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Ultrafast adsorbate excitation probed with sub-ps resolution XAS

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ABSTRACT

We use a pump-probe scheme to measure the time evolution of the C K-edge X-ray absorption spectrum (XAS) from CO/Ru(0001) after excitation by an ultrashort high-intensity optical laser pulse. Due to the short duration of the X-ray probe pulse and precise control of the pulse delay, the excitation-induced dynamics during the first ps after the pump can be resolved with unprecedented time resolution. By comparing with theoretical (DFT) spectrum calculations we find high excitation of the internal stretch and frustrated rotation modes occurring within 200 fs of laser excitation, as well as thermalization of the system in the ps regime. The ~100 fs initial excitation of these CO vibrational modes is not readily

rationalized by traditional theories of nonadiabatic coupling of adsorbates to metal surfaces, *e. g.* electronic frictions based on first order electron-phonon coupling or transient population of adsorbate resonances. We suggest that coupling of the adsorbate to non-thermalized electron-hole pairs is responsible for the ultrafast initial excitation of the modes.

MAIN TEXT

The vibrational, rotational, and translational motions of adsorbates are the fundamental physical processes behind chemical reactions on surfaces. Understanding the details of adsorbate dynamics can therefore be essential for describing heterogeneous catalytic reactions, with the eventual goal of tailoring catalysts for high efficiency and selectivity. In particular, how energy is transferred from the bulk substrate to the adsorbate modes is key to understanding selectivity in catalysis. Since surface chemical reactions proceed through several elementary dynamical steps on different time scales in parallel with competing energy relaxation processes, studying these in a time-resolved manner can give knowledge about the elementary adsorbate-surface interaction and energy transfer processes.

Exciting an adsorbate/substrate system with an intense femtosecond optical pulse and then probing the dynamical evolution of the adsorbate by time-resolved optical techniques, such as sum-frequency generation (SFG) [1–6] and two-pulse correlation (2PC) [7–9] has a modest history. More recently, X-ray absorption and emission spectroscopy (XAS/XES) [10–14], has also become a powerful approach for gaining insight into the transient excitation of adsorbate vibrations and the consequent diffusion, desorption, and final chemical reactions. X-ray spectroscopies are inherently element-specific local probes and thus particularly suited for studying any process that affects the electronic structure and chemical environment of a specific atom [15]. In particular, an X-ray free-electron laser (XFEL) delivering femtosecond X-ray pulses allows probing dynamical changes of the adsorbate electronic structure induced by the femtosecond optical laser. Activation of atomic oxygen [12] and carbon monoxide [10], as well as driving of CO oxidation [13], are examples of reactions that have been studied with this technique.

In this work, we report experimental C K-edge XAS of CO/Ru(0001) during and after pulsed optical laser excitation, probed with a time resolution of ~ 100 fs in a conventional pump-probe setup. This is possible due to the ultrashort duration of XFEL pulses (at DIPROI beamline of the FERMI facility in Elettra, Italy). The XFEL is seeded by a tunable solid state laser system and the photon energy is multiplied through a two-stage high-gain harmonic generation (HG) process [16]. The polarized soft X-ray pulses with less than 50 fs duration are delivered to the sample at a repetition rate of 10 Hz as the “probe”. Meanwhile, a portion of the seed laser is separated and coupled into the experimental chamber collinearly with the X-rays, acting as the “pump” with the same repetition rate. Since the seed and the

pump originate from the same laser pulse, the time jitter between the pump laser and the XFEL X-ray pulse is less than 6 fs [17]. The aforementioned properties allow direct detection of the adsorbate electron structure as influenced by internal excitations before the substrate phonons have been excited, which takes around 1 ps. Since CO chemisorbs on Ru(0001) in an upright geometry at the on-top site (with the C atom coordinated toward the Ru atom) [18], the dipole selection rule gives the strongest moment for the C K-edge $1s \rightarrow 2\pi^*$ transition when the electric field vector (E-vector) is parallel to the surface. Using linearly polarized X-rays and measuring with the E-vector oriented either in-plane (lin H) or out-of-plane (lin V) with respect to the metal surface, we can thus monitor how the electronic structure and molecular orientation temporally changes following the optical laser pumping. The optical laser is p-polarized, precluding any direct optical excitation of the $2\pi^*$. The C K-edge, due to similar intramolecular potential energy curves in the ground and core-excited states, gives only small Franck Condon factors for C-O vibrational excitation in the x-ray transition [19,20], and thereby provides excellent spectral contrast in comparison to the much broader O K-edge used in most previous work [10,11,21].

