Efficient N₂ Formation on Ag(111) by Eley–Rideal Recombination of Hyperthermal Atoms

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Abstract

Using molecular dynamics simulations and potential energy surfaces of *ab-initio* quality, we show that direct pick-up of N adsorbates by gas-phase N is a highly efficient channel for N_2 formation on Ag(111). This recombination process, called Eley-Rideal, was traditionally associated to lighter projectiles and regarded as marginal, but here we obtain reactivities for

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N of $\gtrsim 35\%$ even at incident energies of a few eV. The good agreement found between simulated and published experimental energies of the desorbed N₂ is a fingerprint of this otherwise elusive recombination.

TOC Graphic



Keywords: Gas-surface interactions, Eley-Rideal reactions, scattering, potential energy surfaces, molecular dynamics.

Among the various elementary processes that occur in everyday gas-surface reactions, those resulting in the recombination and desorption of molecules deserve a special attention, as they regulate the replenishment of the surface active sites. One of these processes, the direct Eley-Rideal (ER) recombination has been found to be a marginal phenomenon, only observed experimentally for incoming light atoms (H,D). ^{1–9} Originally postulated as an atom exchange, ^{10–12} the ER process is nowadays assigned to the direct recombination of an incoming gas-phase species that picks up a surface adsorbate to conform a molecular compound that desorbs during the collision. Even if the initial measurements pointed to ER cross sections of about 2-5 Å², ^{4,6,9} subsequent theoretical studies demonstrated that they were around one order of magnitude smaller because most of the recombination events proceed after few collisions of the incoming (H,D) atom with the surface, i.e., through the slower hot-atom (HA) process. ^{13–18}

In the scope of reactive ion scattering (RIS), C abstraction from a graphite surface by N⁺ ions has been characterised by experiments, and found to happen after several collisions with the substrate¹⁹ Recombination of neutral atomic species heavier than (H,D) has been experimentally studied in the context of CO formation and CO oxidation on metal surfaces. In all cases, CO and CO₂ formation is dominated by the Langmuir-Hinshelwood (LH) recombination in which the species involved in the reaction are fully thermalized with the surface.^{20–22} Also in these systems, the direct ER pick-up events are accepted to play a very minor role. Within this context, the recent experiments by Ueta *et al.* suggesting the existence of direct recombination on the N-covered Ag(111) surface between gas-phase and adsorbed N atoms are certainly intriguing.^{23,24} All the more so when recent simulations of N₂ formation on the W(100) surface also exhibit low ER probabilities.²⁵

In this letter we demonstrate that thermal and hyperthermal gas-phase N scattered off Ncovered Ag(111) is indeed a prototype system that fulfills all the requirements to observe a highly efficient direct ER recombination. This outcome poses a new scenario for the study of gas-surface interactions, as it shows that fast direct recombination channels may be active in contrast to the usual assumption of neglecting them, specially for large atomic projectiles. The complex modeling of the ER mechanism has been successfully attained by a careful mixing of the *ab-initio* potential energy surfaces calculated for gas-phase N and N₂ impinging, respectively, on the (1×1) N-covered and clean Ag(111) surfaces. Importantly, our calculations nicely reproduce the recent measurements by Ueta *et al.* using N and N₂ mixed molecular beams reflected off clean and N-covered Ag(111).^{23,24} In these experiments a small amount of molecules is detected at outgoing polar angles $\Theta_f < 40^\circ$ off N-covered Ag(111), but not off clean Ag(111) (see filled and open triangles in Figure 1). The authors cautiously interpreted the existence of those molecules as an indirect evidence of a recombination process taking place on the N-covered surface. The calculations in the present letter prove unambiguously that the origin of those molecules is a surface reaction. Quantitative agreement with the experimental energy angular spectra supports the interpretation. Moreover, we show that at least 1/3 of the impinging atoms result in an ER event. All in all, the present molecular dynamics (MD) study confirms that Ref.²⁴ contains the first experimental fingerprint of N₂ formed by ER reaction, a process that up to now has been exclusively detected for incoming light atoms.

