Femtosecond Thermal and Nonthermal Hot Electron Tunneling Inside a Photoexcited Tunnel Junction

Natalia Martín Sabanés,† Faruk Krecinic,† Takashi Kumagai, Fabian Schulz, Martin Wolf, and Melanie Müller*

ABSTRACT: Efficient operation of electronic nanodevices at ultrafast speeds requires understanding and control of the currents generated by femtosecond bursts of light. Ultrafast laser-induced currents in metallic nanojunctions can originate from photoassisted hot electron tunneling or lightwave-induced tunneling. Both processes can drive localized photo-currents inside a scanning tunneling microscope (STM) on femto- to attosecond time scales, enabling ultrafast STM with atomic spatial resolution. Femtosecond laser excitation of a metallic nanojunction, however, also leads to the formation of a transient thermalized electron distribution, but the tunneling of thermalized hot electrons on time scales faster than electron–lattice equilibration is not well understood. Here, we investigate ultrafast electronic heating and transient thermionic tunneling inside a metallic photoexcited tunnel junction and its role in the generation of ultrafast photocurrents in STM. Phase-resolved sampling of broadband terahertz (THz) pulses via the THz-field-induced modulation of ultrafast photocurrents allows us to probe the electronic temperature evolution inside the STM tip and to observe the competition between instantaneous and delayed tunneling due to nonthermal and thermal hot electron distributions in real time. Our results reveal the pronounced nonthermal character of photoinduced hot electron tunneling and provide a detailed microscopic understanding of hot electron dynamics inside a laser-excited tunnel junction.

KEYWORDS: hot electron tunneling, ultrafast scanning tunneling microscopy, femtosecond electron dynamics, THz near-field sampling, lightwave-induced tunneling, two-temperature model

Hot carriers harvested in solids through optical excitation can drive a wide range of physical and chemical processes, the understanding of which lies at the heart of hot carrier science and technology.1 The ultrafast dynamics of photoexcited hot carriers are intimately related to optoelectronic nanodevice operation. A particular example is laser-excited ultrafast scanning tunneling microscopy (STM), which pursues the ultimate goal to image the ultrafast dynamics of individual atoms and molecules on surfaces with simultaneous Ångstrom spatial and femtosecond temporal resolution. Recent advances in combining ultrashort optical or terahertz (THz) pulses with low-temperature STM demonstrated the potential of light-driven STM to reach that goal.2–6 Attaining ultrahigh spatiotemporal resolution in STM requires the generation of ultrafast, atomically localized tunneling currents. In the weak excitation regime, photoassisted tunneling of hot electrons below the vacuum barrier gives rise to localized ultrafast currents that can be used for imaging in photon-driven STM.6–8 In contrast, lightwave-driven STM operates in the strong-field regime, where an intense localized light field of several V/nm modulates the junction barrier and, hence, the tunneling current on subcycle time scales.2,4,5,9–13 The two regimes are commonly distinguished by the Keldysh parameter, γ = √F l/4πω, relating the work function of the material, ϕ, to the photon energy, E ph = hω, and the light field, F l.14

Received: May 17, 2022
Accepted: August 19, 2022
Published: August 26, 2022
Both processes are distinctly different in terms of the involved electronic excitation: Whereas photoassisted tunneling employs the tunneling of “hot” but nonthermal electrons, lightwave-driven STM is mediated by adiabatic tunneling of thermal but “cold” electrons directly from the Fermi level. Herein, the terms “hot” and “cold” describe whether there is excess energy (e.g., via absorption) stored in the electronic system, whereas the terms “thermal” and “nonthermal” state whether the electronic distribution follows Fermi–Dirac statistics. Focusing ultrashort laser pulses on a metal gives rise to light absorption and the generation of energetic, nonthermal electrons, which rapidly redistribute their energy via electron–electron and electron–phonon scattering, leading to the formation of a thermalized carrier distribution on the 10–100 fs time scale. Hot electrons from both thermal and nonthermal distributions can tunnel through the potential barrier in STM, though with different time scales and energy distributions of the respective tunneling currents. Although the ultrafast dynamics and thermalization of hot electrons in metals has been intensively investigated,15,16 the transient tunneling of thermalized hot electrons is rarely considered as source for ultrafast photocurrents in STM. However, ultrafast thermionic tunneling should coexist with both photon-driven and lightwave-driven tunneling currents and thus accompanies any pulsed laser-driven STM. Moreover, the dynamics of hot electrons determine the initial step of laser heating of STM tips illuminated by ultrashort laser pulses.17,18 It is thus of general interest to understand the role of hot electron dynamics for femtosecond heating and tunneling in photoexcited STM.

Transient thermionic emission has been discussed in the context of ultrafast electron guns using laser-excited nanotips.19–22 It describes the emission of electrons above the vacuum barrier originating from the high-energy tail of a time-dependent Fermi–Dirac carrier distribution at elevated temperatures23,24 and can be described by the Richardson–Dushman equation.25 In contrast to freestanding tips, the nanometer-sized tip–sample gap in STM allows electrons to tunnel through the barrier at all energies down to the Fermi level; therefore, ultrafast thermionic currents can flow at much lower electron temperatures. The situation is similar to thermally enhanced field emission from a high voltage (kV) biased nanotip emitter19,26,27 operating at fields of >108 V/m, which can be described by Murphy’s and Good’s equation for thermo-field emission.28 To understand thermionic tunneling on femtosecond time scales, the ultrafast evolution of the electronic temperature after laser excitation has to be known. Before electron–phonon equilibration has been achieved (>1 ps), different temperatures are assigned to the electronic and phononic subsystems. Their respective time evolutions can be described by the two-temperature model (TTM)15,19 or by solving Boltzmann transport equations16,20 for the coupled electron–phonon system. Whereas photoassisted tunneling and optical-field-driven tunneling are temporally confined to the presence of the laser pulse, thermionic tunneling can occur on time scales longer than the laser pulse duration and might lead to significantly delayed photocurrents in STM due to electron dynamics inside the tip.