Recently, CO desorption from CO/Ru was studied on ps time scales, and a weakly adsorbed precursor state [11,22] was directly identified and its evolution described [14]. In this paper, we focus on the ultrafast dynamics occurring in the CO within the very first picosecond following optical laser excitation of the Ru. The high spectral resolution and controlled pump-probe delay, together with detailed spectrum simulations, allow us to identify the excitation of specific adsorbate modes and their initial excitation times. We find significantly faster energy transfer to the adsorbate than conventional non-adiabatic theoretical approaches suggest, and we postulate that non-thermal electron-hole (e-h) pairs initially created by the optical laser may account for the ultrafast initial excitations of these modes.

The Ru(0001) crystal was first cleaned by standard procedures [14], and a saturated CO adlayer (~ 0.66 ML) on Ru was prepared by pressurizing the UHV chamber with 1.0×10^{-8} torr of CO while cooling the crystal down to liquid nitrogen temperature [23]. The crystal sample was pumped using 400 nm optical laser pulses (fluence of ~ 91 J/m²) at a grazing incidence angle of around 2° with duration of 100 fs. A fresh sample spot for each shot was realized by continuously scanning the sample. A CO background pressure of 1.0×10^{-8} torr was used to ensure a fully recovered surface adlayer once the same spot was revisited after scanning the complete sample. XAS was recorded by detecting X-ray fluorescence, using a previously described detector setup [24], and the incident energy was stepwise changed across the carbon $1s \rightarrow 2\pi^*$ resonance edge in the range of 285–290 eV. The XAS was thus plotted by averaging the total number of emitted photons recorded as a function of the incident energy for every desired pump-probe delay. The definition of time zero delay is discussed in the supplementary information (Suppl. A [25]). Further details of the experimental method can be found elsewhere [14].

The top panel of Fig. 1 shows the lin H XAS spectra evolving with pump-probe delay time. The optical unpumped spectrum (averaged between -300 – 0 fs) depicts a strong C 1s \rightarrow 2 π^* resonance peak centered at around 288 eV. For delays 0 – 100 fs following the laser pump pulse, a sudden decrease of intensity at the peak center and an increase of intensity on the lower energy side of the spectrum is clearly seen, while the main peak energy stays the same. However, in the next 100 fs delay a redshift of about 80 meV of the peak center can be identified. No further rapid changes are found in the following delays, indicating a slower energy equilibration after the initial excitation. The lin V spectra (Fig. 1(b)), on the other hand, indicates that the aforementioned observation in the lin H spectrum at 0-100 fs delay cannot be ascribed to the molecule tilting since no corresponding intensity increasing at 288 eV can be

