Reaching the cold regime: $S^{(1)D} + H_2$ and the Role of Long-Range Interactions in Open Shell Reactive Collisions

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Reactive cross-sections for the collision of open shell $S^{(1)D}$ atoms with ortho- and para-hydrogen, in the kinetic energy range 1–120 K, have been calculated using the hyperspherical quantum reactive scattering method developed by Launay et al. [Chem. Phys. Lett. 169, 473 (1990)]. Short-range interactions, described using the ab initio potential energy surface by Ho et al., were complemented with an accurate description of the long-range interactions, where the main electrostatic ($\sim R^{-3}$) and dispersion ($\sim R^{-6}$) contributions were considered. This allows the comparison with recent experimental measurements of rate constants and excitation functions for the title reaction at low temperatures [Bertelot et al., accepted in Phys. Rev. Lett., 2010]. The agreement is fairly good. The behavior in the considered energy range can be understood on the average in terms of a classical Langevin (capture) model, where the centrifugal barriers determine the amount of reactive flux which reaches the barrierless transition state. Additionally, the structure of the van der Waals well provides temporal trapping at short distances thus allowing the system to find its way to the reaction at some classically-forbidden energies. Interestingly, the cross-section for para-hydrogen shows clearly oscillating features associated to the opening of new partial waves and to shape resonances which may be amenable to experimental detection.

I. INTRODUCTION

Until very recently, most experimental and theoretical studies in reaction dynamics had focused on thermal (or higher) collision energies. With the field of cold molecules emerging as an active interdisciplinary area of research, a great interest in understanding cold and ultracold chemical reactions has risen [1, 2]. Electric and magnetic decelerators, helium buffer gas cooling, and magneto- and photo-association (among other techniques) are opening the access to cold and ultracold samples of molecules [3]. Given the success in controlling interactions achieved during the last two decades in the field of ultracold atomic physics [3], the question is whether a similar level of control is possible for more general processes involving molecules. Very few partial waves contribute at low temperatures. However, reactive processes have been shown to occur rapidly being dominated by tunneling and Feshbach resonances, in a regime where the smallest barrier to reaction surely exceeds the collision energy [5–9]. Remarkably, at low kinetic energies the presence of external fields modifies the dynamics in an essential way, appearing as "knobs" to steer the system to the desired outcome [10]. In addition, the theoretical advances in the coherent control of bimolecular processes achieved in the nineties [11, 12] found in the thermal averaging of the colliding partners a fundamental obstacle. Consequently, the forthcoming access to coherent samples of molecules at very low temperatures has renewed the quest for the control of chemical reactions [13].

Given the success in the production of (ultra-)cold alkali metal dimers, attention has mainly focused in such particular systems [14, 19]. In a landmark experiment, reaction rates for collisions involving fermionic $^{40}K^{87}Rb$, at temperatures less than 1 µK, were recently measured [20]. In contrast, advances in the analysis of other type of reactions are still hampered by the lack of versatile methods to produce molecules at low temperatures and the very low densities achieved. Luckily, emerging experimental techniques like the Crossed Stark decelerators [21] promise to provide detailed information on cold bimolecular collisions. More related with our work, CRESU (Reaction Kinetics in Uniform Supersonic Flow) experiments implemented in Rennes [22, 23], and crossed molecular beam experiments performed in Bordeaux [24, 25], are exploring barrierless reactions between neutral species in conditions approaching the cold regime [26]. In particular, rate constants and cross-sections have been recently obtained for the title reaction at temperatures and kinetic energies as low as $\sim 5K$ (1K $\sim 8.3 \times 10^{-3} KJ/mol$) [27]. We will analyze their results below.

Given the availability of experimental results, the title reaction provides us with a good chance to check and extend our predictive power from the explored thermal regime, where ab initio reaction dynamics is routinely applied, to the low energy one where such methodology should fail due to the lack of accuracy. In the limit, the deep ultracold regime, governed by Wigner laws, has been found amenable to parametrization in terms of a very few inputs (the scattering length, for example) which the experiments may provide soon [20]. Paradoxically, the extreme sensitivity of such parameters to small details of the PES, and the action of the surface as a whole, makes extremely difficult to predict them or, conversely, to deconvolute from their measurement the underlying interactions. Collisions in the range of $\sim 1 K$ thus lie in the limits of what can be done using our conventional theoretical tools. Besides, such collisions are more than an obliged milestone in our route from the thermal to the ultracold regime, having interest on their own. They report the combined game of short-range (SR) and long-range (LR) interactions. While the ultracold scenario privileges LR interactions, leading to

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universality in extreme cases\cite{2}, the thermal regime and its higher kinetic energies make (SR) chemical forces prevail. As we will show, both SR and LR interactions play the game at the concerned kinetic energies. Nevertheless, their roles may be independent enough to allow to associate trends in the behavior to different regions of the PES, thus offering some insight in the underlying dynamics.

The $S(1D) + H_2(X 1 \Sigma^+_g) \rightarrow SH + H$ thermal reaction and its isotopic variants were the subject of detailed analysis in the past\cite{28-39}. Belonging to the class of insertion reactions\cite{40}, the system is characterized by the presence of a deep well (96 kcal mol$^{-1}$) on its singlet ground potential energy surface (the $1A'$ electronic state of $H_2S$) and by a small exoergicity (6.90 kcal mol$^{-1}$). Lee and Liu carried out crossed molecular beam experiments which were able to conclude the insertion mechanism\cite{37,39}. Their measurements motivated a number of theoretical studies. Zyubin et al.\cite{28}, working at a multi-reference configuration interaction (MRCI) level with multi-configuration self-consistent field (MCSCF) reference wave functions, computed the five ($1A'$, $2A'$, $3A'$, $1A''$ and $2A''$) potential energy surfaces (PES) which correlate with the $S(1D) + H_2$ asymptote. The collinear barrier of the first $1A'$ excited PES was found to be 10 kcal mol$^{-1}$ high. The other three surfaces being repulsive, the reaction at moderate energies may be assumed to occur on the $1A'$ PES. An improved ground state, based on the same \textit{ab initio} data set was subsequently produced by Ho et al.\cite{29}. This PES is the one we will use and it will be labeled as S1. Working on it, the quantum calculations of Honvault and Launay\cite{32}, precursors to the statistical model of Rackham and Manolopoulos\cite{44}, the latter were shown to account very well for the QM and the quasiclassical statistical model\cite{45}. In Refs. [33] and [36] the models of Rackham and Manolopoulos\cite{44} and the quasi-classical statistical model\cite{45} were recalled and the conclusions will be given in Section VI.

The S($1D$) + $H_2(X 1 \Sigma^+_g)$ system was also revealed in a recent quantum study performed on the isotopic variant $S(D)$ + $H_2(1S)$ by Maiti \textit{et al.}\cite{30} using trajectory surface-hopping methodology. Their calculations conclude that the electronic quenching process, $S(1D) + H_2 \rightarrow S(P_{0,1,2}) + H_2$, plays a major role in the removal of $S(1D)$ at energies $\sim$ 250K. The significance of non-adiabatic effects in the title system was also revealed in a recent quantum study performed on the isotopic variant S($1D$) + HD\cite{46}. We thus expect a significant contribution of the quenching process in the current regime.

In this work we will show the results of new quantum mechanical calculations for the title collision at very low temperatures. They were obtained starting from an accurate description of the LR interactions, essential in order to describe collisions at low energy\cite{47}. In a regime where the kinetic energy is so small, LR interactions determine the amount of incoming flux which reaches the SR, where rearrangement may occur\cite{20}. In our case the system is characterized by the presence of a significant quadrupole-quadrupole (QQ) contribution, which should lead to important reorientation effects. Therefore, the dynamical methodology has been revisited in order to include the most of the anisotropy that an adiabatic electronic treatment may enable. The theoretical results will be compared with the experimental results from the work by Costes and co-workers and Sims and co-workers\cite{23}.

