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Direct observation of electron dynamics in the attosecond domain

A. Föhlisch¹, P. Feulner², F. Hennies¹, A. Fink², D. Menzel², D. Sanchez-Portal³, P. M. Echenique³ & W. Wurth¹

Dynamical processes are commonly investigated using laser pump-probe experiments, with a pump pulse exciting the system of interest and a second probe pulse tracking its temporal evolution as a function of the delay between the pulses1-6. Because the time resolution attainable in such experiments depends on the temporal definition of the laser pulses, pulse compression to 200 attoseconds (1 as = 10^{-18} s) is a promising recent development. These ultrafast pulses have been fully characterized, and used to directly measure light waves6 and electronic relaxation in free atoms2-4. But attosecond pulses can only be realized in the extreme ultraviolet and X-ray regime; in contrast, the optical laser pulses typically used for experiments on complex systems last several femtoseconds (1 fs = 10^{-15} s)^{1,5,6}. Here we monitor the dynamics of ultrafast electron transfer-a process important in photo- and electrochemistry and used in solid-state solar cells, molecular electronics and single-electron devices-on attosecond timescales using core-hole spectroscopy. We push the method, which uses the lifetime of a core electron hole as an internal reference clock for following dynamic processes9-19, into the attosecond regime by focusing on short-lived holes with initial and final states in the same electronic shell. This allows us to show that electron transfer from an adsorbed sulphur atom to a ruthenium surface proceeds in about 320 as.

When studying electron transfer processes in complex systems, it is of equal importance to address the temporal evolution of the electron wave packet and the question of which atomic centre an electron is localized at before charge transfer to the substrate occurs. This atom specific information cannot be provided in pump-probe experiments in the spectral regime of optical transitions. By adapting an element specific synchrotron based soft X-ray spectroscopy method, namely core-hole clock spectroscopy, we can effectively determine on an attosecond timescale electron transfer dynamics originating from an atomically localized state by making use of extremely fast Coster-Kronig decay processes of core-excited states.

The principle of core-hole clock spectroscopy is to take the corehole lifetime 7 as an internal reference clock for the temporal evolution of a dynamic process under investigation 419. To study charge transfer on the timescale of τ , the dynamics of an electron resonantly excited into an unoccupied state from an atomically localized adsorbate core level (Fig. 1a) is monitored through the autoionization process that accompanies the core-hole decay (Fig. 1b and c). If the initially excited core electron remains in an atomically localized resonance, a linear relation between the energies of the incoming photon and of the outgoing electron in the autoionization is observed (Fig. 1b). This is the so-called Raman autoionization channel at constant binding energy (1). In contrast, if the initial

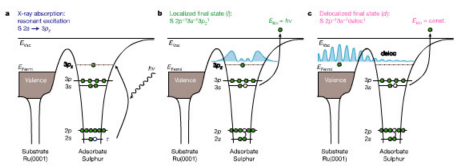


Figure 1 | Core-hole clock spectroscopy-schematic overview. a, Initially, a core electron is promoted by resonant excitation from the \$2s level into a bound resonance localized at an adsorbed sulphur atom (\$2s-13p1) on ruthenium $c(4 \times 2)$ S/Ru(0001) with a core-hole lifetime $\tau = 0.5$ fs. In the autoionization decay processes, Coster-Kronig decay of the \$2s core hole takes place in the presence of this electron, the so-called 'spectator' electron,

leading to two different final states. b, Localized final state S 2p-13s-13p1: state l. The initially excited electron is still localized at the sulphur atom. c, Delocalized final state \$2p^{-1}3s^{-1}deloc1: state d. The initially excited electron has already left the localized resonance. E_{vao} vacuum energy; E_{Bermi}, Fermi energy; Ekin, kinetic energy.

institut für Experimentalphysik, Universität Hamburg, Luruper Chaussee 149, D-22761 Hamburg, Germany. ²Physik Department E20, Technische Universität München, D-85747 Garching, Germany. Centro Mixto CSIC-UPV/EHU "Unidad de Física de Materiales", Donostia international Physics Center (DIPC), and Departamento de Física de Materiales Universidad del País Vasco, Apdo. 1072, 20080 Donostia-San Sebastián, Spain