Here we study the interplay between ultrafast thermal and nonthermal tunneling from a femtosecond laser-excited STM tip. By tracking the temporally delayed response of thermionic currents in real-time we can discriminate between noninstantaneous thermal and instantaneous nonthermal currents, without the need to distinguish their energy distributions as for example in photoemission experiments.20,21,29 Energy information could, in principle, be obtained in STM by varying the bias voltage, but the bidirectional tunneling of photocarriers between tip and sample complicates interpretation of the respective hot carrier distributions. Instead, we measure the temporal response of ultrafast thermionic currents via phase-resolved sampling of single-cycle ultrabroadband THz pulses inside the tip–sample junction.30 Specifically, we measure the time-dependent THz-induced change of photocurrents from the laser-excited STM tip,30–32 which is determined by the instantaneous THz bias $U_{\text{THz}}(\tau)$ and the time evolution of the photocurrent $f_{\text{ph}}(\tau)$. In the case of prompt currents, the THz-field-induced photocurrent modulation yields the tip-enhanced THz near-field waveform. In contrast, delayed photocurrents will suppress higher THz frequencies and modify the measured waveform similar to a low-pass filter that limits sampling bandwidth. Knowledge of the original instantaneous THz waveform combined with numerical simulation of the time-dependent tunneling process allows us to extract the thermionic photocurrent dynamics and relate it to the electronic temperature evolution inside the STM tip and to reveal the competition between thermal and nonthermal photocurrent contributions.

RESULTS AND DISCUSSION

Phase-Resolved THz Sampling of Ultrafast Photocurrents in STM. Figure 1a shows the experimental scheme of our measurement. We excite a grounded tungsten STM tip with 800 nm near-infrared (NIR) laser pulses of 10 fs duration with peak intensities ranging from $0.8 \times 10^{11}$ to $2.5 \times 10^{11}$ W/cm², which corresponds to laser powers between 0.11 and 0.36 mW under our conditions. The tip is placed above the Ag(111) sample at a distance $d$. A positive bias of $U_{dc} = 8$ V is applied to the sample to suppress photocurrents from the sample with opposite sign and to ensure unidirectional flow of photocarriers originating from the tip. Phase-stable single-cycle broadband THz pulses are focused into the STM junction at variable delay $\tau$ with respect to the NIR pulses. The quasi-instantaneous field of the tip-enhanced THz field acts as a transient bias modulating the potential barrier and hence the current on femtosecond time scales. Provided that the THz-induced change of the photocurrent, $\Delta f_{\text{THz}}(\tau)$, is dominated by a process that is instantaneous on the time scale of the THz field, the THz bias can be sampled with a time resolution as determined by the NIR pulse duration. The delay-dependent $\Delta f_{\text{THz}}(\tau)$ reveals the transient THz bias, and the local slope of the photocurrent–voltage curve allows us to calibrate the THz bias amplitude $U_{\text{THz}}(\tau)$.30,32 Precise knowledge of the waveform and amplitude of the THz bias will be essential to extract the time evolution of ultrafast noninstantaneous currents photoexcited inside the metallic junction, which will lead to temporal distortions of the measured THz waveform as illustrated in Figure 1a. We point out the difference from “state-selective” THz-gated tunneling, where the resonant THz-induced tunneling into individual molecular states during the peak of a THz half-cycle yields a time resolution faster than the THz half-cycle. Due to the threshold character of the resonant tunneling process, this regime does not require detailed knowledge of the THz waveform to access subcycle dynamics. In contrast, accessing the dynamics of “nonresonant” systems (i.e., systems with a continuous variation
The different photocurrent mechanisms that can be operative for a photoexcited STM tip are displayed in Figure 1b. At nanometer gap sizes, photoassisted tunneling at energies of the NIR pulse duration, above-barrier photoemission and current channels from the laser-excited STM tip. On the time scale through the barrier from a nonthermal steplike electron temperature photocurrents depending on the time evolution of the electron of thermalized hot electrons (red arrow) can lead to delayed their respective bias dependence and emission times.

