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MCTDH study of CH₄ dissociation on Ni(111)

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Abstract

We present results of quantum simulations using the multi-configuration time-dependent Hartree method for the dissociation of CH₄ on Ni(111) with a potential-energy surface based on density-function calculations. As varied coordinates we include the distance of the methane molecule to the surface, a C-H distance, and the orientation of the CH₄. Taking into account the latter coordinate decreases the dissociation probability. For the range of translational energies we studied, the dissociation occurs via tunneling.

Keywords: Alkanes; Computer simulations; Low index single crystal surfaces; Surface chemical reaction

1. Introduction

The dissociation of CH₄ on transition metals is very interesting for a number of reasons. In steam reforming to produce H₂ the dissociation is the rate limiting step [1]. More general, as methane is the simplest of all alkanes, its dissociation can be viewed as prototypical for C–H activation in catalysis. It is with this in mind that we started some years ago with density-functional theory (DFT) calculations of the dissociation on nickel. Another reason for our interest are a number of molecular beam experiments in which the dissociation probability was measured as a function of translational energy [2–8]. These experiments have contributed much to our understanding of the mechanism of the dissociation. Also some information has been obtained on the role the internal degrees of freedom play. However, our knowledge is still negligible compared to what we know about a well-studied reaction as H₂ dissociation on copper [9].

In a number of previous studies we have calculated the reaction rate constant of CH₄ dissociation on Ni(111) using DFT and transition-state theory (TST) [10–14]. These studies have shown that there is a late activation barrier. We calculated various heights of this barrier, the most accurate one, in our view, being 121 kJ·mol⁻¹. The barrier height could not be obtained reliably either from the experiment, as very different numerical values were obtained depending on the assumed reaction mechanism; precursor or not, tunneling or not. Although our calculations gave results consistent with experiment, they were unsatisfactory, because they yielded only thermally averaged dissociation probabilities, and could therefore not be compared directly with the more detailed molecular beam experiments.

Neither did our calculations yield much informa-
tion on the role of the internal vibrations. It was observed experimentally that vibrationally hot CH₄ dissociates more readily than cold CH₄, with the energy in the internal vibrations being about as effective as the translational energy in inducing dissociation. However, a more detailed assessment of the importance of the internal vibrations could not be made, because of the large number of internal vibrations. For these reasons we decided to do quantum simulations on the potential-energy surface (PES) that we obtained with our DFT calculations, thus obtaining a better assessment of the quality of these latter calculations.

The quantum simulations have been done using the multi-configuration time-dependent Hartree (MCTDH) method [15,16]. This is a rather new method which has been developed to deal with a large number of degrees of freedom and with large grids. The method has been successfully applied to gas phase reactions (photodissociation of NOCl [15], NO₂ [17], CH₃I [18,19] and ICN [20], vibrational predissociation of Cl₂Ne [21] and the hydrogen exchange reaction H₂ + H → H + H₂ [22]), but also to reactions at surfaces (photodissociation CH₃I on MgO [23,24], H₂ dissociation on surfaces [16] and H₂ scattering from LiF(001) [25]). The most advanced work has been done on three coupled 5D potential-energy surfaces [18,19]. Much of the work has, however, been directed at testing the method, by choosing models for which other methods were used as well for comparison. This paper presents one of the first real applications of MCTDH to a surface reaction.

2. Computational details

2.1. The potential-energy surface

Some time ago we did DFT calculations on the dissociation of CH₄ on Ni(111) using cluster models [10–14]. We have based the PES on the results of these calculations. As various clusters were used to model the Ni(111) surface, and the results of these calculations were not all the same, we have first identified common characteristics. These consisted of the structure of the transition state for the dissociation, the vibrational frequencies of the adsorbed products CH₃ and H, and, of course, the vibrational frequencies of the CH₄ in the gas phase. Two important parameters that we have not taken from the DFT calculations are the reaction energy and the height of the barrier. The reaction was endothermic for all DFT calculations, whereas experimentally it is found to be slightly exothermic [5]. We have assumed the reaction to be thermodynamically neutral. Many clusters we used gave more or less the same barrier of about 200 kJ·mol⁻¹, which is about a factor two higher than the highest experimental estimates [5,26]. Only a Ni₁₃ cluster gave a reasonable barrier height of 121 kJ·mol⁻¹. We have decided to fit the barrier height to the experimental dissociation probability at the translational energy of 64 kJ·mol⁻¹. This is roughly in the middle of the range of translational energies in the experiment [5].

