Non-adiabatic couplings and dynamics in proton transfer reactions of \( \text{H}^n+ \) systems: Application to \( \text{H}_2^+ + \text{H}_2 \rightarrow \text{H}^+ + \text{H}_3^+ \) collisions

Cristina Sanz-Sanz, Alfredo Aguado, Octavio Roncero, and Fedor Naumkin

Citation: The Journal of Chemical Physics 143, 234303 (2015); doi: 10.1063/1.4937138
View online: http://dx.doi.org/10.1063/1.4937138
View Table of Contents: http://scitation.aip.org/content/aip/journal/jcp/143/23?ver=pdfcov
Published by the AIP Publishing

Articles you may be interested in
Radiative charge transfer in \( \text{He}^+ + \text{H}_2 \) collisions in the milli- to nano-electron-volt range: A theoretical study within state-to-state and optical potential approaches

Nuclear spin dependence of the reaction of \( \text{H}_3^+ \) with \( \text{H}_2 \). I. Kinetics and modeling

Influence of nuclear exchange on nonadiabatic electron processes in \( \text{H}^+ + \text{H}_2 \) collisions

A time-dependent wave packet quantum scattering study of the reaction \( \text{H}_2 \text{D}^+ ( v = 0 – 3 ; j = 1 ) + \text{He} \rightarrow \text{He}^+ + \text{H} + \text{H}_2 \text{D} + + \text{D} ( \text{H} ) \)

Exact quantum dynamics study of the \( \text{O} + + \text{H}_2 ( v = 0 , j = 0 ) \rightarrow \text{O} \text{H} + + \text{H} \) ion-molecule reaction and comparison with quasiclassical trajectory calculations
Non-adiabatic couplings and dynamics in proton transfer reactions of $H_3^+$ systems: Application to $H_2 + H_2^+ \rightarrow H + H_3^+$ collisions

Cristina Sanz-Sanz,1 Alfredo Aguado,1 Octavio Roncero,2,a) and Fedor Naumkin3,b)
1Departamento de Química Física Aplicada (UAM), Unidad Asociada IFF-CSIC, Facultad de Ciencias C-XIV, Universidad Autónoma de Madrid, 28049 Madrid, Spain
2Instituto de Física Fundamental, CSIC, C/Serrano, 123, 28006 Madrid, Spain
3Faculty of Science, UOIT, Oshawa, Ontario L1H 7K4, Canada

(Received 7 October 2015; accepted 23 November 2015; published online 17 December 2015)

Analytical derivatives and non-adiabatic coupling matrix elements are derived for $H_3^+$ systems ($n = 3–5$). The method uses a generalized Hellmann-Feynman theorem applied to a multi-state description based on diatomics-in-molecules (for $H_3^+$) or triatomics-in-molecules (for $H_3^+$ and $H_3^+$) formalisms, corrected with a permutationally invariant many-body term to get high accuracy. The analytical non-adiabatic coupling matrix elements are compared with ab initio calculations performed at multi-reference configuration interaction level. These magnitudes are used to calculate $H_2(v' = 0, j' = 0) + H_2(v, j = 0)$ collisions, to determine the effect of electronic transitions using a molecular dynamics method with electronic transitions. Cross sections for several initial vibrational states of $H_3^+$ are calculated and compared with the available experimental data, yielding an excellent agreement. The effect of vibrational excitation of $H_3^+$ reactant and its relation with non-adiabatic processes are discussed. Also, the behavior at low collisional energies, in the $1 \text{ meV}-0.1 \text{ eV}$ interval, of interest in astrophysical environments, is discussed in terms of the long range behaviour of the interaction potential which is properly described within the triatomics-in-molecules formalism. © 2015 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4937138]

I. INTRODUCTION

In the interstellar medium (ISM), hydrogen is by far the most abundant atom. $H_2$ is the most abundant molecule and is formed on the surface of dust particles by successive adsorption of H atoms1 and several gas phase processes such as radiative association of H atoms. Another gas phase reaction of interest is

$$H + H_2^+ \rightarrow H_2 + H^+, \quad (1)$$

which is important in the formation of $H_2$ in the early universe. Once $H_2$ is formed, the chemistry in space starts by the formation of the first hydrides. In cold clouds, the most abundant ion is $H_3^+$ which is formed in the collision

$$H_2 + H_2^+ \rightarrow H_3^+ + H, \quad (2)$$

which is exothermic by 1.2 eV and therefore very efficient. This explains why $H_3^+$, formed by the ionization of $H_2$ by cosmic rays, is not very abundant. On the contrary, when $H_3^+$ collides with molecular hydrogen, a proton exchange is produced as

$$H_2 + H_3^+ \rightarrow H_3^+ + H_2, \quad (3)$$

which does not destroy $H_3^+$, but simply transforms it. This last reaction is the main source of ortho/para-conversion of $H_3^+$ and when involving deuterium is the main source of deuterated $H_3^+$.

$H_3^+$ is involved in many processes in astrochemistry with other atoms and polyatomic molecules, electrons and hydrogen, and protons and $H^-$. These processes are described in details in many review articles such as Refs. 2–6 and there are special issues devoted to $H_3^+$.7 Many of these processes are very efficient because they are highly exothermic. In cold molecular clouds, many reactive species are adsorbed on icy particles, and the densities of electrons and cosmic rays are considerably reduced, making possible slow processes such as deuteration through Eq. (3) exchange reaction. Deuterated species are favored by their lower zero-point energy (ZPE), thus shifting the equilibrium in gas phase reactions such as that of Eq. (3),8,9 $H_2^+$, as the universal protonator,2,10–12 plays a central role in the interstellar chemistry, since it is at the origin of the formation of the first hydrides, including deuterated species.

