Kinetic Monte Carlo Studies of Hydrogen Abstraction from Graphite

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We present Monte Carlo simulations on Eley-Rideal abstraction reactions of atomic hydrogen chemisorbed on graphite. The results are obtained via a hybrid approach using energy barriers derived from DFT calculations as input to Monte Carlo simulations. By comparing with experimental data we discriminate between contributions from different Eley-Rideal mechanisms. A combination of two different mechanisms yields, good quantitative and qualitative agreement between the experimentally derived and the simulated Eley-Rideal abstraction cross sections and surface configurations. These two mechanisms include a direct Eley-Rideal reaction with fast diffusing H atoms and a dimer mediated Eley-Rideal mechanism with increased cross section at low coverage. Such a dimer mediated Eley-Rideal mechanism has not previously been proposed and serves as an alternative explanation to the steering behavior often given as the cause of the coverage dependence observed in Eley-Rideal reaction cross sections.

I. INTRODUCTION

Molecular hydrogen is the most abundant molecule in the interstellar medium (ISM), where it serves as an important coolant and as a precursor for the formation of more complex molecules. In the cold and dilute interstellar medium no efficient gas phase routes exist for the formation of H₂. Hence, the most likely route is to form the molecule on the surfaces of small dust particles. This process has been intensely studied over the past years [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12]. The general consensus is that molecular hydrogen can be formed by a diffusive Langmuir-Hinshelwood mechanism, in the very cold regions below ~20 K [13]. At these low temperatures the hydrogen atoms are physisorbed on the dust particles, but can still easily move on the surface. Once two atoms meet, they will react to form molecular hydrogen. If the surface temperature becomes too high, the residence time of atoms on the surface becomes so short that, at the low fluxes in the ISM, the chance of two atoms meeting becomes negligible. Observations show that molecular hydrogen is also formed in warmer areas (> 20 K) like Photon Dominated Regions (PDRs) and post-shock regions. A possible mechanism at these conditions would be through an Eley-Rideal reaction where an incoming hydrogen atom reacts with another H atom that is chemically bound to the surface. Since a considerable fraction of the interstellar grains is expected to consist of carbonaceous material, a popular model system for this process is hydrogen abstraction from graphite. This Eley-Rideal reaction has been studied by Density Functional Theory (DFT) and Quantum Wave packet calculations [14, 15, 16, 17, 18, 19]. Based on these calculations several different mechanisms have been proposed to contribute to the Eley-Rideal abstraction process. These include: Direct Eley-Rideal [14, 15, 16, 17, 18, 19], barrier-less abstraction of one hydrogen atom forming part of a para-dimer configuration [20] and abstraction by rapidly diffusing H atoms in physisorbed states [21].

However, due to computational limitations such detailed calculations are unable to take the full complexity of the H-graphite system into account. Hence, in the calculations the flat surface approximation is often used and generally either none or only a single C atom on the simulated graphite surface were allowed to relax during the Eley-Rideal reaction. These simplifications are potentially problematic. By not allowing for relaxation of the C atoms on the surface zero barrier reaction channels, such as sticking of hydrogen atoms into specific dimer configuration [21, 22], are in some cases artificially closed, making it impossible to evaluate the contributions from different proposed mechanisms to the Eley-Rideal abstraction process. Eg. calculations on the Eley-Rideal reaction by Martinazzo and Tantardini [19] show a higher probability for reaction if the incoming H atom is 0.5 - 1.5 Å away from the target H atom instead of a direct hit on. This distance corresponds to the H-H distance in an ortho-dimer configuration (see Fig. 1) on the graphite surface (1.42 Å). Hence, surface corrugation and competing sticking reactions must be expected to influence the abstraction behavior. In these calculations, however, dimer formation was not a possible route since the flat surface approximation was used and only the carbon atom underneath the original hydrogen atom was allowed to relax. Impact parameters corresponding to the para-dimer (see Fig. 1) distance (2.84 Å) were not included in the study.

Both experimental observations and theoretical calculations show that hydrogen atoms preferentially form adsorbate clusters on the graphite surface. Scanning Tunneling Microscopy (STM) investigations show that at a hydrogen atom coverage of ~ 0.5 % more than 75 % of all surface configurations are clusters [23]. This indicates that ~ 85 % of the hydrogen atoms are part of...
larger clusters. In particular, the existence of hydrogen dimer configurations has been studied theoretically and experimentally. Hornekær et al. identified two stable hydrogen dimer configurations on the graphite surface, an ortho-dimer and a para-dimer (see Fig. 1). DFT calculations show that formation of the para-dimer is barrierless and that formation of the ortho-dimer has a reduced barrier, which makes dimer formation a competing channel to Eley-Rideal abstraction at non-zero impact parameter.

One set of experiments on the Eley-Rideal abstraction reaction forming HD has been reported. In these experiments a high cross section for the abstraction reaction was observed which varied from 17 Å² at low coverage to 4 Å² at high coverage. This variation in the cross section with coverage has been ascribed to a steering effect. However, other mechanisms could also contribute to the high cross section at low coverage. Options include the barrier-less abstraction of one of the H atoms in a para-dimer configuration and abstraction by fast diffusing H atoms. We propose a further possibility, namely abstraction via a hydrogen dimer state, where the incoming atom is not immediately thermalized and some of its excess initial energy is used to overcome the barrier to H₂ formation and desorption. The finding that adsorption into the hydrogen para-dimer state is barrierless makes this a strong competing reaction channel.