The intensity at which this transition occurs is independent of the number of electrons per pulse (see Figure S.2), implying that space charge effects are not responsible for the decreasing slope. Similar data from another tip are shown in Figure S.3, where pronounced waveform changes are observed purely in the multiphoton photoemission regime. The observed waveform changes thus emerge in both the weak- and strong-field regimes of photoemission. In both regimes, the photoemission process is prompt, and the emitted current will be temporally confined to the laser pulse width. Hence, photoemission cannot explain the observed low-pass filter behavior, which requires a “slow” photocurrent contribution (i.e., a delayed carrier response). We conclude that an additional delayed photocurrent component, which coexists with instantaneous weak- and strong-field photocurrents, is responsible for the observed waveform changes. As will be discussed below, the observed waveform changes can be assigned to delayed ultrafast thermionic currents from the STM tip, whose electronic subsystem is heated on the femtosecond time scale.
like distribution and (ii) thermal electrons that follow a Fermi−Dirac distribution, as illustrated by the blue and red distributions in Figure 1b, respectively. Photoexcitation by an ultrashort laser pulse initially promotes electronic single-particle excitations, leading to a nonthermal electron distribution that thermalizes via electron−electron scattering.
with a rate \( \alpha E / \epsilon_{2-\omega} \) which takes place on time scales typically of a few 10 fs in transition metals.\(^{15,16,43}\) On longer time scales of \( \sim 1 \) ps, electron–phonon scattering leads to energy transfer from the electronic to the phononic subsystem with a rate \( \alpha E / \epsilon_{2-\omega} \). Due to the different time scales of \( \epsilon_{2-\omega} \) and \( \epsilon_{2-\omega} \), it is reasonable to assume that electron–phonon coupling only leads to cooling of the thermal electron distribution, but it does not remove energy from the nonthermal part of the electronic system. In addition, ballistic transport and diffusion of hot electrons into the bulk can influence the electron distributions at the tip surface.\(^{15,44}\) Both nonthermal and thermal electron distributions can give rise to a photocurrent from the tip, which we write as the sum

\[
 j_{\text{ph}}(t) = j_{\text{th}}(t) + \delta(t - t_0) j_{\text{nonth}}
\]

of an ultrafast time-dependent thermionic current \( j_{\text{th}}(t) \) and an instantaneous nonthermal current \( j_{\text{nonth}} \), where \( \delta(t - t_0) \) is a delta function located at the laser pulse center \( t_0 \). The separation in eq 1 and the assumption of an instantaneous nonthermal current is reasonable here considering the limited time resolution of 10s of fs determined by the finite THz bandwidth, which is insufficient to resolve the electron thermalization process via electron–electron scattering.

The ultrafast thermionic current through the THz-modulated potential barrier is given by the total charge \( Q_{\text{th}} \) emitted per unit area per pulse due to a thermalized hot electron distribution,\(^{28,45}\)

\[
 j_{\text{th}}(\tau) \propto Q_{\text{th}}(\tau)
\]

\[
 = e \int D(W, F(t - \tau)) N_{\text{th}}[E, T_{\text{el}}(t)] \, dW \, dE \, d\tau
\]

where \( \tau \) is the delay between THz and NIR pulse, \( F(t) = E_k + F_{\text{THz}}(t) \) is the combined dc and THz electric field, and \( D \) is the time-dependent transmission probability, where \( W = E - (p_z^2 + p_r^2) / 2m_e \) is the kinetic energy of the electron in the positive z-direction. \( N_{\text{th}} \) is the time-dependent thermal electron occupation at energy \( E \) and temperature \( T_{\text{el}} \) which follows the Fermi–Dirac distribution

\[
 N_{\text{th}}(E, T_{\text{el}}(t)) = \frac{1}{\exp[(E - E_F)/k_B T_{\text{el}}(t)] + 1}
\]

The THz-modulated transmission probability \( D \) is calculated by solving the one-dimensional Schrödinger equation

\[
 -\frac{\hbar^2}{2m_e} \frac{d^2 \psi}{dz^2} + V(F, z) \psi = W \psi
\]

for the time-dependent quasi-static potential

\[
 V(t, z) = \begin{cases} 
 0 & z \leq 0 \\
 E_F + \phi - \frac{\hbar^2}{16\pi\epsilon_0 e^2} - eF(t)z & z > 0 
\end{cases}
\]

where \( \hbar \), \( \epsilon_0 \), and \( m_e \) are the reduced Planck constant, vacuum permittivity, and electron mass, respectively, \( E_F \) is the Fermi level, \( \phi \) is the work function of the tip, and the third term in eq 5 describes the effect of image charges. Due to the high positive sample bias we can neglect current contributions originating from the sample in the present calculations. The time evolution of the electronic temperature \( T_{\text{el}}(t) \) is obtained by solving the two-temperature model (TTM) in three dimensions inside the tip including ballistic and diffusive electron transport using COMSOL Multiphysics (details are described in section 4 of the Supporting Information). The TTM source term \( S_\text{el} \) for electronic heating is obtained from the spatially inhomogeneous absorbed power density, convoluted with the mean free path of electrons in tungsten\(^{15,46}\) to account for the initial fast redistribution of energy due to ballistic transport. The left panel in Figure 3b shows the spatial profile of \( S_\text{el} \) inside the tip after ballistic energy redistribution, which on the nanometric scale of the tip occurs instantaneously on the time scale of the THz field. For a given tip–sample geometry, we obtain the ultrafast electronic temperature evolution, the locally enhanced optical field as well as the dc field along the surface of the tip using COMSOL simulations. The middle and right panel in Figure 3b show the electronic temperature distribution inside the tip at \( \tau = 20 \) fs and \( \tau = 1 \) ps, respectively. \( T_{\text{el}} \) initially resembles the profile of \( S_\text{el} \) but becomes blurred on longer time scales due to diffusive hot carrier transport. Finally, the experimental calibration of the THz bias allows us to fix the ratio of dc and THz field. As we know the waveform of the THz bias, we can thus calculate the THz-induced change of the ultrafast thermionic current by calculating the emitted charge \( \Delta Q_{\text{th}}(\tau) = Q_{\text{th}}(\tau) - Q_{\text{th}}(\tau \ll 0) \) in the presence of the instantaneous THz field using eq 2.