In the reactive cycle of pure hydrogen species of Eqs. (1)–(3), there are several electronic states, corresponding to electron holes in each of the equivalent hydrogen atoms. This makes necessary to use a multi-state description of the potential energy surfaces (PESs), such as the diatomics-in-molecules (DIM) frequently used to describe these systems.13–16 The DIM description is however not enough to describe accurately $H_3^+$ fragments, and several many-body (MB) terms have to be added. For $H_3^+$, this was done for ground singlet17 and triplet18 states by adding a three-body term to the DIM PES so that an accuracy of about 1-5 cm$^{-1}$ was reached on global surfaces designed to describe the reactive collisions. This was later improved by adding the long range (LR) terms, charge-induced dipole and charge-quadrupole, on the diagonal

---

a)Electronic address: octavio.roncero@csic.es
b)On sabbatical at IFF-CSIC.
terms of the DIM Hamiltonian matrix. Another approach was adopted adding three-body terms on the non-diagonal matrix elements of the DIM matrix. For $H_2^+$ and $H_3^+$, the DIM approach does not describe the different fragmentation channels because many of them correspond to triatomic $H_3^+$. In order to describe these channels, a generalization of the DIM approach was used, the triatomics-in-molecules (TRIM) approach, which is essentially exact at the asymptotes but not in the interaction region. To get the desired accuracy, many-body terms were added to describe the ground state. With this approach, accurate 6 and 9 dimension PESs were obtained for $H_2^+$ and $H_3^+$.

In this work, we generalize these approaches to describe several electronic states of $H_2^+$, with special emphasis on analytic non-adiabatic coupling matrix elements (NACMEs) and potential derivatives, using the generalized Hellmann-Feynman theorem. The analytical derivatives of the ground PES are tested with quasi-classical trajectories (QCTs) on $H_2 + H^+$, $H_2 + H_2^+$, and $H_2 + H_3^+$ reactive collisions, getting a significant speedup and better accuracy as compared to the numerical ones previously used for $H_2^+$ and $H_3^+$.

For $H_2^+$ with a deep insertion well, the reaction mechanism is essentially statistical, and QCTs underestimate the cross section because this method is unable to describe tunneling for high total angular momenta. For $H_3^+$, the exchange reaction of Eq. (3) has been treated with statistical approaches, using asymptotic long range interaction potentials between reactants. When considering the full potential of Ref. 22 and 26–28 for high total angular momenta. For $H_3^+$, the exchange reaction of Eq. (3) has been treated with statistical approaches, using asymptotic long range interaction potentials between reactants. When considering the full potential of Ref. 22 and 26–28 for high total angular momenta.

In this work, we focus on the non-adiabatic dynamics of the reaction of Eq. (2) using the classical molecular dynamics with quantum transitions (MDQTs) of Tully. The MDQT is a variant of the commonly known surface hopping applied to study the non-adiabatic reactive $H^+ + H_2$ collisions by Krenos et al. and supported by the experiments of Ochs and Teloy. In this reaction, there is a curve crossing in the entrance channel very close to the ZPE of the reactants, and here, we shall analyze the effect of non-adiabatic transitions as a function of translational energy and vibrational excitation of the $H_3^+$ reactants. The results are compared with the available experimental data and with the previous surface hopping results. These last results were obtained with DIM PESS, which do not include long-range interaction terms and are not accurate to describe $H_2^+$ fragments. In the present approach, these two problems are solved.

The organization of the paper is as follows. In Section II, the theoretical methods used are presented: the generalized Hellmann-Feynman approach used to obtain analytical potential derivatives and non-adiabatic matrix elements applied to the TRIM+many body (TRIM+MB) method. Also, some details of the numerical implementation of the MDQT developed in this work will be briefly described. In Section III, the results obtained for the reaction of Eq. (2) will be shown and discussed. Finally, Section IV is devoted to extract some conclusions.

II. THEORETICAL METHODS

A. Generalized Hellmann-Feynman theorem

We will consider $N$ adiabatic eigenstates, $\phi_n^a$, which are obtained as solution of the electronic Schrödinger equations

$$H(R)\phi_n^a = W_n(R)\phi_n^a,$$

where $W_n$ is the electronic eigenvalue at a nuclear configuration given by the nuclear coordinates $R$. These eigenfunctions are represented in a basis set, $\varphi_m$, as

$$\phi_n^a = \sum_m T_m \varphi_m^d.$$

This basis set, hereafter called diabatic, does not depend on the nuclear configuration $R$ and is assumed to be complete and orthonormal. The procedure can be easily generalized to the non-orthonormal case. In this diabatic basis, the Hamiltonian operator $H$ takes a matricial form of elements $H_{mn}^d$.

Differentiating Eq. (4) with respect to $\alpha \equiv R$, a component of the hypervector determining the nuclear configuration,

$$\frac{\partial H}{\partial \alpha} \phi_n^a + H \frac{\partial \phi_n^a}{\partial \alpha} = \frac{\partial W_n}{\partial \alpha} \phi_n^a + W_n \frac{\partial \phi_n^a}{\partial \alpha},$$

and premultiplying by $\phi_n^a$ and integrating over electronic coordinates, a generalized version of the Hellmann-Feynman theorem is obtained as

$$\langle \phi_n^a | \frac{\partial H}{\partial \alpha} | \phi_n^a \rangle = (W_n - W_n) \langle \phi_n^a | \frac{\partial \phi_n^a}{\partial \alpha} \rangle + \delta_{n,n'} \frac{\partial W_n}{\partial \alpha},$$

where the NACMEs, $\langle \phi_n^a | \frac{\partial \phi_n^a}{\partial \alpha} \rangle$, are non-diagonal.

To evaluate the derivatives of the electronic Hamiltonian it is convenient to represent the adiabatic eigenfunctions in the diabatic basis, so that using $\frac{\partial \phi_n^a}{\partial \alpha} = 0$, two cases can be distinguished as the following:

1. Diagonal with $n = n'$, in which NACMEs are zero, giving the derivative of the adiabatic potential as

$$\frac{\partial W_n}{\partial \alpha} = \sum_{mn'} (T_m^a)^d \frac{\partial H_{m'n}}{\partial \alpha}.$$

An equivalent formalism was previously employed, in particular, for obtaining derivatives of DIM potential energy surfaces of $Rg_n^a$ (Rg = He, Ne, Ar) and $Ar_n^a$ used further for their geometry optimizations.