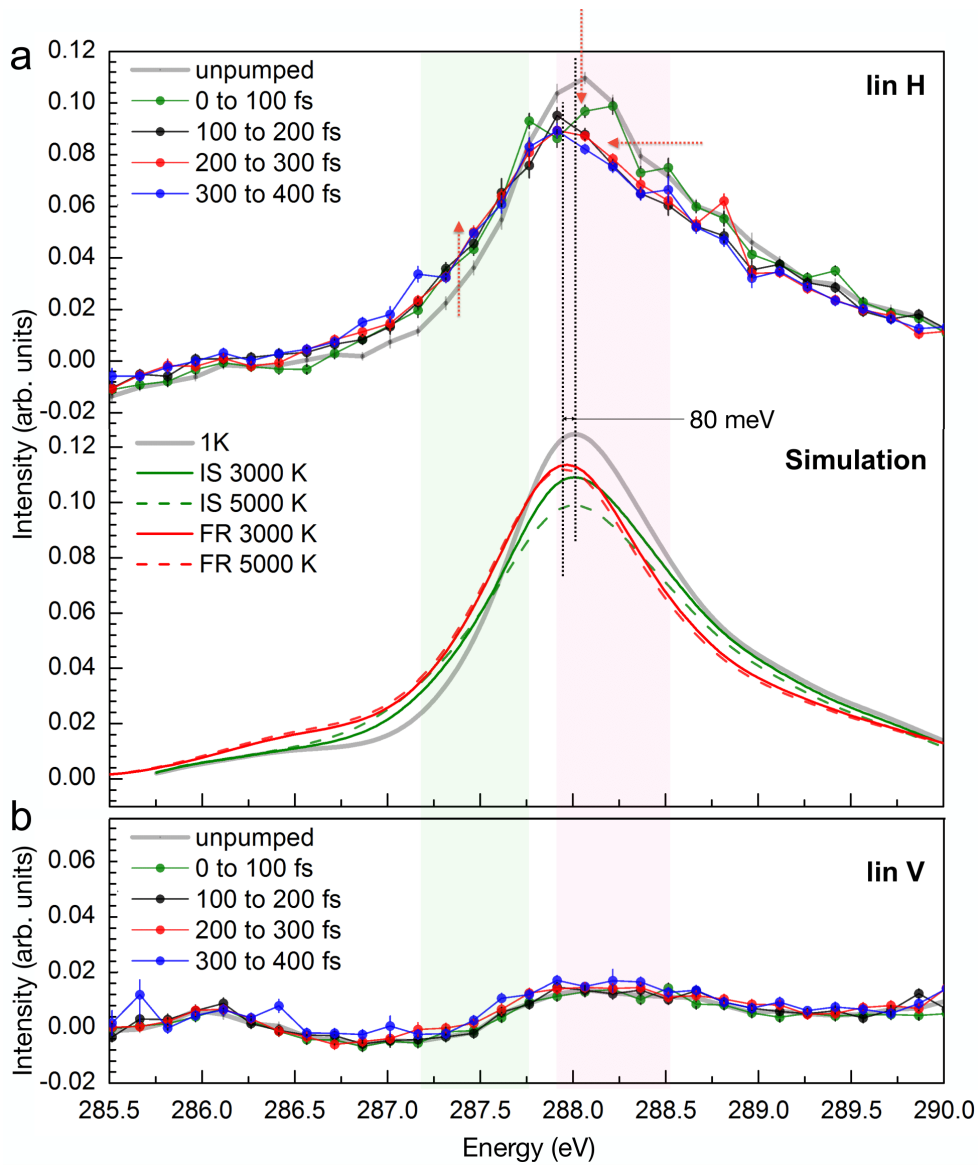


FIG. 1. (a) Measured lin H spectra, averaged over the indicated pump-probe delay ranges, together with calculated spectra with excited internal stretch (IS) and frustrated rotation (FR) modes. (b) Measured lin V spectra. (The negative intensity is due to background subtraction.)

seen in the lin V spectrum. Instead, a mild increase of overall intensity in lin V spectra can be seen in the following delays, up to about 1 ps.

To study the time evolution in detail, the intensities are integrated over two energy regions (main peak region in the pink area and lower energy region in the green area) chosen in order to give a clear view of the time-dependence. The time evolution of these integrated intensities is shown in Fig. 2. In addition, the lin H and lin V spectra at longer time delays are shown in Suppl. B [25]. We first focus on the main peak region. Fig. 2(a) shows a sudden decrease of intensity in the first 0-100 fs and 100-200 fs delays of the lin H spectrum due to the main peak intensity dropping and peak shifting, respectively (cf. Fig. 1(a)), followed by a gradually decreasing intensity after 200 fs and a corresponding increasing intensity in the lin V spectrum with time. On the lower energy side (Fig. 2(b)), a sudden increase of intensity in the first 0-100 fs delay followed by a gentle increase of intensity after 100 fs can be observed in the lin H spectra, while for lin V spectra no obvious changes in 0-100 fs delay can be seen, yet a similar intensity increasing trend as in the lin H spectra shows up after 100 fs. We can thus distinguish two processes

