possible for nonrotationally symmetric systems, allowing practical designs to be simulated with high accuracy; (2) mechanical fabrication tolerances have advanced materially in the past 15 years; (3) high stability electronic components have become available in the past 10 years, allowing the packaging of many, very high stability, computer-controlled power supplies in a small space; and (4) high-speed small computers are now available for real-time processing of the shadow map Rochigram data to obtain aberration parameters.

We believe aberration correction technology combined with the inherent power of the STEM instrument opens the way for lower voltage, smaller, ‘smarter’, more easily managed instruments that will be capable of routinely imaging and analysing materials at sub-
Ångstrom resolution, using several different crystal orientations. Specialized instruments, perhaps using some submillimetre-sized electrostatic imaging elements, may reach the diffraction limit for the imaging electrons. Multiple corrector systems will allow aberration control of both probe size and detector field of view, in a manner somewhat similar to confocal light microscopy. Finally, detailed control of the amplitude and phase of the incident and scattered electron wavefunction will be possible, allowing specific specimen inelastic transitions to be accessed. Thus aberration correction marks a shift towards precise optical control that will allow routine atomic level characterization of defects and interfaces within bulk materials.

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ligands. For this dye the upper limits on the electron transfer time have been measured\(^\text{5–9}\). As a model interface for this important system we have chosen a monolayer of dcb adsorbed on a rutile TiO\(_2\)(110) surface\(^\text{20}\). The structure of the interface is illustrated in Fig. 1. The carboxylic groups bind to the substrate in a 2M-bidentate fashion, that is, with each of the carboxylic oxygen atoms attached to one substrate Ti-atom. This geometry is similar to the geometry surmised for bi-isonicotinic acid on anatase TiO\(_2\)(101)\(^\text{21}\) and on anatase TiO\(_2\) nanoparticles grown in situ (I.S. et al., manuscript in preparation) as well as to one of two suggested geometries of the original cell\(^\text{11–13}\). The bandgap of rutile TiO\(_2\) (3.05 eV)\(^\text{22}\) is close to that of the nanostructured anatase used in time have been measured\(^\text{5–9}\). As a model interface for this important ligands. For this dye the above upper limits on the electron transfer

Relevant processes. Figure 2

(3.05 eV)\(^\text{22}\) is close to that of the nanostructured anatase used in the photoelectrochemical cell (3.20 eV)\(^\text{23}\). The high structural qual-

The electronic structure of the interface is presented in Fig. 3. The bandgap of the substrate is located between 5 and 8 eV binding energy. We have studied the first three unoccupied states of the adsorbate. The lowest unoccupied molecular orbital (LUMO) is situated almost completely within the bandgap. This is an effect of the core hole (Fig. 2b) produced in XAS, which leads to energy shifts to higher binding energies\(^\text{24}\), similar to the case of valence excitations\(^\text{25}\). The other two orbitals (LUMO+1 and LUMO+2), in contrast, lie within the substrate conduction band. The calculations confirm these fundamental experimental results. These findings imply that in the presence of the core hole an electron residing in the LUMO cannot be transferred into the substrate, while for an electron in the LUMO+1 and LUMO+2 a transfer is energetically feasible. We now use RPES to show that the latter cases indeed lead to a very rapid electron delocalization into the substrate.

Figure 4 shows the RPES and contains the XAS from Fig. 3 for comparison. The large peak at 398.6 eV in the RPES shows that the excited electron promoted to the LUMO is localized on the molecule, in agreement with the results for the density-of-states. In contrast, the signals observed in the XAS for the LUMO+1 and LUMO+2 are completely quenched in the RPES, indicating ultra-

fast delocalization of the excited electron into the substrate. In Fig. 4 we indicate the noise level \(N\) of the RPES and the XAS signal intensity \(S\). \(N\) is at most 10% of the XAS intensity, that is, \(N/S < 0.1\). A possible RPES signal for the LUMO+2 hidden by the noise could thus at most amount to 10% of the XAS intensity \(S\), that is \(I_{\text{RPES}}/I_{\text{XAS}} < 0.1\) in terms of the intensities in the RPES and the XAS. Using the equation given in the Methods section—that is, \(\tau_T = \tau_C (I_{\text{RPES}}/I_{\text{XAS}})/[C = (I_{\text{RPES}}/I_{\text{XAS}})]\) where \(\tau_C = 6\) fs and \(C = 1/3\)—determined from the data in the Supplementary Information—this result corresponds to an upper limit on the electron transfer time, \(\tau_T < 2.5\) fs, much faster than previous studies could resolve. For the LUMO+1 the noise level is slightly higher, leading to an upper limit \(\tau_T < 3\) fs, which with little loss of precision we adopt for both the LUMO+1 and LUMO+2 states.