2. Non-diagonal with $n \neq n'$, to evaluate the NACME as

$$\langle \phi_n^a | \frac{\partial \phi_n^a}{\partial \alpha} | \phi_n^a \rangle = \delta_{n,n'} \frac{\partial W_n}{\partial \alpha} = \frac{1}{W_n - W_{n'}} \sum_{mn'm} (T_m^a)^d \frac{\partial H_{m'n}}{\partial \alpha}.$$
B. TRIM+MB method for $H^+_3$

In this work, we use the DIM and/or TRIM representation, improved by including many-body interactions, generally called TRIM+MB, described in detail previously.\textsuperscript{21,22}

The basis set used is an antisymmetrized product of an orbital and a spin part, $\varphi_m = \mathcal{A} \psi_m \mathcal{S}_f$, with $m = i l$, corresponding to linear combinations of Slater determinants or configurations, $\mathcal{A}$ being the anti-symmetrization operator. When applied to $H^+_3$, we use for the orbital part the minimum basis set of 1s orbitals in each hydrogen atom, which for the $H^+_4$ case is written as

$$\phi_i = s_i(1) s_j(2) s_k(3)$$

(10)

corresponding to an electron hole on nuclei $i$, and one electron in each 1s function of the rest of nuclei $(i,j,k,l = 1,2,\ldots$ cyclic and $j = i + 1, k = j + 1, l = k + 1)$.

We use a complete set of spin functions for 2-4 electrons to describe $H^+_3(1^1A', 1^3A'), H^+_4(1^2A'),$ and $H^+_5(1^2A)$. For $H^+_3$ either one singlet or one triplet function is needed. However, for $H^+_4(2^2A)$ and $H^+_5(1^2A)$, two spin functions are needed to describe the doublet and singlet states, respectively. Thus, the total basis is composed by 3, 8, and 10 functions for $H^+_3$, $H^+_4$, and $H^+_5$, respectively.

The electronic Hamiltonian matrix is partitioned as

$$H = H^{TRIM} + H^{MB},$$

(11)

where $H^{TRIM}$ is the TRIM electronic matrix (DIM matrix for $H^+_3$), and $H^{MB}$ is the many-body term matrix added to get the desired accuracy.

$H^{MB}$ matrix elements are constructed\textsuperscript{15,46} as permutationally invariant linear combinations of products of Rydberg polynomials, depending on the internuclear distances $R_{ij}$ between atoms $i$ and $j$. The parameters of these polynomials are fitted to minimize the difference between the energies obtained in this representation and \textit{ab initio} points calculated at the higher accuracy. In the present implementation, $H^{MB}$ is a diagonal matrix in which all $H_{mm}$ terms are equal, corresponding to many-body terms added to reproduce the ground state of each of the systems considered.

The $H^{TRIM}$ is the total electronic Hamiltonian rewritten as a sum of in triatomic and diatomic fragment terms, and is formally exact. In the case of $H^+_4$ for the electronic configuration corresponding to an electron hole at nuclei $i$, it can be written as\textsuperscript{21}

$$H^{TRIM}_i = \sum_{n,i,o,n} H^+_{ino}(n-i,o-i) - \sum_{p>i} H^+_{ip}(p-i)$$

(12)

(please note in Eq. (4) of Ref. 21 that the sign of the monoatomic $\hat{H}^+_ip(p-i)$ Hamiltonians, the second term in the rhs, is wrong and should be negative). For $H^+_5$, it is written as\textsuperscript{22}

$$H^{TRIM}_i = \sum_{n,i,o,n} H^+_{ino}(n-i,o-i) - 2 \sum_{p>i} H^+_{ip}(p-i).$$

(13)

In these expressions, $H^+_{ip}(p-i)$ are monoatomic Hamiltonians of $H^+_3$ fragments and $H^+_{ino}(n-i,o-i)$ are bielectronic Hamiltonians (for electrons $n-i,o-i$) describing $H^+_5$ for the $ino$ nuclei. The $i,n,o,p$ indexes are defined in a cyclic way and runs up to 4 and 5, for $H^+_4$ and $H^+_5$, respectively.

The evaluation of the matrix elements of $\hat{H}^+_ip(p-i)$ describing $H^+_3$ is simple, since they can be directly deduced from \textit{ab initio} points obtained for few electronic states, the ground state on each symmetry box (see the Appendix of Ref. 22). The usual method is to fit these points to analytical functions, allowing a simple way to evaluate the derivatives as a function of internuclear distances. This is the same as it is usually done in the DIM approach.

The matrix elements of the triatomic Hamiltonians,

$$\langle \phi_i | \hat{H}^+_{ino}(n-i,o-i) | \phi_i \rangle,$$

(14)

correspond to diagonal and non-diagonal matrix elements involving linear combination of spin adapted diabatic matrices of $H^+_3$, of the first three electronic states of singlet and triplet symmetry. A good description of these triatomic matrix elements is what allows an accurate description of $H^+_3$ and $H^+_5$ systems, specially at long distances needed to properly describe collisions at low energies. For the evaluation of the matrix elements of the triatomic Hamiltonian, we use a generalization of DIM+MB method,\textsuperscript{17} which is formally analogue to the TRIM and which is briefly described below.

C. DIM+MB method for $H^+_3$

The electronic Hamiltonian in the DIM approach\textsuperscript{13-16} is expressed as a sum of atomic and diatomic fragment terms, and for $H^+_3$, it is written as\textsuperscript{22}

$$H^{DIM}_i = \sum_{n>i,o>n} \hat{H}_{ino}(n-i,o-i) + \sum_{p>i} \hat{H}_{ip}(p-i),$$

(15)

where $\hat{H}_{ino}(n-i,o-i)$ are the bielectronic Hamiltonians of $H_2$. This Hamiltonian is formally exact. The approximation is in the use of a small basis set that introduces inaccuracies. To improve the accuracy, 3-body terms are added, $H^{MB} \equiv H^{3B}$, as indicated in Eq. (11). Thus, the triatomic electronic Hamiltonians involved in Eqs. (12) and (13) can be rewritten as

$$\hat{H}^+_{ino}(n-i,o-i) = H^{DIM}_i(n-i,o-i) + H^{3B},$$

(16)

The triatomic DIM 3 x 3 matrices of $H^+_3$ are given by Ref. 17 for the singlet and by Ref. 47 for the triplet state. The $H^{3B}$ here is also considered as a diagonal matrix, whose elements are all equal, as described above.