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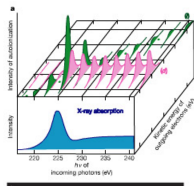


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excitation involves an electronic state delocalized over many atomic centres (that is, the excited atomic resonance is coupled to a continuum), we obtain independently of the incident photon energy autoionization at constant kinetic electron energy (Fig. 1c). This is the charge transfer channel of autoionization (d). Owing to this different dispersive behaviour, the Raman (1) and charge transfer (d) channels of autoionization can be spectroscopically separated (Fig. 2a), and the ratio of Raman to charge transfer intensity is related to the degree of atomic localization in the excited state on the timescale τ of the core-hole decay. This can be translated into a dynamic picture of an electron residence time, or alternatively as the charge transfer time, τ_{CD} of electron hopping to the substrate. As spectral intensities are compared, the \sqrt{N} uncertainty (where N is the number of events) allows a statistically significant analysis only as long as the intensities of the spectroscopic channels are less than one order of magnitude apart. Thus a temporal range of charge transfer times between $0.1\tau \le \tau_{CT} \le 10\tau$ becomes accessible. The typical core-hole lifetimes of inner shell vacancies lie at oxygen KLL ($\tau = 4 \, \text{fs}$; ref. 20), nitrogen KLL ($\tau = 5 \, \text{fs}$; ref. 20), carbon KLL $(\tau = 6 \text{ fs}; \text{ ref. } 20), \text{ and argon } L_3 M_{4/5} M_{4/5} \ (\tau = 6 \text{ fs}; \text{ refs } 13, 18).$ We note that the core-hole lifetimes depend only weakly on the chemical environment. In a comparison between atomic and molecular systems, variations of the order of roughly 20% have been



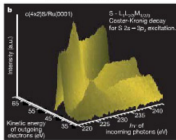


Figure 2 | Core-hole clock spectroscopy—the spectroscopic signatures. a, Diagram of the spectroscopic autoionization signatures leading to a localized final state (l) with linear dispersion and a delocalized final state (d)at constant kinetic energy, and their relation to resonant excitation by X-ray absorption. b, Experimental sulphur L1L2/3M1/2/3 Coster-Kronig autoionization spectra of c(4 × 2)S/Ru(0001) as a function of incident

To access dynamic processes in the attosecond range reliably, shorter core-hole lifetimes are required. Our approach is to perform attosecond charge transfer core-hole clock spectroscopy in the soft X-ray region by monitoring Coster-Kronig autoionization channels

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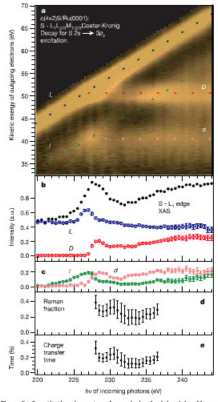


Figure 3 | Quantitative charge transfer analysis of sulphur L₁L_{2/3}M_{1/2/3} Coster-Kronig autoionization spectra of c(4 × 2)S/Ru(0001) as a function of photon energy. a, Experimental intensities as a function of incoming photon energy and kinetic energy of the outgoing electrons. Lighter colours correspond to higher autoionization intensity. Shown are Raman channels with linear dispersion for localized final states L $(2p^{-1}3p^{-1}3p_z^1)$ at 170.7 eV binding energy and $I(2p^{-1}3s^{-1}3p_z^1)$ at 181.7 eV binding energy, and charge transfer channels with delocalized final states D(2p-13p-1deloc1) at 50.8 eV kinetic energy and d (2p-13s-1deloc1) at 40.6 eV kinetic energy. b, Sum of spectral intensities representing the S-L1 edge X-ray absorption spectrum. Also shown are separate intensities of the spectral channels (L, D) from curve fitting with lorentzians of 3.3 eV FWHM. Error bars show the standard deviation of each fit. c, Separate intensities of the spectral channels (I, d) from curve fitting with lorentzians of 3.3 eV FWHM. Error bars show the standard deviation of each fit. **d**, Raman fraction f = l/(l+d) as a function of photon energy. Error bars are derived from the standard deviation of the fits (see c). e, Charge transfer time obtained from the Raman fraction as $\tau_{CF} = \tau f(1 - f)$ and the S2s core-hole lifetime $\tau = 0.5$ fs. Error bars are derived from the standard deviation of the fits (see c).