The DFT calculations have optimised four coordinates describing the dissociating CH₄, and vibrational frequencies have been determined explicitly for six coordinates [13]. Our quantum simulations have only been done for three coordinates. Three reasons can be given for this. First, at this moment it is not possible to do a fully quantum simulation with six coordinates for CH₄ dissociation on Ni(111). Second, the DFT calculations provided us with information of the PES only for three small regions in configuration space for CH₄; namely, far from the surface, around the top of the barrier of the PES, and for CH₃ and H on the surface. As the quantum simulations need a complete PES, we had to interpolate, which proved to be taxing even for just three coordinates. Third, it is not clear how important the various coordinates are. We have included the distance of CH₄ to the surface, the distance between the H atom that splits off and the CH₃ that is formed, and the orientation of the CH₃/CH₄. The first two coordinates are evident. The reason for including the third coordinate is that the DFT calculations showed that the orientation of the CH₃ that is formed at the transition state always corresponds with a tilting vibration with a very low frequency (only a few wavenumbers), whereas on the Ni, cluster the CH₃ orientation even dominates the reaction coordinate. (On all other clusters the
CH$_3$–H stretch corresponds to the reaction coordinate. We have also done a few simulations with a fourth coordinate representing a lattice mode, very much as has been done in the simulations of CH$_4$ dissociation on Pt(111) [27–29]. In agreement with experimental findings, that showed no [5] or only a small [26] effect of the substrate temperature for Ni(111), we found that these 4D simulations gave similar results as the 3D simulations, and we did not continue with them. We like to note that in this decision computer time was only a minor issue, as for this system the 4D MCTDH simulations took only twice as long as the 3D simulations. No results of the 4D simulations will be presented.

The PES consists of two parts,

$$V(Z, s, \tau) = V_{\text{stretch}}(Z, s) + V_{\text{tilt}}(s, \tau),$$

where $Z$ is the distance between the centre-of-mass of CH$_4$ and the plane through the nuclei of the atoms in the first layer of the Ni(111) surface, $s$ is the distance between the H atom that splits off and the centre-of-mass of the CH$_3$ that is formed, and $\tau$ is the angle between the threefold axis of this CH$_3$ and the surface normal. The second term is given by

$$V_{\text{tilt}}(s, \tau) = AS\left(\frac{s - s_1}{s_2 - s_1}\right)(\cos \tau - 1),$$

with $A = -308$ kJ·mol$^{-1}$, and $S$ is a switch function,

$$S(x) = \begin{cases} 
0, & \text{if } x \leq 0, \\
1, & \text{if } x \geq 1, \\
10x^3 - 15x^4 + 6x^5, & \text{if } 0 \leq x \leq 1,
\end{cases}$$

which is continuous and has continuous first and second derivatives everywhere. The constants $s_1 = 0.169$ nm and $s_2 = 0.287$ nm correspond to the top of the barrier and the CH$_3$–H distance where the dissociation is complete, respectively. These values, and also the value of $A$, have been obtained directly from the DFT calculations. Note that there is no $\tau$-dependence up to the top of the barrier. This is reasonable for the CH$_4$ prior to dissociation as CH$_4$ is spherical to a good approximation. But also at the top of the barrier the DFT calculations show only a very weak $\tau$-dependence.