In the $H^{3B}$, the long range interactions are included, describing the interaction of the $H^+$ with the induced electric dipole and quadrupole of $H_2$,

$$V_{H^+\cdot-H}(R,r) = Q_2(r)P_2(\cos \theta)R^{-3} - \left[ \frac{1}{2} a_0 + \frac{1}{3} (a_\parallel - a_\perp) P_2(\cos \theta) \right] R^{-4},$$

(17)

where $a_0, a_\parallel, a_\perp$ are the average, parallel, and perpendicular polarizabilities and depend on the internuclear distance $r$ and $\cos \theta = \mathbf{R} \cdot \mathbf{r}/R$. The interaction of $H^+_3$ with $H$ is considered as two $H^+ - H$ interactions,

$$V_{H^+\cdot-H}(R) = -\frac{9}{4} R^{-4} - \frac{15}{2} R^{-6},$$

(18)

which are already included in the mono-electronic terms that correspond to $H^+_3$ fragments, $H^+_{ip}(p-i)$.\textsuperscript{19} Being included in
the diatomic terms, the $H^+_2 + H$ long range terms are included in all the $H^+ + H_2$ asymptotes, while it should not and has to be subtracted properly. Moreover, there are three equivalent rearrangement channels all tending to the same long range interactions so that it cannot be repeated either.

Let us consider a triangular configuration characterized by the internuclear distances $R_i = P_{i+2} - P_{i+1}$ ($i = 1, 2, 3$ cyclic), where the charge is in atom $i$ and $P_{i+1}$ is the position vector of atom $i + 1$. The weighting function $w_i = \frac{e^{-2R_i^2}}{\sum_j e^{-2R_j^2}}$ is defined to include the LR interaction as

$$V_{12}^L(R_1, R_{1+2}) = w_i \left\{ V^{H_2+H}(R, R_i) - V^{H^+H}(R_1) - V^{H^+H}(R_{1+2}) \right\},$$

(19)

where the $R = [R_{i+1} + R_{i+2}] / 2$ is the corresponding Jacobi vector. This form is slightly different to that used previously in Ref. 19 and has the advantage that its derivatives are continuous. Also, the Jacobi vector $R$ can only reach the value of zero when the corresponding internuclear $R_i$ distance is very long, and in this situation, the weight $W_i$ becomes zero, i.e., this function acts as a damping function.

Including the long range term as a contribution in the $H^{1B}$ allows to describe very accurately the long range behavior in $H^+_3$. In addition, in the TRIM+MB method, several triatomic terms are considered, and these long range terms are involved in each of them, thus allowing to describe the delocalization of the charge in $H^+_2$ or $H^+_3$ when accounting for the $H^+_2 + H_2$ or $H^+_3 + H_2$ long range interactions.

For example, in Fig. 1, we show the long range dependence of the $H_2 + H^+_2$ interaction potential, for distances $R > 10$ a.u. For $R > 20$ a.u., the TRIM+MB potential fits very well the long range potential of Eq. (17). However, for $10 < R < 20$ a.u., the potential is better fitted using $C/R^{16}$. This is due to the contribution of two triatomic terms, each one with a charge in one of the two atoms of $H^+_2$. At these intermediate distances, the potential does not yet reach its long range behavior, what has some influence in the collision dynamics, as described below.

The error of the potential in $10 < R < 20$ a.u. region is of the order of 1 cm$^{-1}$, when compared with the $ab$ initio points. In some configurations, the $ab$ initio points are better fitted by Eq. (17). Thus, we conclude that some more work has to be done to check the influence on the dynamics described below. Some work in this direction is currently under way.

The derivative of all the $\tilde{H}^+_{ino}(n-i,o-i)$ and $\tilde{H}^+_{ip}(p-i)$ can be evaluated analytically, and hence those of the full diabatic matrices, $\frac{\partial H^H_{ino}}{\partial \alpha}$, required in Eqs. (8) and (9). These analytical derivatives were implemented to evaluate the derivative of the potential of the ground states of $H^+_n$ and compared with the numerical ones, giving excellent results. In addition, they were used in classical trajectory calculations, giving lower errors in conserving the total energy than the numerical ones.

The minimum energy paths (MEPs) of the first states of $H^+_3$, $H^+_4$, and $H^+_5$ are shown in Fig. 2. For $H^+_4$ and $H^+_5$, the ground state potential crosses with the first excited state at intermediate distances, due to the charge transfer between the fragments. The three codes, for $n = 3–5$, are available upon request.

**D. MDQT**

The non-adiabatic dynamics is studied in the adiabatic representation with the MDQT method as developed and described by Tully. Here, only some details on the numerical implementation made in this work are described.

An electronic wave-packet is expanded as $\Psi(R|t) = \sum_{\eta} \phi_{\eta}(t) \phi_{\eta}^*$, with $\phi_{\eta}(t = 0) = \delta_{\eta n}$ determining the initial conditions. Its evolution in time is governed by

$$i\hbar \dot{\phi}_{\eta}(t) = \sum_{kl} \left[ W_{\eta k} \delta_{nl} - i\hbar R \cdot \dot{d}_{kl} \right] \phi_{\eta}(t).$$

(20)

These coupled equations are integrated in time using a split operator propagator, along a classical trajectory $R(t)$. To avoid numerical problems at conical intersections, where the non-adiabatic couplings diverge, the NACMEs are truncated so that $-i\hbar R \cdot \dot{d}_{kl} \leq E_{max}$. In these expressions, $R$ is the multidimensional velocity vector, corresponding to the $3 \times N$

**FIG. 1.** Interaction potential for the $H_2$ ($r_1 = 1.4$ a.u.) + $H^+_1$ ($r_2 = 2$ a.u.) interaction as a function of the distance $R$ between their centers of mass. The two molecules are in the plane, with $r_1$ and $r_2$ being parallel and perpendicular to $R$. The internuclear distances $r_1$ and $r_2$ approximately correspond to their equilibrium distances.

**FIG. 2.** Minimum energy paths (in eV) of the lowest electronic states of $H^+_3$, $H^+_4$, and $H^+_5$. In each system, a different zero of energy is chosen to plot all of them together.
Cartesian coordinates of the position vectors of the $N$ atoms.