First, we have simulated the scattering dynamics of N₂ molecules off clean Ag(111). The interaction between N₂ and Ag(111) has been modeled by a six-dimensional (6D) potential energy surface (PES), which has been constructed from interpolation of *ab-initio* energies evaluated for a dense set of $\{x, y, z\}$ coordinates of the N-atoms and frozen Ag positions. Those *ab-initio* energies have been calculated within the density functional theory using ultrasoft pseudopotentials²⁶ and the PW91 generalized gradient approximation²⁷ for the exchange and correlation functional. The interpolation has been carried out using the corrugation reduction procedure method.²⁸ We find that the molecule-surface interaction is, overall, strongly repulsive. Further details on the calculations can be found in Ref.²⁹

Classical 6D MD simulations have been performed with molecular projectiles that are set to match the experimental settings, including incoming polar angles $\Theta_i = 60^\circ$ and a broad distribution of translational kinetic energies with average value $\langle E_i \rangle = 5.6 \text{ eV}$ and fwhm = 6.3 eV, i.e. we use an *effusive beam*.²⁴ Initially, the N₂ molecules are not rotationally or vibrationally excited.

An optimal description of the dynamics is achieved when including an energy loss channel into Ag lattice vibrations by means of the Generalized Langevin Oscillator (GLO) model^{30–33} using the same implementation of Refs., ^{34–36} where the surface vibration frequencies are taken from experimental values for Ag(111).³⁷ The agreement between theory and experiment is exceedingly good for the resulting angular distribution of final-to-initial translational energy ratios, $\langle E_f \rangle / \langle E_i \rangle$, as shown in Figure 1. The analysis of the angular distribution of scattered molecules together with additional calculations for monoenergetic beams explain the distinctive V-shaped experimental distributions, nicely reproduced here by theory. Since the molecules in the beam with higher E_i can access more corrugated regions of the PES, they produce broader angular distributions than the less energetic ones, as shown in the middle panel of Figure 1. For this reason, on average, the effusive molecular beam has a $\leq 40^{\circ}$ spread and, importantly, the tails of the distribution are dominated by the more energetic molecules. As a result the final-to-initial translational energy ratio increases the more the scattered N₂ departures from specularity. In agreement with the experiment, there is no hint of dissociative sticking in the simulations and no molecules are detected at $\Theta_f < 40^{\circ}$.

On the N-covered surface, however, a small amount of N₂ molecules leave the N-covered surface at $\Theta_f < 40^\circ$ with intensities about 50 times smaller than the specular peak.²⁴ The off-specular signal is consistent with a corrugated PES, as it is the case for N-atoms but not for N₂ molecules, which suggests an underlying molecule formation on the surface. In this respect, MD studies on the (1×1) N-covered Ag(111) surface with a 3D PES have shown that every adsorption trajectory ends up in a N(g)-N(a) bond, where N(g) and N(a) denote gas-phase N atoms and N adsorbates, respectively. Despite the limitations of a 3D description, which keeps N(a) at frozen coordinates, this theoretical result also points to N₂ formation.³⁶ The progressive cleaning of the N(a) via N₂ formation also explains the experimental observation that the N-atom scattering data on N-covered Ag(111) resemble that of the clean surface. Those recombination processes compete in turn with the replenishment of clean Ag(111) patches by N(g). Thus, a fundamental question that raises here is the efficiency of the recombination processes. We show next that the ER mechanism, which sets a lower boundary value of the total N₂ yield, is unusually efficient.

