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Origin of Power Laws for Reactions at Metal Surfaces Mediated by Hot Electrons

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A wide range of experiments have established that certain chemical reactions at metal surfaces can be driven by multiple hot-electron-mediated excitations of adsorbates. A high transient density of hot electrons is obtained by means of femtosecond laser pulses and a characteristic feature of such experiments is the emergence of a power law dependence of the reaction yield on the laser fluence $Y \sim F^n$. We propose a model of multiple inelastic scattering by hot electrons which reproduces this power law and the observed exponents of several experiments. All parameters are calculated within density functional theory and the delta self-consistent field method. With a simplifying assumption, the power law becomes exact and we obtain a simple physical interpretation of the exponent *n*, which represents the number of adsorbate vibrational states participating in the reaction.

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Hot-electron-induced femtochemistry at surfaces (HEFatS) is a class of chemical reactions, where the energy to overcome the reaction barrier is provided by energetic ("hot") electrons. These electrons are often photoelectrons emitted from the surface when struck by intense laser pulses, but other hot-electron sources are possible, such as scanning tunneling microscope tips and metal-insulatormetal nanodevices [1]. For these reactions, the reaction rate usually scales as a power law with the electron flux. For example, in a pioneering study of NO on Pd(111) [2], it was shown that femtosecond laser pulses could induce desorption of NO, and a mechanism by multiple electronic excitations was identified [3]. The yield depended on the fluence as $Y \propto F^n$ with $n \sim 3.3$. Subsequently, desorption induced by femtosecond lasers has been demonstrated for several other adsorbate systems [4-7] showing nonlinear vields, which can be fitted to power laws with exponents 3 < n < 8. It has also been shown that femtosecond laser pulses can induce surface hopping [8] and oxidation reactions [9–11], all of which can be characterized by power laws.

A popular theoretical approach to the interaction of adsorbates with a high density of hot electrons is the electronic friction model [12] where the excited electrons are assumed to equilibrate rapidly to a thermal distribution with electronic temperature T_e . For sufficiently large time scales the propagation of the adsorbate is well approximated by semiclassical Langevin dynamics with an electronic friction η_e depending on T_e as well as the adsorbate coordinates. While the friction model has certainly been successful in reproducing various experimental observations, it cannot account for the physical origin of the power law exponent *n*. Furthermore, measurements on the distribution of hot electrons in gold excited by subpicosecond laser pulses implies thermalization times up to 1 ps [13], which is on the order of reaction times, and there are

examples of laser induced surface chemistry where the assumption of a thermal distribution of hot electrons is in direct conflict with observations [5,11,14].

In this Letter we introduce a general model for firstprinciples calculations of the rates of HEFatS processes, regardless of the source of hot electrons. The central object of the model is a nonadiabatic Hamiltonian, which is used to calculate the hot-electron-induced vibrational transition probabilities of adsorbates. All parameters in the Hamiltonian are obtained from density-functional theory (DFT) and we show that the model reproduces experimentally observed power laws. Finally, we will make a simple approximation for the transition probabilities and show that the power law then becomes exact in the limit of large electron flux and that the exponent is given by $n = E_R/\hbar\omega$ where E_R is the energy barrier of the reaction and $\hbar\omega$ is the energy quantum of the vibrational mode dominating the energy transfer.

To analyze HEFatS rates involving multiple hot electrons, we consider a quadratic potential energy surface of an adsorbate with vibrational states $|n\rangle$ coupled to a localized electronic resonance $|a\rangle$. The probability for a hot electron with energy ε to inelastically scatter on the localized state and induce a vibrational transition $m \rightarrow n$ is denoted $P_{mn}(\varepsilon)$ and we assume a constant flux J_0 of hot electrons incident on the adsorbate. It is further assumed that each vibrational quantum has a fixed lifetime $T_{\rm vib}$ and that there exists a maximum quantum number n_R such that a reaction will proceed immediately if $n \ge n_R$. The probability that one vibrational quantum survives the time interval $\Delta t = 1/J_0$ between subsequent scattering events is then $e^{-\Delta t/T_{\text{vib}}}$. Each incoming electron will thus change the distribution of adsorbate vibrational states Q(n) until an equilibrium is reached between decay and reexcitation.