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with attosecond core-hole lifetimes. Here the initial and final state vacancies are in the same electronic shell (same principal quantum number n); the probability for these transitions is higher and the corresponding core-hole lifetimes shorter than in the case of decay

processes involving different values of n. With the c(4 × 2)-S/Ru(0001) surface21,22, we obtained the S L₁L_{2/3}M_{1/2/3} Coster-Kronig autoionization spectra shown in Fig. 2b as a function of the incoming photon energy hν tuned across the $S2s^{-1}3p_{i}^{3}$, core level resonance at $h\nu = 227.5 \,\text{eV}$. On resonance, the excited electron's energy lies 1.68 ± 0.1 eV above the Fermi level. We observe resonant enhancement and branching of decay channels at this absorption resonance. The data are converted to a colourcoded plot in Fig. 3a, where higher intensity corresponds to lighter colour. We can directly discern spectral features (l, L) at constant binding energy, which branch into charge transfer spectral features (d, D) with constant kinetic energy. Let us assign the spectral features: starting from the electronic ground state (GS), the S2s-13p1 coreexcited state can autoionize through Coster-Kronig channels with and without participation of the excited electron ('participator' and 'spectator' channels, respectively). The participator channel, involving the core-excited 3pt in the decay, leads to the spin-orbit split 2p-1 final state, identical to the main lines of photoionization, with $161.5 \,\text{eV} \, (2p_{3/2}^{-1})$ and $162.6 \,\text{eV} \, (2p_{1/2}^{-1})$ binding energy. Its spectral features lie outside the range of Figs 2b and 3a.

The spectral features shown in Figs 2b and 3a are thus associated with spectator channels, in particular the 2p-13s-13p1 (1) and

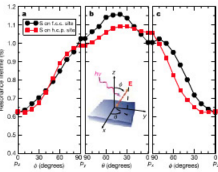


Figure 4 | Theoretical charge transfer time for S in f.c.c. and h.c.p. hollow sites computed as \$ 3p resonance lifetime. Theory predicts a strong dependence of the charge transfer time on the symmetry of the initial wave packet, which translates to a strong dependence on the light polarization. The coordinates x, y, z shown in the diagram in b correspond to the crystallographic directions [100], [010], [001]. The circles and squares correspond respectively to sulphur atoms in f.c.c. and h.c.p. sites of the surface. a, Resonance lifetime as a function of the angle ϕ of the electric field vector with respect to the surface normal in the y-z plane. Polarization of the synchrotron light along the z axis ($\phi = 0$, the experimental geometry) produces an initial excited state with p_symmetry. b, Resonance lifetime as a function of the angle θ of the electric field vector with respect to the x axis in the x-y plane. In this geometry, p_x and p_y symmetries and combinations of them would be obtained with in-plane polarization. c, Resonance lifetime as a function of the angle ϕ of the electric field vector with respect to the surface

2p-13p-13p, (L) final states at 181.7 eV and 170.7 eV binding energy,

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$$G_{2} = \frac{m_{1} - m_{2}}{m_{1} - m_{2}} + \sum_{i \neq j} \frac{m_{1} - m_{2}}{m_{2} - m_{2}} + \sum_{i \neq j} \frac{1}{2^{i} - 1} \frac{1}{2^{i}} + \sum_{i \neq j} \frac{m_{1} - m_{2}}{m_{2} - m_{2}} + \sum_{i \neq j} \frac{m_{2} - m_{2}}{m_{2}} + \sum_{i \neq j} \frac{m_{2} - m_{2}}{m_{2}} + \sum_{i \neq j} \frac{m_{2}}{m_{2}} + \sum_{i \neq j}$$

The $2p^{-1}3s^{-1}3p_z^1$ final state l can only be reached via autoionization, whereas the $2p^{-1}3p^{-1}3p^{1}$ final state L can in addition be reached as a photoionization satellite, obeying the monopole selection rule of photoionization shake-ups. Thus, L is present at all photon energies, whereas I is a pure autoionization feature observed only when a core hole has been present. We therefore base all further analysis on the autoionization channel l $(2p^{-1}3s^{-1}3p_{\perp}^{l})$ and the related charge transfer feature d $(2p^{-1}3s^{-1}deloc^{1})$, which branches off at 40.6 eV constant kinetic energy. The final states l and d are shown schematically in Fig. 1b and c.