This result is semiquantitatively explained by the calculations, which show quite broad LUMO+1 and LUMO+2 bands (compare Fig. 3), with a width of the order of 1.5 eV. Interpreted as band-

widths, and assuming that no vibrational or other broadening effects exist, this corresponds to electron transfer times much shorter than 3 fs. Closer agreement on the resonance energy has been achieved for a simpler system\(^\text{26}\), and a similar level of agreement for systems of the present complexity is a challenge for the near future. A key aspect of the present study is that the soft-X-ray excitation frequencies used here are much higher than the fastest possible electron transfer rate. Application of these short electron transfer times to the optical excitations relevant to solar cell applications requires care, because the timescales of both processes are comparable in such cases. Because of this, optical excitation and...
transfer of the electron cannot be disentangled. Hence it is more appropriate to discuss the adsorbate–substrate interaction in terms of coupling strength, that is, in terms of an electron transfer bandwidth (for example, ref. 27), rather than in terms of a well-defined electron transfer time. From the measured electron transfer time of less than 3 fs for a core excitation we can conclude that the system is characterized by strong interfacial electronic coupling. Together with previous calculations24 the present results show that this extremely rapid transfer relies on the overlap of ring, carboxylic group and substrate electronic states. The weak-coupling Marcus theory commonly used to model electron transfer reactions (for reviews and discussion see, for example, refs 2, 4 and 17) is inappropriate to describe the electronic coupling at these interfaces. This is confirmed by the observation that the electron transfer rate is not limited by the timescale of vibrational motion in this case, although vibrations are excited in the N 1s as in optical transitions, here easily observed as the dominant contribution to the width of the LUMO resonance. Similar vibronic coupling is expected for the LUMO+1 and LUMO+2 states. This observation is consistent with at least three models15,28,29. Furthermore, the electron transfer takes place on a much faster timescale than charge redistribution30,31 and thermalization12–14 in N3 and similar systems.

Aromatic rings such as those in dcb are well known to facilitate charge delocalization. Here we establish that the carboxylic acid group, which forms a covalent bond to the oxide substrate, completes an efficient conduit of electric current from the N atoms into the substrate; the coupling is comparable to the strongest observed for CO on metal surfaces15. The large distance introduced by the carboxylic spacer group could be expected to lengthen the electron transfer time28. The present results show in contrast that the ligand presents a almost negligible barrier, helping to rationalize the observed ultrafast electron transfer for the N3 dye itself 5–8. The observed ultrafast electron transfer rate is appropriate to discuss the adsorbate–substrate interaction in terms of the occupied electronic density-of-states. In the case presented here an electron from an occupied electronic state of a sample, that is, it images the occupied electronic density-of-states. In the present case (Fig. 2b), RPES monitors only decay processes in which the excited electron takes part. For very fast transfers the RPES vanishes and can often be mapped onto electronic density-of-states ground state calculations. Here we call the adsorbate state at lowest binding energy the HOMO (highest occupied molecular orbital), the next one HOMO – 1, and so on. The energy scale has its origin at the vacuum level E_{vac} (ref. 16). Clean crystal and monolayer PES spectra were mutually aligned using the O 2s level.

X-ray absorption spectroscopy
XAS maps the unoccupied electronic density-of-states. In the present case (Fig. 2b), a valence electron with well-defined energy hv promotes an N 1s core electron to a bound unoccupied state with p-character. We term this electron the excited electron. By varying the photon energy different unoccupied states can be reached, and the small spatial extent of the 1s orbital ensures that adsorbate states are probed. The first observed state is referred to as the lowest unoccupied molecular orbital (LUMO), followed by the LUMO+1, and so on. The process of removing an electron from a core state, that is, the creation of a core-hole, can quantitatively be viewed as an increase of the positive nuclear charge by one unit16. Owing to this the observed energies are lower than the calculated ground state energies, especially for the LUMO. The XAS was placed60 onto the same energy scale as PES.

Density-of-states and XAS calculations
The theoretical curve for the density-of-states is a periodic pseudopotential calculation with gradient corrected functionals of the total ground state density-of-states and was performed using version 3.3 of the CPMD code (for more detail see http://www.fz-juelich.de/nic-series/Volume3/marx.prl). It has been aligned to the PES data of clean TiO2(110), with the gap artificially adjusted to the measured optical one22. The XAS has been modelled by calculating the p-like density-of-states with an excited N 1s electron, which is directly comparable to the experimental data31.