The quasi-instantaneous photocurrent due to nonthermal electrons can be written as a superposition of currents through each multiphoton channel \( n \)

\[
 j_{\text{nonth}} \propto Q_{\text{nonth}} = C_0 \sum_n Q_n
\]

with

\[
 Q_n = D_n N_n = D_n [(C_{\text{ex}})^n I_{\text{IR}}^{2n}]
\]

where \( Q_n \) is the charge emitted per unit area per pulse through channel \( n \), \( I_{\text{IR}} \) is the tip-enhanced laser field, \( C_0 \) is a constant scaling factor, and \( D_n \) and \( N_n \) are the electron density and transmission probability at the surface at energy \( E_n = E_F + n\hbar \omega \) with \( n = [1, 2, 3, ...] \), respectively. The excitation constant \( C_{\text{ex}} \ll 1 \) is a free parameter whose dependence on \( n \) accounts for the decreasing probability for excitation into higher multiphoton channels. However, the relative contribution from higher channels increases with increasing laser intensity as expected due to \( I_{\text{IR}}^{2n} \). Equation 6 thus follows the intensity scaling and \( n \)-dependent excitation probability expected for multiphoton absorption in the perturbative regime.\(^{27} \) The THz-induced change of the nonthermal charge density, \( \Delta Q_{\text{nonth}}(\tau) \), is calculated from eq 6 taking into account the time dependence of the transmission probability \( D_n \) due to the instantaneous THz field at the tip surface. Finally, the simulated waveforms are obtained from the sum \( \Delta Q_{\text{THz}}(\tau) = \Delta Q_{\text{th}}(\tau) + \Delta Q_{\text{nonth}}(\tau) \). The free parameters of our model are the incident laser intensity, the constants \( C_0 \) and \( C_{\text{ew}} \), and the tip–sample geometry including the gap distance \( d \).

At high laser fields approaching \( \gamma_{\text{IR}} \sim 1 \), above-threshold photoemission becomes significant. Yet, as those channels are relatively insensitive to the potential barrier and hence the THz bias, we neglect them here and restrict our calculations to
channels below or close to the top of the barrier \((n \leq 3\) in our case). At even higher laser fields, entering the optical tunneling regime at \(h_{\text{IR}} \ll 1\), the laser field is significantly stronger than the applied THz and dc fields. Hence, optical tunneling should be almost unaffected by the weak THz fields applied here; therefore, the THz-induced change of optical-field-driven tunneling currents is expected to be very small. We simulated NIR-lightwave-tunneling from the THz-biased tip by adding the NIR laser field to the total field \(F(t)\) in eq 5. The results confirmed that optical tunneling contributes insignificantly to the THz-induced change of the photocurrent due to its low bias sensitivity in the parameter range used here. We can thus neglect strong-field effects and optical tunneling in our waveform simulations. We further note that the measured photocurrent is NIR-cycle-averaged and integrated over energy. In contrast to energy-resolved measurements, which record the final kinetic energy of the photoelectrons after propagating through the oscillating THz field,\(^\text{31}\) our measurement is not sensitive to electron propagation but only to the instantaneous THz field at the tip surface. Since \(E_{\text{dc}} \gg F_{\text{THz}}\), we can also neglect THz streaking of photoelectrons back into the tip.

**Simulation of Ultrafast Thermionic Currents and THz Waveforms from a Photoexcited STM Tip.** Figure 3c shows calculated THz waveforms obtained at the apex of a tip with work function \(\phi = 4.5\) eV, tip radius \(R_{\text{tip}} = 5\) nm, and shaft opening angle \(\alpha_{\text{tip}} = 4^\circ\) (a SEM image of the STM tip used in this work is shown in Figure S.5). The simulations with incident laser intensities close to those estimated from our experimental conditions reproduce very well the measured waveform broadening and temporal shift. The exact laser intensities required to observe electronic heating in the simulated waveforms vary with tip size and shape, which is consistent with the experimental variation of the laser power we require to observe waveform deformations for different tip conditions. The corresponding time dependence of the electron temperature and thermionic current are shown in Figure 3d,e, respectively. Electronic temperatures of several 1000 K are reached inside the tip on ultrafast time scales due to the small heat capacity of electrons in metals.\(^\text{48}\) It is important to note that thermionic currents alone do not sufficiently describe the waveforms, especially at low laser intensities at which \(\Delta Q_{\text{th}}(t)\) does not yield the original THz waveform (pure "thermionic waveforms" are shown in Figure S.4.2). To reproduce the laser power dependence of the waveforms in Figure 2b precisely, the combination of thermal and nonthermal currents is required. Whereas nonthermal currents, at far distances predominantly emitted through \(n = 3\), dominate and reproduce the original THz waveform at low laser intensities, delayed thermoionic currents are responsible for the waveform distortions at higher laser intensities. Figure 3f shows the calculated power scaling of the THz-induced change of the thermionic charge density \(\Delta Q_{\text{th}}\) through channel \(n = 3\) compared to the power scaling of the nonthermal charge density \(\Delta Q_{\text{nth}}\) through channel \(n = 3\). The slope of \(\Delta Q_{\text{th}}\) is much steeper than that of \(\Delta Q_{\text{nth}}\) and continuously decreases at higher laser intensities, as opposed to the constant slope \(n = 3\) of the nonthermal photocurrent. Whereas the scaling of \(\Delta Q_{\text{nth}}\) is determined solely by the excitation into states at \(E_n = 3\hbar \omega_0\), which increases as \(I_{\text{IR}}^2\), the power scaling of \(\Delta Q_{\text{th}}\) is determined by the increasing electron temperature and its decreasing sensitivity to the barrier, and hence THz field, at high \(T_{\text{el}}\). Overall, the relative contribution of \(\Delta Q_{\text{th}}\) is insignificant at low intensities, but it increases rapidly with increasing laser intensity. It has been discussed previously that noninteger nonlinearities, which are frequently measured for photoemission from nanotips, can be caused by long-lived hot electron distributions inside the tip,\(^\text{49,50}\) which is supported by our results. We note that we can exclude that the delayed currents originate from laser-driven rescattering\(^\text{51}\) and delayed re-emission of electrons from the tip,\(^\text{52}\) as the waveform distortions are observed also (and for some tips solely) in the weak-field regime, in which laser-driven electron scattering is negligible (section S3 of the Supporting Information). The results in Figures 2 and 3 thus confirm that femtosecond laser heating of the STM tip can lead to ultrafast thermal currents persisting on time scales of several 100 fs (i.e., much longer than the exciting laser pulse duration).