Hence, in total 5 different abstraction mechanisms have been proposed to contribute to Eley-Rideal abstraction of hydrogen on graphite:

I. direct Eley-Rideal
II. direct Eley-Rideal with steering
III. preferred direct Eley-Rideal with hydrogen atoms part of a para-dimer
IV. a dimer mediated reaction where the incoming atom is first adsorbed into a dimer configuration and, before thermalizing to the substrate temperature, reacts with the other atom in the dimer to form H₂ and desorb.
V. Eley-Rideal reactions by fast diffusing H atoms in the physisorption state

However, due to the simplifications needed in the complex DFT and quantum wave packet calculations a quantitative comparison between experimental data and theory has not been possible. In this paper we employ a hybrid approach in which we include the findings of ab-initio DFT calculations in a Monte Carlo simulation program and then simulate Eley-Rideal abstraction experiments on a more realistic surface area with more complex hydrogen adsorbate configurations than what is possible in the DFT and wave packet calculations. Through this quantitative approach we aim to discriminate between the contributions of the different underlying mechanisms for hydrogen abstraction.

II. MONTE CARLO MODEL

The Monte Carlo simulation program is a so-called lattice-gas simulation program, where the atoms are confined to an adaptive grid depending on the initial coverage. The hydrogen atoms can chemisorb to the graphite surface at the sites directly on top of the carbon atoms as indicated in Figure 2. The puckering of the carbon atom upon chemisorption is included indirectly by using the barriers for chemisorption and binding energies from DFT calculations that allow this motion. Interaction between two hydrogen atoms is accounted for in a similar way as will be discussed in the following sections. The hydrogen atoms can physisorb to the surface without a barrier. Since the potential energy surface for physisorption is rather flat, there are probably no specific physisorption site, however since the lattice-gas model forces us to choose specific sites, we confine physisorption to the sites directly above the carbon atoms and an additional site at the center of the ring (Figure 2).

The simulation starts with a clean graphite surface. The first event will be a deposition attempt of the first atom. The time at which this attempt will occur is

\[ t_{\text{dep}} = -\frac{\ln(X)}{f} + t \]  

where \( \sigma \) is the density of sites, \( X \) is a random number between 0 and 1, \( f \) is the hydrogen flux in atoms per time per area, and \( t \) is the current time. Deposition times for subsequent sticking events are determined by
the same expression (Eq. 1). How the atoms will bind to the surface is determined by another random number and depends on the barrier for chemisorption in that specific position. If the site is already occupied by another hydrogen atom, an abstraction reaction is considered by comparing a Boltzmann factor including the abstraction barrier against a random number. Upon reaction both atoms will leave the surface in the form of H₂, else the incoming atom is deflected. The abstraction channel can be closed by making the barrier infinitely large.

Once hydrogen atoms populate the surface, they can diffuse, desorb, or recombine with other atoms to form H₂. For each surface hydrogen the time at which they will undergo one of these events is determined by

\[ t_i = \frac{\ln (X)}{R_{\text{dif}}^i + R_{\text{des}}^i + R_{\text{rec}}^i} + t \]  

(2)

with \( R_{\text{dif}}^i \), \( R_{\text{des}}^i \), and \( R_{\text{rec}}^i \) the rate for diffusion, desorption, and reaction of atom \( i \), respectively. Another random number determines which of the three events occur according to their relative probability of occurrence. Reaction between two chemisorbed atoms can only occur if they form a dimer configuration. The rates are given by

\[ R = \nu \exp \left( -\frac{E}{kT} \right), \]  

(3)

where \( \nu \) is the attempt frequency which is assumed to be \( 10^{12} \) Hz for physisorbed atoms and \( 10^{13} \) Hz for chemisorbed atoms [24] and \( T' \) is the ‘temperature’ of the atom that is involved.

In the experiments that we aim to reproduce, the atoms arrive at the surface at normal incidence and at a temperature around 2000 K which is much higher than the surface temperature. Furthermore, the atoms will gain energy due to the high binding energy if they chemisorb. The atoms will not be thermalized instantaneously, but will most likely gradually lose their energy to the substrate. The dissipation of the excess energy into the substrate is expected to be exponential [30]. To reduce the computation complexity of the model we emulate this exponential energy loss by a simpler expression and use the following function to describe the temperature

\[ T'(t) = \max \left( T_s, \frac{T_{\text{start}}}{(1 + B(t - t_a))^2} \right) \]  

(4)

where \( T_s \) and \( T_{\text{start}} \) are the surface and starting temperature respectively, \( t_a \) is the time at which the atom has adsorbed on the surface. The sensitivity of the model to the exact functional form was checked and found to be small. For most cases we use \( T_{\text{start}} = 2000 \) K, but also higher values of 7000 K and 10,000 K are considered, matching the binding energy of a monomer. The parameter \( B \) can be chosen freely. Fig. 3 studies the effect of this parameter. It displays the result of series of 10,000 simulations of one deposition event for a particular value of \( B \). The top panel indicates the percentage of deposition attempts that resulted in sticking. The sticking fraction slowly approaches the monomer sticking barrier \( B \). The top panel indicates the percentage of deposition attempts that resulted in sticking. The sticking fraction slowly approaches the monomer sticking barrier.