The first term in Eq. (1) is given by a generalisation of a LEPS form [30],

$$V_{\text{stretch}}(Z, s) = \frac{1}{2}Q_1(Z) + Q_2(s) - Q_2(Z)Q_1(s),$$

with

$$Q_1(Z) = \varepsilon_1(\varepsilon_1 \varepsilon_2 e^{-2\gamma_1(Z-Z_0)} + \beta Z e^{-\gamma_1(Z-Z_0)}),$$

$$Q_2(s) = \varepsilon_2[(1 - \varepsilon_2) e^{-2\gamma_2(Z-Z_0)} - (2 + \beta_Z e^{-\gamma_2(Z-Z_0)}],$$

and similar expressions for $U_s$ and $Q_s$. In the asymptotic limits $Z \to \infty$ and $s \to \infty$ the potential $V_{\text{stretch}}$ becomes the Morse potential $U_s + Q_s$ and $U_s + Q_s$, respectively. The parameters are given by $\varepsilon_1 = -480$ kJ·mol$^{-1}$, $Z_0 = 0.206$ nm, $s_0 = 0.115$ nm, $\gamma_1 = 11.0$ nm$^{-1}$, $\gamma_2 = 17.4$ nm$^{-1}$, $\varepsilon_2 = 0.6596$, $\varepsilon_2 = 1.3867$, $\beta_Z = -0.6999$ and $\beta_s = -0.2281$. The parameters $\varepsilon_1$ and $\varepsilon_2$ are obtained, using DFT, from the dissociation energy for CH$_4$ in the gas phase and the assumption that the reaction on Ni(111) is thermodynamically neutral. The parameters $s_0$ and $Z_0$ are obtained from the optimised geometries we determined with DFT. The parameter $\gamma_2$ is derived by fitting to the vibrational frequencies of the products as determined by the DFT calculations. Analogously, the parameter $\gamma_1$ is obtained from the CH$_3$–H stretch vibration in the gas phase. The other parameters are chosen to give a top of the barrier of the PES at $Z = 0.222$ nm and $s = 0.169$ nm, and a barrier height of 93.2 kJ·mol$^{-1}$. The position is once again determined by DFT calculations, but we choose the height to obtain a dissociation probability equal to the experimental one at a translational energy of CH$_4$ of 64 kJ·mol$^{-1}$. In fact, the number of parameters is overcomplete to determine just the barrier. The remaining freedom had to be used to avoid a deep physisorption minimum at low $Z$ and low $s$. Fig. 1 shows a plot of $V_{\text{stretch}}$. In agreement with the experimental observations of increased dissociation probability by vibrational excitation we find a very late barrier.
2.2. Initial and final states

The initial states in the simulation are given by
\[
\Psi(Z, s, \tau) = \psi^{(tr)}(Z) \psi^{(stretch)}(s) \psi^{(rot)}_m(\tau).
\] (7)

In this expression \(\psi^{(tr)}\) represents the initial translational part. It is a Gaussian wavepacket,
\[
\psi^{(tr)}(Z) = \left(\frac{2\pi \sigma^2}{-1} \right)^{-1/4} \exp \left( -\frac{(Z - Z_0)^2}{4\sigma^2} + iP_Z Z \right),
\] (8)

where \(Z_0\) is the initial position of the wavepacket (we used \(Z_0 = 0.582\) nm), \(\sigma\) is the width, which was chosen large to have a small spread in momentum and energy (we used \(\sigma = 0.0468\) nm), and \(P_Z\) is the initial momentum. The \(s\)-dependence of \(\Psi\) is given by \(\psi^{(stretch)}\), which is an eigenstate of the Hamiltonian for the CH\(_3\)-H stretch in the gas phase. The potential in this Hamiltonian is given by \(U + Q_v\). The index labels the various eigenstates; \(v, = 0\) for the ground state, and \(v, = 1, 2, 3, \ldots\) for the excited states. The \(\tau\)-dependence is given by
\[
\psi^{(rot)}_m(\tau) = \begin{cases} 
(1/\sqrt{\pi}) \cos(m\tau), & \text{if } m \geq 0, \\
1/\sqrt{2\pi}, & \text{if } m = 0, \\
(1/\sqrt{\pi}) \sin(|m|\tau), & \text{if } m < 0.
\end{cases}
\] (9)

As there is no \(\tau\)-dependence for \(Z \rightarrow \infty\) in the PES, it would be more natural to choose \(\exp(im\tau)\). But as \(\exp(im\tau)\) and \(\exp(-im\tau)\) are degenerate, the definition above is also valid. It has the advantage that it is adapted to the mirror symmetry in \(\tau\) of the PES.