**Gap Size Dependence.** To examine the role of thermalized hot electrons for photoinduced tunneling in STM, we measure the THz waveform versus tip–sample distance as plotted in Figure 4a. The NIR power is set to a value that yields significant waveform deformations due to delayed thermal photocurrents at large tip–sample distances. The zero position \(\Delta z = 0\) nm is defined by the STM set point of \(U_{\text{dc}} = 10\) V and \(j = 1\) nA, at which the current is dominated by static tunneling. The relative gap distance is varied in the range between \(-0.2\) and 6 nm (negative values mean smaller gap size). We can roughly estimate the absolute tip–sample distance by fitting the decay of the static current (dashed line in Figure 4b) and extrapolating to the quantum conductance \(G_0\) which yields \(d \approx 2.9\) nm for \(\Delta z = 0\) nm, which is reasonable at our conditions. As demonstrated previously, the THz bias does not depend on gap size;\(^\text{11,50}\) therefore, the THz bias measured at large gap distance (gray dashed line) applies to all distances. We note that “static” rectified THz currents are negligible due to the small THz bias amplitude and the comparably large gap distance when operating at 10 V bias. As discussed in the Supporting Information, our measurements are not affected by quasi-static or transient (pulse-to-pulse) thermal expansion of the STM tip due to pulsed laser illumination. Starting from a distorted waveform at \(\Delta z = 6\) nm, the waveform continuously transforms such that it reproduces the original waveform of the THz bias at the smallest gap size. Complete vanishing of the thermal waveform distortions correlates with the onset of static tunneling, as evident from the distance dependence of the current plotted in Figure 4b. This is surprising, as naively one would expect a slower current decay due to increased tunneling at lower energies, for which the occupation and current decay times are larger than at the high-energy tail of the Fermi distribution. In addition, ultrafast heating will be enhanced at nanometer distances, because the optical field, and hence the absorbed power, are inversely proportional to the gap size. The dependence of the waveform deformations on the gap distance, which is distinctly different from the dependence on laser power shown in Figure 2b, thus clearly indicates that instantaneous current contributions must exceed thermionic tunneling at reduced gap size close to the static tunneling regime. Figure 4c shows the effective nonlinearity extracted from current-distance curves measured at varying laser pulse energy (details are described in section S6 in the Supporting Information).
Figure 4. Gap size dependence of measured THz waveforms. (a) THz near-field waveforms measured against relative gap distance $\Delta z$ from the set point. The THz waveform gradually transforms into the original THz bias waveform at short gap distances. (b) Gap size dependence of the total current (left y-axis) and THz-induced change of the photocurrent (right y-axis, peak value). Recovery of the original THz bias waveform correlates with the onset of static tunneling (dashed line). (c) Effective nonlinearity $n_{th}$ of the photocurrent versus relative gap distance (set point 1 nA and 10 V, laser power 2.1 mW).

Information). Whereas at large tip–sample distances the photocurrent exhibits a nonlinearity of $n_{th} \sim 3$, the nonlinearity continuously decreases with decreasing $\Delta z$, indicating that photoassisted tunneling of lower orders dominates at small gap sizes.