In the brief moment after a deposition, the rates in Eq. 3 are time dependent due to the decreasing temperature and Eq. 2 cannot be used. Instead we use the method by Jansen et al. [31] to determine \( t_i \). This method makes use of

\[ -\ln (X) = \int_t^{t'} R_{\text{dif}}^i (t') dt + \int_t^{t'} R_{\text{des}}^i (t') dt + \int_t^{t'} R_{\text{rec}}^i (t') dt, \]  

(5)

that transforms to Eq. 2 if all rates are time dependent. To obtain \( t_i \) the expression has to be solved. Using

\[ \Omega(t^i) = \int_t^{t'} \nu \exp \left( -\frac{E(1 + B(t - t_a))^2}{kT_{\text{start}}} \right) dt \]  

(6)

\[ = \frac{\nu}{2B} \sqrt{\frac{\pi T_{\text{start}}}{E}} \exp \left( \frac{1 + B(t - t_a)}{T_{\text{start}}} \right) \]  

with \( \text{erf} \) the error function, Eq. 5 becomes

\[ -\ln (X) = \Omega_{\text{dif}}(t^i) + \Omega_{\text{des}}(t^i) + \Omega_{\text{rec}}(t^i). \]  

(8)
This can be solved numerically to $t_i$ using the Newton-Raphson method \cite{32}, since both $\Omega$ and $\frac{d\Omega}{dt}$ decrease monotonically. Notice that different functions for $T'$ will result in a different expression for $\Omega(t')$. As the order in $t$ increases, solving $\Omega(t_i)$ becomes more computationally expensive.

### III. INCLUDED REACTIONS AND ENERGY BARRIERS

The previous section described the general Monte Carlo algorithm. For all processes energy barriers are needed to determine the corresponding transition probabilities. These barriers are taken from independent DFT calculations \cite{33} of the binding energies of the different configurations that are considered, sticking trajectories and diffusion trajectories. Since many possible configurations are formed during the simulations, especially for high coverages, including all these different possibilities explicitly would make the Monte Carlo program very slow and would require a huge set of barriers that all have to be calculated independently. To overcome this problem a number of simplifications are introduced resulting in the following sets of energy barriers:

#### A. Physisorbed atoms

For the physisorption binding energy, the value of $\sim 40$ meV is used based on results from selective adsorption experiments \cite{34}. For diffusion an activation energy of 4 meV taken from \cite{21} was used. We will come back to this diffusion rate at the discussion of mechanism IV. For physisorbed atoms $T'(t) = T_s$ was used.

#### B. Sticking

Numerous different barriers for sticking of an isolated hydrogen atom (a monomer) into the chemisorption site on the graphite surface have been given. Jeloaica and Sidis found a barrier of $\sim 0.2$ eV using the coronene molecule as a model of a graphite surface \cite{35}. Sha et al. obtained a barrier slightly above $0.2$ eV using a slab super cell with 4 layers each containing 8 carbon atoms \cite{16}. Hornekær et al. \cite{25} found a barrier for chemisorption into a monomer of 0.15 eV using a single layer super cell containing 32 carbon atoms. The influence of adding a second carbon layer was investigated and found to be negligible. Using the same model surface a sticking barrier of 0.1 eV into the ortho-dimer and 0 eV into the para-dimer configurations were found. All possibilities for sticking into a trimer state, starting from the para-dimer, were seen to have non-zero sticking barriers between 0.1 and 0.15 eV. Adding a fourth atom resulting in a triple para-dimer configuration again did not exhibit a barrier \cite{25}. Barrierless sticking into the para-dimer state was also found by Rougeau et al. \cite{26}.

Even though there is a clear dependence on the local configuration of the impact site, we decided only to include variations in sticking barriers for dimers in the simulation. The program determines if the incoming atoms can form a dimer ignoring the larger configuration it might be part of and determines the barrier for sticking accordingly. This assumption will cause deviations between simulations and experimental results at high coverage, since it overestimates the formation of trimer configurations that contain para-dimers. No sticking barrier was used for physisorption of H atoms. The barriers for sticking used in the simulation are summarized in Table I.

Section VI tests the different mechanisms to their sensitivity to various input parameters. The monomer sticking barrier is one of them.

#### C. Binding energies

The binding energies for different configurations of chemisorbed H atoms are displayed in table II. The different configurations are displayed in Fig. 4. These values are based on DFT calculations reported in \cite{25, 33}.