To quantify the relative strength of the Raman (1) and charge transfer (d) channels, we performed a curve fit of the spectra at all photon energies with fixed line shapes, where each channel was described phenomenologically by a lorentzian of 3.3 eV full-width at half-maximum (FWHM), and only the intensities were varied. The charge transfer peaks (D, d) were kept at constant kinetic energy and the Raman peaks (L, l) at constant binding energy with varied photon energy. In Fig. 3b and c, these four contributions (D, L, d, l) and the standard deviation of the fit at each photon energy are shown together with their sum. The latter yields the S-L₁ edge X-ray absorption spectrum (XAS) (Fig. 3b) with 3 eV FWHM dominated by the S2 s core-hole lifetime of $\tau = 0.5$ fs (ref. 23). From this fit, as a function of photon energy, the relative strength of the Raman contribution expressed as the Raman fraction f = l/(l + d) and the charge transfer time $\tau_{CT} = \tau f / (1 - f)$ using the S 2s core-hole life time ($\tau = 0.5 \, \text{fs}$) is derived and displayed in Fig. 3d and e,

For photon energies below the S2s-13pt absorption resonance $(h\nu = 227.5 \text{ eV})$, we observe intensity in the Raman channel l only, as charge transfer below threshold is energetically forbidden, equivalent to an infinitely long charge transfer time. Just above the \$2s-13p. resonance at $h\nu = 228$ eV, we determine $\tau_{CT} = 0.32 \pm 0.09$ fs. For higher photon energies, shorter charge transfer times down to 0.11 ± 0.06 fs at $h\nu = 234$ eV are found. An energy dependence of the electron transfer time has been observed before 15,18. This energy dependence is most probably due to the detailed nature of the projected band structure of the substrate, and thus the number and character of the final states available, as well as their overlap with the initial adsorbate state.

We have compared our experimental results with firstprinciples computations of the charge-transfer dynamics in the $c(4 \times 2)$ S/Ru(0001) system. The initial electron wave packet is constructed as a linear combination of the \$3p orbitals projected onto the unoccupied bands of the combined system, that is, the Ru substrate with the adsorbed sulphur atoms, at the resonance energy. We find that the excitation into a resonance with predominantly 3p, character yields a charge transfer time of 0.63 ± 0.15 fs (Fig. 4a). This corresponds to the state that is excited in the experiment. In comparing experiment to theory, the theoretical time constant confirms that the charge transfer process takes place well below a femtosecond timescale. In particular, the agreement with the experimental value of 0.32 ± 0.09 fs is very satisfactory, taking into account that the core vacancy is not described explicitly in the theoretical ground state calculation; and the theoretical resonance position at -2 eV above the Fermi level is shifted relative to the experimental absorption resonance, which is at 1.68 ± 0.1 eV above the Fermi level. Furthermore, theory predicts a detailed dependence of the charge transfer time on the symmetry of the initial excited state,





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which is summarized in Fig. 4. For excitation into 3px or 3pv-like resonances (in plane) (Fig. 4b) with a smaller overlap to the substrate, a significantly larger charge transfer time of up to 1.15 ± 0.15 fs is calculated. This theoretical result indicates that different polarizations of the light favour different initial excited states, with different symmetries and overlaps with the states of the substrate, thus leading to different transfer times.

The demonstration that soft X-ray spectroscopy can be used as a tool to study the motion of electrons on attosecond timescales opens a possible way to interesting new research areas. Potentially the method is suited to the investigation of electron transfer in complex molecular systems. In such investigations, the ability to excite individual atomic centres (even atoms of the same element in chemically different environments) by exploiting core level shifts should be of particular importance. A second future application could be the investigation of spin-dependent electron transfer processes, which are important in spintronics. Here core level excitation using circularly polarized light could be used to create spin-polarized excitations.