The observations in Figure 4 suggest that at close distances the instantaneous tunneling of nonthermal electrons competes with delayed thermionic tunneling from the laser-excited STM tip. To corroborate this assumption, we simulate THz waveforms for varying gap distances $d$. With decreasing $d$, the dc field, THz field, and optical field at the tip apex increase, as do the absorbed power and hence the electronic temperature inside the tip. Figure 5a,b shows the distance scaling of $T_{th}$, $E_{dc}$, and the optical field enhancement at the tip apex, together with the decay time $\tau_{th}$ of the ultrafast thermionic current. The longer decay times $\tau_{th}$ at reduced gap distances result from the higher electron temperatures and an increasing contribution from electrons tunneling at lower energies. Figure 5c compares the gap size dependence of $\Delta Q_{THz}$ due to thermal and nonthermal electrons of orders $n = [1, 2, 3]$, respectively. The THz waveforms calculated from the superposition of those currents are plotted as a function of $d$ in Figure 5d, which reproduces well the experimental observations in Figure 4a. We can thus explain the gap size dependence of the measured THz waveforms by the competition between delayed thermal and prompt nonthermal tunneling from the photoexcited STM tip. The distinctive THz waveform changes can be understood from the distance-dependent mixture of thermal and nonthermal photocurrents, whose relative contributions vary with gap size according to their specific tunneling probabilities and nonlinear dependence on the local laser field, as reflected in the different slopes in Figure 5c. Whereas the nonthermal contribution from $n = 3$ increases mainly due to its nonlinear dependence on local laser intensity, the lower order contributions $n = 2$ and, in particular, $n = 1$ increase rapidly at reduced gap size due to their steeply increasing tunneling rates. In contrast, the source term of electronic heating in the TTM scales linearly with laser intensity. Moreover, the thermionic current is integrated over energy and thus exhibits an overall reduced sensitivity to the barrier compared to low-order nonthermal tunneling channels. This becomes apparent from the energy distribution of $J_{th}$ plotted in Figure 5e together with the corresponding thermal electron occupation $N_{e}$ and transmission probability $D$ for three gap sizes at $\tau = 20$ fs. Even though the peak of the thermionic current distribution shifts to lower energies and develops a significant low-energy tail at nanometer gap distances, a majority of thermalized electrons are emitted at energies $\sim 2$ eV above $E_F$ and, hence, experiences a reduced sensitivity to the THz field compared to the nonthermal electrons excited into channel $n = 1$. It is the nonlinear laser intensity dependence of channels $n > 1$ on the one hand and the higher sensitivity of low-order channels to the tunneling barrier on the other hand that cause the instantaneous nonthermal currents to exceed the delayed thermionic current at nanometer gap sizes. Our findings clearly show the pronounced nonthermal and prompt character of hot electron-induced tunneling from a photoexcited STM tip.

Finally, we note that our simple model can reproduce the data reasonably well without calculation of the full nonthermal steplike distribution. The experimental data, however, shows a waveform change, within a distance variation of only 1 Å close to the set point, more rapid than that observed in the simulations. This indicates that our model might underestimate nonthermal hot electron tunneling close to the Fermi level, which exhibits an even higher sensitivity to the tip–sample distance. Further improvement of the agreement between the experimental and simulated results could be obtained by extending our model by (i) the simulation of the electron thermalization process, (ii) the inclusion of the full nonthermal distribution, and (iii) replacing the Fowler–Nordheim barrier by the Simmons barrier, \cite{Simmons} which is more appropriate at nanometer gap sizes. We expect that more detailed spectroscopic information about the nonthermal distributions might be obtained in future experiments by variation of the dc and THz bias. Moreover, it may be possible to measure the electron thermalization process and its effect on tunneling by using shorter THz transients to increase the time resolution.

CONCLUSIONS

In conclusion, we investigated the competition between ultrafast thermal and nonthermal photocurrents inside a
laser-excited STM junction via phase-resolved sampling of broadband THz transients. Our results reveal that hot electron tunneling from a laser-excited STM tip is dominated by nonthermal electron distributions, accompanied by delayed tunneling of thermalized hot electrons. At large gap sizes, at which photoinduced tunneling through the barrier is negligible, thermalized hot electrons dominate the THz-induced photocurrent from a tungsten tip excited at high laser intensities. However, at small gap distance, this current contribution is exceeded by nonthermal currents as soon as the probability for photoassisted tunneling at low energies below the vacuum level increases appreciably. We infer the markedly nonthermal character of the photoexcited electron distribution from the distinctly different durations of the thermal and nonthermal photocurrents. Prompt nonthermal tunneling is followed by a thermionic photocurrent that can be delayed by several 100 fs, as determined by the time at which the laser-excited STM tip reaches its peak electronic temperature, which can increase up to several thousand Kelvin. The weak THz field serves as a highly sensitive and selective probe for photoinduced tunneling currents in ultrafast laser-excited STM, which is a unique capability of THz-gated STM. Detailed information about the phase and amplitude of the THz bias allows one to extract the ultrafast dynamics of hot electrons on time scales much shorter than a single THz cycle even in nonresonant systems like metals or semimetals. Our results provide detailed microscopic insight into the nonequilibrium dynamics of hot electrons in a laser-excited STM tip, which is of general relevance to experiments that aim to study photophysical or photochemical processes on ultrafast time scales with pulsed-laser excited STM. We envision that extension of our approach by the imaging capabilities of STM will allow for the spatiotemporal investigation of ultrafast carrier dynamics at photoexcited surfaces on Angstrom length and femtosecond THz-subcycle time scales.