The classical trajectories $\mathbf{R}(t)$ are obtained solving the Hamilton equations of motion in the $6N$ generalized coordinates using a step adaptive Adams-Bashforth-Moulton predictor-corrector method. At each time step, the switching probability, $g_{i,f}$, from electronic state $i$ to each $f$ is calculated as

$$g_{i,f} = \frac{\Delta t}{|\mathbf{C}|^2} \left[ 2 \Re(\mathbf{c}_i^* \cdot \mathbf{R} \cdot \mathbf{d}_f) \right],$$

(21)

so that the sum of all these probabilities is 1. These probabilities are then considered consecutively in a vector as

$$\mathbf{g}_i = [g_{i,1}, g_{i,2}, \ldots, g_{i,f}, \ldots, g_{i,N}],$$

where $\mathbf{g}_i$ is the probability vector of all transitions from state $i$. The sum of all these probabilities is 1.

In order to get the reaction cross section for the reaction

$$\text{H}_2^+(v, j = 0) + \text{H}_2(v' = 0) \rightarrow \text{H}_3^+ + \text{H},$$

(24)

a Monte Carlo sampling of initial conditions is performed. The method used is completely analogous to that introduced by Karplus et al. In order to distinguish between $\text{H}_2$ and $\text{H}_3^+$ in a single electronic state, we take the inner classical turning point for $\text{H}_2$ and the outer turning point for $\text{H}_3^+$ in their respective initial vibrational states, for a reason which will become clear below. More than 10^5 trajectories are run for each energy and initial state. The energy conservation in all of them is better than 0.001 cm$^{-1}$.

In order to get an approximate relationship between the non-adiabatic transitions and the non-adiabatic matrix elements, we shall use the inverse of the Massey parameter defined as

$$M_{ij}^{-1} = \frac{\mathbf{R} \cdot \mathbf{d}_{ij}}{2(W_i - W_j)},$$

(25)

III. APPLICATION TO H$_3^+$

A. Non-adiabatic couplings

As shown in Fig. 2, the first two electronic states are separated by $\approx 0.5$ eV in the $H_2 + H_2$ entrance channel, which is close to the sum of the vibrational zero-point energies of the reactants. These two states cross at configurations where the two diatomic molecules have the same internuclear distances, as can be seen in the top panel of Fig. 3. In a pure adiabatic representation, the positive charge is not located in any of the two diatomic fragments, but it is delocalized. When the two internuclear distances are the same, the charge has an equal probability of being in either one of the two first lower electronic states giving rise to two degenerate states, i.e., to a crossing.

In the middle panels of Fig. 3, some cuts of the potential energy for the first two adiabatic states are shown as a function of the internuclear distance $r_1$ of the first diatomic fragment. The other diatom is frozen for three values, $r_2 = 1.4, 1.7$, and $2$ a.u. All the other coordinates are frozen, with the two bonds parallel in a plane. Three different energies are plotted: the $ab$ initio points (calculated at Multi-Reference Configuration Interaction (MRCI) level using the MOLPRO package as described in Ref. 21), results obtained with the TRIM+MB (labeled TRIM) described here, and finally the energies obtained with the DIM approach. The DIM energies are rather good in the $H_2^+ + H_2$ entrance channel but already show some disagreements with the $ab$ initio ones. The TRIM+MB results for the ground state are excellent, since the MB term is adjusted for that, showing some discrepancies for the excited electronic state. Nevertheless, for the collision energies of interest, below 1 eV, these results are considered to be accurate enough, giving a rather good description of the crossings.

The NACMEs obtained with these three methods are shown in the bottom panels of Fig. 3. Note that the TRIM+MB method gives DIM results when removing the MB terms in the triatomic fragments and in the $H_2^+$ potential. The $ab$ initio results are obtained with the MOLPRO package, using a first order difference method with an interval of 0.01 a.u. For the square geometry, $r_2 = 2$ a.u., the crossings and the NACME obtained with the three methods are in very good agreement. As $r_2$ distance decreases, TRIM and DIM results remain very close but their differences with respect to the $ab$ initio results increase. These differences are considered to be rather small and are neglected here.

The TRIM NACMEs are only slightly better as compared to the DIM results. The reason is that the adiabatic wave-functions are obtained from the approximated $H^{TRIM}$ Hamiltonian, in whose triatomic terms and the diagonal $H^{BB}$ term are all the same. Under this condition, the DIM and TRIM NACMEs are very similar. The inclusion of the “exact” $H^{BB}$ terms in the $H^{TRIM}$ method would improve the description of the crossings and the accuracy of the NACME. Work in this direction is in progress.

Summarising, the TRIM+MB improves significantly the accuracy of the ground adiabatic state, including the long-range interactions and triatomic fragments, as described previously. In addition, it improves the position of the curve crossing in the entrance channel with respect to the DIM approach. The NACMEs of the TRIM method with the approximations made so far are very similar to those obtained with the DIM approach, and both are pretty similar to the $ab$ initio results.

These results are analogous to all the geometries and distances analyzed in the entrance channel. In the $H + H_2^+$ products channels, the ground and excited states separate, and non-adiabatic effects become negligible. We therefore conclude that the present approach is a significant improvement with respect to the previous DIM method used to study the non-adiabatic dynamics in this system.
FIG. 3. Top panel: Energy diagram of the lowest electronic states of H$_4^+$ showing curve crossings as a result of charge transfers between equivalent subunits. Middle panels show the potential energy curves and bottom panels the NACME, obtained for two parallel H$_2$ fragments in a plane at fixed values of $R = 2$ a.u. (the distance between the two centers of mass) and $r_2 = 2$ a.u. and by varying the internuclear distance of the first H$_2$ subunit, $r_1$. The NACME is calculated in Cartesian coordinates, and here, only one is shown, along the z component of $r_1$. The DIM, TRIM, and MRCI label the method used to obtain the energies or NACMEs.

B. Reactive collision cross sections

This reaction has 10 rearrangement channels

\[ AB^+ + CD \rightarrow \begin{array}{cccc}
AB^+ + CD & \text{inelastic} \\
\rightarrow AB + CD^+ & \text{electron transfer} \\
\rightarrow (A + BCD^+) & \text{proton hop} \\
\rightarrow (B + ACD^+) & \text{atom hop} \\
\rightarrow (ABC^+ + D) & \text{exchange}.
\end{array} \]

The electron transfer channel is included within the first adiabatic electronic state, as discussed later, in which the charge is in either of the two fragments and can only be distinguished in the dynamical calculations by the vibrational frequency when the distance between the two diatomic fragments is long.