The states \(\psi^{(stretch)}_v\) and \(\psi^{(rot)}_m\) are eigenstates for \(Z \rightarrow \infty\), and we will be interested in the population of these eigenstates for the scattered part of the wavefunction. The dissociated part of the wavefunction we will project on \(\psi^{(norm)}(Z)\) and on \(\psi^{(tilt)}_v(\tau)\). Here \(\psi^{(norm)}\) is an eigenstate of the Hamiltonian for the vibration of the dissociated CH\(_4\) as a whole normal to the surface (potential \(U_Z + Q_Z\)), and \(\psi^{(tilt)}_v\) is an eigenstate of the Hamiltonian for the CH\(_3\) tilt motion (potential \(A(\cos \tau - 1)\)). We have \(v, v, = 0, 1, 2, \ldots\), where higher indices mean higher excitation levels.

2.3. The multi-configuration time-dependent Hartree method

As this method has been described extensively elsewhere we will give here only an outline. We have used MCTDH with equations of motion for the natural single-particle functions [16]. The total
wavefunction $\Psi$ is written as

$$\Psi(q_1, \ldots, q_D; t) = \sum_{n_1 \ldots n_D} c_{n_1 \ldots n_D}(t) \times \psi_{n_1}^{(1)}(q_1; t) \cdots \psi_{n_D}^{(D)}(q_D; t),$$

(10)

where $\psi_{n_i}^{(i)}$ are the natural single-particle functions, which are defined as an eigenstate of the reduced density operator

$$\rho_j(q_j, q_{j-1}) = \int dq_1 \cdots dq_{j-1}dq_{j+1} \cdots dq_D$$

$$\times \Psi(q_1, \ldots, q_{j-1}, q_j, q_{j+1}, \ldots, q_D)$$

$$\times \Psi^*(q_1, \ldots, q_{j-1}, q_j, q_{j+1}, \ldots, q_D).$$

(11)

The equations of motions for the natural single-particle functions can be obtained by differentiation of the eigenvalue equation. This gives us

$$ih \frac{\partial}{\partial \tau} \psi_{n_j}^{(j)} = h_j \psi_{n_j}^{(j)} + \sum_{m_j} B_{n_j m_j}^{(j)} \psi_{m_j}^{(j)}$$

$$+ \frac{\langle \psi_{n_j}^{(j)} | V | \Psi \rangle}{\langle \psi_{n_j}^{(j)} | \psi_{n_j}^{(j)} \rangle} - \sum_{m_j} \frac{\langle \psi_{n_j}^{(j)} | V | \psi_{m_j}^{(j)} \rangle}{\langle \psi_{n_j}^{(j)} | \psi_{m_j}^{(j)} \rangle} \psi_{m_j}^{(j)},$$

(12)

where

$$B_{n_j m_j}^{(j)} = \frac{\langle \psi_{n_j}^{(j)} | \psi_{m_j}^{(j)} | V | \Psi \rangle - \langle \Psi | V | \psi_{n_j}^{(j)} \psi_{m_j}^{(j)} \rangle}{\langle \psi_{n_j}^{(j)} | \psi_{n_j}^{(j)} \rangle - \langle \psi_{m_j}^{(j)} | \psi_{m_j}^{(j)} \rangle}$$

(13)

and

$$\tilde{\psi}_{n_j}^{(j)} = \sum_{n_1 \ldots n_{j-1}} \sum_{n_{j+1} \ldots n_D} c_{n_1 \ldots n_D}$$

$$\times \psi_{n_1}^{(1)} \cdots \psi_{n_{j-1}}^{(j-1)} \psi_{n_{j+1}}^{(j+1)} \cdots \psi_{n_D}^{(D)}.$$  

(14)

The Hamiltonian $H$ is given by

$$H = \sum_{j=1}^D h_j + V,$$

(15)