Figures 5. Simulation of photocurrent channels and THz waveforms versus tip–sample distance. (a) Distance scaling of electron peak temperature (left) and decay time of the thermionic current (right). (b) Distance scaling of optical field enhancement (left) and dc field (right). (c) Dependence of the THz-induced charge density $\Delta Q_{\text{THz}}$ on gap size for thermionic tunneling (red) and nonthermal photoassisted tunneling at energies $E_n = n \hbar \omega$ (dashed) at the peak of the THz pulse. (d) THz waveforms calculated for the combined photocurrent channels shown in (c) at varying gap distance. (e) Energy distribution curves of the thermionic current (solid red, left y-axis) at delay $\tau = 20$ fs after photoexcitation for three gap distances 1 $\mu$m (bottom), 6 nm (middle), and 2 nm (top). The black dashed curves show the corresponding Fermi–Dirac distributions ($N_{\text{th}}$, log-scale) with peak temperatures of $T_{\text{th}} = 5722$ K at 1 $\mu$m, $T_{\text{th}} = 7466$ K at 6 nm, and $T_{\text{th}} = 7901$ K at 2 nm. The black solid curve shows the transmission probability $D$ in the absence of the THz field. The blue vertical lines mark the positions of the nonthermal photocurrent channels $n = \{1, 2, 3\}$. The gray shaded area shows the Fermi-Dirac distribution at room temperature.

METHODS

Measurements are recorded with a customized STM from Unisoku (USM-1400 with Nanonis SPM controller) operated at room temperature and under ultrahigh vacuum conditions (base pressure of $<5 \times 10^{-10}$ mbar). The spring-loaded STM platform is equipped with two off-axis parabolic mirrors (PM, 1 bare Au and 1 protected Ag, 1 in. diameter, 35 mm focal length) for illumination with a broadband NIR laser and broadband THz pulses, respectively. Both beams are incident to the tip axis at an angle of 68° and are polarized along the tip axis. The THz beam enters the UHV chamber through a 500 $\mu$m thick diamond window and is focused by the Au PM. The Ag PM is used to focus the NIR pulses for photoexcitation of the STM junction. Precise focus alignment on the tip apex is ensured by precise
positioning of the PMs in UHV, which are motorized and can be moved in the x-, y-, and z-directions (Attocube GmbH). The tip position is fixed, and the sample is moved for coarse motion and scanning. The dc bias is applied to the sample and the current is collected from the grounded tip. The current preamplifier (Femto DLP-CA) is operated at a gain of 10^4 V/A at 1 kHz bandwidth. A mechanical chopper operated at a frequency of 607 Hz modulates the power of the NIR laser beam used for THz generation, and lock-in detection is used to detect the THz-induced current. Repeated cycles of Ar+ sputtering and annealing up to 670 K were conducted to clean the Ag(111) sample before measuring. Electrochemically etched tungsten tips are transferred to UHV immediately after etching.

The laser system is a broadband optical parametric chirped-pulse amplifier (Venteon OPCPA, Laser Quantum) delivering NIR laser pulses of 10 fs duration at 800 nm center wavelength and with 3 μJ energy (2 μJ are available for the THz-STM setup) at f_rep = 0.5 MHz repetition rate. Part of the laser power is used for the generation of single-cycle THz pulses from a spintronic THz emitter (STE; 5.8 nm thick W/CoFeB/Pt trilayer on 500 μm sapphire substrate) excited at normal incidence in transmission geometry. A motorized translation stage is used to control the delay between the THz pulses and the NIR laser pulses for photoexcitation of the STM. Part of the NIR laser pulses can be overlapped collinearly with the THz beam for THz pulse characterization using electro-optic sampling, as well as for precise THz beam alignment inside the STM. Peak laser intensities are calculated assuming a sech^2 laser pulse profile, for which the peak intensity is determined as I_peak = 0.88 × \( \Phi_{\text{peak}} / f_{\text{rep}} \), where \( \Phi_{\text{peak}} \) is the fwhm laser pulse duration and \( \Phi_{\text{peak}} = E_p/(n w^2/2) \) is the peak fluence in the center of the Gaussian beam with waist \( w \), and \( E_p = P_p / f_{\text{rep}} \) is the pulse energy at a given laser power \( P_p \). A beam waist of \( w = 4 \) μm was used in the calculation, which was measured inside the STM under ambient conditions after the same alignment procedure we follow under normal operation conditions.

Numerical simulations are carried out using COMSOL Multiphysics 5.6, which provides the required values of the dc field, tip-enhanced optical field, and electronic temperature evolution inside the tip. The photoinduced thermal and nonthermal currents and their THz-induced change (THz waveforms) are simulated using custom Python scripts. Details about the model are described in the Supporting Information.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nano.2c04846.

Calibration of the THz bias, bias dependence of photocurrent-power scaling at large gap size, THz waveforms for a tip operating solely in the weak-field regime, theoretical model, SEM image of STM tip, power-dependent current-distance curves, discussion of thermal expansion (PDF)

AUTHOR INFORMATION

Corresponding Author
Melanie Müller — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany; orcid.org/0000-0002-3269-2107; Email: m.mueller@hhi-berlin.mpg.de

Authors
Natalia Martin Sabanes — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany; IMDEA Nanoscience, 28049 Madrid, Spain; orcid.org/0000-0003-4704-0408

Faruk Krecinic — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany
Takashi Kumagai — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany; Center for Mesoscopic Sciences, Institute for Molecular Science, 444-8585 Okazaki, Japan
Fabian Schulz — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany; orcid.org/0000-0002-1359-4675
Martin Wolf — Department of Physical Chemistry, Fritz Haber Institute of the Max Planck Society, 14195 Berlin, Germany

Complete contact information is available at: https://pubs.acs.org/doi/10.1021/acsnano.2c04846

Author Contributions
N.M.S. and F.K. contributed equally to this work. M.M. conceived the experiment. N.M.S. and M.M. carried out the measurements and analyzed the data. F.K., N.M.S., and M.M. carried out the numerical simulations. F.K. developed the python codes. T.K., M.M., and M.W. designed the STM instrumentation. N.M.S., F.S., and M.M. constructed the experimental setup. M.M. wrote the manuscript. All authors contributed to the discussion and commented on the manuscript, and have given approval to the final version of the manuscript.