The paper is structured as follows. In the next section, we will briefly describe the new LR calculations, while further details will be given in the Appendix. In section III, the hyperspherical approach to the dynamics\cite{48} will be recalled and small methodological improvements will be explained. The results from the dynamical calculations will be shown in section IV and they will be discussed and compared with the experimental data in Section V. Finally, a summary of the work and the conclusions will be given in Section VI.

II. LONG-RANGE INTERACTIONS

A. Formalism

The LR interactions between the sulfur atom $S(1D)$ and hydrogen molecule $H_2(X 1 \Sigma^+_g)$ have been described following perturbation theory up to second order, using a multi-polar expansion of the electrostatic interaction operator $\hat{H}_e$. To account for the 5-fold degeneracy of the open-shell atom $S(1D)$, we defined a set of asymptotically degenerate diabatic states, taken as a product of the atomic $|LA^{(A)}\rangle$ and diatomic $|0^{(B)}\rangle$ unperturbed electronic states. The quantum number $\lambda = 0, \pm 1, \pm 2$ is the projection of the atomic orbital angular momentum $L$ along the Body-Fixed (BF) $z$-axis, chosen along the intermolecular vector $\mathbf{R}$, and the projection $\lambda = 0$ of the diatomic orbital angular momentum relates to the diatom axis $\mathbf{r}$, chosen to be in the BF $xy$-plane. Following previous works on open-shell systems\cite{49,52}, the matrix elements of the first-order and second-order perturbation operators in the $|LA^{(A)}\rangle|0^{(B)}\rangle$ basis set lead to LR potentials that depend on the internal coordinates $(R, \theta)$, with $R$ being the intermolecular separation and $\theta$ the angle between $\mathbf{R}$ and $\mathbf{r}$. The dependence of the LR potentials on $r$, the internuclear distance of $H_2$, has been neglected in the present case. We have fixed $r$ at its vibrationally averaged value in the ground state, $(r)_{v=0}$.

The anisotropy of the LR $S(1D)$-$H_2$ interaction can induce transitions between the fragment states $|LA^{(A)}\rangle|0^{(B)}\rangle$ associated with different values of the projection quantum number $\lambda$, whereas $L$ is assumed to be conserved throughout the collision for large $R$. We can therefore drop the quantum number $L$ and label by $|\lambda\rangle$ the electronic diabatic basis of the $S(1D)$-$H_2$ interacting system. For linear geometries ($\theta = 0$), there is one-to-one correspondence between the states $|\lambda\rangle$ and the adiabatic states of $\Sigma$, $\Pi$ and $\Delta$ symmetry of $S(1D)$-$H_2$. Those adiabatic states are labeled according to the value of the quantum number $\Lambda = 0, \pm 1, \pm 2$, the projection of the total electronic orbital angular momentum $L$ along the BF $z$-axis, with $L$ being de-
fined with respect to the triatomic center-of-mass BF frame. For non-linear geometries, the diabatic states \(|\lambda\rangle\) are coupled through LR interactions, and one has to deal with a \(5 \times 5\) LR potential matrix \(V_{1,5}(R, \theta)\). Since the issue of non-adiabatic couplings is not addressed in present work, we chose to keep the (complex) signed-\(\lambda\) diabatic basis \(|\lambda\rangle\) to express the LR potential matrix elements. The diagonalization of the \(5 \times 5\) LR potential matrix yields the adiabatic potentials associated with the three \(A'\) and two \(A''\) adiabatic states correlating with \(S(1D)-H_2\).

In the present study, we have considered the electrostatic interaction between the permanent quadrupole moments of \(S(1D)\) and \(H_2\), leading to electrostatic energies varying as \(R^{-3}\), and the dispersion interaction between the dipole-induced moments of the two species, giving rise to dispersion energies varying as \(R^{-6}\). The contribution of induction effects to the LR potentials has been neglected, since the dominant induction contribution involves interactions between permanent quadrupole moments and dipole-induced moments, which are proportional to \(R^{-8}\). According to Eqs. [13] and [15] of the Appendix, the LR potential matrix elements write:

\[
V_{1,5}(R, \theta) = \frac{1}{R^3} C_{1,5}^{\lambda \lambda} C_{2,5-\lambda}^{\lambda \lambda}(\theta, 0) - \frac{1}{R^5} \sum_{k=0,2} C_{6,k}^{\lambda \lambda} C_{k,5-\lambda}^{\lambda \lambda}(\theta, 0)
\]

where the angular functions \(C_{lm}(\theta, \phi)\) are normalized spherical harmonics [76], and the interaction coefficients \(C_{1,5}^{\lambda \lambda}\) and \(C_{6,k}^{\lambda \lambda}\) stands for the electrostatic (quadrupole-quadrupole) and dispersion (dipole-induced dipole-induced) contributions, respectively.

### B. Quadrupole moments and dipole polarizabilities

The evaluation of the LR electrostatic \(C_{1,5}^{\lambda \lambda}\) and dispersion \(C_{6,k}^{\lambda \lambda}\) coefficients require to determine the permanent quadrupole moments and dynamic dipole polarizabilities of \(S(1D)\) and \(H_2\). For \(H_2\), we have selected accurate values from the literature, corresponding with a \(H_2\) geometry at its vibrationally averaged value, \(\langle r \rangle_{v=0} = 1.449\ a_0\). We used the permanent quadrupole value \(\langle 0|\bar{Q}_{20}|0\rangle = +0.481\ a_0\) of Ref. [53], obtained from full Configuration Interaction (CI) calculations with a large Gaussian-type orbitals basis set, including bond-centered polarization functions. For the dynamic dipole polarizabilities \(\alpha_{uu}(\omega)\) and \(\alpha_{vv}(\omega)\), we chose the values tabulated in Ref. [54], obtained from a sum-over-states formalism with explicitly electron-correlated wavefunctions. The corresponding static polarizabilities, \(\alpha_{uu}(0) = 6.721\ a_0\) and \(\alpha_{vv}(0) = 4.739\ a_0\), are in good agreement with other literature values [53].

For \(S(1D)\), we performed \textit{ab initio} calculations by means of the Dalton quantum chemistry code [55]. The permanent quadrupole moments have been calculated as the expectation values of cartesian quadrupole moment operators, \(\Theta_{uu}\), with \(u = \{x, y, z\}\), using \textit{ab initio} MCSCF electronic wavefunctions. The MCSCF wavefunctions were generated by distributing six electrons among 13 orbitals (3s, 3p, 3d, 4s and 4p), the inner-shell orbitals being kept frozen in their ROHF form, and the Sadlej-\(pVTZ\) basis set was used. The resulting value of the quadrupole moment \(\langle L_0|\bar{Q}_{20}|L_0\rangle = +2.075\ a_0\) was employed to get the whole set of nonzero matrix elements from the Wigner-Eckart theorem. Notice that the latter value is identical to the one already reported in the literature [56, 57] for the same level of calculation. The dynamic dipole polarizabilities of \(S(1D)\) have been determined by means of MCSCF linear response calculations, using the MCSCF wavefunctions previously defined. For each substate of definite symmetry, we obtained a set of Cauchy moments, each of them corresponding with a given cartesian component of the electric dipole operator \(\mu_{uu,n} = \langle uu|\hat{e}_n|uu\rangle\). The evaluation of the LR electrostatic contribution is described in previous works on alkali atoms [58, 59], and we get for the isotropic and anisotropic dispersion contributions \(\alpha_{uv}(\omega)\) following the procedure described in Ref. [49].