Differentiation of

$$c_{n_1 \ldots n_D} = \langle \psi_{n_1}^{(1)} \cdots \psi_{n_D}^{(D)} | \Psi \rangle$$

(16)

gives us

$$ih \frac{d}{dt} c_{n_1 \ldots n_D} = \langle \psi_{n_1}^{(1)} \cdots \psi_{n_D}^{(D)} | V | \Psi \rangle$$

$$- \sum_{j=1}^D c_{n_1 \ldots n_{j-1} n_j \ldots n_D} B_{n_j m_j}^{(j)},$$

(17)

Eqs. (12) and (17) are to be solved numerically. More general equations of motion can be derived starting from a time-dependent variational principle [15]. It has explicitly been shown that Eqs. (12) and (17) are recovered by imposing appropriate restrictions [31,32]. We find that these equations are somewhat more convenient to analyse results than the more general equations of motion. In particular, it is easy to determine the convergence of the simulations with respect to the number of terms (configurations) in Eq. (10). However, other equations of motion may be numerically more advantageous by allowing a larger step size in the numerical integration. This is very much still an open question. We have used the variable-order variable-step Adams method as implemented in routine D02CBF of the NAG library for the numerical integration [33,34]. The length of each simulation was 360 fs.

The grid that we have used spans the region $0.132 \leq Z \leq 0.807$ nm, $0.053 \leq s \leq 1.405$ nm, and $-\pi \leq \tau \leq \pi$ in configuration space. There are 256 grid points in $Z$, 512 in $s$, and 64 in $\tau$. The large range in $s$ was necessary as the H atom that splits off moves very fast, and very far, over the surface. The grid has also to be dense, because of the high mass of CH$_4$. Such a large grid presents no problem at all for MCTDH. Our system is very close to the limiting case where the computer time used by MCTDH scales linearly with the number of grid points for one coordinate. The other limiting case is where MCTDH scales exponentially with the number of natural single-particle functions for one coordinate [15]. The number of configurations we used was $7 \times 7 \times 4$, which means seven natural single-particle functions for $Z$, seven for $s$, and four for $\tau$. Calculations with more natural single-particle functions showed only a negligible difference for the range of translational energy that we looked at.

3. Results and discussion

Fig. 2 shows the dissociation probability for a ground state CH$_4$ molecule as a function of translational energy. We see that it scales approximately exponentially with the translational energy, except for large probabilities, which is in agreement with
the experiment [5]. This has been interpreted as an indication that the dissociation occurs via tunneling. We will show below that this is indeed correct. We have fitted the barrier height of the PES to the experiment, but we see that the curve of the simulation is much steeper than the experimental one. This implies that the barrier of the PES that we used is too wide and too low. To obtain some idea of the error in our barrier height and width we estimate an improved height and width as follows. The experimental results have been interpreted using [35] for the dissociation probability $P_{\text{diss}}$ as a function of the translational energy $E_T$, the height $E_{\text{bar}}$ of the barrier, and the mass $\mu$ corresponding to the reaction coordinate. The barrier height of our PES is $E_{\text{bar}} = 93.2 \text{ kJ} \cdot \text{mol}^{-1}$, which gives us for the width $W = 0.0546 \text{ nm}$ using $P_{\text{diss}} = 3.66 \times 10^{-4}$ at $E_T = 64 \text{ kJ} \cdot \text{mol}^{-1}$, and using for $\mu$ the reduced mass of the H atom that splits off. Using $E_{\text{bar}} = 108 \text{ kJ} \cdot \text{mol}^{-1}$ and $W = 0.04 \text{ nm}$ gives the same dissociation probability at $E_T = 64 \text{ kJ} \cdot \text{mol}^{-1}$, but these latter values fit the experimental dissociation probability also for other energies [5]. We see that the barrier should be higher and narrower. Eq. (18) is really only valid for a particular 1D model of tunneling. If we use all our simulation results and fit them with Eq (18) we find $E_{\text{bar}} = 84.1 \text{ kJ} \cdot \text{mol}^{-1}$ and $W = 0.0725 \text{ nm}$. The difference with the real barrier height is a consequence of the fact that our simulations are 3D and not 1D. One consequence of this is that zero-point vibrations perpendicular to the reaction coordinate changes the effective barrier height (see below). Also the twelve degrees of freedom of CH$_4$ we do not include in our simulations will have an effect on the effective barrier height. There is a whole range of experimental values for the barrier height, 33.0 [8], 52.7 [36], 108.7 [5] and about 130 kJ mol$^{-1}$ [26], but there is a fair agreement between values based on a dynamical interpretation of experimental results [5,26] and our value of 93.2 kJ mol$^{-1}$ and also the value of 121 kJ/mol$^{-1}$ we calculated with DFT.