A good formulation of the problem should contain both the large potential corrugation brought by the N-coverage and N(a) mobility to account for eventual desorption events. With this in mind, our model PES uses energies of *ab-initio* quality and is constructed as sketched in Figure 2 by merging (i) the 3D PES for N(g) approaching the (1×1) N-covered Ag(111) for N(g)-N(a) distances $d > R_s$ and (ii) the 6D PES for the N₂ molecule approaching the clean surface for $d < R_s$. When N(g) crosses the boundary from the 3D to the 6D region, N(a) is allowed to move and may become an atom of a N₂ molecule. The choice $R_s = 2.2$ Å for the boundary ensures negligible discontinuities when integrating the equations of motion. Details on the PES continuity and the settings of the MD simulations performed on the merged PES can be found in the SI.²⁹

Figure 3 shows the fraction of atomic trajectories that result in N_2 formation by ER, where we observe that $\gtrsim 35\%$ of the projectiles in the experimental effusive beam used in the calculation succeed to react with surface species. It is a fast process with typical simulation times below <0.4 ps for the effusive atomic beam, and therefore the rigid surface approximation is well justified. Inclusion of GLO in the dynamics has a non-negligible impact, albeit small, in the reaction probability for $E_i < 2 \text{ eV}$. For those E_i values, energy exchange with the Ag(111) lattice reduces the ER efficiency by $\sim 10\%$. This uncommonly large efficiency, specially at incident energies as high as $E_i = 7 \text{ eV}$, is a non-trivial outcome of the dynamics of the system. Slower N(g) with $E_i < 2 \text{ eV}$ increase further the reaction probability. Low-energy projectiles visit lower energy regions of the PES, so that the dynamics is dictated by the highly exothermic and barrierless character of the reaction (9.8 eV are dissipated by a single N_2 formation, whilst only just 2.01 eV are needed to desorb each N(a)³⁸). On similar grounds, CO formation by ER at C-covered Pt(111) with $E_i(O) < 0.5 \text{ eV}$ was predicted to proceed with 97% efficiency by early MD simulations.³⁹ However, later molecular beam experiments showed that LH was the dominant mechanism for CO recombination in this system.²⁰ On N-covered W(100), the existence of an energy barrier makes N_2 formation by ER less efficient than on N-covered Ag(111), despite the reaction being also exothermic.^{25,40} In the case of $H_2/Cu(111)$, the H-H interaction is repulsive when the atoms lie close to the metal, and most recombinations are driven by HA.¹⁴ In other words, the efficiency of the reaction depends on the balance between several interactions: atom-surface, atom-atom, etc. The nature of the recombination mechanism relies thus in the chemical interaction between the neutral atoms on the surface. Effective ER with heavy ions has been measured in RIS. However, the ER scenario for ions in RIS is different to the one with neutral atoms, since the recombination mechanism is due to the electrostatic ion-adsorbate attraction (ion-dipole) and it is effective only for physisorbed atoms.^{41,42}

The angular dependence of the molecular translational energy is accounted for in the molecular beam experiments by Ueta *et al.*,²⁴ and it is in reasonable agreement with the theoretical value from MD simulations that use the atomic effusive incident beam, as shown in Figure 1 for $\Theta_f < 40^\circ$ inplane directions. At $\Theta_f > 40^\circ N_2$ molecules originated by ER contribute only to $\leq 3\%$ of the total N₂ intensity, and therefore those $\langle E_f \rangle / \langle E_i \rangle$ values are dominated by N₂ reflection. This is due to the fact that most formed N₂ molecules leave the surface at out-of-plane directions, and thus they are not to be observed in the experimental spectra. The angular distribution of inplane N₂ molecules formed by ER is shown at the top panel of Figure 1. These intensities are about two orders of magnitude smaller than the intensities of in-plane scattered N₂ (middle panel), in reasonable agreement with the experimental observations.²⁴ The theoretical $\langle E_f \rangle / \langle E_i \rangle$ values at $\Theta_f < 40^\circ$ are somewhat larger than the experimental ones because secondary recombination processes are not captured by the current PES model. These would yield, if any, slower products as the HA longer surface residence times allow for further energy dissipation into the substrate. The good agreement in $\langle E_f \rangle / \langle E_i \rangle$ strongly indicates that most of those observed N₂ molecules at $\Theta_f < 40^\circ$ result from a ER reaction.