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**Table I: The sticking barrier for different configurations**

| Configuration | $E_{\text{stick}}$ [meV] |
|--------------|--------------------------|
| para-dimer   | 0.0                      |
| ortho-dimer  | 0.1                      |
| other chemisorption site | 0.15              |
| physisorption | 0.0                      |

**Table II: The total binding energy for different configurations**

| Configuration$^1$ | $E_{\text{bind}}$ [meV] |
|-------------------|--------------------------|
| monomer           | -0.8                     |
| para-dimer (G)    | -2.9                     |
| ortho-dimer (A)   | -2.8                     |
| meta-dimer (E)    | -0.8                     |
| trimer (A, G)     | -3.9                     |
| trimer (G, I)     | -3.7                     |
| trimer (D, G)     | -3.5                     |
| trimer (B, G)     | -3.5                     |
| trimer (F, G)     | -3.3                     |
| tetramer (A, G, H)| -5.9                     |
| tetramer (A, B, E)| -5.9                     |
| tetramer (A, D, G)| -5.7                     |
| tetramer (A, B, D)| -5.5                     |
| tetramer (B, E, G)| -5.1                     |
| tetramer (A, B, C)| -5.0                     |
| tetramer (A, B, G)| -4.8                     |
| tetramer (A, B, F)| -4.5                     |

$^1$ See Fig. 4.
FIG. 4: Schematic guide to obtain the configurations used in Table II. A configuration is made up from hydrogen atoms positioned on top of the black carbon atom and the atoms indicated by the characters given in Table II. In this way, an ortho-dimer is represented by the character (A).

For several dimer and tetramer configurations the individual binding energies and the transition barriers between some of the configurations were determined via DFT calculations. The binding energy of the individual dimer and tetramer configurations are given in Table II. The tetramer binding values and diffusion barriers are taken from [33]. Figure 5 plots the diffusion barrier as a function of the energy difference of the configurations. As the figure clearly shows there is a linear relation between the two quantities and dimers and tetramers follow the same relation. A least-squares fit resulted in

\[ E_{\text{diff}} = 0.5\Delta E + 1.04 \text{ eV}. \] (9)

In order to obey detailed balance, or microscopic reversibility, transition probabilities for diffusion should fulfill

\[ \frac{P_{ij}}{P_{ji}} = \exp \left( \frac{\Delta E_{ij}}{kT} \right) \] (10)

Using Eq. 9 it can be shown that the empirically found relation, Eq. 9 follows this requirement.

E. Desorption

Finding a general expression for the desorption energy of the individual atoms from a configuration is less straightforward. The binding energies are determined for the configuration and not for the individual atoms. Again calculating all possible desorption pathways would not be feasible. We base the desorption on the total binding energy of the configuration of \( n \) atoms, \( E_{\text{bind}} \). The desorption energy is then

\[ E_{\text{des}} = \frac{E_{\text{bind}}}{n} + E_{\text{stick}} \] (11)

where \( E_{\text{stick}} \) is the barrier for sticking in the same position. \( E_{\text{stick}} \) and \( E_{\text{bind}} \) can be found in Tables I and II.

F. Thermally activated H\(_2\) formation

Formation of H\(_2\) via reaction between two chemisorbed hydrogen atoms to gaseous molecular hydrogen occurs from the dimer states and has to overcome barriers of 2.49 eV for the ortho-dimer and 1.4 eV for the para-dimer state [24]. Reaction barriers from the isolated dimer states are used for all configurations. This approximation will again lead to inaccuracies for simulations at high coverage.

G. Abstraction

Five different Eley-Rideal abstraction mechanisms are considered in the model:

I. direct Eley-Rideal with all hydrogen atoms regardless of their local configuration.

II. a simple version of Eley-Rideal with steering where a direct Eley-Rideal reaction is allowed not just for H atoms impinging on an already occupied site but also for H atoms impinging on adjacent sites.

III. direct Eley-Rideal with only hydrogen atoms part of a para-dimer.
FIG. 6: The amount of HD molecules per time leaving the surface in ML/s as a function of time for different pre-exposures of D. At $t = 0$ the hydrogen beam is switched on. For (I) mechanism I: Direct Eley-Rideal ($E_{\text{ER, mono}} = E_{\text{ER, dimer}} = 9$ meV), (II) mechanism II: Direct Eley-Rideal with steering ($E_{\text{ER, mono}} = E_{\text{ER, dimer}} = 9$ meV, $s = 1$), (III) mechanism III: Dimer Eley-Rideal ($E_{\text{ER, mono}} = \infty; E_{\text{ER, dimer}} = 0$ meV), (IV) mechanism IV: Dimer mediated Eley-Rideal ($B = 1 \times 10^9$ s$^{-1}$), and (V) mechanism V: Direct Eley-Rideal with fast diffusion of physisorbed atoms ($E_{\text{ER, mono}} = E_{\text{ER, dimer}} = 9$ meV, $R_{\text{diff}} = 5 \times 10^{13}$ s$^{-1}$).

IV. a dimer mediated reaction where the incoming atom is first adsorbed into a dimer configuration and, before thermalizing to the substrate temperature, forms H$_2$ and desorbs.

V. direct Eley-Rideal together with a high diffusion rate of the atoms in the physisorption state.

The influence of the different Eley-Rideal mechanisms can be controlled by using different barriers for the direct Eley-Rideal reaction and different values of the thermalization parameter $B$. Furthermore, a simple version of steering is implemented with an adjustable parameter $s$ as will be described below. The values of the parameters used for simulating the different mechanisms are given in Table III.