The resulting LR interaction coefficients for the electrostatic \(C_{1,5}^{\lambda \lambda}\) and dispersion \(C_{6,k}^{\lambda \lambda}\) contributions are tabulated in the electronic supplementary information (ESI). To the best of our knowledge, no LR coefficients had been determined so far for the \(S(1D)-H_2\) system, and thus it is not possible to compare with the results of previous works. By summing the tabulated values over the diagonal matrix elements \(\lambda = \lambda'\), we can retrieve state-averaged interaction coefficients. The state-averaged electrostatic contribution vanishes, and we get for the isotropic and anisotropic dispersion contributions \(C_{6,0} = 40.338\ a_0\) and \(C_{6,2} = 4.146\ a_0\), respectively.

### III. DYNAMICAL METHODOLOGY

#### A. The hyperspherical approach

The quantum methodology used to carry out the dynamical calculations was described in previous works on alkalis at ultracold energies [53, 60] and elsewhere in the context of thermal reactive scattering [61]. In fact, most of the convergence parameters used in this work are the same which were used in the study of the title collision at thermal energies [31]. Let us simply recall that in the hyperspherical quantum reactive scattering method developed by J. M. Launay [48] the configuration space is divided into inner and outer regions. The positions of the nuclei in the inner region are described in terms of hyperspherical democratic coordinates. The logarithmic derivative of the wavefunction is propagated outwards on a single adiabatic PES. At a large enough value of the hyper-radius the former is matched to a set of suitable functions, called asymptotic functions, to yield the scattering S-matrix.
matching.

The asymptotic functions provide the collisional boundary conditions. When working at thermal energies they are familiar regular and irregular radial Bessel functions which account for the presence of the centrifugal potential at large intermolecular separation. Thus, they were properly modified to include the effect of an isotropic dispersion interaction (\( R^{-6} \)), thus enabling the study of ultracold collisions of alkalis\[59, 60\]. In the current work further changes are required. Asymptotic functions must account also for the presence of the anisotropic electrostatic and dispersion LR interaction, introducing reorientation effects which may take place while the reagents approach. More details about their definition are given in the following section.

B. Introducing the anisotropy in the outer region: the asymptotic functions

Using the set of Jacobi coordinates \((R, r, \theta)\) corresponding to the S+H\(_2\) arrangement, the total Hamiltonian of the system can be expressed as

\[
\hat{H} = \frac{\hbar^2}{2\mu} \left( \frac{\partial^2}{\partial R^2} + \frac{1}{2} \frac{\partial^2}{\partial r^2} + \frac{1}{2} \frac{\partial^2}{\partial \theta^2} \right) + \hat{H}_{el} (2)
\]

where \(\hat{H}_{el}\) is the electronic Hamiltonian, \(I\) is the orbital angular momentum of the atom with respect to the center of mass of the diatom, and \(J\) is the rotational angular momentum of the latter. The total angular momentum of the nuclei (conserved in an adiabatic approach) is given by \(J = j + I\). In the current study, we chose an adiabatic treatment of the dynamics, assuming that the collision occurs only on the ground adiabatic PES. Hence, among the five adiabatic PESs correlating with \(S(1D)+H_2\) (three \(A\) and two \(A'\) singlet states), we consider only the lowest one, associated with the ground \(1 \, ^1A'\) electronic state. Hereafter, the lowest eigenvalue of \(\hat{H}_{el}\) will be labeled by \(V^0(R, r, \theta)\). The diagonalization of the LR potential matrix of Eq.1 provides a value for such energy, accurate at large intermolecular separations, with \(r\) fixed to its vibrationally averaged value \(\langle r \rangle_{v=0}\).

A convenient basis in order to expand the nuclear wavefunction in the LR region is the one characterized by quantum numbers \((J, M, v, j, l)\), with \((v, j)\) the rovibrational quantum numbers of the diatom, \(l\) the relative orbital angular momentum and \((J, M)\) the total angular momentum and its projection on the Space-Fixed Z axis. We will call this basis B1, and represent it as \(\phi_{v,j,M}^J\). Such a basis, well adapted to handle Coriolis couplings, is used in the hyperspherical approach to expand the asymptotic wavefunctions, which are matched with the SR information obtained in hyperspherical coordinates.

If the system approaches collision with quantum numbers \((J, M, v_0, j_0, l_0)\), we will assume that (in addition to \(J\) and \(M\)) the rovibrational quantum numbers, \((v_0, j_0)\), remain well conserved in the LR region. This is well justified in the present case given the large energy gap between different rovibrational states relative to the considered collision energy range. Within this approximation the total electro-nuclear wavefunction, \(\Psi_{v_0,j_0,l_0}^{JM}\), can be expanded in the LR region as

\[
\Psi_{v_0,j_0,l_0}^{JM} = \sum_l \frac{F_l^{v_0}(R)}{R} \phi_{v,j,M}^J (1 \, ^1A')
\]

where \(1 \, ^1A'\) is the ground adiabatic electronic state and all the "conserved" quantum numbers have been suppressed in the notation of the radial coefficients, \(F_l^{v_0}(R)\). Introducing the expansion (3) into the time-independent Schrödinger equation associated with a total energy \(E, \psi = \psi V\), and using the Hamiltonian in Eq.2, it is straightforward to obtain the following system of coupled radial equations

\[
\left[ -\frac{\hbar^2}{2\mu} \frac{\partial^2}{\partial R^2} + \frac{l(l+1)\hbar^2}{2\mu R^2} - E_c \right] F_l^{v_0}(R) = - \sum_{l'} \langle \psi_{l'} \hat{F}_l^{v_0}\rangle F_{l'}^{v_0}(R) (4)
\]

where \(E_c\), the collision energy, is given by \(E - E_{v,j}\), with \(E_{v,j}\) the internal energy of the diatom. \(\langle \psi_{l'} \hat{F}_l^{v_0}\rangle\) designs the matrix elements in basis B1 of \(V^0(R, r, \theta) - V_{H_2}(r)\), with \(V_{H_2}(r)\) the asymptotic \(H_2\) diatomic potential. By inwards integration of Eq.4 we obtain the "regular" \((F_l^{v_0})\) and "irregular" \((F_{l_0}^{v_0})\) asymptotic radial wavefunctions, corresponding to an incoming \((J, M, v_0, j_0, l_0)\) channel. They are defined as the ones which behave as

\[
F_l^{(1)} \xrightarrow{R \to \infty} \sin(kR - l\pi/2)\delta_{l_0,l}/k^{1/2} (5)
\]

\[
F_{l_0}^{(2)} \xrightarrow{R \to \infty} -\cos(kR - l\pi/2)\delta_{l_0,l}/k^{1/2}
\]

where \(k\) is the wavenumber of the considered channel. Such radial functions account for the correct LR behaviour to match with the inner-dynamics in the presence of an anisotropic \(V^0(R, r, \theta)\) potential. The coupled-equation version of the method of De Vogelaere [62] is used to solve for them.