Reactive trajectory tracking unveils the ER mechanism at the atomic scale. For a N(g)-N(a) pair, a ER reaction impact parameter, *b*, is defined as sketched in Figure 2. For a given target atom N(a), Figure 4 shows the initial impact parameter distributions of all the incident atoms, $P_0(b)$, and the reactive ones (i.e. those that will result in an ER event), $P_r(b)$. The potential outside the shaded region in Figure 2 is able to steer the N(g) projectile, deflecting it towards a target other than the one it was initially set to collide with. If the potential far from the surface were negligible, the N(g)

atoms would move along straight trajectories and would impinge on N(a) with impact parameters $R_{th} \leq b \leq R_s$ due to the shadowing effect of neighbouring adsorbates. The steering effect comes through as an E_i -dependent shift towards larger b values in the distributions, so that they start at $b > R_{th}$ and take values $b > R_s$. The lower the E_i , the stronger the steering, which evidences the long range attraction exerted by a dense N(a) coverage. The distribution $P_0(b)$ is maximal at $b \simeq 2$ Å, under the current incidence geometry. $P_r(b)$ is double peaked, and the peak position and width changes with E_i . The peaks in $P_r(b)$ can be assigned to two distinct recombination mechanisms. At large b values, the N(g) momentum component along the N(g)-N(a) direction is small, and therefore the net attractive force between both atoms acts long enough as to produce the adsorbate *abstraction*. At low b values, N(g) approaches N(a) fast enough to enter the repulsive region of the PES. A collision occurs that destabilizes N(a) from its equilibrium adsorption site (it retracts downwards by an average $\Delta z = 0.2$ and 0.4 Å for $E_i = 1$ and 4 eV, respectively) and the recoiling N(g) can easily drag it along. This peak height is smaller at larger E_i , since the recoil may be too fast for N(a) to follow the projectile. If we consider the quotient $P_r(b)/P_0(b)$ between the distributions of the reactive and all the incoming N(g)-atoms, it turns out that the collision (abstraction) mechanism has an approximate maximum efficiency of 75% (100%) at all E_i values. Nevertheless, the global likelihoods of these two mechanisms in the actual dynamics are very different, as only a few N(g) atoms are beamed onto the surface at small b initial conditions [see $P_0(b)$ curves in Figure 4]. Double peaked distributions have been reported in the formation of HCl on Au(111) by ER,¹⁵ too. In contrast, a small b mechanism is not possible at W(100), since repulsion between N-atoms exists at vertical distances 2.0 - 2.5 Å that deflects N(g) towards the N(a) sides.²⁵

The decrease in efficiency when energy exchange with the lattice is included in the model using GLO (see Figure 3) is mainly associated to the collision reactive channel. As shown in Figure 4 the low-*b* peak height is decreased upon inclusion of GLO, while the high-*b* peak of $P_r(b)$ remains unaffected. Since the lattice is allowed to move, a fraction of the energy released in the collision is transferred to the substrate and N(a) does not acquire enough energy to become unbound from the

hollow site. The N-N bond formation on Ag(111) is exothermic by $\sim 7 \text{ eV}$, thus large amounts of energy are released that are shared between the product and the substrate. On the one hand, we have checked that the formed N₂ products leave the molecule at high rovibrationally excited states. On the other hand, we can extract information on the average energy transferred to the substrate upon molecule formation by comparing the formed N₂ average kinetic energies in GLO and adiabatic frozen surface simulations. We find that this energy transfer is 1.57 eV for an incident effusive atomic beam of $\langle E_i \rangle = 4.3 \text{ eV}$. For a monoergetic beam of $E_i = 7.0 \text{ eV}$, 2.42 eV can be transferred to substrate lattice vibrations. This energy is large enough to induce desorption of N adsorbates and create hot atoms, increasing N₂ production by indirect mechanisms that are beyond the scope of the present work.