Proton hop and atom hop mechanisms both yield to H$_3^+$ products and have relatively similar cross sections, as shown in Fig. 4. The proton hop mechanism has a slightly higher cross section, probably because the H$_2^+$ fragment with a larger equilibrium distance is easier to brake.

The exchange mechanism is about 100 times lower. This mechanism involves two atoms or proton hops or a simultaneous exchange. This reaction is rather direct, and the reaction occurs without giving time to a second atom/proton transfer. This could explain why the exchange mechanisms present lower cross sections.

The proton and atom hop mechanisms cannot be separated, and only two products can be distinguished, H$_2^+$ + H$_2$ and H$_3^+$ + H, because all the hydrogen atoms are

FIG. 4. Cross sections (in Å$^2$) for the H$_2(v'=0,j'=0) + H_3^+(v=0,j=0)$ collisions for the different arrangement channels indicated in Eq. (26) as calculated here with the MDQT method.
indistinguishable. This is a fundamental principle of quantum mechanics, and in order to distinguish among some channels, it is necessary to substitute one or several hydrogen atoms by an isotope.

In this work, we present MDQT calculations for the H₂(v′ = 0, j′ = 0) + H₂(ν, j = 0) reactions, always keeping the reactants in their ground rovibrational state, except in Section III C, where the effect of the initial vibrational excitation of H₂(ν) is analyzed. Unless otherwise indicated, the first two adiabatic electronic states have been included.

In Fig. 5, we compare the total reaction probabilities, the sum of proton hop and atom hop, with previous experimental and theoretical results. In all cases, the agreement is very good. The merged beam results of Glrewinkel-Meyer and Gerlich were obtained with H₂⁺ ions produced in a storage ion source. The relative error is due to the calibration of the target density and, therefore, constant over the energy range shown. Thus, our MDQT results are all within the experimental error bars.

The previous results of Eaker and Schatz for collision energies higher than 0.3 eV are also in good agreement with the results presented here. Those results were obtained using the DIM potential of Stine and Muckerman, using also a variation of a surface hopping method. This indicates that at this intermediate energies, the DIM approach is rather good in describing the total reaction cross section. The quantum mechanical results of Baer and Ng are lower than the present results, and this may be explained by their use of a lower dimensional model needed to perform their calculations.

The Langevin model gives results significantly lower than the results obtained here, which means that the interaction potential averaged over the H₂(j′ = 0) rotation does not vary as −α/2R². At the collision energies of 1, 10, and 50 meV, the maximum impact parameter for the reaction is found to be 30, 15, and 10 a.u., respectively. As discussed above, for these “intermediate” distances, the averaged potential does not follow the C/R⁴ long range behaviour but it is better fitted by adding a −C/R⁴ term (see Eq. 1). In this reaction, once the H₂⁺ + H product channels are reached, there is no return back to the entrance channel, and the capture models are then applicable here. According to these models, the cross section takes the form

\[ \sigma(E) = \pi q(s) (C/E)^{2/s} \]  

with \( q(s) = \frac{s/2}{(s-2)^{(s-2)/s}}. \)

The MDQT cross section of Fig. 5 is perfectly fitted by this expression for E < 0.1 eV, with C = 11.51 Å² eV and s = 4.16. This value of s is in agreement with the dependence of the potential in the region 10 < R < 20 a.u. To get the Langevin results, much lower collision energies have to be considered, involving longer distances, at which the average interaction potential varies as \( R^{-1} \). This can be explained knowing that in this system, the positive charge is not punctual, but it is distributed in H₂⁺.

For collisions of H₂ with atomic cations, it is expected that at these energies of 1-100 meV, the asymptotic Langevin limit is already achieved, since the charge can be considered punctual. This is, for example, the case of H₂ + O⁺ in experimental and as well as in theoretical simulations. However, to check this finding in H₂ + H₂⁺ collisions, a better fit of the 10 < R < 20 a.u. region, with much lower errors, should be obtained.

C. Effect of initial vibrational excitation

The possibility of generating H₂⁺ in different vibrational states made possible many experimental and theoretical studies for its effect. Here, we analyze the effect of H₂(ν = 0, 1, 2, 3) initial vibrational states in the formation of H₂⁺ in the 0-1 eV translational energy range, as shown in Fig. 6. The behaviour of all of them is very similar, being nearly parallel as a function of energy, clearly demonstrating that the capture model is valid in all the cases.

The small decrease of the cross section of the H₂⁺ formation as ν increases is not explained by the increase of the exchange cross section, in the right panel of Fig. 6. When the two mechanisms are added, the resulting plot is very close to that shown in the left panel. This exchange

![Graphs showing cross sections for reactions](image-url)
TABLE I. Experimental\textsuperscript{12} and simulated cross sections (Å\textsuperscript{2}) for the \( \text{H}_2(\nu'=0, j'=0) + \text{H}_2^*(\nu, j=0) \rightarrow \text{H}_2^* + \text{H} \) reaction as a function of the vibrational excitation of \( \text{H}_2^* \) for 0.1 eV of translational energy.

<table>
<thead>
<tr>
<th>( \nu )</th>
<th>Experiment\textsuperscript{12}</th>
<th>Simulated</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>51 ± 2</td>
<td>59.9</td>
</tr>
<tr>
<td>1</td>
<td>45 ± 3</td>
<td>59.9</td>
</tr>
<tr>
<td>2</td>
<td>46 ± 3</td>
<td>52.2</td>
</tr>
<tr>
<td>3</td>
<td>42 ± 3</td>
<td>49.4</td>
</tr>
</tbody>
</table>

mechanism has a much lower cross section, but its increase with \( \nu \) is rather important.