If the adsorbate is initially in the ground state, the distribution of vibrational states induced by the first elec-

tron with energy ε_1 is

$$Q_1(n;\varepsilon_1) = P_{0n}(\varepsilon_1). \tag{1}$$

The probability of the adsorbate being in the *n*th vibrational state after the second electron has scattered is

$$Q_2(n;\varepsilon_1,\varepsilon_2) = \sum_{m=0}^{n_R-1} p_1(m;\varepsilon_1) P_{mn}(\varepsilon_2), \qquad (2)$$

where $p_1(m)$ is the probability that the adsorbate is in the state *m* after the time interval Δt given by

$$p_1(m;\varepsilon_1) = \sum_{l=m}^{n_R-1} Q_1(l;\varepsilon_1) {l \choose m} (e^{-\Delta t/T_{\text{vib}}})^m \times (1 - e^{-\Delta t/T_{\text{vib}}})^{l-m}.$$
(3)

We exclude terms with $m \ge n_R$ since such excitations would have led to a reaction by assumption. Proceeding like this, the probability $Q_3(n; \varepsilon_1, \varepsilon_2, \varepsilon_3)$ of being in the *n*th excited state after the third scattering event can be expressed in terms of $Q_2(n; \varepsilon_1, \varepsilon_2)$ and so forth. Since all vibrational states $n \ge n_R$ lead to a reaction, the reaction probability of the *k*th electron is

$$P_k^R = \sum_{n=n_R}^{\infty} Q_k(n).$$
(4)

For large k this will approach a limiting value, P_R .

To calculate the vibrational transition matrix $P_{mn}(\varepsilon)$ we consider a Newns-Anderson-type Hamiltonian with substrate states $|k\rangle$ and a resonant state $|a\rangle$ linearly coupled to a number of vibrational modes with creation operators for vibrational quanta b_i^{\dagger} [15,16]:

$$H = \varepsilon_0 c_a^{\dagger} c_a + \sum_k \epsilon_k c_k^{\dagger} c_k + \sum_k (V_{ak} c_a^{\dagger} c_k + V_{ak}^* c_k^{\dagger} c_a)$$
$$+ \sum_i \hbar \omega_i \left(b_i^{\dagger} b_i + \frac{1}{2} \right) + \sum_i \lambda_i c_a^{\dagger} c_a (b_i^{\dagger} + b_i).$$
(5)

The model essentially describes a harmonic oscillator, which is displaced when the state $|a\rangle$ is occupied and the coupling V_{ak} to the metallic states introduces a finite lifetime of $|a\rangle$. As previously published [17], the transition probabilities $P_{mn}(\varepsilon)$ can be calculated exactly in the wideband limit where the density of states projected on the localized state $|a\rangle$ is a Lorentzian centered at ε_0 with full width at half maximum given by $\Gamma = 2\pi \sum_{k} |V_{ak}|^2 \delta(\varepsilon_0 - \varepsilon_0)$ ϵ_k). The probabilities $P_{mn}(\varepsilon)$ depend on the dimensionless parameters, $\hbar \omega_i / \Gamma$ and $g_i = (\lambda_i / \hbar \omega)^2$, and the reaction probability P^R also depends on these parameters in addition to the reaction quantum number $n_R \sim E_R/\hbar\omega$. The quantities E_R and ω can be calculated within standard density-functional theory and Γ is estimated from the projected density of states. The resonance energies as well as the nonadiabatic coupling parameters λ_i are obtained from excited state potential energy surfaces which are calculated with the method of linear expansion delta self-consistent field (Δ SCF). The method is a generalization of standard delta self-consistent field designed to handle molecular orbitals hybridized with metallic states and calculates the expectation values of excited states which are not eigenstates of the Hamiltonian but involve an occupied resonance. Details on the method and comparison with experiment can be found in Refs. [17,18].

As an example we calculate the fluence dependent transfer of energy from hot electrons to a CO molecule adsorbed on Cu(111), mediated by excitation of an unoccupied π^* orbital. CO adsorbs with the molecular axis perpendicular to the surface and the symmetry of the adsorbed molecule thus only allows a linear coupling to the center of mass (c.m.) and internal stretch vibrations of CO. The parameters for CO chemisorbed at a Cu(111) bridge site were calculated with the code GPAW [19,20], which is a realspace density-functional theory code using the projector augmented wave method [21]. We modeled the surface by a three layer (4×4) supercell with the top layer relaxed, a grid spacing of 0.2 Å, and a 4×4 surface K-point sampling. With the revised Perdew-Burke-Ernzerhof [22] functional, we find $\varepsilon_0 = 2.4$ eV, $\hbar\omega = 231$ meV, and $\lambda =$ -118 meV for the internal stretch vibration, and $\hbar\omega =$ 42 meV and $\lambda = -4$ meV for the c.m. vibration. The internal mode completely dominates the transfer of energy from hot electrons to the molecule since it has a much larger coupling λ , and the quantum of energy is 5 times larger than for the c.m. mode. Figure 1 shows the density of an excited top site molecule obtained with Δ SCF DFT relative to the ground state density and one clearly sees the excited π^* orbital. In Fig. 2 we have used (1)–(5) with hot electrons at $\varepsilon = 2.0$ eV corresponding to the laser frequency used in [4] to calculate reaction rates, which require energies corresponding to 3, 4, 5, and 6 internal vibrational excitations. In the nonlinear regime corresponding to reactions induced by multiple scattering

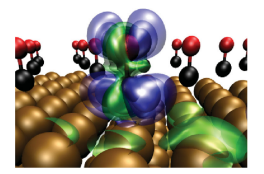


FIG. 1 (color). The difference between excited and ground state densities for CO adsorbed on Cu(111). Black balls are carbon atoms and red balls are oxygen atoms. Blue contours is excess density in excited state and green contours is excess density in ground state. The excited state is constructed by occupying a π^* orbital of CO. For clarity we only show the density difference in a single supercell.

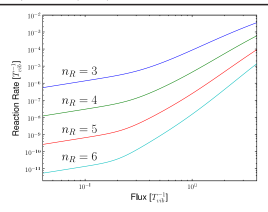


FIG. 2 (color online). The reaction yield of CO on Cu(111) as a function of hot-electron flux for four reaction energies. Note the initial linear dependence corresponding to single electron reactions.

events, the rates are very well approximated by power laws with n = 2.8, n = 3.6, n = 4.3, and n = 5.1. Varying the parameters in the model reveals that, in general, one always obtains very good power law fits with exponents $n \sim$ n_R . Increasing the energy of the hot electrons above the resonant energy ε_0 tends to decrease n.

Table I shows the calculated power law exponents for five systems compared with corresponding experiments. The DFT parameters used were the same as for CO on Cu(111). We have assumed that a reaction occurs when an energy of 0.2 eV in excess of the reaction barrier has been transferred, which is consistent with measurements of the kinetic and internal energy distributions of desorbed molecules [7], and we have used hot-electron energies $\varepsilon =$ $\varepsilon_f + \hbar \Omega$ where Ω is the laser frequency. The agreement is very good except for CO on Cu(100). This could be due to the role of frustrated rotations, which has previously been found to couple strongly to metallic electrons in this particular system [23]. In general, however, the frustrated rotations have a low energy compared with the internal stretch mode and we expect their effect to be neglectable. Because of symmetry the frustrated rotations cannot couple linearly to the resonant electron and one would

TABLE I. Power law exponents obtained from (1)–(5). The systems involving NO, CO, and O_2 are desorption experiments whereas O/Pt(111) is hot-electron-induced diffusion of atomic oxygen from a step to a terrace hollow site. For O_2 /Pt(111), the exponent depends on the laser frequency.

System	Experimental n	Calculated n	E_R
NO/Pd(111) [2]	3.3	3.7	1.0 eV
CO/Cu(111) [4]	3.7	3.6	0.4 eV
CO/Cu(100) [7]	8.0 ± 1	3.7	0.5 eV
$O_2/Pt(111)$ [5]	3.0/5.6	2.5/5.5	0.4 eV
O/Pt(111) [8]	15	12	0.8 eV

need a quadratic term like $\lambda_2 c_a c_a^{\dagger} (b + b^{\dagger})^2$ in (5) to include these in the model [24].

A particularly interesting case is the hot-electroninduced desorption of O₂ from Pt(111) where a power law with exponent $n \sim 5.6 \pm 0.7$ was observed using a photon energy of 2.0 eV and an exponent of $n \sim 3.0 \pm$ 0.5 using a photon energy of 4.0 eV [5]. The fact that the power law exponent depends on the photon energy contradicts the picture of a thermalized hot-electron gas interacting with the molecule, which is the basic assumption in models based on electronic friction. In contrast, the model (1)–(5) naturally gives rise to a decrease in the power law exponent when the energy of hot electrons is increased.

The transition matrix $P_{mn}(\varepsilon)$ has a very complicated structure and it is hard to extract the physics of the power law using these probabilities and (1)–(4). However, the magnitude of $P_{mn}(\varepsilon)$ is largely governed by the prefactor $g^{n-m}/(n-m)!$ [17] and in the following we will examine the consequences of assuming transition probabilities of the form

$$P_{mn} = e^{-\alpha} \frac{\alpha^{n-m}}{(n-m)!}, \qquad n \ge m, \tag{6}$$

where α is a dimensionless coupling constant, which describes the coupling of hot electrons to the adsorbate vibrational states. Repeated use of the algorithm (1)–(4) with these probabilities then reveals that to leading order in α one has

$$Q_k(n) = \frac{\alpha^n}{n!} \left(\sum_{j=0}^{k-1} e^{-j\Delta t/T_{\rm vib}} \right)^n.$$
(7)

We then consider a large flux $e^{-\Delta t/T_{\rm vib}} \sim 1 - \Delta t/T_{\rm vib}$, sum up the geometric series, take the limit $k \to \infty$ corresponding to steady state, and get

$$Q(n) = \frac{\alpha^n}{n!} (T_{\rm vib} J_0)^n, \tag{8}$$

where $\Delta t = 1/J_0$. Thus, for small α the reaction probability (4) will be dominated by such a term with $n = n_R$.

The power law emerges from summing up the detailed combinatorics of all possible ways of rising through the vibrational states in the potential well. In Fig. 3(a) we show the reaction yields for three values of n_R and they are seen to approach power laws of the form $Y \propto J_0^{n_R}$ for large fluxes. Even if α is not small, Eqs. (1)–(4) tend to conserve the power law although the exponent becomes reduced from the value of n_R when terms beyond leading order are not vanishing. In Fig. 3(b) we show the yield when $n_R = 15$ for $\alpha = 0.2$, $\alpha = 0.5$, and $\alpha = 1.0$. At large fluxes the yields are well approximated by power laws with exponents 14, 12, and 10, respectively. One might worry that the fixed time interval Δt between scattering events is too crude an approximation for the random nature of hot electrons interacting with the adsorbate. However, a sequence of time intervals $\{t_k\}$ with an average of Δt would

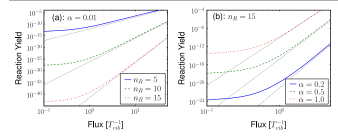


FIG. 3 (color online). The yield as a function of electron flux obtained using the transition probabilities in Eq. (6) for different parameters (see text).

lead to the replacement $j\Delta t \rightarrow \sum_{i=1}^{j} t_{k-i}$ in (7), which is well approximated by $j\Delta t$ for large *j*. We have repeated the calculations leading to Fig. 3 but with the time intervals randomly drawn from an exponential distribution $p(t) \sim e^{-t/\Delta t}$ and the power law is conserved on average.

The interpretation of the power law exponent as the number of contributing vibrational states can be used to identify the reaction channel of a given adsorbate system. For example, in the study of hot-electron-mediated desorption of NO from Pd(111) [2] a power law with exponent $n \simeq 3.3$ was found. The internal stretch vibration corresponds to an energy of $\hbar\omega \simeq 210$ meV whereas the other modes have vibrational energies $\hbar \omega \leq 70$ meV. Since the adsorption energy is $E_a \simeq 1.0$ eV we conclude that the power law exponent $n_R \sim E_a/\hbar\omega$ has to arise from sequential excitation of the internal stretch vibration and subsequent anharmonic energy transfer to the desorption coordinate. In contrast, the study of hot-electron-induced surface diffusion of atomic oxygen on Pt(111) [8] gave rise to a power law with exponent $n \simeq 15$, which fit very well with an experimental diffusion barrier of $E_d \simeq 0.8$ eV and vibrational modes on the order $\hbar \omega \sim 50$ meV.

In summary, we have presented a theory of multiple inelastic scattering with transition probabilities calculated in a nonadiabatic Newns-Anderson model, which lead to the ubiquitous power law of HEFatS and reproduce experimentally found exponents. The interpretation of the exponents as the number of contributing vibrational states is a very useful tool to identify the reaction channel of a given system and also indicates that a classical treatment of the adsorbate motion is insufficient. However, the model can only treat quadratic potentials and is thus not able to treat the anharmonic effects, which are expected to play an important role in the transfer of internal vibrational energy to the reaction coordinate. In the case of molecular desorption the model applies because the internal stretch mode, which is well approximated by a harmonic potential, dominates the energy transfer. Furthermore, while our assumption of constant hot-electron energy $\varepsilon = \varepsilon_f + \hbar\Omega$ gives the right dependence of the exponents on Ω , the hot electrons proceeding a laser pulse will undergo scattering and produce some distribution of electron energies. The true (time-dependent) distribution lies somewhere between the present assumption and a thermalized hot-electron gas and the model should thus be regarded as complementary to a statistical approach based on electronic friction and Langevin dynamics, which assumes a thermalized hotelectron gas.

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