The tunneling mechanism has recently been questioned in the case of H$_2$ dissociation on copper. Instead, the dissociation probability has been explained classically by the hole model [37]. This model uses a distribution of barrier heights. At higher energy the hole in configuration space through which the system can pass from reactant to product is bigger than at low energy. It has been remarked, however, that it would be unlikely that a distribution of barriers would produce the exponential scaling [5]. It is absolutely clear that in the simulations we are seeing tunneling, as the incoming methane molecule simply does not have enough energy (translational and 15.9 kJ mol$^{-1}$ zero-point vibrational energy) to get over the barrier. This does not mean that tunneling is important for the thermally averaged dissociation probability. The contribution from high energies has a small Boltzmann factor, and at low energies there is a small dissociation probability. It is still an open question whether the largest contributions come from energies below or above the activation energy. The results from Beebe et al. [26] seem to suggest the former, whereas our TST results suggest the latter [11–14].

Fig. 2 also shows the effect that including the
rotation of methane has. We see that the dissociation probability in the 2D simulations, without the coordinate \(z\), is about a factor of five larger than in the 3D simulations. One way to understand this is by using a model with 1D effective potentials [38, 39]. We first define a reaction path as a steepest-descent path from the top of the barrier. We do this in the configuration space of mass-weighted coordinates as otherwise the reaction path cannot be defined unambiguously [40]. Next we define an effective potential as a function of the reaction coordinate as the sum of the value of the PES on the reaction path plus an eigenvalue corresponding to the vibrations perpendicular to the reaction path. The results are shown in Fig. 3. In the entrance channel there is no difference between the 2D and the 3D case for \(m_r = 0\). In the exit channel the effective potential is 9.8 kJ·mol\(^{-1}\) higher in the 3D case due to the zero-point energy of the tilt vibration of the CH\(_3\) fragment. This makes the barrier a little bit wider on the exit side, which has a large effect on the dissociation probability, because of the tunneling. Note that when we would treat the reaction coordinate classically that there would be no difference for the two effective potentials, because the heights of both is the same. For a fully classical 2D and 3D simulation there would be a difference in the dissociation probability for translational energies above the minimum barrier height, because the distribution of barrier heights is different. As the zero-point energy of the CH\(_3\) tilt is not large, yet has a large effect on the dissociation probability, the inclusion of other degrees of freedom may have a substantial effect too.

Fig. 2 also shows some results for dissociation with a singly excited CH\(_3\)-H stretch. As is to be expected for a late barrier, the dissociation increases enormously. This too can be explained using 1D effective potentials (see Fig. 4). The effective barrier height decreases rapidly with increasing vibrational excitation level, and vanishes already at \(v_s = 3\). The effective barrier height is 79.4 kJ·mol\(^{-1}\) for \(v_s = 0\) and 60.0 kJ·mol\(^{-1}\) for \(v_s = 1\). This is very similar to what has been found for H\(_2\) model studies [38]. The additional vibrational energy of 33.3 kJ·mol\(^{-1}\) is very efficient in inducing dissociation; we can see in Fig. 2 that the dissociation probability curve of \(v_s = 1\) is about 22 to 26 kJ·mol\(^{-1}\) lower in energy than the one of \(v_s = 0\).