Regarding the calculation of the potential matrix elements \(\langle \psi_{l'} \hat{F}_l^{v_0}\rangle\), it is convenient to define an intermediate nuclear basis, B2, labelled by the projection \(\Omega_j\) of \(j\) and \(J\) on the BF \(z\)-axis. The basis set B2 is given by

\[
\phi_{v,j,M}^{\Omega_j} = \frac{\chi_{v,j}(r)}{r} \sqrt{\frac{2l + 1}{4\pi}} D_{l l_0}^{\Omega_j}(\alpha, \beta, \gamma) Y_{l_0}(\theta, 0), (6)
\]

where \(D_{l l_0}^{\Omega_j}\) designs a Wigner rotation matrix element and \((\alpha, \beta, \gamma)\) are the Euler angles corresponding to the transformation between SF and BF frames. The change of basis between B1 and B2 simply involves \(3j\) symbols. The matrix elements of the ground PES in basis B2 are given by

\[
\langle \psi_{\Omega_\alpha, \Omega_\gamma}\rangle (R) = \delta_{\Omega_\alpha, \Omega_\gamma} 2\pi \int_{0}^{2\pi} \chi_{v,j}(r)^2 Y_{l_0}^2(\theta, 0) V^0(R, r, \theta) \sin \theta \; dr \; d\theta (7)
\]

The potential does not couple states with different \(\Omega_j\), although reorientation effects in the BF system happen due to the Coriolis term. Assuming in Eq.7 that the variations with \(R\) of the mean value of \(V^0(R, r, \theta)\) in the rovibrational state \(\chi_{v,j}(r)\) are well approximated by the variations of the potential at the averaged bond distance, \(r = \langle r \rangle_{v,j}\) (assumption which has been proven valid for the current PESs), we reach the convenient expression

\[
\langle \psi_{\Omega_\alpha, \Omega_\gamma}\rangle (R) = \delta_{\Omega_\alpha, \Omega_\gamma} 2\pi \int_{0}^{2\pi} \chi_{v,j}(r)^2 Y_{l_0}^2(\theta, 0) V^0(\langle r \rangle_{v,j}, \theta) \sin \theta \; dr \; d\theta
\]
Once the potential matrix elements are calculated in basis B2 we change to basis B1, thus obtaining the elements \( \langle V \rangle_{\Omega,\Omega'}(R) \) involved in Eq. \[4\]

\[
\langle V \rangle_{\Omega,\Omega'}(R) = \langle \Omega \Delta \rangle \int \frac{Y^2_{\alpha_0}(\theta, 0) V^0(R, \{r\}, \theta) \sin \theta d\theta. }
\]

Let us finally note that \( \langle V \rangle_{\Omega,\Omega'}(R) + l_0(l_0 + 1)\hbar^2/2\mu R^2 \) has the meaning of the effective potential felt by the colliding partners at a distance \( R \) when approaching in the state \( \phi_{\nu,\lambda} \). Such meaning will be used below.

### C. The potential energy surface

In our adiabatic approach we consider that the collision takes place on the ground PES. The global fit for the \( ^1A' \) PES performed by Ho et al. \[29\] (surface S1) was widely used in the past and its SR region was tested by comparison with experiments at higher energies \[31\]. Nevertheless, the surface does not pay the necessary attention to the description of the LR region, that being the reason why we performed our own calculations of the LR potentials (see Sec. \[II\]). To combine the SR interactions given by S1 with our description of the LR region (the lowest eigenvalue of the LR potential matrix in Eq. \[1\]), we performed a smooth switching from one to the other. Thanks to the approximation in Eq. \[8\] only the LR potentials for fixed \( r = \langle r \rangle_{\nu, \lambda} \sim \langle r \rangle_{\nu, 0} \) are required, and only the switching for such internuclear distance is considered. The switching was performed around the central value \( R_0 = 12.5 \text{ a.u.} \), within a range of intermolecular distances given by \( \Delta R = 3 \text{ a.u.} \). The angle-independent switching function \( f(R) = \{\cos[\pi(R + \Delta R/2 - R_0)] + 1\}/2 \) was used. In contrast, note that an intermolecular separation of \( \approx 10 \text{ a.u.} \) was considered for the matching of the inner dynamics with the external asymptotic functions.

It is worth mentioning that a new global \( ^1A' \) PES for the title system was recently published \[63\]. A term to account for the \( \sim R^{-6} \) dipole-induced dipole-induced (DD) dispersion interaction was included in the global fit. Nevertheless, the quadrupole-quadrupole (QQ) electrostatic interaction \( \sim R^{-5} \), which dominates the LR behavior, was not explicitly considered. As such term is relevant to describe the reactivity of ortho-hydrogen at low collision energies, we have preferred to employ the PES obtained from the switching procedure just described.

### D. The collision with \( j=0 \) para-hydrogen

The collision of \( S(1^D) \) with \( \text{para-hydrogen} \) in its ground rotational state deserves special attention. Within a non-adiabatic treatment the effective contribution of the anisotropic terms in \( V_{\lambda\lambda'}(R, \theta) \) (Eq. \[1\]) is found to vanish. This is not the case when using an adiabatic approach instead. In order to show this we define a set of electro-nuclear basis functions in terms of the diabatic electronic basis defined in Section \[II\]

\[
\phi_{\nu,\lambda, M}(r) = \frac{X_{\nu,0}(r)}{r} \sqrt{\frac{2J_T + 1}{4\pi}} \frac{I_{M_T, M}}{I_{\nu,\lambda}} (\alpha, \beta, \gamma) Y_{\nu,\lambda}(\theta, 0) |L, \lambda \rangle (10)
\]

where \( J_T \) designs a total angular momentum, \( J_T = j + l + L \), which in contrast with Eq. \[6\] includes the atomic electronic angular momentum, and \( M_T = (\lambda + 1) \) and \( M_T \) label its projection on the BF and SF z-axis, respectively. The use of this basis in combination with the whole interaction matrix given by Eq. \[II\] would efficiently account for the exchange of angular momentum among electrons and nuclei (\( J_{T} \) being conserved) and reorientation effects appearing while the reagents approach \[64\]. Potential couplings between two different channels in this non-adiabatic treatment would involve the integration in \( \theta \) of the product of two spherical harmonics (originating from Eq. \[10\]) and one matrix element \( V_{\lambda\lambda'}(R, \theta) \). The latter consists for spherical harmonics, \( Y_{k,m}, \) with \( k = 0, 1, 2 \) \( (C_{k,m} \sim Y_{k,m} \text{ in Eq. } 1) \). When considering the case of \( \text{para-hydrogen} \) with \( j=0 \), such integral is found to vanish for \( k \neq 0 \). The contribution of QQ and dispersion anisotropic terms disappears, and thus the \( R^{-5} \) dependence. Only the isotropic dispersion term, \( \sim R^{-6} \), is found to play a role. This is not the case when the adiabatic approach is used. The diagonalization of the LR \( V_{\lambda\lambda'}(R, \theta) \) matrix yields a ground PES with a \( R^{-5} \) dependence, which is not cancelled through the integral in Eq. \[8\]. In order to correct such a drawback of the adiabatic treatment we artificially matched S1 with a pure isotropic dispersion term, \( -C_6/R^6 \), when dealing with \( j=0 \) para-hydrogen. In this way we introduced in the adiabatic treatment the effective LR interaction which the colliding partners should experience. The \( C_{6,0} \) coefficient, the lowest isotropic one, was chosen for that purpose.

