Accurate Probabilities for Highly Activated Reaction of Polyatomic Molecules on Surfaces Using a High-Dimensional Neural Network Potential: CHD$_3$ + Cu(111)

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Supporting Information

ABSTRACT: An accurate description of reactive scattering of molecules on metal surfaces often requires the modeling of energy transfer between the molecule and the surface phonons. Although ab initio molecular dynamics (AIMD) can describe this energy transfer, AIMD is at present untractable for reactions with reaction probabilities smaller than 1%. Here, we show that it is possible to use a neural network potential to describe a polyatomic molecule reacting on a mobile metal surface with considerably reduced computational effort compared to AIMD. The highly activated reaction of CHD$_3$ on Cu(111) is used as a test case for this method. It is observed that the reaction probability is influenced considerably by dynamical effects such as the bobsled effect and surface recoil. A special dynamical effect for CHD$_3$ + Cu(111) is that a higher vibrational efficacy is obtained for two quanta in the CH stretch mode than for a single quantum.

A ccurately describing molecule–surface reactions is of vital importance for the understanding of heterogeneously catalyzed processes such as the Haber–Bosch and steam reforming processes. Unfortunately, the complexity of the interaction between molecules and metals limits the accuracy of theoretical studies on these kinds of processes. Often, chemically accurate results are obtainable at high computational cost with ab initio molecular dynamics (AIMD) combined with the so-called Specific Reaction Parameter (SRP) approach. However, the investigation of reactions with low reactivity (<1%) remains challenging due to the need for a large number of trajectories in combination with a large computational cost. Therefore, neural network approaches have recently been employed in order to obtain results with the accuracy of AIMD using density functional theory (DFT), but with a considerably smaller computational cost. So far, these studies either involved diatomic molecules or they neglected the movement of surface atoms. Very recently, a high-dimensional neural network potential (HD-NNP) has been developed for a system in which a linear triatomic molecule interacts with a metal surface, i.e., CO$_2$ + Ni(100), also including surface atom motion. The neglect of surface motion can limit the accuracy of these studies due to the neglect of energy exchange between the molecule and the surface. This lack of energy exchange represents a severe approximation for the dynamics of polyatomic molecules reacting on metal surfaces due to their high mass. A modified Shepard interpolation method has also been used to describe the potential of a polyatomic molecule reacting on a metal surface but again with the neglect of surface motion. Reactive force field fits have been made that do include surface motion, but the quality of these fits remains unclear. However, no neural network potential has been employed so far for nonlinear polyatomic molecules interacting with surfaces that explicitly includes the effect of surface motion as well.

In this work, we focus on the dissociative chemisorption of CHD$_3$ on Cu(111) because the system exhibits a low reactivity, making reactive AIMD studies untractable for most incidence energies achievable in molecular beams. Moreover, high-quality graphene can be synthesized using methane dissociation on copper, and this warrants additional study of the rate-controlling step, namely, the breaking of the first CH bond. The Eley–Rideal reaction of D with CD$_3$ preadsorbed on Cu(111) has also been studied. The methane + Cu(111) system shows interesting dynamics in that the low reactivity of methane on Cu(111) is not only caused by a high barrier (167 kJ/mol) but also by specific features of the potential energy surface (PES) such as the curvature of the minimum energy path (MEP). For all of these reasons, we apply the neural network Behler–Parrinello approach for the first time to a nonlinear polyatomic molecule reacting on a metal surface, which makes accurate simulations feasible while including surface motion, using CHD$_3$ + Cu(111) as an example.

In the HD-NNP, the total energy is evaluated as a sum of atomic contributions that are dependent on their energetically relevant local environment, which is described by many-body interactions.
atom-centered symmetry functions.\textsuperscript{44} In total, 38 000 DFT data points were used to train the HD-NNP, of which 14 000 points were taken from an AIMD study.\textsuperscript{12} Other data points included structures sampling the van der Waals well and transition state regions and the molecule’s vibrational modes. Finally, dynamically important structures missing from the data set were identified by running molecular dynamics (MD) on the (incomplete) HD-NNP and then added to the data set in a procedure described in ref \textsuperscript{14}. For the neural network, two hidden layers were used, each with 15 nodes. The training was carried out using the RuNNer code\textsuperscript{45-47} and the MD was performed with LAMMPS.\textsuperscript{48,49}