The height of the abstraction barrier is somewhat uncertain due to the limitations in the accuracy of DFT calculations. Morisset et al. [18] found a barrier for direct abstraction of a monomer just below 10 meV. Others also found low barriers to abstraction [17, 19]. Following Morisset et al. we employ a barrier of 9 meV but later investigate the effect of changing the value of the barrier. The same group also found that the abstraction reaction with one of the hydrogen atoms in a para-dimer can proceed without barrier [20].
TABLE III: Model parameters for the different abstraction mechanisms

| Mechanism      | $E_{ER, \text{mono}}$ [meV] | $E_{ER, \text{dimer}}$ [meV] | $B$ [s$^{-1}$] | $T_{\text{start}}$ [K] | $s$ | $R_{\text{d}}$ [s$^{-1}$] |
|----------------|-----------------------------|-----------------------------|---------------|-------------------------|-----|--------------------------|
| I (Fig. II)    | 9                           | 9                           | $10^{12}$     | 2000                    | 0   | $7.2 \times 10^{12}$     |
| II (Fig. III)  | 9                           | 9                           | $10^{12}$     | 2000                    | 1   | $7.2 \times 10^{12}$     |
| III (Fig. III) | $\infty$                   | 0                           | $10^{12}$     | 2000                    | 0   | $7.2 \times 10^{12}$     |
| IV (Fig. IV)   | $\infty$                   | $10^9$                      | 2000          | 0                       | 0   | $7.2 \times 10^{12}$     |
| V (Fig. V)     | 9                           | 9                           | $10^{12}$     | 2000                    | 0   | $5.0 \times 10^{13}$     |

IV. COMPARISON WITH EXPERIMENTS

The Monte Carlo simulation results are compared with the experimental data on the Eley-Rideal reaction presented by Zecho et al. [28]. These experiments consist of two phases. First, a graphite substrate was exposed to a normal incidence deuterium beam with a flux of $3.8 \times 10^{15}$ atoms cm$^{-2}$s$^{-1}$ at 150 K. The exposure time was varied to give a range of initial coverages. The temperature of the atom beam was 2000 K. During the second phase the pre-exposed substrate was exposed to a normal incidence hydrogen beam and the formed HD molecules were measured using a mass spectrometer. The cross section of the deuterium abstraction with hydrogen, $\sigma$, was determined from these spectra using

$$\frac{d[H_2D_2]}{dt} = \Phi [D_{\text{ad},0}] \exp(-\sigma \Phi t),$$

with $\Phi$ the H flux and $[D_{\text{ad},0}]$ the initial D coverage. For the derivation of this equation, we refer to [28]. The expression is obtained assuming only direct Eley-Rideal. Since the reaction mechanism is implicitly included in the cross section, it is hard to directly interpret the results obtained in this way, but we will use the method to compare the experimental data with our simulation results.

The hydrogen adsorbate configurations found in the Monte Carlo simulations are also compared to the adsorbate configurations observed in STM experiments. In particular the fraction of hydrogen atoms in dimer configurations or larger clusters in simulation and experiment are compared.

V. RESULTS

Several simulation runs were performed using a similar set of pre-exposures as used by Zecho et al. [28]. The H and D fluxes were $4.8 \times 10^{13}$ cm$^{-2}$s$^{-1}$ at a temperature of 150 K. 250 $\times$ 125, 500 $\times$ 250, and 750 $\times$ 500 chemisorption sites were used depending on the initial D coverage. This corresponds to an array of 250 $\times$ 250, 500 $\times$ 500, and 750 $\times$ 750 to accommodate the extra physisorption sites. If the noise was still considerable for arrays of 750 $\times$ 500 multiple runs with different random seeds were made. In order to test to what extent the five proposed abstraction mechanisms contribute to the Eley-Rideal abstraction process comparisons to the experimental results, with different parameter sets (see Table III) were performed.

Figure 6 shows $d[H_2D_2]/dt$ for the five different abstraction mechanisms. Panel (I) only includes the direct Eley-Rideal with a barrier of 9 meV independent of the configuration on the surface, panel (II) investigates the influence of steering, panel (III) only allows an Eley-Rideal reaction, with a zero barrier, if the surface atom is part of a para-dimer configuration, panel (IV) uses the slower thermalization with $B = 1 \times 10^8$ s$^{-1}$ to test the dimer mediated mechanism, and panel (V) uses very fast diffusion. As $d[H_2D_2]/dt$ is plotted on a logarithmic scale, the curves should be linear according to Eq. (12) with the slope the cross section times the flux. The flux has the same value throughout the simulations.