### IV. RESULTS

Quantum reactive cross-sections for collisions of \( S(1^D) \) with \( j=0 \) para-hydrogen (p-H\(_2\)) and \( j=1 \) ortho-hydrogen (o-H\(_2\)), in the energy range 1–120 K, were calculated using the methodology described in the previous section. In Fig. \[I\] the obtained results are shown. A small dependence with the rotational state and some interesting oscillations, more apparent...
FIG. 1: Quantum reaction cross-sections for the collision of $S(1D)$ with $j=0$ $p$-$H_2$ and $j=1$ $o$-$H_2$ at low kinetic energies.

in the case of $p$-$H_2$, are relevant features. The cross-sections were averaged using the room-temperature ratio 3:1 of $o$-$H_2$ to $p$-$H_2$, so-called normal-hydrogen ($n$-$H_2$), which is expected to be valid in the conditions of the experiment by Costes and co-workers [27]. They were also convoluted with a Gaussian distribution of collision energies to account for the velocity and crossing angle spread in the experiment [27]. In Fig. 2 the experimental excitation functions (in arbitrary units) were scaled to minimize the sum of the squares of the differences between every experimental result and its corresponding theoretical value. Both sets show very similar decreasing trends as a function of the collision energy.

Some additional calculations for particular collision energies in the range 150–1500K were performed in order to allow interpolations and the calculation of thermally-averaged rate constants to compare with the experimental results by Sims and co-workers [23, 27]. Again, the room-temperature ratio 3:1 was considered. It was estimated that such proportion is not perturbed by the rapid cooling to 77 K and the subsequent adiabatic expansion to 5.8 K in the experiment. By using a Maxwell-Boltzmann distribution the population of higher rotational states was estimated to be negligible for all the experimental temperatures except for the highest one, $T=300K$. Given the independency of the sets of rotational states with even and odd quantum number and the big rotational constant of $H_2$ ($\sim 85K$), only in the case of $p$-$H_2$ and for such high temperature the population of $j=2$ was found to be similar to the population of the ground state. Accordingly, we calculated the cross-sections for $j=2$ at some particular collision energies and concluded a difference of $\sim 5\%$ between the cross-sections for $j=0$ and $j=2$. The contribution of $j=2$ was not considered in the calculation of thermally averaged rate-constants and a value somewhat lower than the one in Fig. 3 is expected for $T=300K$.

Interestingly, the theoretical rate constants approximate fairly well the experimental ones, remaining below them with the exception of the value for the lowest experimental temperature (5.8K). This temperature is also the most challenging from an experimental point of view. The 300 K experimental result of Black and Jusinski [65] is also shown for comparison, in perfect agreement with the recent measurements. Let us remark that a multiplicative factor 1/5 was included in the calculation of rate constants, as only one of the five asymptotically degenerate adiabatic surfaces is assumed to lead to the products in the considered energy range.

V. DISCUSSION

A. The significance of the long-range interactions

In order to stress the need for accuracy in the description of LR effects while working at low collision energies, the sensitivity to such effects is shown in Fig. 3. Cross-sections for the collision with $p$-$H_2$, calculated using an uncorrected PES S1 [29], are compared to the ones obtained by matching it with a pure isotropic $R^{-6}$ dispersion term at LR (see Section III D). As the figure shows, for energies below 20K the accurate inclusion of the LR interactions appears as essential. The use of S1 without corrections leads to differences of the order of 30% for energies around 1K.

Note that the LR behaviour of an accurate ab initio surface for the ground electronic state contains a non-negligible $R^{-5}$ character in the title system. As argued above, such contri-
with accurate treatment of the LR non-adiabatic dynamics seems forces the idea that for asymptotically degenerate systems an approach (which considered the 5 asymptotically degenerate surfaces and all the non-adiabatic couplings). Nevertheless, it does not cancel while using an adiabatic approach (that being the reason to force a LR behaviour $R^{-6}$ in the calculations with $p$-H$_2$). The need for such artificial modifications reinforces the idea that for asymptotically degenerate systems an accurate treatment of the LR non-adiabatic dynamics seems essential. To the best of our knowledge, non-adiabatic effects in low energy collisions have not been explored in depth up to the date.

B. The electronic quenching

In view of the results of Maiti et al.\textsuperscript{36} and Tian-Shu et al.\textsuperscript{46} the electronic quenching is expected to play a role in the title collision. Since the experimental rate constants measured by Sims and co-workers\textsuperscript{27} probe the total removal of $S(1D)$ by $n$-H$_2$, they should comprise the disappearance of $S(1D)$ due to the reactive process (formation of SH+H) or to the electronic quenching (relaxation to $S(^3P_{0,1,2})$). The excitation functions measured by Costes and co-workers\textsuperscript{27} monitor only the first mechanism. As our theoretical calculations do not include the possibility of quenching some comments are pertinent in order to justify mutual comparisons. What should we expect from the use of an electronically-uncoupled (EU) formalism to describe the dynamics when an electronically-coupled (EC) one would be more appropriate? A reasoning according to the statistical model of Rackham and Manolopoulos\textsuperscript{44}, applied in the past to the system at thermal energies\textsuperscript{33, 42}, may help to clarify the limits and validity of our approach. If the collisional process is assumed to be mediated by a long-lived complex (due to the deep well in the PES) we can distinguish the step of formation from the step of statistical decomposition of the complexes. Since the singlet-triplet crossing occurs inside the well we will consider the quenching as an alternative way of decomposition. Besides, we affirm that our calculation describes correctly the first step, the way the probability is captured. Regarding the second step, in a scenario where some complexes would decompose to give back the reactants (I), some to give reaction products (R), and the rest would suffer quenching (Q), our electronically-uncoupled (EU) calculation does not allow this latest possibility. By making use of the principles of the model, and designing by $\sigma^E_C$, $\sigma^E_R$, $\sigma^E_Q$ and $\sigma^E_{EU}$, $\sigma^E_{EC}$ the cross-sections for each possible outcome in an hypothetical EC (unprimmed) or in our EU (primmed) calculation, we will show that

$$\sigma^E_R + \sigma^E_Q > \sigma^E_{EU} > \sigma^E_{EC}$$

(11)

As a consequence, the theoretical rate constant calculated using $\sigma^E_{EU}$ constitutes a lower bound for the experimental total removal rate of $S(1D)$. The latter should be given by the sum of both reactive and quenching contributions in an exact EC calculation. Besides, $\sigma^E_R$ is itself an upper bound for the real reaction cross-section, monitored in arbitrary units by the measured excitation functions. These statements should be valid even if $\sigma^E_{EU}$ is far from $\sigma^E_{EC}$. Below we will show how to reach these conclusions.

According to the statistical model\textsuperscript{44}, the fraction of complexes which decompose into a particular channel is related to the capture probability of forming the complex starting from that particular channel. Let us denote with $\Lambda_i^R(E)$, $B_i^R(E)$ and $C_i^p(E)$ the energy dependent capture probabilities starting from reagent, product and quenching sides, cor-
responding to particular channels \( n, n', n'' \), respectively. \( J \) labels the total angular momentum. Let us define the sum in open channels of the capture probabilities associated to each possible outcome, \( A^i(E) = \sum A^i_n(E) \), \( B^i(E) = \sum B^i_n(E) \) and \( C^i(E) = \sum C^i_n(E) \). Following the prescriptions of the model, the reaction probability corresponding to the reagents colliding in channel \( n \), \( P^I_{R,n}(E) \), is given by \( P^I_{R,n}(E) = A^i_n(E)B^i(E)/\left(A^i(E) + B^i(E)\right) \) in our EU calculation and by \( P^I_{R,n}(E) = A^i_n(E)B^i(E)\left(A^i(E) + B^i(E) + C^i(E)\right) \) in an hypothetical EC calculation. Interestingly, the former is larger. Now, essentially by summing over \( J \), one obtains cross-sections and it follows that \( \sigma^E_{EC} > \sigma^E_{R} \), which is the second inequality in expression (11). A similar reasoning leads to the probability of total removal, \( P^I_{R,n}(E) + P^Q_{R,n}(E) = A^i_n(E)\left(B^i(E) + C^i(E)\right)\left(A^i(E) + B^i(E) + C^i(E)\right) \), to the first inequality. In agreement with it, the thermally averaged theoretical rates in Fig. 3 remain below the experimental ones, with the exception of the value for the lowest temperature.