First, the accuracy of the HD-NNP is tested by comparing the 2D elbow plot of methane on Cu(111) in which methane is fixed in all molecular coordinates according to its transition state geometry, as depicted in Figure 1a, except for Z and r (the distance between the carbon and surface and the length of the dissociating CH bond). The HD-NNP is compared directly with DFT calculations in Figure 1b. Here we see that the HD-NNP reproduces the DFT data remarkably well, even though points from the 2D cut are not included in the data set. When the methane is relaxed in all degrees of freedom other than r and Z (Figure 1c), the MEP lies slightly closer to the surface than to the MEP of the constrained methane. Again, the HD-NNP reproduces the direct DFT calculations quite well. Moreover, both the electronic and mechanical coupling\textsuperscript{39,50} are in good agreement with DFT (see Figure 1d,e), which means that changes in the barrier height and geometry with respect to the motion of the surface atom below the dissociating molecule are described correctly. Furthermore, using 90% of the DFT data set as the training set and 10% as the test set, the root-mean-square error (RMSE) is 1.7 kJ/mol for the test set, which is well within chemical accuracy (4.2 kJ/mol). (Note that all errors reported in this work are with respect to the full system, i.e., the total energy.) The high fitting accuracy is also observed in Figure 2, where the distributions of the absolute error for the training and test set are shown and the vast majority of the errors falls within chemical accuracy. The total energy for all of the structures in the training and test set obtained with the HD-NNP and direct DFT calculations is also shown in Figure S1. Moreover, the RMSE for the forces in the test set is 2.3 kJ/mol/Å. The RMSE of 1.7 kJ/mol that we obtain here on the basis of 38 000 DFT points for CHD\textsubscript{3} + Cu(111) with the surface atoms allowed to move compares well with the RMSE of 1.5 kJ/mol obtained for a recent 15D NN static surface PES for CHD\textsubscript{3} + Ni(111) on the basis of 200 000 DFT points.\textsuperscript{22,30} We also note that the approximate modified generalized Langevin oscillator method used in ref \textsuperscript{30} to effectively add surface atom motion to the problem may run into problems if the molecule-surface interaction depends on more than just one surface atom coordinate, as for instance is the case for H\textsubscript{2}O + Ni(111)\textsuperscript{50} and may be the case for methane interacting with stepped metal surfaces.\textsuperscript{30}

The goal is to make a HD-NNP that is capable of accurately evaluating the energy and forces on the fly during MD simulations. Therefore, not only are incidence energies with low reaction probabilities (<1%) investigated but also regimes with higher reaction probabilities that are obtainable with AIMD in order to test the validity of the results obtained with the HD-NNP. Figure 3 shows the results obtained for the dissociative chemisorption of CHD\textsubscript{3} on Cu(111) with MD using the HD-NNP and with AIMD\textsuperscript{12} by simulating a molecular beam for the rovibrational ground state and under laser-off and laser-on conditions. Under laser-off conditions, the molecular beam’s vibrational state population is sampled according to the nozzle temperature, and under laser-on conditions, the CH stretch mode $\nu_3$ is excited with one or two
In order to describe the reaction probability with good statistics, 10,000–110,000 quasiclassical trajectories were computed per incidence condition. Here we see that at high incidence energy and for vibrationally excited methane, for four sets of initial conditions resulting in reaction probabilities obtained with AIMD, good agreement exists between AIMD and MD performed with the HD-NNP (see also Table 1 and the statistical analysis in the SI). Moreover, reaction probabilities as low as $5 \times 10^{-5}$ have been computed with the HD-NNP, which was previously not possible using accurate methods. It is observed that at the highest incidence energy (181 kJ/mol) the laser-off simulation yields a similar reaction probability as the $\nu_1 = 1$ simulation, which is caused by the high amount of vibrational excitation in the laser-off beam due to the high simulated nozzle temperature ($T_n = 1000$ K). However, it should be noted that sticking probabilities computed for laser-off conditions and nozzle temperatures higher than 650 K may be unreliable due to intramolecular vibrational energy redistribution (IVR) among vibrational states in which CD bends and stretches are excited.

Our dynamical simulations show that the reaction of methane is promoted by both translational and vibrational energy. Plotting the reaction probability as a function of the total energy (vibrational + translational energy) shows that putting vibrational energy into the reaction is almost equally or more efficient than increasing the translational energy, depending on the amount of quanta in the $\nu_1$ CH stretch mode (see Figure 3b). The vibrational efficiency is equal to or larger than 0.8, which can be expected for such a late barrier system$^{19}$ combined with a MEP of the shape shown in Figure 1c, causing incoming molecules to have to react over considerably higher barriers because they run off of the MEP (the “bobbed effect”$^{32,53}$). This could play a large role at catalytic conditions, where graphene is produced from methane using very high temperatures (>1200 K)$^{36,41}$ and thus vibrational excitation is prevalent. Interestingly, the vibrational efficiency$^{34,35}$ for the excitation from the $\nu_1 = 1$ to 2 overtone ($\eta_{\nu_1=2,1} = 1.7$) is considerably higher than that for the excitation from the ground state to $\nu_1 = 1$ ($\eta_{\nu_1=1,0} = 0.8$).