The key results of the simulations are summarized in Table IV and compared to the experimental findings. Each entry consists of nine individual simulations. The top twelve rows represent the runs that allow only a single mechanism. The second section summarizes simulation series of combinations of mechanisms and the third part gives the results with a different sticking barrier and will be discussed in Section VI. The final entries give the experimental results for comparison. The difference in cross section between high and low coverage, the saturation coverage, and the linearity of the initial signal in initial coverage are indicators of the agreement with the experiment. The latter linearity is measured by the Pearson correlation coefficient that gives one for perfect correlation and zero for no correlation. For a better comparison between the different simulation runs all cross sections at low coverage are given for $\sim 0.013$ ML which is generally achieved after an exposure of 0.03 ML (the lowest considered pre-exposure). Simulations including mechanism IV (dimer mediated reaction) need a longer exposure to reach 0.013 ML whereas simulations with mechanism V (fast diffusion) reach it faster than 0.03 ML. The cross sections at high coverage are taken at the highest considered pre-exposure of 2.9 ML. The tenth column gives the corresponding coverages. In almost all cases the saturation coverage has been reached at this point. The last column of the table indicates the fraction of atoms that are part of a dimer configuration or larger cluster at a coverage of 0.5 % at conditions similar to the ones used by Horneck at al. [24].
Mechanism I: Direct Eley-Rideal

The five plots in Fig. 6 all show a very different behavior. Panel (I), which tests the direct Eley-Rideal mechanism, shows a linear relation between $\ln \left( \frac{d[HD]}{dt} \right)$ and the deposition time over the whole time range. The slope appears to have some dependence of the initial D coverage. This would result in a cross section that is dependent of initial coverage. Figure 7 plots, among other quantities, the cross section which indeed shows some coverage dependence. It is however much less than seen in the experiments [28]. The cross section and the value of the initial signal is obtained by fitting the curves shown in Figure 6-I to Eq. 12. Since for some parameter choices only the first section is linear, only these points are included in the fit. The number of points included in the fit can have a strong effect on the final result, especially if the linear part is very short. This is reflected by the error bars. Various runs with different initial seed showed that this error should at least be 0.5 Å². The initial signal depends linearly on the deuterium coverage, which agrees with Figure 3c in [28]. Figure 7 also displays the total HD yield. Since this value is comparable with the initial D coverage it is clear that only a negligible fraction of D atoms remains on the surface after hydrogen exposure. This is in good agreement with the experimental findings [28]. The dimer fraction is very low in contrast with the experiments as can be seen in the first row in Table IV.

Mechanism II: Eley-Rideal with steering

A simple version of Eley-Rideal abstraction with steering is implemented by allowing atoms that land on empty sites close to chemisorbed hydrogen atoms to attempt a reaction with these atoms. For the reaction barrier the same $E_{ER, \text{mono}}$ is used as for the direct hit. Eley-Rideal via a direct hit (mechanism I) is also allowed. If an atom lands on an empty site, the possibility of steering is considered by comparing a random number between 0 and 1 against parameter $s$. If the random number is smaller than $s$, then steering is activated. In the case of steering, the three neighbouring sites are checked for reacting species. If two or more different atoms are found, another random number determines with which of the atoms the incoming H atom will undergo the reaction attempt. Simulations for $s = 1$ and 0.5 were performed. The results of these two simulation runs are summarized in Fig. 8 and the rows indicated by II in Table IV and the $s = 1$ simulation run is shown in Figure 6-II. Fig. 8 shows the initial coverage and yield after 200 seconds of H exposure as function of the initial D pre-exposure. Notice that the yield is given in ML in these graphs and not as a percentage. It further gives the cross section and initial signal as a function of the initial coverage. For $s = 1$, a reasonably high cross section at low coverage is obtained (14.7 ± 1.4 Å²), but the saturation coverage is very low (0.10 ML) and also the cross section at this coverage is too high. The series which simulates steering with $s = 0.5$ gives somewhat lower cross section resulting in better agreement with the experimental findings at high coverage and too low a cross section at low coverage. Furthermore, an almost linear decrease in cross section is found as a function of coverage in disagreement with the experimental findings. Again the dimer ratio is too low.

Mechanism III: Dimer Eley-Rideal

The curves in panel (III) are not linear and show a maximum at later times for the low coverages. Since this panel presents the results with the dimer Eley-Rideal mechanism, this maximum indicates the time at which a maximum of D containing dimers is formed. This behavior contradicts the experiments [28] where the [HD] signal decreases over time and is therefore not likely to be the primary mechanism involved in the abstraction.
Mechanism IV: Dimer mediated abstraction

Panel (IV) presents the results for the dimer mediated mechanism. Here a clear linear dependence can be observed for very early times with a decreasing slope for increasing initial coverage. To obtain this graph large arrays up to 1500 × 3000 sites were used due to the low sticking rate at low coverage. Notice that the y-axis range of this panel is different from the others. Figure 7 studies this set of simulations more closely. The upper two panels show that not all D is converted into HD, but that a substantial amount of deuterium still resides at the surface after the 200 seconds exposure. This is in contrast with the experimental finding that all deuterium is abstracted after a 3 ML dose. Figure 9 also shows the cross section determined from Fig. 6-II. Again the bars indicate the uncertainties in obtaining the slope. Since the linearity is much less than for mechanism I, the error bars are larger than in Fig. 2. In contrast with this latter figure, Figure 9 shows a strong dependence of the cross section on the initial deuteron coverage. This corresponds with the trends observed by Zecho et al. The values are also in the correct range, although the cross section is too high at low coverage and too low for high coverage. Both the cross section and the initial signal indicate that the dimer mediated mechanism becomes very inefficient for high coverages. The initial signal even decreases for increasing surface coverage indicating that at high coverage it is hard to make dimers due to the decreasing number of available sites. As the coverage increases, either by H or by D atoms, the mechanism becomes less efficient and the deuterium atoms will not be abstracted. This also results in a very high saturation coverage. The dimer fraction is higher as compared to the other mechanisms, although still much too low. The increase in dimer fraction is due to the fact that many of the chemisorbed atoms desorb due to the initial energy. Since atoms in dimer position are less likely to desorb because of their higher binding energy, this results in an elevated dimer ratio.