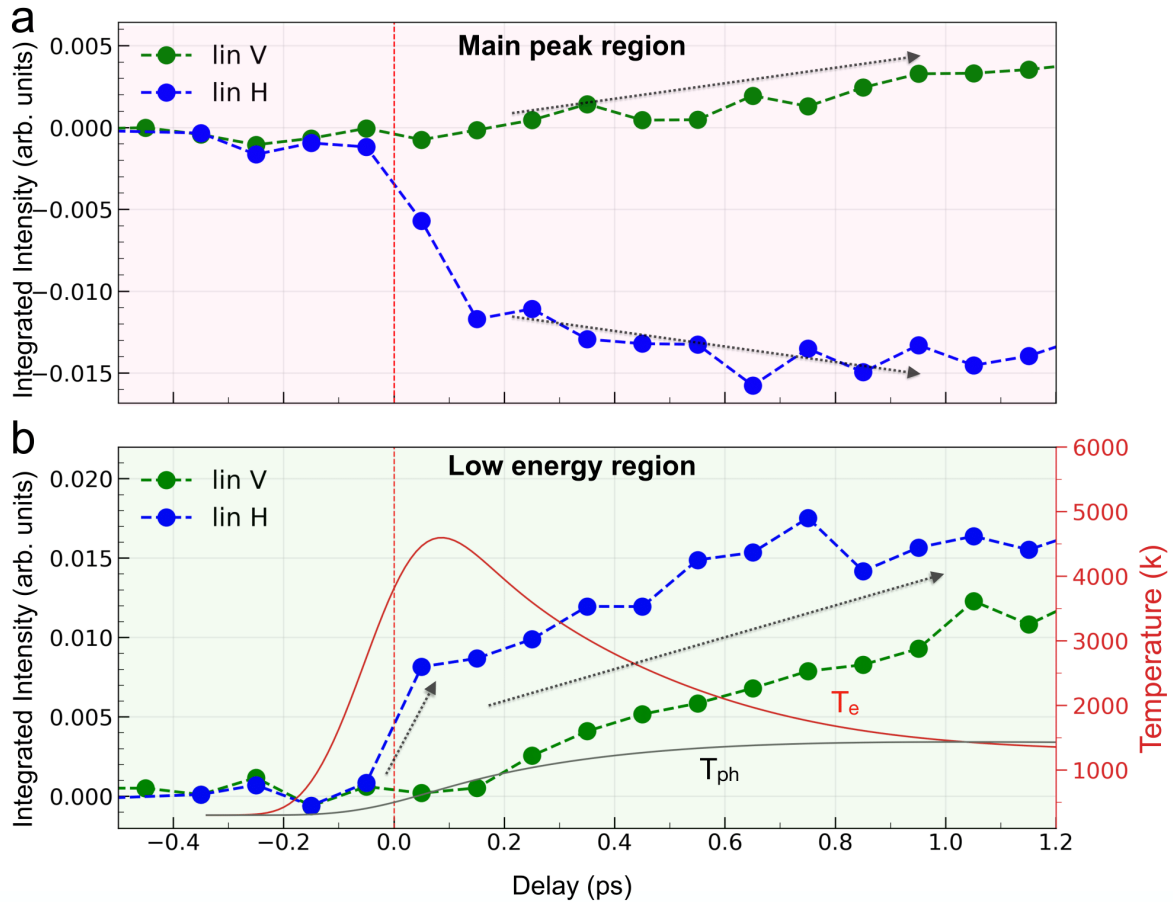


FIG. 2. Time evolution of the integrated spectral intensity, minus the unpumped intensity, in (a) the main peak region and (b) low energy region, respectively (cf. Fig. 1). Also shown in (b) are the electron (T_e) and phonon (T_{ph}) temperatures from the two-temperature model.

occurring on different time scales, one very rapid which is completed after around 200 fs and mainly affects the lin H spectrum, and one slower affecting both polarizations.