C. The Langevin-type capture dynamics

Both the theoretical and the experimental excitation functions, in Fig. 2 show no threshold, consequence of the attractive character without barrier of the reaction on the ground electronic state. Remarkably, both follow very similar trends as a function of the kinetic energy, in spite of the absence of quenching channels in the theoretical calculations. As a first guess, such proportionality would result from the real reaction probabilities following the entrance channel capture probabilities, \( A^i_n(E) \), given that the capture is well described in our approximate calculation. Moreover, if our guess were correct the experimental reaction cross-sections should have a Langevin-capture character (see note [67]).

Such qualitative reasoning can be put in solid grounds by making use again of the statistical model. Given the exothermicity of both the reactive and quenching processes, many internal states are open even at low collision energies in the corresponding arrangements. In particular, \( B^i(E) \) may be approximated by the number of open channels, thus being much larger than \( A^i(E) \). The value of \( C^i(E) \) is more difficult to estimate as the capture from the quenching side requires a non-adiabatic transition, of unknown probability, to occur. Regardless, it is reasonable to accept that \( B^i(E) \), and thus \( B^i(E) + C^i(E) \), are much bigger than \( A^i(E) \) in the energy and angular momentum ranges which contribute, thus following that most of the complexes are expected to form SH+H or to suffer quenching. Besides, such figures are expected to be essentially constant, \( B^i(E) \sim B, B^i(E) + C^i(E) \sim B + C \). Accordingly, in we neglect the contribution of \( A^i(E) \) in the denominator of \( P^I_{R,n}(E) \) (see above) we find a result which is essentially equal to a reactant capture probability, \( A^i_n(E) \). Proceeding similarly we find that \( P^I_{R,n}(E) \) is a constant portion of the capture probability, \( P^I_{R,n}(E) \sim \left(B^i(B + C)A^i_n(E)\right) \), and thus proportional to \( P^I_{R,n}(E) \). Finally, the sum \( P^I_{R,n}(E) + P^Q_{R,n}(E) \) is again equal to the capture probability \( A^i_n(E) \) and therefore to \( P^I_{R,n}(E) \). By summing over \( J \) to obtain cross-sections, the previous results are found equivalent to the following statements: i) \( \sigma^E_{EC} \) is essentially a capture cross-section which accounts for the total number of complexes formed in the collision, ii) \( \sigma^E_{EC} \) should be found proportional to \( \sigma^E_{R} \), and iii) \( \sigma^E_{EC} + \sigma^E_{Q} \) and \( \sigma^E_{EC} \) must be similar to each other. We will check their validity below.

Regarding the first statement, the quantum cross-section averaged for \( n-H_2 \) is compared in Fig. 5 with an averaged Langevin capture estimate and a good agreement is found. A capture dispersion potential \(-C_5/R^6 \) was assumed for \( p-H_2 \), what leads to an analytical Langevin cross-section given by \( \sigma^E_{Lam}(E) = 3\pi(C_6/4E)^{1/3} \). A QQ electrostatic potential, \(-C_5/R^3 \), and the corresponding cross-section, \( \sigma^E_{EC}(E) = 5\pi(C_5/6\sqrt{E})^{2/5} \), were assumed for \( o-H_2 \) instead. The values \( C_6 = C_6^{0,0} = 38.9 \) a.u. and \( C_5 = 4.08 \) were respectively taken. The former is the lowest isotropic dispersion coefficient and the latter results from an angular average of the ground eigenvalue of the QQ contribution in Eq. 1. Double weight is assigned to the perpendicular approach, \( C_5(\theta = 90) = \frac{\sqrt{39}}{2} \langle \hat{Q}(20)\rangle\langle \hat{Q}(20)\rangle_{H_2} \), with respect to the linear one, \( C_5(\theta = 0) = 6 \langle \hat{Q}(20)\rangle\langle \hat{Q}(20)\rangle_{H_2} \), the factor 4.08 resulting from such weighted average. The second statement explains the similar trends found between theoretical and experimental excitation functions, in spite of the non-inclusion of quenching in the calculations. Finally, the third statement justifies the similarity of the experimental and theoretical rates, shown in Fig. 5. The bigger discrepancy between the experimental and theoretical rate coefficients at 5.8 K, the latter being larger, may be related with the neglected non-adiabatic couplings between the three singlet \( A' \) states which correlate with the reactants. In fact, the discrepancy can be interpreted as the theoretical simulations capturing more probability in the complex than they should, and neglecting the couplings to repulsive states could be a reason for such an effect. The effect of such couplings should increase dramatically when decreasing the kinetic energy, while the contribution from the neglected singlet-triplet couplings is expected to be less energy dependent. Regardless, such low kinetic energies should be more sensitive to possible inaccuracies of the considered PES.

To outline this section, let us remark that our theoretical cross-sections are essentially capture cross-sections and they must account well for the total number of complexes formed in the collision. The number of complexes which eventually react is found a constant portion of the total number. This explains the experimental reaction cross-sections being proportional to the theoretical cross-sections. Finally, as the experimental total removal rates include the sum of reaction and quenching, they are monitoring again the total number of complexes, so they are well approximated by our theoretical results.

D. The structures in the cross-sections

The theoretical cross-sections show interesting oscillations superimposed to the overall Langevin behaviour (Fig. 6).
FIG. 5: In the figure, the theoretical cross-section averaged using the 3:1 proportion of \( o-H_2 \) to \( p-H_2 \) present in the experiment by Costes and co-workers \cite{27} is compared with the averaged analytical Langevin estimate \cite{66} (see text).

They can be observed for both \( p-H_2 \) and \( o-H_2 \), although their presence seems to be washed out by the sum in partial waves in the latter case. In a naive approach, zero-order resonances present in the case of \( p-H_2 \) would split in the case of \( o-H_2 \) due to the different values of \( \Omega \). Their abundance and mutual overlap would justify their being more difficult to observe in the latter case.

The observed structures, which can be attributed to particular partial waves and channels have a double origin: some of them display the effective opening of a partial-wave \( l \); others seem to be due to the presence of shape resonances. Regarding the former, the Wigner threshold laws provide a behaviour for the \( l \) cross-section below the centrifugal barrier given by \( E^{l+1/2} \). If, after the barrier were overcome, the reaction probability reached quickly the limit value of one, the behaviour should follow a \( 1/k^2 \) dependence, that is \( \sim E^{-1} \) \cite{67–69}. This would result as a “bump” in the cross-sections. A bump of this type should happen at energies over the height of the centrifugal barrier. On the contrary, oscillations for high partial waves stemming from shape resonances would occur at energies below the centrifugal barriers, the associated effective potentials providing some kind of trapping.