Mechanism V: Fast diffusion

As discussed above, the barriers to diffusion in the chemisorbed state are quite high making this process essentially negligible at least at low coverage. Diffusion in
the physisorbed state is, however, a completely different matter. If a simple expression for an activated process is assumed, then an atom thermalized to a surface temperature of 150 K will make around 20 hops, which corresponds to a distance of 5 Å before desorbing. However, as Bonfanti et al. [21] pointed out diffusion is not an activated process, since the activation barrier of 4 meV is negligible at these temperatures. This asks for a different estimation of the moving rate. As a first approach we took the diffusion coefficient obtained by Bonfanti et al. [21]. This results in a moving rate of $1.3 \times 10^{13}$ s$^{-1}$. The upper limit for diffusion will be free movement in two dimensions according to the average gas phase velocity. This results in a diffusion rate of $5.0 \times 10^{13}$ s$^{-1}$ at 150 K. We will perform simulations using both rates. The results for the highest rate and $E_{ER,\text{mono}} = 9$ meV are shown in Figures 6 and 10. Results of other simulation series with different combinations of $E_{ER,\text{mono}}$ and the diffusion rate are summarized in Table IV. A clear increase in the dimer fraction can be observed for increasing diffusion rate, although still slightly too low compared with experiments. The cross section curve as a function of the coverage has the correct dependence although it is still not high enough at low coverage. Furthermore the initial signal as a function of coverage does not have a linear dependence but exhibits a kink around 10 %. At this coverage the number of trimers starts declining in favour of the number of tetramers. Most of the configurations (60 %) are now a configuration of three or higher. This non-linearity could well be an artifact of the simplified treatment of reaction and sticking for these configurations which makes the model inaccurate at high coverage as discussed above.

**Combined mechanisms**

The experiments cannot be explained by one of the five mechanisms separately. Mechanism I is not efficient enough at low coverages, mechanism II leads to very low saturation coverages, mechanism III shows a maximum at $t > 0$, mechanism IV is not efficient at high coverages. Moreover, all these four mechanisms show very low dimer fractions compared with experiments. Mechanism V has an increased number of dimers and a higher cross section, but again this is not high enough to match the experimental observations. We therefore investigate different combinations of the five mechanisms.

Again the results are summarized in Table IV. The table clearly shows that a combination of mechanisms IV and V gives the best results, both in terms of dimer ratios and cross section. The best agreement is obtained for the high diffusion rate and $E_{ER,\text{mono}} = 9$ meV (in bold face). The detailed analysis of these simulations is given in Figure 11. The cross section has the correct coverage dependence, both in trend and in value. Also the saturation coverage of 0.4 ± 0.2 ML found experimentally is reproduced. The major discrepancy is again the initial HD signal which does not depend linearly on the initial cross section, but as mentioned earlier this can be an artifact due to the simplified assumptions made for sticking and reaction for complex configurations, which make high coverage simulations inaccurate. The dimer ratio is still below the experimental values, but much higher than all previous series. The dimer ratio includes both ortho and para dimers, but the main contribution is from the para dimers (40 times more at 0.1 %), since these sites have no barrier for sticking. The residence time of the physisorbed atoms at ortho sites is not long enough for the atoms to easily enter the chemisorbed state. As the dimers have been observed in more equal ratios, the rate for crossing this barrier is probably higher, possibly due to a lower barrier or higher pre-exponential factor than used in the present model or due to a longer residence time at the ortho site because of the irregularity in the lattice caused by the first hydrogen atom. The reason that this combination of mechanisms give higher cross section at low coverage is because more reaction events go via a dimer state. In this state an atom gains some energy due to the strong binding energy and this extra binding energy is used to overcome the barrier for reac-
VI. SENSITIVITY OF THE MODEL ON PARAMETERS

By comparing the results listed in Table IV it is clear that a combination of mechanisms IV and V reproduces the experimental findings quite well. This section will discuss the influence of the parameter settings on the final result. The table already showed four different parameter choices varying the diffusion rate and the barrier for abstraction. The diffusion rate has, as mentioned before, a clear influence on the number of formed dimers, but it appears to have very little effect on the obtained cross sections for the mechanisms IV and V. Lowering it to a very low value of 7.2 · 10^{11} \text{s}^{-1}, which turns it into a combination of mechanisms I and IV, results in a lower cross section at low coverage. We found that reducing the diffusion rate even further eventually leads to a completely flat cross section and a very high Pearson correlation coefficient. Also for a pure mechanism V a higher diffusion rate gives a stronger coverage dependence of the cross section as was suggested by Bonfanti et al. [21].