Experiments. The experiments were performed at beamline I311, MAX-lab in Lund, Sweden. At 5 × 10⁻¹¹ torr base pressure, a clean Ru(0001) surface was prepared by cycles of Ar 2-ion sputtering, oxygen-exposure and annealing. The c(4 × 2)S/Ru(0001) surface, with sulphur atoms chemisorbed in hexagonal close packed (h.c.p.) and face-centred cubic (f.c.c.) hollow sites21,22, was prepared by dissociative adsorption of 400 Langmuir H₂S (1 Langmuir = 10⁻⁶ torr s) at 550 K and annealing to 850 K. The surface quality was checked by core electron spectroscopy (XPS) and low-energy electron diffraction (LEED). At 7° grazing incidence, the electric field vector of the incident radiation was 7º off the surface normal, exciting preferentially into the S3p, orbital oriented normal to the surface. The electron spectrometer (Scienta SES 200) was in the polarization plane at 45° to the incident radiation. Narrow bandwidth excitation and high spectral resolution are prerequisites for separating charge-transfer from noncharge-transfer states. Thus the bandwidth of the incident radiation and the ΔE of the electron analyser were both set to 100 meV.

Electronic structure calculations. The density functional calculation of the electronic structure of c(4 × 2)S/Ru(001) has been performed using the SIESTA code^{14,25}. We used a symmetric slab containing 7 Ru layers and the surface geometry known from LEED21,23. Approximately 5 eV below the Fermi energy a strong S3p density of states is found. Above the Fermi level we also find a broad resonance with a large weight on the S3p orbitals, although strongly hybridized with Ru states in an anti-bonding S-Ru interaction. The resonance maximum lies -2 eV above the Fermi level, which is marginally higher than the experimentally observed resonance maximum at 1.68 ± 0.1 eV above the Fermi

Calculation of the charge transfer times. The charge transfer dynamics are computed using the electronic hamiltonian obtained in the density functional calculations previously described. The initial electron wave packet $|\phi_R\rangle$ is constructed as a projection of a linear combination of the S3p orbitals onto the unoccupied bands at the energies of the resonance region. The wavefunction of the resonance depends on the excitation process. Since the electron is excited from a state of s-symmetry, the admixture of p_x , p_y and p_z components is given by the direction of the electric field vector of the incoming radiation E. Therefore we take $|\phi_R\rangle = |\phi(t=0)\rangle \propto \sum_i E_i |p_i\rangle$. From the time evolution of the wave packet we can calculate the probability of finding the electron in the initial state $P(t) = |A(t)|^2$, where $A(t) = \langle \phi_R | \phi(t) \rangle$ is the so-called survival amplitude. The Fourier transform A(t) is directly related to the projection of the Green function onto the initial state $\tilde{A}(\omega) \propto \langle \phi_R | \tilde{G}(\omega) | \phi_R \rangle = G_{RR}(\omega)$ (ref. 26). The characteristic decay time of the resonance population τ is then computed using two procedures. Either the width of the resonance Δ is directly estimated from $G_{RR}(\omega)$ and $\tau = \hbar \Delta^{-1}$, or P(t) is transformed into real time and τ is defined such that $P(t) \le 1/e$ if $t \ge \tau$. Both methods produce very similar results. However, we prefer the second method since 7 is obtained directly and it is not necessary to assume the lorentzian line-shape. We should point out that $G_{RR}(\omega)$ has to be calculated for the semi-infinite system, that is, we have to get rid of the finite size effects associated with the slab calculations. This is instrumental in getting reliable resonance widths and lifetimes. This is done by combining the information from a slab calculation with the ab initio hamiltonian obtained for bulk Ru and using recursive techniques to calculate $G_{R0}(\omega)$. We have assigned an error bar of 0.15 fs to our theoretical values. This reflects both the presence of two

non-equivalent sulphur atoms in the surface and the numerical accuracy of our

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