Since it is difficult to distinguish experimentally HA from ER products, ^{43,44} MD studies come in as a valuable tool in the study of gas-surface reaction pathways. They have shown that the ER cross-section for HCl formation on Au(111) is $\sigma < 1 \text{ Å}^2$ (about 10% of the HCl yield), ¹⁶ instead of the initially reported experimental value $\sigma = 2 \text{ Å}^2$.⁴ For (H,D) reacting on Cu(111), theory also yields smaller cross-sections ($\sigma < 0.5 \text{ Å}^2$, equivalent to 3.2-4.4% of the H₂ or HD products)^{13,14} than experiments $\sigma=5\,\text{\AA}^{2,2,5}$ Less than 5% of the N_2 formed on W(100) follows from ER with cross-sections $\sigma = 0.4 \text{ Å}^2$.²⁵ Considering this background, it is remarkable to find that the single collision ER route towards N_2 on Ag(111) with efficiency $>35\%~(\sigma>2.5\,\text{\AA}^2)$ as shown in Figure 3, even with the use of much more energetic projectiles than in the aforementioned reactions. A competingly efficient N-atom sticking process on clean Ag(111) is present for $E_i \leq 1.5 \text{ eV}$, but drops fast to negligible values for $E_i \ge 2 \text{ eV}$.³⁸ The ER efficiency obtained in this work is a lower boundary value to the total N2 yield, that may also result from HA or LH. Therefore, the surface cleaning may be significant also for projectiles below 2 eV and a fast adsorbate removal from the N-covered Ag(111) is foreseen that would explain the experimental observation that reflected Natom distributions on clean and N-covered Ag(111) are almost identical²⁴ despite the significant differences in their PES topographies.³⁶ An accurate description of the cleaning-replenishment cycles would require knowledge of the kinetic constants of the individual events. Nevertheless, since

 N_2 recombination probabilities are larger than N sticking probabilities on Ag(111),³⁸ it seems plausible that cleaning dominates as the experiment is being conducted on the covered surface.

In summary, we have shown that impinging N projectiles of ~eV energies on N-covered Ag(111) can result in the efficient formation of N_2 by fast pick-up events, also known as Eley-Rideal (ER) processes. We find that at least 35% of the incident N will recombine following this mechanism. This result sharply contrasts with what is known for other gas-surface systems where ER has a lesser weight and is typically restricted to light projectiles. The high N_2 yield found here is mainly due to the relatively long-ranged attractive and barrierless nature of the N-N interaction potential in the presence of the Ag(111) surface. This finding establishes the necessary conditions for efficient ER recombination and, importantly, poses a new scenario for many surface reactions, where now fast direct pick-up should be seen as a relevant product desorption channel.

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Supporting Information Available

The Supporting Information contains: (i) the settings used in the construction of the 6D PES of a N_2 molecule that interacts with the Ag(111) surface and (ii) a test on the trajectory continuity at the boundary R_s between the 3D and 6D PESs (see Figure 2). This material is available free of charge via the Internet at http://pubs.acs.org.

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Figure 1: Top panel: angular distribution of N_2 molecules formed by ER upon atomic N scattering on N-covered Ag(111), which leave the surface at in-plane directions in GLO simulations. Middle panel: angular distribution of N_2 reflected in-plane off clean Ag(111) in an adiabatic frozen surface simulation. In these two panels, the solid (dashed) line corresponds to an effusive (monoenergetic) beam. Bottom panel: angular distribution of the final-to-initial energy ratio of in-plane detected N_2 . Large open and filled triangles correspond to the experimental results of Ref.²⁴ on clean and N-covered Ag(111). Small symbols are theoretical results from MD using effusive incident beams. Small circles and squares are obtained from adiabatic frozen surface and GLO simulations, respectively, of N_2 reflected off clean Ag(111), and small diamonds correspond to N_2 formed by ER of N scattering on N-covered Ag(111). The inset shows the in-plane scattering geometry.



Figure 2: Schematic view of the incidence geometry used to simulate the ER events. The spheres with radius R_s denote the range around each surface adsorbate where the 6D PES is to be used. The impact parameter *b* is defined as the minimum distance between the N(g) incident direction at time t = 0 and the target N(a). The curved trajectory describes a case of projectile steering in the V^{3D} region.