To interpret the time evolution of the spectra, we perform density functional theory (DFT) calculations using the Quantum Espresso [26,27] code, with spectra calculated using the xspectra [28–30] code. The general method for spectral calculation and treatment of the core-hole has been described elsewhere [31]. A $4\times 4\times 4$ periodic slab was used for all spectrum calculations with a single CO molecule adsorbed at an on-top site, corresponding to $\frac{1}{4}$ ML coverage instead of the experimental 0.66 ML. However, we have checked that our spectroscopy simulations are not sensitive to the absolute coverage (see Suppl. F [25]). For further computational details, see Suppl. C [25].

Previous theoretical studies of the first order electron-phonon coupling of CO adsorbed on transition metal surfaces, *e.g.* CO/Cu(100), CO/Ni(111) and CO/Pt(111), suggested that the internal C-O stretch (IS) and the frustrated CO rotation (FR) modes are the modes most likely to be significantly excited directly by thermal metal electron-hole pair excitations [32]. Our theoretical studies of the first order electron-phonon coupling for CO/Ru (see Suppl. K [25]) confirm this pattern, with the IS exhibiting the strongest coupling followed by that for FR. To calculate the XAS for internal stretch vibrationally excited states of the CO molecule at the surface, we determine the potential energy curve for the electronic ground state by varying the C-O distance and calculating the DFT total energy for each point. We then fit a Morse potential to the data, which gives the eigenstates and an analytical representation of the vibrational eigenfunctions. The potential for the internal stretch mode with core-excited C was determined in the same way, with the C pseudopotential containing an explicit core-hole. The unit cell was kept neutral by adding an electron at the Fermi level, since the metal electrons screen the core-hole very rapidly [33]. The Franck-Condon factors for the C $1s \rightarrow 2\pi^*$ transition can then be calculated analytically. We calculate the spectra at several displacements and average them together with the appropriate weight for the vibrational eigenstate to get the spectrum for each vibrational state (for details, see Suppl. D and E [25]). The frustrated rotation mode was treated similarly, but with a harmonic potential assumed for the description of the mode. Finally, the spectra for the different vibrational states were added together, with weights given by the Boltzmann distribution for a desired temperature.

The calculated spectra for different vibrational modes at various thermal excitations are shown in Fig. 1. We first note that the agreement of the ground state theoretical spectrum with the experimental spectrum at negative delay is excellent (see also Fig. S4), giving confidence to the spectrum simulations. The calculated spectra for elevated temperatures of the internal stretch (IS) mode show a broadening of the peak and decrease of the maximum intensity, however the peak position remains unchanged. High temperature in the frustrated rotation (FR) mode however gives an 80 meV redshift, in excellent quantitative agreement with the experimental results for ~ 200 fs delay. We therefore identify the IS

excitation as inducing the initial decrease of the peak and the FR excitation as causing the redshift of the peak. It might be expected that FR motion would also give a strong signal in the vertical polarization direction, since tilting the molecule means the $C\ 1s \rightarrow 2\pi^*$ dipole moment will have nonzero projection in the vertical direction. However as seen also in Fig. S5 [25] the actual response is quite small. During pure FR excitation the molecular center-of-mass remains fixed, and there is no simple direct relation between the two polarization signals (Fig. S5). In addition, the slow increase we see in the vertical signal over the first ps is consistent with the anticipated complicated dynamics en route to overall thermalization of the system. For comparison, the results of the commonly used two-temperature model is also shown in Fig. 2(b) (see also Suppl. G [25]). It is immediately clear that the initial response in the lin H signal (blue dotted line) is faster than the rise of the phonon temperature (black line), and thus must be electron-driven, while the lin V signal (green dots) increase is due to more complicated dynamics involving both electrons and phonons (see discussion later). After ~ 1 ps, many phonons are excited and the surface geometry becomes much less well-defined [3]. There is therefore a large number of adsorption sites and rotational states available, giving a slow increase of resonance intensity projected on the vertical direction (lin V) and a complementarily decreasing intensity in lin H spectra in the main peak region at 0.2 - 1 ps delay (Fig. 2(a)). The spectral changes at 1 ps can be qualitatively reproduced by simulating spectra from *ab initio* molecular dynamics at a temperature of 1300 K (see Suppl. H [25]), suggesting that near-equilibration between all modes occurs by ~ 1 ps. Desorption starts to be visible in the spectrum after a few ps [14], beyond the time range studied in this paper.