![Fig. 3. Effective 1D potentials (in kJ·mol\(^{-1}\)) showing the effect of including the rotational degree of freedom. The fat line is the value of the potential-energy surface along the reaction path. The dashed line is the effective potential for the 2D simulation with CH\(_3\) in the ground state. The solid lines are effective potentials for CH\(_4\) in the ground state and in a state excited in \(r\). The number on the right indicates \(v_s\) (\(v_s = 0\) correlates with \(m_s = 0\), and \(v_s = 2\) with \(m_s = 1\), because of symmetry). Only potentials for states even in \(r\) are shown.](image1)

![Fig. 4. Effective 1D potentials (in kJ·mol\(^{-1}\)) showing the effect of excitation of the CH\(_3\)-H stretch. The fat line is the value of the potential-energy surface along the reaction path. The solid lines are effective potentials for CH\(_4\) in the ground state and in states excited initially in \(s\). (They become excited in \(Z\) to the right of the barrier.) The numbers on the left indicate \(v_s\).](image2)
Fig. 5. Dissociation probability as a function of translational energy (in kJ·mol$^{-1}$) for CH$_4$ initially in the ground state (circles), and rotationally excited states $m_z = 1$ (triangles pointing upwards), $m_z = -1$ (triangles pointing downwards), and $m_z = 2$ (squares). The symbols are simulation results, and the solid lines are guides for the eye.

Fig. 5 shows the dissociation probability for rotational excitation. We see that the initial states even in $\tau$ have about the same dissociation probability, but that the one odd in $\tau$ has a much lower one. This indicates that the even-$\tau$ states do not follow an effective 1D potential. For if they did we would find that the $m_z = 1$ initial state would become $v_z = 2$ in the exit channel, and $m_z = 2$ would become $v_z = 4$. The barriers for these processes are much wider than for the lowest effective potential (see Fig. 3), and a drastically reduced dissociation probability would be found. Instead we see that terms that couple the effective potentials are important [38]. Indeed, we hardly find any vibrational excitations of the CH$_3$ tilt. For the initial state $m_z = -1$ the effective potential leads to $v_z = 1$. In this case no coupling exists with the lowest effective potential, because it is symmetry forbidden. Consequently, a wider barrier is felt, and the dissociation probability becomes much lower.

That there are substantial coupling terms between the effective potentials can also be seen from the excitations of the CH$_3$ tilt. Only at the highest translational energy that we looked at (80 kJ·mol$^{-1}$) did we find an occupation of the $v_z = 2$ level in the exit channel comparable to the $v_z = 0$ (0.084 and 0.047 being the fractions of the total wavefunction at the end of the simulation ending up in $v_z = 0$ and $v_z = 2$, respectively). Below 72 kJ·mol$^{-1}$ vibrational excitation of the CH$_3$ tilt accounts for less than 1% of the wavefunction in the exit channel. On the other hand, as is to be expected for a late barrier, the vibration perpendicular to the surface of the dissociated CH$_4$ as a whole is highly excited. The effect is stronger than for H$_2$, as the frequency is lower due to the higher mass of CH$_4$ [38]. For low translational energies the occupation of the levels scales exponentially with translational energy, as does the dissociation probability. The levels $v_z = 0$, $v_z = 1$ and $v_z = 2$ have also about the same exponential factor as the dissociation probability.

Fig. 6 shows scattering probabilities. The excitation $v_z = 1$ of the CH$_3$-H stretch seems to scale exponentially, and resembles rather closely the curve of the total dissociation probability. This is, we think, related to the fact that for the reaction path coming from the entrance channel the CH$_3$-H stretch coordinate stays around its gas phase value for a long time, and only just before the top of the barrier the CH$_3$-H bond stretches. Consequently, at low translational energies almost all CH$_4$ is scattered from the surface without really feeling

Fig. 6. Scattering probabilities as a function of translational energy (in kJ·mol$^{-1}$) for vibrational and rotation excited levels. Circles indicate excitation to $v_z = 1$, squares to $v_z = 2$, diamonds to $m_z = 1$, and triangles to $m_z = 2$. The symbols are simulation results, and the solid lines are guides for the eye.
the bend in the reaction path (see Fig. 1). Although the excitation energy of the CH$_4$ rotation is even much smaller than for the CH$_3$–H stretch, the rotation is even less excited. The reason for this is that the PES becomes only $\tau$-dependent at the exit side of the barrier. However, once the rotation is excited it is not restricted to just small $|m_\tau|$ values. At $E_T = 80 \text{ kJ} \cdot \text{mol}^{-1}$ we have found scattering probabilities of 0.83, 0.025, 0.011, $3.8 \times 10^{-3}$ and $1.0 \times 10^{-3}$ for $m_\tau = 0, 1, 2, 3$ and 4, respectively.