Resonant photoemission spectroscopy
In XAS the system is left in an excited state with an additional electron in a bound unoccupied level and a vacancy (a missing electron) in a core (N 1s) level. This core-excited system will decay exponentially via Auger processes (99.9%), with a time constant given by the core-hole lifetime \( \tau_c \) (6 fs for N 1s). As shown in Fig. 2c, a valence electron (or the excited electron itself) fills the core hole, while another valence electron (or the excited electron) takes up the energy released in that process and is ejected into the vacuum and measured. RPES monitors only decay processes in which the excited electron takes part. These processes are termed resonant photoemission. Because the remaining system is left in the same state as for PES (compare Fig. 2a), RPES measurements are obtained by integrating the valence PES containing the resonant photoemission signal for the photon energies studied in XAS, that is, one probes the effect of placing the excited electron in different unoccupied states. The RPES intensity is then compared to the XAS intensity for each unoccupied state, and results in a particular state-dependent ratio that we call C for the case of an isolated molecule (in our case, \( C = 1/3 \), see Supplementary Information). C describes the sensitivity of RPES compared to XAS. If the particular unoccupied orbital to which the excited electron was promoted is coupled to unoccupied substrate states, the excited electron delocalizes into the substrate in an exponential process (to first order) with the characteristic electron transfer time \( \tau_C \). Thus the transfer process competes with the decay process characterized by \( \tau_C \) and monitored in RPES (Fig. 2c and d). If the two timescales are comparable the RPES intensity serves as a fingerprint for the extent of excited electron localization. In principle, this enables RPES to probe the electron transfer dynamics within roughly one order of magnitude (up or down) of the core-hole lifetime18,19. The transfer time of the excited electron into the substrate can be estimated from the XAS and RPES intensities \( I_{XAS} \) and \( I_{RPES} \) for the particular unoccupied state from the equation \( \tau_C = I_{RPES}/I_{XAS} \). For very fast transfers the RPES vanishes completely for the particular state. This simple model can be applied because the XAS excitation occurs on the 10-attosecond (10^{-12}s) timescale, far shorter than attainable electron transfer times and also far faster than optical excitations. The RPES, the best of several runs, contains a linear, slowly decreasing, direct photoemission background.

Methods

Photoemission spectroscopy
PES maps the energies of the occupied electronic states of a sample, that is, it images the occupied electronic density-of-states. In the case presented here an electron from an occupied valence state is excited by absorption of a photon with a well-defined photon energy hv into the vacuum (Fig. 2a) to kinetic energies \( E_{k} \) of approximately 100 eV. A so-called electron binding energy \( E_{B} \) in the adsorbate–substrate system is derived via delocalization. For the LUMO+1 and LUMO+2, we estimated the maximum possible RPES signal consistent with the noise level \( N \), and compared it to the XAS signal \( S \), determining the upper limits in the text. The XAS has a 12:1 signal-to-noise ratio at the LUMO+2.

\[ E_B = h\nu - E_F \]

\[ t_C = I_{RPES}/I_{XAS} \]

\[ C = (I_{RPES}/I_{XAS}) \]

\[ \tau_C = I_{RPES}/I_{XAS} \]

\[ \tau_C = I_{RPES}/I_{XAS} \]
Ecosystem carbon loss with woody plant invasion of grasslands

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The invasion of woody vegetation into deserts, grasslands and savannas is generally thought to lead to an increase in the amount of carbon stored in those ecosystems. For this reason, shrub and forest expansion (for example, into grasslands) is also suggested to be a substantial, if uncertain, component of the terrestrial carbon sink[1-14]. Here we investigate woody plant invasion along a precipitation gradient (200 to 1,100 mm yr⁻¹) by comparing carbon and nitrogen budgets and soil ¹³C profiles between six pairs of adjacent grasslands, in which one of each pair was invaded by woody vegetation, with drier sites gaining, and wetter sites losing, soil organic carbon. Losses of soil organic carbon at the wetter sites were substantial enough to offset increases in plant biomass carbon, suggesting that current land-based assessments may overestimate carbon sinks. Assessments relying on carbon stored from woody plant invasion to balance emissions may therefore be incorrect.

One-third to one-half of the Earth’s land surface has been transformed by human action[2,4]. Many continuing transformations exchange woody and herbaceous plants[15-18], including deforestation, desertification, and woody plant invasion (the expansion of woody species into grasslands and savannas). Shifting dominance among herbaceous and woody vegetation alters primary production, plant allocation, rooting depth and soil faunal communities, potentially metres underground[15,18], in turn affecting nutrient cycling and climate storage[19,20]. The two carbon pools most likely to change are woody plant biomass and soil organic matter, the dominant pool of carbon and nitrogen in grasslands[10,20-23]. New woody biomass stores carbon in amounts that depend on the age, productivity and density of the stand. Changes in soil organic carbon (SOC) over the entire rooting zone are much harder to predict, and have the potential to enhance or offset biomass carbon gains, complicating projections of ecosystem carbon storage.

On the basis of a global analysis of more than 2,700 SOC profiles[23], we examined where increased biomass C in woody plants might be offset by SOC losses (see Methods). Across the global data set, the slope of the relationship between SOC and precipitation was 2.6 times higher for grassland vegetation than for shrublands/woodlands (P = 0.001; see Supplementary Information). Whereas grassland SOC was statistically indistinguishable from values for woody plants at 200 mm mean annual precipitation, woodlands had 43% less total SOC than grasslands at 1,000 mm (P < 0.01). Although suggestive, this analysis lacked a direct test of vegetation change independent of other covarying factors, including soil properties. So we also examined the direct effect of vegetation change at six paired grassland and invaded woody sites along a