Figure 3: The open symbols denote the efficiency of the ER process as a function of the incident N(g) kinetic energy E_i in simulations of atomic N scattering on N-covered Ag(111). Filled symbols account for the efficiencies obtained when an effusive atomic beam of $\langle E_i \rangle = 4.3 \text{ eV}$ is used. The ER cross-section, σ , can be formulated as the efficiency times the area per adsorbed atom.



Figure 4: Initial impact parameter distributions of all the incident N(g) atoms, P_0 (solid line), and the ones that result in ER reactions of atomic N on N-covered Ag(111), P_r , obtained from GLO (dashed line) and adiabatic frozen surface calculations (dotted line). The vertical lines indicate the geometrical threshold values (see text) for the PES boundary choice made in the present work, $R_s = 2.2$ Å.

Supporting information for: "Efficient N₂ Formation on Ag(111) by Eley-Rideal Recombination of Hyperthermal Atoms"

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Construction of the 6D PES

The interaction energy of the N₂ molecule with the Ag(111) surface is described with a full adiabatic six-dimensional (6D) potential energy surface (PES) that depends on the center of mass position $\mathbf{R}_{CM} = (X_{CM}, Y_{CM}, Z_{CM})$ and the internal coordinates (r, θ, φ) of the molecule. The PES is constructed from the interpolation of 6578 *ab-initio* energies corresponding to 11 values for *r*, 23 for Z_{CM} and 26 for different configurations of $(X_{CM}, Y_{CM}, \theta, \varphi)$.

All *ab-initio* data are obtained using the DFT "Vienna *Ab initio* Simulation Program" (VASP) code ¹ that operates with a plane-wave basis set. The exchange and correlation energy is calculated with the generalized gradient approximation (GGA) and the Perdew-Wang energy functional (PW91) .² The electron-core interaction is described with ultra-soft pseudopotentials .³ Preliminary calculations have been performed to ensure that all *ab initio* energies are calculated to a prescribed accuracy. Thus, the energy cutoff in the plane-wave expansion is 453 eV; the fractional occupancies are determined through the broadening approach of Methfessel and Paxton⁴ with *N*=1 and σ =0.2; and, given the surface's hexagonal geometry, the Brillouin-zone integration is performed with a 5x5x1 Gamma Centered Grid of special k points.⁵

The theoretical lattice constant obtained from a bulk calculation is a=4.17 Å. This leads to an interlayer spacing of d=2.408 Å. The Ag(111) surface is modeled by a 5-layer slab with a supercell vector along the normal to the surface (*OZ* axis) of 10*d*. Under these conditions and keeping the third layer fixed, the interlayer distance is relaxed to get the surface equilibrium geometry. These relaxation corrections are however found to be almost negligible (around 0.1%) for both the first and second interlayer distances. Once obtained, the relaxed geometry of the slab is kept frozen for the calculations that follow. The *ab-initio* energies for the N₂/Ag(111) system are obtained using a (2 × 2) surface structure (0.25 coverage). The energies are referred to that of the N₂ molecule located parallel to the surface and midway between the two slabs, which is the simulated asymptotic region.

Once the 6578 point energy grid is obtained, the PES is built by interpolating the *ab-initio* data. The Corrugation Reducing Procedure⁶ is used to obtain an auxiliary, smooth, and easy

to interpolate function F(X,Y,Z), which is spline interpolated along the Z_{CM} and r coordinates and 2D Fourier interpolated within the $X_{CM}Y_{CM}$ plane and the angular internal coordinates. The interpolated value of F(X0,Y0,Z0) can easily be converted back to an interpolated system energy E(X0,Y0,Z0). Note that the 2D Fourier expansion is here chosen over a 2D Periodic Spline in order to profit from the fact that it analytically respects the surface hexagonal symmetry. The accuracy of the constructed 6D PES has been checked by comparing its output to a set of *ab-initio* off-grid values. The errors are found to be of just a few tens of meV.