Some structures have been selected and they are shown in Fig. 6 to be associated to particular partial waves and channels, \( \phi_{M}^{JH} \). The corresponding centrifugal barriers are also displayed as vertical lines. They were calculated using effective potentials, \( \langle V \rangle_{l}(R) + l(l+1)\hbar^2/2\mu R^2 \), where the average \( \langle V \rangle_{l}(R) \) was defined in Eq. 9. Panel (a) corresponds to \( j=0 \) \( p-H_2 \) and panel (b) to \( j=1 \) \( o-H_2 \). Given the shape and location (above the centrifugal barrier) of the peaks associated to \( p-H_2 \), with entrance channels \( J = l = 7 \) and \( J = l = 12 \), and to \( o-H_2 \) with entrance channel \( J = 9, l = 8 \), we conclude that their origin is the opening of a new partial wave. In contrast, the peaks labelled A and C for \( p-H_2 \), associated to the \( J = l = 15 \) and \( J = l = 16 \) contributions, are found below barrier. The same happens for the peaks A, B, C and D in the lower panel, associated to the entrance in channels \( (J = 16, l = 15), (J = 16, l = 16), (J = 17, l = 16) \) and \( (J = 17, l = 17) \), respectively. As we will explain below, we attribute them to the presence of shape resonances. Note that peaks A and C are found at similar energies for \( o-H_2 \) and \( p-H_2 \).

The presence of multiple resonances in the reaction probability of the title system for zero total angular momentum was reported in the past \cite{50}. They manifest the indirect mechanism through the formation of long-lived complexes. If the peaks selected in the current work were resonances, the associated trapping should happen in an intermediate range of radial distances. On the one hand, they appeared in calcula-
tions where the LR interaction was not well accounted for (see Fig. 4), so they cannot be located at very large distances. On the other, their location at very short range would hamper, due to the high anisotropy of SR chemical forces, any rotational adiabaticity which may lead to similar effective potentials, and thus to the similar structures observed, for \( j = 0 \) and \( j = 1 \). In good agreement with such qualitative reasoning, the PES shows a shallow well in the entrance valley at distances of \( \sim 7 \) a.u. [42]. With a depth of \( \sim 500 \)K for collinear approaches, and appearing as a small secondary minimum opening the access to the big well for insertion paths, this complex is the probable origin of many of the structures in the cross-sections. As a result of such well, the effective potentials of the channels associated to the peaks show a double maximum structure (see Fig. 7). The local minimum inbetween may support shape resonances at precisely the energies where the structures appear. Remarkably, the effective potentials corresponding to channels \((J = 16, l = 15)\) and \((J = 17, l = 16)\) of \(\alpha\)-H\(_2\) are respectively very similar to the ones corresponding to \(J = l = 15\) and \(J = l = 16\) of \(p\)-H\(_2\). This justifies the appearance of peaks A and C at similar energies in both cases. The approaching partners would be trapped in this preliminar well until they tunneled into the big one, thus finding the way to reaction, or they tunneled backwards, thus giving the reagents back. The quasibound states seem to prefer “prereaction” though, increasing reactivity at the associated energies. Note finally that Fano-type profiles, which indicate quick changes in the phase of the elastic \(S\)-matrix elements, are found in the elastic cross-sections at similar energies, thus reinforcing the hypothesis about the nature of the peaks as resonances.

Let us remark that the role of van der Waals wells in reactive collisions at very low energy was recently stressed [70]. As stated, the supported quasibound states may lead to a temporal trapping of the system at short distances, and to an increase of tunneling through the barrier. Such effects were mainly discussed for tunneling-dominated reactions like \(\text{Li}+\text{HF} [71]\) and \(\text{Cl}+\text{HD} [70]\), where the reaction would be closed otherwise. In addition to the considered shape resonances, Feshbach-type ones may occur for zero angular momentum and can dramatically modifying ultracold reaction cross-sections and zero-temperature rate constants [72].

VI. CONCLUSIONS

In this work, the \(\text{S}^1(1D) + \text{H}_2\) collision was analyzed in the kinetic energy range \(1–120\)K. Within an adiabatic treatment, working on the lowest surface which correlates with the reagents, cross-sections and thermally averaged rate constant were obtained using the quantum reactive scattering method developed by Launay et al. [48]. The resulting picture of the process is the following: the behavior can be understood on the average in terms of a classical Langevin (capture) model where centrifugal barriers determine the amount of reactive flux which reaches the barrierless transition state; additionally, the structure of the van der Waals well provides temporal trapping at short distances, thus helping the system to find its way to reaction for high partial waves. Comparison with landmark experimental data obtained using the crossed molecular beam machine with variable beam intersection-angle [24] (Bordeaux) and the CRESU technique [22, 23] (Rennes) shows a fairly good agreement. Accurate electronic calculations of the LR interactions were required in order to properly describe collisions at such low kinetic energy. In particular, the anisotropic QQ term of the interaction appears to be important to describe the collision with \(\alpha\)-H\(_2\), although it does not contribute in the case of \(p\)-H\(_2\).

Previous theoretical works at higher energies indicated that the electronic quenching process, and thus the intersystem crossing, may play a significant role in the total removal of \(\text{S}^1(1D)\) [56, 46]. Nevertheless, a good agreement of our theore-
the experimental total removal rates has been found in spite of the adiabatic treatment of the collision. Moreover, the resealed experimental excitation functions follow very well the theoretical ones. An small contribution of the quenching must not be concluded though. The current theory accounts well for the total number of complexes formed in the collision, this amount being what our theoretical rates and cross-sections essentially monitor. Now, as most of the complexes decompose to give either reaction or quenching, and experimental removal rates sum up both contributions, a good agreement of theoretical and removal rates follows. Besides, the ratio of reaction to quenching is not expected to significantly change in the small considered energy range. The amount of complexes which ’react’ at one particular energy appears then a constant fraction of the total number of complexes captured at such energy. A proportionality of the ’real’ reaction cross-section to our theoretical cross-sections automatically follows, and therefore the good agreement with the experimental excitation functions. It must be noted that both discussed agreements seem to follow independently to the proportion of quenching and, in our opinion, the question remains open. Future measurements of the branching ratio reaction/quenching would be thus desirable.

Interestingly, the theoretical cross-sections showed oscillating features. We conclude that some of them are associated to the presence of a small van der Waals well in the entrance valley of the ab initio PES S1.20 Far from being an artifact of the fit, this well seems to be real, appearing in some ab initio calculations we have performed to verify its existence. Nevertheless, we have found a much smaller depth. Although the current experimental results do not allow conclusive confirmation of the presence and nature of the oscillating structures, the isolation of the contribution of p-H2 in the cross-beam experiment or the complementary measurement of differential cross-sections at thermal energies has proven elusive.23. If they were resonances, they would be very sensitive to the potential energy surface and they would give us important information about it. Note that due to the high degree of averaging into orientations, energies and directions of approach, direct observation of resonant structures in reaction cross-sections at thermal energies has proven elusive.24. Up to the date it was only successful in the F+HD system, and thanks to the concurrence of a set of favorable dynamical conditions. The current experimental access to the low-temperature regime, charaterized by small partial-wave averaging, may lead to a different scenario.

Finally, our work reinforces the idea that an accurate treatment of non-adiabatic dynamics at LR for asymptotically degenerate systems seems fundamental at low energies. Previous studies at thermal energies stressed the redistribution of asymptotic flux onto the different PES which may result from such degeneracy while the reagents approach.64. The role of such non-adiabatic connections at low energies has not been explored yet. A better description of the title process including non-adiabatic effects is in progress.

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VII. APPENDIX : LONG-RANGE ELECTROSTATIC AND DISPERSION ENERGIES

We consider below the case of LR interactions between an open-shell atom, A, and a closed-shell diatom, B. To account for the n-fold degeneracy of the atom, we define a set of asymptotically degenerate diabatic states, taken as a product of the atomic |Lλ⟩(A) and diatomic |0⟩(B) unperturbed electronic states. The quantum number λ = −L,...,L is the projection of the atomic orbital angular momentum L along the Body-Fixed (BF) z-axis, chosen along the intermolecular vector R, and the projection λ = 0 of the diatomic orbital angular momentum refers to the diatomic axis r, chosen to be in the BF xz-plane.