Funding
Open access funded by Max Planck Society.

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors thank A. Paarmann, E. Ernstorfer, and T. Kamprath for valuable discussions. The authors further thank Unisoku Inc. for support and discussions in the development of the STM instrumentation. We thank T. Kamprath, T. Seifert, G. Jakob, and M. Kläui for providing us with the spintronic THz emitter, H. Kirsch for tip preparation support, and A. Hammud for recording SEM images of the tip. N.M.S. acknowledges the MSCA program MSCA-IF-2019-892667.

ABBREVIATIONS

STE spintronic THz emitter

REFERENCES

(1) Brongersma, M. L.; Halas, N. J.; Nordlander, P. Plasmon-Induced Hot Carrier Science and Technology. Nat. Nanotechnol. 2015, 10, 25–34.
(2) Cocker, T. L.; Peller, D.; Yu, P.; Repp, J.; Huber, R. Tracking the Ultrafast Motion of a Single Molecule by Femtosecond Orbital Imaging. Nature 2016, 539, 263–267.
(3) Li, S.; Chen, S.; Li, J.; Wu, R.; Ho, W. Joint Space-Time Coherent Vibration Driven Conformational Transitions in a Single Molecule. Phys. Rev. Lett. 2017, 119, 176002.
(4) Garg, M.; Kern, K. Attosecond Coherent Manipulation of Electrons in Tunneling Microscopy. Science 2020, 367, 411–415.
(5) Yoshida, S.; Arashida, Y.; Hirori, H.; Tachizaki, T.; Taninaka, A.; Ueno, H.; Takeuchi, O.; Shigekawa, H. Terahertz Scanning Tunneling Microscopy for Visualizing Ultrafast Electron Motion in Nanoscale Potential Variations. ACS Photonics 2021, 8, 315–323.

(6) Wu, S. W.; Ho, W. Two-Photon-Induced Hot-Electron Transfer to a Single Molecule in a Scanning Tunneling Microscope. Phys. Rev. B 2010, 82, 085444.

(7) Schröder, B.; Bunjes, O.; Wimmer, L.; Kaiser, K.; Traeger, G. A.; Kgotzott, T.; Ropers, C.; Wenderoth, M. Controlling Photocurrent Channels in Scanning Tunneling Microscopy. New J. Phys. 2020, 22, 033047.

(8) Garg, M.; Martin-Jimenez, A.; Luo, Y.; Kern, K. Ultrafast Photon-Induced Tunneling Microscopy. ACS Nano 2021, 15, 18071–18084.

(9) Cocker, T. L.; Jelic, V.; Gupta, M.; Molesky, S. J.; Burgess, J. A. J.; Reyes, G. D. L.; Titova, L. V.; Tsui, Y. Y.; Freeman, M. R.; Hegmann, F. A. An Ultrafast Terahertz Scanning Tunneling Microscope. Nat. Photonics 2013, 7, 620–625.

(10) Yoshioka, K.; Katayama, I.; Minami, Y.; Kitajima, M.; Yoshida, S.; Shigekawa, H.; Takeda, J. Real-Space Coherent Manipulation of Electrons in a Single Tunnel Junction by Single-Cycle Terahertz Electric Fields. Nat. Photonics 2016, 10, 762–765.

(11) Jelic, V.; Iwaszkusz, K.; Nguyen, P. H.; Rathje, C.; Hornig, G. J; Sharam, H. M.; Hoffman, J. R.; Freeman, M. R.; Hegmann, F. A. Ultrafast Terahertz Control of Extreme Tunnel Currents through Single Atoms on a Silicon Surface. Nat. Phys. 2017, 13, 591–598.

(12) Yoshioka, K.; Katayama, I.; Arashida, Y.; Ban, A.; Kawada, Y.; Konishi, K.; Takahashi, H.; Takeda, J. Tailoring Single-Cycle Near Field in a Tunnel Junction with Carrier-Envelope Phase-Controlled Terahertz Electric Fields. Nano Lett. 2018, 18, 5198–5204.

(13) Ammerman, S. E.; Jelic, V.; Wei, Y.; Breslin, V. N.; Hassan, M.; Everett, N.; Lee, S.; Sun, Q.; Pignedoli, C. A.; Ruffieux, P.; Fasel, R.; Cocker, T. L. Lightwave-Driven Scanning Tunneling Spectroscopy of Atomically Precise Graphene Nanoribbons. Nat. Commun. 2021, 12, 6794.

(14) Keldysh, L. Ionization in the Field of a Strong Electromagnetic Wave. Sov. Phys. JETP 1965, 20, 1307.

(15) Hohlfeld, J.; Wollershoffer, S.-S.; Gudde, J.; Conrad, U.; Jähnke, V.; Matthias, E. Electron and Lattice Dynamics Following Optical Excitation of Metals. Chem. Phys. 2000, 251, 237–258.

(16) Rethfeld, B.; Kaiser, A.; Vicanek, M.; