Femtosecond-laser induced dynamics of CO on Ru(0001): New insights from a HOT-ELECTRON, ELECTRONIC FRICTION MODEL INCLUDING SURFACE MOTION

Robert Scholz^{1,2}, Gereon Floß¹, Peter Saalfrank¹, Gernot Füchsel³, Ivor Lončarić⁴, and J. I. Juaristi^{4,5,6}

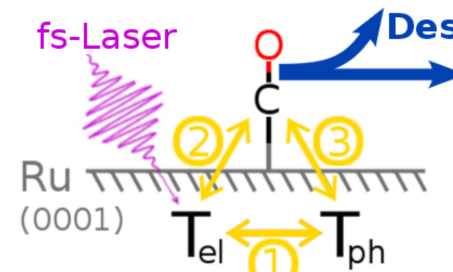
¹Institut für Chemie, Universität Potsdam, Karl-Liebknecht-Str. 24-25, D-14476 Potsdam, Germany ²Fritz-Haber-Institut der Max-Planck-Gesellschaft, Faradayweg 4-6, D-14195 Berlin, Germany ³Universiteit Leiden, Gorlaeus Laboratories, Einsteinweg 55, 2333 Leiden, The Netherlands ⁴Centro de Física de Materiales CFM/MPC (CSIC-UPV/EHU), Paseo Manuel de Lardizabal 5, 20018 Donostia-San Sebastián, Spain ⁵Departamento de Física de Materiales, Facultad de Químicas, Universidad del País Vasco (UPV/EHU), Apartado 1072, 20080 San Sebastián, Spain ⁶Donostia International Physics Center DIPC, P. Manuel de Lardizabal 4, 20018 San Sebastián, Spain

Introduction

Motivation

- research on small molecules adsorbed to metals is important for:
- -catalytic applications
- -fundamental understanding of bonding
- femtosecond(fs)-lasers are a valuable tool for such research as they
- -allow for investigations on small timescales
- open up new processes compared to heating (femtochemistry)
- -may enable specific control over catalytic reactions (photocatalysis)
- specific motivation for system CO/Ru(0001)
- -experimentally well studied regarding fs-laser irradiation, e.g. [1, 2]
- -fulldimensional ab-initio potential recently developed in our group [3] -details of this indicate interpretation of experiment [2] may be wrong

How does fs-laser-irradiation affect metal surfaces?



Desorption

Diffusion (and possibly Reactions)

(1) Electron-phonon coupling

....low T_{el}

t/ps

∠ 5000 -

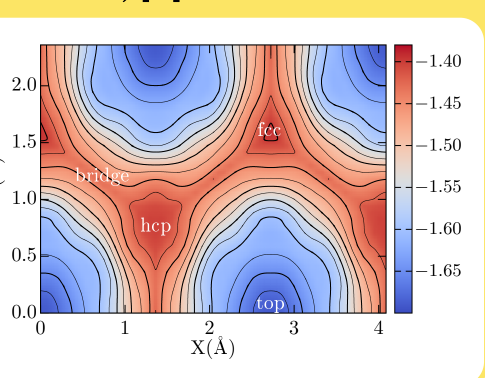
- (2) Electronic friction
- (3) Phonon-adsorbate interaction
- metals: ion lattice plus quasi-free electron gas
- visible light is absorbed only by the electrons
- produced electron-hole pairs thermalize quickly \Rightarrow "hot" Fermi-Dirac-distribution (after $\sim 10 \text{ fs}$)
- electrons transfer part of energy to ion lattice, via **1** electron-phonon coupling
- (phonons = lattice vibrations; quasi-particles)
- -electrons couple to phonons as their fast movement causes "shockwaves" in ion lattice -equilibration process completes after $\sim 1 \text{ ps}$
- \Rightarrow Thus, with fs-lasers, two different temperatures:
 - $-T_{\rm el}$ electron temperature
 - $-T_{\rm ph}$ phonon temperature
- can be simulated using a Two-Temperature Model (2TM)[4] (see right)

Models and Methods

Six-dimensional Potential Energy Surface (6D PES)[3]

- Basis for dynamics: precomputed PES from DFT (rPBE + D2)
 - all 6 dimensions of the adsorbate

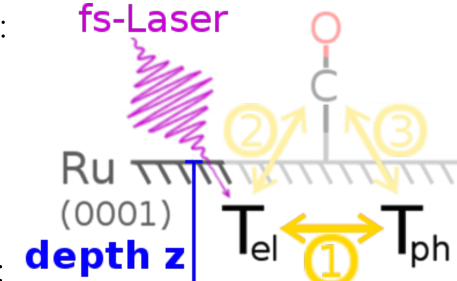
 - \bullet analytical PES and gradients \Rightarrow very fast
 - ⇒ number and length of trajectories can be large • downsides: - surface atoms frozen \Rightarrow no phonons
 - had to be constructed first



Two-Temperature Model (2TM)[4]

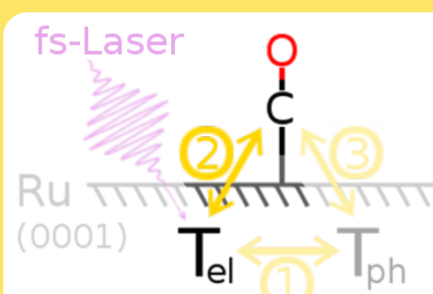
• describes interaction of metal with laser, using two differential equations:

$$C_{\rm el} \frac{\partial T_{\rm el}}{\partial t} = \frac{\partial}{\partial z} \kappa \frac{\partial}{\partial z} T_{\rm el} - g(T_{\rm el} - T_{\rm ph}) + S(z, t),$$
$$C_{\rm ph} \frac{\partial T_{\rm ph}}{\partial t} = g(T_{\rm el} - T_{\rm ph}).$$



- \Rightarrow get $T_{\rm el}$ and $T_{\rm ph}$ as f(z,t) from laser parameters and material properties: -laser wavelength λ (affects penetretion depth into material) -electron and phonon heat capacities $C_{\rm el}$ and $C_{\rm ph}$
 - (effective) absorbed fluence F (energy/area) - electron heat conductivity κ
 - -pulse duration τ (all three appear in the "source term" S(z,t)) -electron-phonon coupling constant g

Electronic Friction: Langevin Dynamics[5] and Local Density Friction Approximation (LDFA)[6]

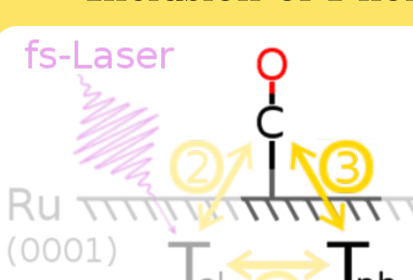


• Langevin equation of motion, a stochastical differential equation:

 $= -\underline{\nabla}_k V(\underline{r}_1, \underline{r}_2) - \eta_{\mathrm{el},k}(\underline{r}_k) \frac{d\underline{r}_k}{dt} + \underline{R}_{\mathrm{el},k}(t).$ Random force Friction force Force on Force due Atom *k* to PES slows movement from e-h pairs

- describes movement of CO on the PES and interaction with electron-hole pairs (friction and excitation)
- Local Density Friction Approx. (LDFA): most simple model to calculate friction coefficients $\eta_{el,k}$ -Atom k embedded in free electron gas with density of bare surface at current position \underline{r}_k
- Random forces $\underline{R}_{el,k}$: gaussian white noise, dependent on both $\eta_{el,k}$ (from LDFA) and T_{el} (from 2TM) -justified by the 2. fluctuation dissipation theorem [10], which relates friction and thermal movement

Inclusion of Phonons: Generalized Langevin Oscillator(GLO)-model[7, 8, 9]



- models influence of phonons in an effective way (augments frozen surface)
- entire surface understood as 3D oscillator (coordinates \underline{r}_s , mass $m_s = m_{\rm Ru}$)
- coupling to molecule via shifting: $V_{\text{GLO}}(\underline{r}_{\text{C}},\underline{r}_{\text{O}};\underline{r}_{s}) = V(\underline{r}_{\text{C}} \underline{r}_{s},\underline{r}_{\text{O}} \underline{r}_{s})$
- additionally coupled to ghost oscillator \underline{r}_q to model influence of the bulk
 - -ghost oscillator is subject to friction $\eta_{\rm ph}$ and random forces $R_{\rm ph}(T_{\rm ph})$

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