The experimental measurements indicate this trend for the formation of \( \text{H}_2^* \), as listed in Table I. The absolute value 51 ± 2 Å\textsuperscript{2} of the cross section of Koyano and Tanaka\textsuperscript{32} is lower at 0.1 eV than the present result of 59.9 Å\textsuperscript{2}. The experimental results of Glenewinkel-Meyer and Gerlich\textsuperscript{35} in Fig. 5 is also lower, being 48 ± 13 Å\textsuperscript{2} at 1.0 eV. The systematic deviation of the cross sections on the collision energy may be due to rovibrational excitation of the ions. However, the error bars in these last experiments were estimated to be larger, of the order of 15 Å\textsuperscript{2}, and the simulated values lie within these larger error bars. For this reason, we consider the present values rather good in describing the reaction cross section, noting that there is a qualitative agreement of the trends with increasing \( \nu \) between the experimental and theoretical values. In addition, the adequacy of the classical method used here should be tested by doing quantum dynamical simulations.

What is the origin of the decrease of the \( \text{H}_2^* \) formation cross section as vibrational excitation increases? One possibility is the non-adiabaticity due to the curve crossing with the first excited state, as shown in Fig. 3. To check this, we have performed the same calculations but including only the ground electronic state, i.e., pure QCT calculation, shown in Fig. 7. The QCT cross section shows a similar trend as compared to those shown in Fig. 6 obtained with 2 electronic states. However, the \( \text{H}_2^* \) formation cross sections for different \( \nu \)’s are closer and with a different ordering. This may be taken as an indication that the small differences observed in the left panel of Fig. 6 are due to non-adiabatic effects.

The MDQT and QCT exchange cross sections shown in the right panels of Figs. 6 and 7 are very similar for collision energies below 0.1 eV. However, for larger energies, the oscillations differ considerably, and these differences are attributed to non-adiabatic effects.

Experimentally, there are no results studying the effect of the neutral \( \text{H}_2(\nu') \) vibrational excitation. However, since the variations are due to non-adiabatic effects (as discussed in more detailed below), we do expect a similar qualitative behaviour when exciting either of the two diatomic reagents, \( \text{H}_2^* \) described above or \( \text{H}_2 \).

D. Non-adiabatic dynamics

The lowest two adiabatic states of \( \text{H}_2^* \) cross at relatively low energies in the \( \text{H}_2 + \text{H}_2^* \) entrance channel and then separate in the \( \text{H}_2^* + \text{H} \) products channel, as can be seen in the middle panel of Fig. 2. The origin of this crossing is that at asymptotic distances, there are two equivalent diabatic states, \( \text{H}_2 + \text{H}_2^* \) and \( \text{H}_2^* + \text{H}_2 \), which combine to form the adiabatic state in which the charge is delocalized between the two fragments. This situation is similar to that of \( \text{H}_2^* \), in which two diatomic states are obtained, of gerade and ungerade symmetries.

In the ground adiabatic state of \( \text{H}_2^* \), at long distances between \( \text{H}_2 \) and \( \text{H}_2^* \) center of masses, \( R \), when one of the diatomic fragments is frozen at the \( \text{H}_2 \) equilibrium distance, the other fragment presents a potential diabatic curve similar to \( \text{H}_2^* \), and vice versa. However, the adiabatic states present crossings with cusps as represented in Fig. 8. If the two fragments are kept in their ground vibrational state, the two \( \text{H}_2 \) and \( \text{H}_2^* \) internuclear distances never get the same value at the asymptote simply because their turning points are slightly separated. This is a classical effect, since the quantum wave-functions penetrate in the potential and overlap.

FIG. 7. Same as Fig. 6 but considering only the ground electronic state, thus being denoted as QCT results.

FIG. 8. Potential energy cuts obtained by freezing \( r_{34} \) internuclear distance to the equilibrium value of \( \text{H}_2^* (r_3 \approx 2 \text{ a.u.}) \) and \( r_{12} \) to the equilibrium value of \( \text{H}_2 (r_1 = 1.4 \text{ a.u.}) \) and varying the other.
When H$_2^+$ gets vibrationally excited, its inner classical turning point is shorter than the outer one of H$_2$, see Fig. 8. In this situation, even at long distances among the diatoms, $R$, the diatomic internuclear distances, $r_1$ and $r_2$, can get the same value. At this configuration, there is a crossing at which the non-adiabatic couplings, shown in Fig. 3, are large and may induce non-adiabatic transitions. As the vibrational excitation increases, the possibility that $r_1 = r_2$ increases and therefore the probability of electronic transitions.

All these are illustrated in Figs. 9 and 10, in which we show several quantities associated to two trajectories leading to H$_3^+$ products, one starting in H$_2^+(v = 0)$ and the other in H$_2^+(v = 2)$. The initial conditions for the two trajectories are very similar, except in the initial H$_2^+$ internuclear distance, $r_2$, corresponding to the outer turning points for $v = 0$ and 2, respectively. In Fig. 9, the two internuclear distances (in the second panel from the bottom) never cross for $R > 10$ a.u. (bottom panel), and as a consequence, there is no electronic transition. The quantum wave-packet (top panel) always remains in the ground electronic state, and the inverse Massey parameter, Eq. (25), only gets significant values for $R < 20$ a.u.

When starting in H$_2^+(v = 2)$, in Fig. 10, the situation changes dramatically. The amplitude of the H$_2^+$ vibration increases, and in this case, $r_1$ and $r_2$ may get the same value. For time <200 a.u., it can be seen that the red and blue curves exchange between them, i.e., an electron transfers between the two diatoms. This is already surprising since it happens at distances $R > 30$ a.u. and in one single adiabatic electronic state. Such exchange of charge occurs when $r_1 \approx r_2$.

When the distance $R$ decreases, and $r_1 \approx r_2$, the non-adiabatic couplings become larger, as illustrated by the inverse Massey parameter in the third panel from bottom in Fig. 10. This makes that electronic transitions may take place even at relatively long distances, $R \approx 30$ a.u., promoting the system to the first excited electronic state. In this state, the motions of the two diatoms become in perfect phase, as shown in the second panel of Fig. 10 between 200 < time < 300 a.u. This result is observed in all trajectories analyzed. The Massey parameter is always large, being able to induce a transition back to the ground state.