The 2D simulations show about the same vibrational excitation in the scattered part of the wavefunction as the 3D simulations. More interesting are the scattering results for excited initial states. The scattering of a CH$_4$ molecule with a singly excited CH$_3$–H stretch vibration is predominantly adiabatic as long as the dissociation probability is smaller than about 0.1. For energies with a higher dissociation probability there is a strong de-excitation in the scattered part. At $E_T = 56 \text{ kJ} \cdot \text{mol}^{-1}$ a fraction of 0.573 of the total wavefunction is scattered and a fraction of 0.456 is scattered into $v_\nu = 0$. Also $v_\nu = 1$ (0.059) and $v_\nu = 2$ (0.047) have some population, but other vibrational levels are hardly populated. At this energy there is also substantial excitation of the CH$_4$ rotation; 0.170 for $m_\tau = 1$ and 0.095 for $m_\tau = 2$. Apart from a shift in the translational energy equal to a vibrational quantum, due to microreversibility the de-excitation probability $v_\nu = 1 \rightarrow 0$ is equal to the excitation probability $v_\nu = 0 \rightarrow 1$ [41]. We have used this to increase the number of data points in Fig. 6. If the initial state is rotationally excited then the scattering is adiabatic for the range of translational energies we have looked at.

We would finally like to make a few remarks on the correlation between the excitations. There is a strong correlation in the scattered part of the wavefunction between the rotational and the vibrational excitation. In the rotationally excited levels the vibration is excited more than in the rotational ground state. The reason for this is obviously that the rotational excitation originates from parts of the wavefunction that have extended even beyond the top of the barrier, which leads also to vibrational excitation. As the translational energy becomes higher the correlation becomes stronger, and also more vibrationally excited states become populated. A reverse effect is seen for the rotational ground state. For this state the vibrational ground state is slightly preferred.

There is much less correlation in the dissociated part of the wavefunction. The only clear effect is that the ground state of the CH$_3$ tilt vibration is a little bit more excited in the vibration perpendicular to the surface than is to be expected based on the population of the perpendicular vibration for the whole dissociated part.

4. Conclusions

We have done quantum simulations for a 3D model of CH$_4$ dissociation on Ni(111) using the MCTDII method. The PES was developed based on information of DFT calculations. It has a very late barrier and a height of 93.2 kJ $\cdot$ mol$^{-1}$.

The dissociation probability scales exponentially with the translational energy of the methane molecule. From the total energy and the height of the barrier it could be deduced that the discussion occurs via tunneling. The inclusion of the rotation in the simulation of CH$_4$ decreases the dissociation probability. This is due to the fixed orientation of the CH$_3$ on the surface, which makes the barrier for the dissociation effectively wider. Because of the late barrier a CH$_4$ molecule with an excited CH$_3$–H stretch vibration dissociates much easier. There is a strong tendency to de-excite a rotationally excited CH$_4$, which has as a consequence that there is only a small effect on the dissociation probability. Only when for symmetry reasons this de-excitation is not allowed, the CH$_3$ tilt will be excited, which reduces the dissociation probability.

Excitation of the CH$_3$–H in scattered CH$_4$ resembles the dissociation probability as both require enough translational energy so that the wavefunction can sample the region of the PES around the top of the barrier, where the C–H bond is stretched. The rotation of scattered CH$_4$ is only excited at the highest translational energy we looked at ($80 \text{ kJ} \cdot \text{mol}^{-1}$), even though the excitation energy is very low. This is because the PES has no rotational dependence on the entrance side of the barrier. When the rotation does become excited the excited rotational levels are also more
vibrationally excited than the rotational ground state.

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