main difference with the formulas reported here will be the change of the atomic state from $P \equiv |n, \ell, \lambda \rangle$ to $P_j \equiv |n, \ell, j, \lambda_j \rangle$ (with $j = 1/2$ or $3/2$), where the state $P_j$ is written as the appropriate superposition of atomic states with different $\lambda$ and spin projections through a unitary transformation. We note, however, that for most of the alkali-metal atoms (from Na to Cs) the fine structure is much larger than the magnitude of the long-range atom-dimer interaction. Therefore, the related $C_5$ coefficients will result from linear combinations of the coefficients of Table II and will not modify the main statement of our study concerning the range of validity of our approach. In contrast, the case of a lithium atom will be remarkable as its $\lambda$-dependence will account for the fine structure of the excited atom, which will induce even more complexity in the present conclusions. This work is currently in progress. A similar discussion obviously holds for the hyperfine interaction of our study concerning the range of validity of our approach.

The present formalism can be generalized to PA of dipolar dimers and atoms, like KRb with K or KRb with Rb. If the atom is in the excited state $n^2 P$ and the heteronuclear dimer in an excited rotational state $j > 0$, the long-range dimer-atom interaction is dominated by a dipole-quadrupole term varying as $C_3/R^4$. The long-range interaction between two identical dipolar molecules can also be treated in the same way, as the leading term will be the usual van der Waals $C_6/R^6$ term if both molecules are in their lowest rotational level $j = 0$ or dipole-dipole $C_3/R^3$ term if one of them is rotationally excited. One could thus investigate the PA of two identical heteronuclear KRb ground-state molecules in their lowest vibrational level using a laser field with a frequency red detuned with respect to $j = 0 \rightarrow j = 1$ transition. In this respect, PA of two dimers is very similar to the PA of two identical alkali-metal atoms, except that the laser frequency is much smaller for the two-dimer PA.

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