To our knowledge, a higher vibrational efficiency for an overtone has not been observed before.$^{34–38}$ In Figure S4, it is observed that when the incidence energy decreases, for $\nu_1 = 2$, reacted trajectories follow the MEP more closely. In Figure 4a, an increase of vibrational energy causes trajectories going on to react to follow the MEP more closely. The dynamical effect (see Figure 4b) is that, because a higher incidence energy is
needed to overcome the barrier for a low $\nu_1$, for low $\nu_1$, the carbon atom smashes into the repulsive wall. The hydrogen moves out while the carbon is still close to the surface, and therefore, a higher barrier needs to be overcome (see Figure 4). Hence, a higher vibrational efficacy is observed for $\nu_1 = 2$ because the bobsled effect will be less prominent and thus lower barriers need to be overcome.

We already noted that the reaction probabilities at high incidence energy obtained with the HD-NNP are in good agreement with AIMD. However, the amount of quasiclassical approximation for the low reaction probabilities needs to be tested by comparison to experiment due to the possibility of quantum effects and potential problems with zero-point energy violation. Even though it has been shown that at elevated surface temperature the reaction of methane happens in a “classical over the barrier fashion” with assistance of surface atom motion and without the need for tunneling.

The main goal of applying the Behler–Parrinello method to polyatomic molecules is to be able to explicitly include surface motion. Therefore, to evaluate the effect of surface motion, reaction probabilities for $\nu_1 = 2$ have also been computed using a static surface model, where the surface was kept in its ideal relaxed static configuration (note that the lattice expansion corresponding to a surface temperature of 550 K was kept). This effectively removes energy transfer between the molecule and the surface in barrier heights and positions related to surface motion. Reaction probabilities for this frozen surface are a factor 2 higher than those when surface motion is allowed (see Figure 3c). Furthermore, when the distortions of a hot surface are included while still excluding surface motion, i.e., modeling a static thermally disordered surface (similar to the so-called static corrugation model), reaction probabilities are increased by 50% compared to the frozen ideal surface at low incidence energies.

At high incidence energies, no difference is observed between the results for the static ideal and the distorted surface, with the latter including the effect of the electronic coupling (or the so-called $\beta$-coupling). Our observation that explicitly including surface motion at these high incidence energies lowers the reaction probabilities suggests that the reaction probabilities are decreased due to energy transfer to the surface as the molecule first impacts on the surface (Figure 4b) and possibly also due to surface recoil (mechanical coupling).

Because the surface recoil effect (which is due to surface atom vibrational averaging) tends to be small, we suspect that the energy transfer is most important. This effect can only be addressed with explicit modeling of the surface motion and not by the sudden and energy averaging methods typically used with quantum dynamics simulations.

To summarize, in this work, the Behler–Parrinello approach was used to develop a HD-NNP that describes a polyatomic molecule reacting on a mobile metal surface, i.e., CHD$_3$ + Cu(111). The HD-NNP was found to be in good agreement with DFT, which means that MD can be performed with the accuracy of AIMD but with a considerably lower computational effort. Using this HD-NNP, reaction probabilities as low as $5 \times 10^{-5}$ were obtained, which was untractable for previous accurate methods such as AIMD, while including surface motion. It was found that vibrational excitation plays a major role in the reactivity, where the overtone has a higher vibrational efficacy than the fundamental vibrational excitation. Moreover, allowing energy transfer from the molecule to the surface considerably reduces the overall reactivity. Hence, surface motion needs to be included explicitly in simulations in order to obtain quantitative results for molecular beam simulations of methane reacting on copper. More work is still required to investigate the effect of surface temperature on the reaction of CHD$_3$ on Cu(111) as we addressed only one surface temperature (550 K). Finally, the quasiclassical approximation needs to be tested for low reaction probabilities by comparison to experiments due to the possibility of quantum effects and zero-point energy violation. However, this would not be an intrinsic problem of the HD-NNP as good agreement with DFT has been shown.

**Computational Method**

The DFT calculations used the same computational setup that was used in an earlier AIMD study, which is summarized here. A $3 \times 3$ Cu supercell with five layers and 13 Å vacuum distance is used, where the bottom two layers are fixed and the metal atoms in the other layers are allowed to move in order to simulate a surface temperature of 550 K. Furthermore, a plane wave cutoff of 350 eV and a $6 \times 6 \times 1$ Γ-centered k-point grid are used. All calculations are performed with the Vienna Ab-initio Simulation Package (VASP version 5.3.5) with the SRP32-vdW functional.
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ASSOCIATED CONTENT

Supporting Information
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Description of the training data set and fitting error, employed symmetry functions for the HD-NNP, initial conditions of the molecular beam, elbow plots, and statistical analysis (PDF)

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Notes
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