The mechanism behind the very fast adsorbate initial IS and FR excitation, which occurs within 200 fs, is unclear. First order electron-phonon coupling calculations of the frictional damping of thermalized metal electron-hole pairs suggest vibrational lifetimes $\tau \sim 2 - 4$ ps for the IS and FR modes for CO/Ru (see Suppl. K [25]). A simple one dimensional classical master equation approach indicates that a coupling strength of $\geq 30\text{ cm}^{-1}$ (corresponding to $\tau \sim 200$ fs) would be needed to obtain an energy transfer fast enough to describe the observed initial excitations in these modes (Suppl. I [25]). We suggest that the most likely reason for the rapid initial excitation is that the highly excited non-thermal e-h pair distribution, initially created in the substrate by laser absorption, could give a stronger coupling to adsorbate motion than friction models based on first order electron-phonon coupling where a thermalized e-h pair bath is assumed to excite the adsorbate. Possible evidence for the contribution of non-thermalized e-h pair to adsorbate excitation has previously been suggested for oxygen activation in a O/CO/Ru system [34], where the non-thermalized electron population right after optical excitation coupled more efficiently to an unoccupied adsorbate resonance (LUMO) compared to a thermalized hot Fermi-Dirac distribution, which does not necessarily form within the first 100 fs [35].

The LUMO $2\pi^*$ resonance for CO/Ru is found around 5 eV above the Fermi level [36], while the optical laser energy is only 3.1 eV (i.e., 400 nm). Even though the $2\pi^*$ resonance is broad and a remnant of its resonance could extend to 3.1 eV, significant direct population of this resonance - as in ion-resonance models responsible for Desorption Induced by (Multiple) Electronic Transitions [37] (DIET/DIMET) - is highly unlikely to give efficient energy transfer. Even if we assumed that the adsorbate LUMO $2\pi^*$ resonance could be populated (e.g., by photo-excitation of initially created hot electrons in the laser pulse) we find that a simple dynamical model gives only little energy transfer into the motion of the adsorbate (Suppl. J [25]), thus making resonant population of the adsorbate LUMO an inefficient excitation mechanism for CO/Ru at 400 nm. However, the non-thermal e-h pair distribution could still excite adsorbate modes more efficiently than thermal e-h pairs, since e-h pair states far from the Fermi level may couple more strongly than close to it.

In this paper, we only discuss the anomalous fast initial excitations of the IS and FR modes observed in the C XAS spectra compared to those predicted by first order electron-phonon coupling, and the description after full thermalization. We make no attempt to describe the route between the two as that is driven by complicated coupled electron and phonon dynamics. Theoretical studies of the temperature dependence of the CO/Cu(100) IS lineshape and its evolution following fs optical laser excitation even imply that first order electron-phonon coupling is not sufficient to describe the temporal dynamics of the coherently excited $\nu = 1$ state of IS [38,39]. Extending the electron-phonon coupling to second order opens up an entirely new important class of energy transfer processes, so-called electron-mediated phonon-phonon couplings (EMPPC). While we have no definitive argument that these EMPPC processes are as important for CO/Ru as for CO/Cu(100), we believe that they likely are so that a complete understanding of the dynamics occurring between initial excitation and the fully thermalized system will ultimately require a multidimensional dynamical model including all of the EMPPC and direct phonon couplings. However, it is unlikely that their addition can account for the ultrafast initial mode excitations responsible for the changes in the XAS. For CO/Cu(100), the fastest decay processes for the wavevector $q \sim 0$, $\nu = 1$ state of IS are due to the first order electron-phonon coupling and dephasing of the $q \sim 0$ into other q states of the same IS vibration (denoted as IS \rightarrow IS) [39]. However, unlike photon excitation of an adsorbate vibrational mode, e-h pair scattering excites the adsorbate modes with all q . In addition, since XAS is a local probe at the C, the XAS spectrum is independent of the phase of vibrations contributing to the lineshape so that the IS \rightarrow IS process does not contribute to the dynamics observed. Therefore, if EMPPC in CO/Ru are similar to those in CO/Cu, inclusion of EMPPC does not rationalize the ultrafast initial changes to the XAS compared to those predicted by first order electron-phonon coupling. Since optically created electron-hole pairs take ~ 100 fs to thermalize in Ru [35], we instead suggest that non-thermalized electron-hole pairs within the first ~ 100 fs have a stronger coupling to the adsorbate and preferentially excite the IS and FR modes within this time scale.

We summarize our main observations in Fig. 3. We can tentatively identify an initial excitation of the IS within 100 fs of the laser pulse, followed by excitation of the FR mode during the subsequent 100 fs. Thermalization of the substrate phonons, and heating of the adsorbate translational modes, proceeds over the first ps, consistent with the commonly used two-temperature model. The initial excitations are significantly faster than what can be accounted for in non-adiabatic friction models with theoretically obtained friction coefficients, pointing to a lack of current theoretical descriptions of the coupling between highly excited electron distributions and adsorbate dynamics. Our observation is however in line with what has been inferred previously from experimental SFG [2,3,5,6] and 2PC [9] studies. A detailed explanation poses a challenge for future theoretical and experimental work.

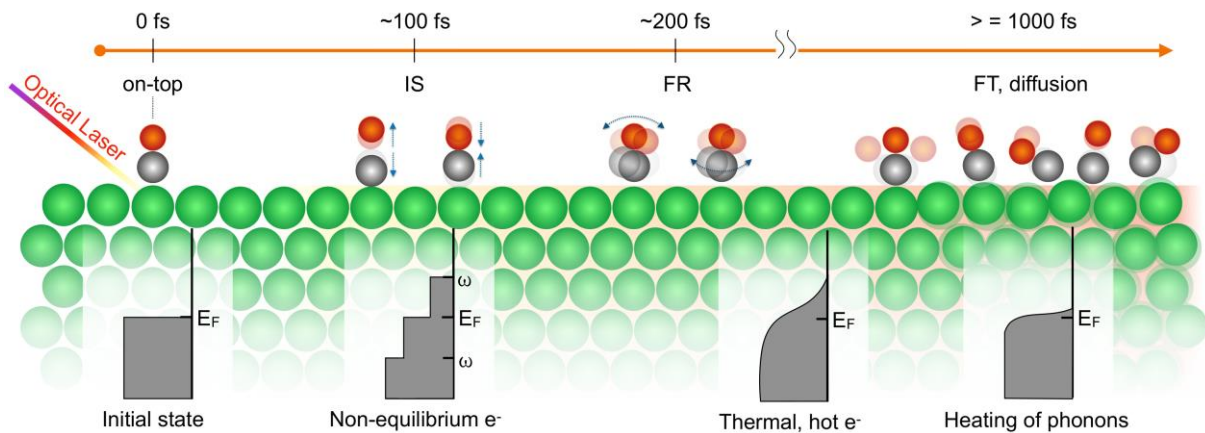


FIG. 3. Sketch of the system evolution during the first picosecond. Atom colors: O red, C grey, Ru green.

This study shows the possibilities of the sub-ps time resolution allowed by the well-controlled pump-probe delay in a seeded FEL setup. Compared to the 2PC technique, where sub-ps dynamics can only be inferred indirectly by its influence on subsequent dynamics, and to SFG, where one can only probe a specific degree of freedom (typically the IS mode at $q \sim 0$), this technique gives uniquely detailed information about the immediate response of the adsorbate local electronic structure to the driving pulse. We have thus been able to directly measure the evolution with time as it occurs, and together with DFT spectral calculations this allows a detailed analysis of the initial excitation and fully thermalized state. We expect that similar methods could be used to investigate the questions raised in this work regarding the role of non-thermal electrons and the seemingly strong frictional coupling.

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