Derivative continuity checks at the boundary between the 3D and the 6D PES

Eley-Rideal (ER) recombination of gas-N, N(g), and adsorbed-N on Ag(111) at saturation coverage, N(a), has been simulated making use of two PESs calculated *ab-initio*. In each MD trajectory, the N(g) projectile departs from an initial height z_i =6 Å over the Ag(111) topmost layer and (x_i, y_i) coordinates generated by a conventional Monte Carlo (MC) sampling. Its incidence direction is defined by a fixed polar angle Θ_i =60°, measured with respect to the surface normal, and a random azimuthal angle Φ_i generated by MC sampling, too. The hyperthermal kinetic energies have been analysed in the present work are E_i =0.3, 1.0, 2.0, 4.0, 7.0 eV, plus an effusive beam of N(g) gas atoms that matches the experimental atomic beam energy distribution, with average energy $\langle E_i \rangle$ =4.3 eV and fwhm=5.4 eV.^{7,8} For each energy, N_{traj} =8 × 10⁵ trajectories are simulated that will be used to extract statistical averages. Classical molecular dynamics are performed by integrating the equations of motion with a time interval $h \leq 10^{-2}$ fs.

As explained in the manuscript, the simulation space is divided in two parts:

(i) Previously to the collision, at distances $R > R_s$ from the N(a) species, the N(g) dynamics are dictated by the 3D PES on the (1×1) N-covered Ag(111) surface, V^{3D} , which is described in detail in Ref.⁹ At this stage of the MD, surface atoms are kept fixed at their adorption sites.

(ii) When the projectile approaches the surface and the distance between the N(g) atom and a N(a) atom on the surface is $R < R_s$, the latter atom is allowed to move, and the interaction between these two atoms evolves under a 6D PES, V^{6D} , namely that of a N₂ molecule interacting with a clean Ag(111) surface. At this stage, recombination can take place.

Since the influence of neighbouring adsorbates is neglected at stage (ii) of the MD, we must dismiss from the analysis channels other than N_2 formation by an ER process with fast interaction with the surface. For the effusive atomic beam, we find average reaction times smaller than 0.4 ps. Other channels, such as adsorption or trapping, may prelude molecule recombination by hot atoms or Langmuir-Hinshelwood mechanisms at longer reaction times after interaction with several adsorbates.

The methodology above will constitute a reliable model for the ER reaction as long as N(g) experiences a small discontinuity when crossing the boundary between the two PESs at $R = R_s$. After a first stage evolving under V^{3D} , N(g) arrives at the boundary with position \mathbf{x}_s and velocity $\dot{\mathbf{x}}_s$. These are the initial conditions for the second part of the simulation with V^{6D} . The amount of energy not conserved because of switching to another PES, $\delta E(R_s)$, can be estimated from the discontinuity in the force experienced by N(g) at the boundary using:

$$\delta E(R_s) \approx h \dot{\mathbf{x}}_s \cdot \left[\nabla V^{3D}(\mathbf{x}_s) - \nabla V^{6D}(\mathbf{x}_s) \right] + \mathcal{O}(h^2)$$
(1)

Fig. Figure 1 shows the distributions of $|\delta E|$ for the studied trajectories, calculated for the largest used time step, h = 0.01 fs, and several values of E_i and R_s . In all the cases $|\delta E|$ is of the order of a few meV, much smaller than the incident energies. Distributions are broader for larger E_i and for smaller R_s . The choice made in the present work, $R_s = 2.2$ Å, ensures that the energy discontinuities will be $\leq 2\%$ of the E_i value with very few exceptions. Typically, the trajectories with large $|\delta E|$ are those that enter the $R < R_s$ region at small z values, i.e. they experience a stronger interaction with neighbouring N(a) in region (i) that is abruptly switched off when they enter region (ii).



Figure 1: Energy discontinuity distributions at the boundary between the 6D and 3D PES regions for E_i values studied in the present work. Each panel accounts for two locations R_s of the boundary.

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