The abstraction barrier has no effect on the dimer ratio, but it influences the cross section at low coverage. It generally gives a lower cross section for a higher barrier. For a pure mechanism I also a barrierless abstraction was considered (see Table IV). The cross section at low coverage did not raise to the values found in the experiments in this case.

The parameter $B$, which controls the thermalization of the sticking atoms, has a very narrow parameter range if only mechanism IV is considered. Lower values than 1 · 10^9 \text{s}^{-1} result in very low sticking at low coverage and higher values will have only a limited effect on the cross section. If mechanism IV is used in combination with mechanism V the sticking will be increased because of the diffusion of the physisorbed atoms and also lower values than 1 · 10^9 \text{s}^{-1} can be used leading to a higher cross section for low deuterium exposure and high dimer ratios. The linearity in the initial HD mass spectrometer signal is however decreased.

Adding mechanism III as a third mechanism in the simulations had a negligible effect on the results. The cross section and initial signal curves remained unchanged within their uncertainties.

The parameter, $T_{\text{start}}$, which controls the initial energy of hydrogen atoms, has a different effect on the final results for the pure mechanism IV and the combination of IV and V. This parameter cannot be controlled independently of $B$, since the sticking probability is constrained within a certain window (see Fig. 3). For a pure mechanism IV a increased value of $T_{\text{start}}$ results in a cross section which is too low and only weakly dependent on coverage. The correlation coefficient for the initial HD signal is higher as compared to $T_{\text{start}} = 2000$ K. This parameter combination further shows an extremely high dimer ratio. Only combinations of mechanism were found to yield similar high values. It appear that the high initial energy is only used to remove monomers and not for dimers to react or to desorb. The relaxation time is probably too fast for these events to occur. If a more elaborate implementation of this mechanism was considered, where the initial energy of atoms in dimers is high, because of their higher binding energy, the reaction from dimers would go up and the dimer ratio go down. The results would then probably closer resemble the $T_{\text{start}} = 2000$ K results. For a combination of mechanisms $T_{\text{start}}$ has less of an effect. Here only the saturation coverage shows a large change.

Finally we checked the influence of increasing the monomer sticking barrier to the value of 0.25 eV which is an upper value for the sticking barrier reported by other authors [12, 17, 18]. This results in a higher cross section at low coverage and a higher dimer ratio in all simulated cases. However, the correlation coefficient of the initial mass spectrometer signal as a function of coverage goes down. In general this increase in the sticking barrier results in a better agreement with the experiments. Although the agreement of the pure mechanism V is improved much, the combination of mechanism IV and V remains the best option for both low and high values of $E_{\text{stick}}$.

VII. CONCLUSION

The Eley-Rideal abstraction of atomic hydrogen chemisorbed on the graphite surface has been studied via a hybrid approach using energy barriers derived from DFT calculations as input to Monte Carlo simulations. Through comparison with experimental data we discriminate between the contributions from different proposed Eley-Rideal mechanisms. Good quantitative and qualitative agreement between the experimentally derived and the simulated Eley-Rideal abstraction cross sections are found if two different Eley-Rideal abstraction mechanisms are included. One is a direct Eley-Rideal reaction with very fast diffusion of physisorbed H atoms leading to the formation of hydrogen dimer configurations, while the other is a dimer mediated Eley-Rideal mechanism with increased cross section at low coverage. Such a dimer mediated Eley-Rideal mechanism has not previously been proposed.

The effect on abstraction of fast diffusing physisorbed H atoms was first considered by Boufanti et al. [21], who suggested that diffusion of the physisorbed atoms could explain the high coverage dependence of the cross section. We tested this mechanism and it was indeed found that diffusion plays an important role, not only for the abstraction reaction but also to explain the high occurrence of dimers found by STM measurements. However,
very little is known about the exact energetic landscape in the vicinity of a chemisorbed hydrogen atom, especially when the second atom is only weakly physisorbed. Since these weak interactions appear to be very important, further study would be desirable.

Furthermore, we investigated the effect of steering [16, 17] as a possibly alternative route to reproduce the experimental findings. The results presented here show that a simple steering mechanism which just results in increased cross sections for abstraction reactions is not sufficient to reproduce the experimental results. This finding does, however, not rule out steering as an important mechanism. It has for example been suggested that a coverage dependent steering effect could result from the creation of and interaction with electron-hole pairs on the surface [36]. Further investigations of steering should, however, consider the presence of preferred binding sites in the vicinity of adsorbed hydrogen atoms. In the present case, sticking in a dimer position with low or no barrier is a competing reaction to the steered Eley-

![Table IV: Simulation results for different parameter settings](image)

![Diagram](image)
Rideal reaction. Hence, the described dimer mediated Eley-Rideal mechanism offers a competing mechanism to steering. Full quantum mechanical calculations where both channels, dimer formation and hydrogen abstraction, are accessible, are needed to settle this case.

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