The multipolar expansion of the first-order, Ĥ(1), and second-order, Ĥ(2), perturbation operators depend on multipole moments 〈Qλ⟩ and 〈Qλ⟩(B), that are defined in frames centered on A and B, with axes parallel to a global Space-Fixed (SF) frame. To treat the atom-diatom case, we introduce multipole moments 〈Qλ⟩ that are defined relative to a local reference frame on B,

\[ \hat{Q}^{(B)}_{M,m_{\lambda}} = \sum_{m_{\lambda}} \hat{Q}^{(B)}_{m_{\lambda}} \left[D_{M,m_{\lambda}}^{(B)}(\hat{\Omega}_B)\right]^\dagger, \]

(12)

where 〈ΩB collects the polar angles of the diatom axis r relative to the SF frame, and D(M,mλ) is a Wigner rotation matrix. By rotating the SF frame such as the z-axis coincides with R, and r lies in the xz-plane, the Wigner rotation matrix reduces to [D(M,mλ)(\hat{\Omega}_B)]\dagger = d(M,mλ)(\theta), with \theta being the angle between R and r.

Following previous works on open-shell interacting systems [49, 52], the matrix elements of Ĥ(1) in the |Lλ⟩(A)|0⟩(B) basis set describe the electrostatic energies and, according to the chosen orientation for the global frame, they can be written as:

\[ V_{L_1L_2}(R,\theta) = \sum_{l_1l_2m} \frac{1}{R_{l_1l_2+1}} V_{l_1l_2}^{L_1L_2} C_{l_2-m}(\theta,0) \]

(13)

where the angular functions C_{l,m}(\theta, \phi) are normalized spherical harmonics, and the electrostatic interaction coefficients are given by:

\[ V_{l_1l_2}^{L_1L_2} = (-1)^{l_2} \left(\frac{2L_1}{2l_1}\right)^{1/2} \langle l_1ml_2 - m | L_20 \rangle \times \langle L_20 | \hat{Q}^{(A)}_{l_1l_2} | L_10 \rangle \]

(14)
where $L_{12} = l_1 + l_2$. From the Wigner-Eckart theorem, it follows that the multipole moments matrix elements $(L|\hat{Q}_{\lambda}^{(A)}|L')$ and $(0|\hat{Q}_{\lambda}^{(B)}|0)$ vanish unless $m = \lambda - \lambda'$ and $\mu_2 = 0$, respectively. For the sake of clarity, the electrostatic interaction coefficient $V_{\lambda\mu}^{\lambda\mu}$ will be substituted hereafter by the label $C_{\lambda\mu}$, since it is commonly used in the context of LR interactions.

The matrix elements of $H_{\lambda\mu}^{(2)}$ in the $(|L\rangle^{(A)}|0\rangle^{(B)}$ basis set describe the dispersion energies, and write as [49,51]:

$$V_{\lambda\mu}^{\lambda\mu}(R, \theta) = - \sum_{l_1, l_1', k_1, k_1'} \frac{1}{R_{l_1+2, l_1'+2}} \sum_{k_2 M} V_{l_1 l_1', k_1 k_1'}^{l_2 l_2', k_2 M} C_{k_2 M}(\theta, 0)$$

where the dispersion interaction coefficients are given by:

$$V_{l_1 l_1', k_1 k_1'}^{l_2 l_2', k_2 M} = \sum_{k_1 k_2} h_{l_1 l_1', k_1 k_2} (k_1 M k_2 - M | k_0) \chi_{l l', k k', k_1 k_2}^{l l', k k', k_1 k_2}$$

The angular coupling coefficient $h_{l_1 l_1', k_1 k_2}$ is defined as in Eq. 21 of Ref. [50], and the quantity $\chi_{l_1 l_1', k_1 k_2}^{l_1 l_1', k_1 k_2}$ is a coupled form of Casimir-Polder integral:

$$\chi_{l_1 l_1', k_1 k_2}^{l_1 l_1', k_1 k_2} = \frac{1}{2 \pi} \int_0^{\infty} \lambda_{l_1 l_1', k_1 k_2}(i\omega) \alpha_{l_1 l_1', k_1 k_2}(\omega) d\omega$$

From the Wigner-Eckart theorem, it follows that the coupled dynamic polarizabilities $\lambda_{l_1 l_1', k_1 k_2}$ vanish unless $M = \lambda - \lambda'$ and $Q_2 = 0$, respectively. Furthermore, in the particular case of dipole-dipole polarizabilities $(l_1 = l_2 = 1)$ and a diatomic state $(|L\rangle = |L'|)$, the matrix elements $\lambda_{l_1 l_1', k_1 k_2}^{l_1 l_1', k_1 k_2}$ vanish unless $Q_2 = \lambda - \lambda'$ and $k_2$ is even. Again, for the sake of clarity, we substitute hereafter the dispersion interaction coefficient $V_{l_1 l_1', k_1 k_2}^{l_2 l_2', k_2 M}$ by the more commonly used label $C_{l_1 l_1', k_1 k_2}^{l_2 l_2', k_2 M}$.

In present work, we have considered the quadrupole-quadrupole interactions ($l_1 = l_2 = 2$) between an open-shell atom $A$ and a closed-shell diatomic $B$. Hence, the quadrupole moments $(L|\hat{Q}_{2\lambda}^{(A)}|L')$ and $(0|\hat{Q}_{2\lambda}^{(B)}|0)$ are the necessary ingredients to compute the LR electrostatic coefficient $C_{2\lambda\lambda}$ of Eq. [14]. Besides, we have considered the dipole-induced dipole-induced interactions ($l_1 = l_1' = l_2 = l_2' = 1$) between the two species. In such a case, the LR dispersion coefficient $C_{l_1 l_1'}^{k_1 k_2}$ vanishes unless $k_1$, $k_2$, and $k$ take even values, and the necessary ingredients are the coupled dipole-dipole dynamic polarizabilities $\alpha_{2\lambda\lambda}^{l_1 l_1', k_1 k_2}$ and $\alpha_{2\lambda\lambda}^{l_1 l_1', k_1 k_2}$ with $k_1 = 0, 2$ and $k_2 = 0, 2$. Notice that the matrix elements are here defined in terms of spherical components of the $\hat{Q}_{2\lambda\lambda}$ moment and $\hat{Q}_{2\lambda\lambda}$ dynamic polarizability operators, together with a basis set of (complex) signed-$\lambda$ electronic states. Nonetheless, we can benefit from the Wigner-Eckart theorem to consider only the diagonal matrix elements, which have one-to-one correspondence with the quantities derived from quantum chemistry calculations.

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[74] The normalized spherical harmonics \( C_{lm} (\theta, \phi) \) relate to the associated Legendre polynomials \( P_l^m (\cos \theta) \) as follows:

\[
C_{lm} (\theta, \phi) = (-1)^m \frac{(l-m)!}{(l+m)!} P_l^m (\cos \theta)
\]

for \( m \geq 0 \), and 

\[
C_{lm} (\theta, \phi) = (-1)^m C_{lm} (\theta, \phi) \quad \text{for} \quad m < 0.
\]

[75] The Langevin capture model assumes that the system is always captured in the complex when the kinetic energy overcomes the centrifugal barrier and, once the complex is formed, that it always leads to products. The former means that capture probabilities, \( A_l^m (E) \), change from zero to one when overcoming the centrifugal barrier. The latter identifies capture probabilities with reaction probabilities, thus assuming the upper limit for the proportion of complexes which decompose into products.