In reactive trajectories leading to H$_3^+$, the energy difference between the two electronic states in the products channel gets apart and the non-adiabatic couplings decrease. Thus, for these trajectories, the electronic transitions do not occur once the H$_3^+$ is formed. The four H$_3^+$ + H product channels are separated...
by a barrier of $\approx 0.25$ eV which hinders the exchange among them in low temperature $H_3^+$ + H collisions.\textsuperscript{57,58}

For those trajectories leading to diatomic fragments, either in the inelastic or the exchange channels, the two electronic states remain close and the coupling is large. Therefore, the maximum number of transitions happens for these kind of trajectories.

The behavior of the two trajectories shown in Figs. 9 and 10, with similar initial conditions, is quite analogous, both leading to the same $H_3^+$ product. A careful analysis of many other trajectories leads us to conclude that the trajectories do not vary very much due to electronic transitions, and this is the reason why the reaction cross section does not vary significantly. The most affected trajectories are those leading back to any $H_2^+$ + $H_2$ channel, and this explains why vibrational excitation affects more significantly this channel.

Once the $H_2^+$ is vibrationally excited, the number of electronic transitions increases with decreasing collision energy, simply because it takes longer time for the two fragments to approach and therefore, $r_1 = r_2$ occurs many times. For $v = 0$, there are no electronic transitions even at $E = 1$ eV, with energy enough to allow transitions to excited electronic states. Thus, the collision dynamics of $H_2(v = 0) + H_2^+(v = 0)$ can be studied on a single electronic state.

E. Isotopic effect

$H_3^+$ is the universal protonator in cold molecular clouds. It is also responsible for the high deuterium fractionation of many hydrides in space, much higher than the natural abundance of the D with respect to H. It is therefore important to analyze how the reaction changes whenever one deuterium substitutes one hydrogen atom in either $H_2$ or $H_3^+$ reactants. Here, we present results for the $H_2D^+$ system, where the D is initially either in the ion or the neutral. As demonstrated above, considering the ground rovibrational state of the two reagents, the dynamics proceeds in a single adiabatic surface, and the usual QCT approach is used.

The total cross sections to form triatomic products, either $H_3^+$ or $H_2D^+$, are compared in Fig. 11. The cross sections for the three reactions considered are very similar, with the same dependence on energy, which can be explained by the capture model of Eq. (27).\textsuperscript{54} There are only small differences at low and high energies.

In the two cases, there are three channels leading to $H_2D^+$ fragments and only one to $H_3^+$. Thus, assuming unbiased statistical scrambling, the cross section for the formation of the deuterated $H_2D^+$ is approximately 3 times that of $H_3^+$ in $H_2 + HD^+$ or $H_2^+ + HD$ collisions. The role of quantum effects, especially at low energies, may change this.

The differences arise when considering the possible mechanisms, i.e., the jump of neutral H or D or of a $H^+$ or a $D^+$, indicated in different channels of Eq. (26). These mechanisms are distinguished in Fig. 12 and are not equivalent when there is a deuterium. The channel which presents larger differences is that involving the hop of a deuterium, charged or neutral, for which at $E > 0.1$ eV is always smaller. This may be explained by an impulsive mechanism, in which heavier atoms are more difficult to deviate.

![FIG. 11. Total reaction cross sections for the isotopic variants: $H_2 + H_3^+(v = 0)$, $HD + H_3^+(v = 0)$, and $H_2 + HD^+(v = 0)$ collisions.](image1)

![FIG. 12. Cross section for the 4 different rearrangement channels leading to triatomic products, in Eq. (26) for the $HD + H_3^+(v = 0)$ and $H_2 + HD^+(v = 0)$ collisions.](image2)
For low collision energies, however, the situation changes. In HD + H$^+_2$ collisions, the 4 channels present similar cross sections, while for H$_2$ + HD$^+$ collisions, the hop of a proton always prevails, probably because it combines the lower mass with the most elongated bond.

IV. CONCLUSIONS

In this work, we have developed analytical potential derivatives and non-adiabatic couplings for H$^+_n$ systems for TRIM-MB multi-state model potentials, allowing the accurate description of H$^+_2$ fragments and the inclusion of long range interactions. This procedure has been applied to the study of H$_2$(v' = 0, j' = 0) + H$^+_2$(v, j = 0) reactive collisions, considering the effect of collision energy, vibrational excitation of H$^+_2$, and non-adiabatic and isotopic effects, using an implementation of the MDQT method of Tully. The calculations have been restricted to the ground rotational levels and therefore do only explore the charge-induced dipole term of the long range interactions. The calculated cross sections are in good agreement with the available experimental data and are slightly larger than those obtained with a Langevin model. The reason of that is that the effective average potential does not vary as $R^{-4}$ for distances between 10 and 20 a.u., because the charge is distributed in the H$^+_2$ molecule and cannot be centered at its center of mass.

There are small changes in the cross section when increasing the vibrational excitation of H$^+_2$ reactants, in agreement with the experimental data of Koyano and Tanaka, which are attributed to non-adiabatic transitions to the first excited electronic state. These electronic transitions occur in the entrance channel when the two internuclear distances of the two diatomic species (H$_2$ and H$^+_2$) are equal, leading to a charge transfer between them, but with rather low cross sections at the energies studied here.

We conclude that for energies below 1 eV and starting in the ground vibrational states of reactants, the non-adiabatic effects are negligible. Then, pure QCT calculations are performed to study the isotopic effects in the HD + H$^+_2$ and H$_2$ + HD$^+$ collisions. Again, the cross sections are governed by the long range interactions. The different mechanisms (proton or atom hops, H or D) are distinguished and the D or D$^+$ hop mechanism is found to be the one with lower cross sections at E > 0.1 eV.

ACKNOWLEDGMENTS

F.N. acknowledges UOIT for the support of a sabbatical stay at IFF-CSIC, Spain. We acknowledge the support of Ministerio de Economía y Competitividad (Spain), for Grant Nos. CSD2009-00038 and FIS2014-52172-C2. The calculations have been performed in CESGA and CSIC computing centers, which are acknowledged. We also thank the support from the European Research Council under the European Union’s Seventh Framework Programme (No. FP/2007-2013)/ERC Grant Agreement No. 610256 (NANOCOSMOS). We also acknowledge the COST Action No. CM1401 “Our Astrochemical History.”


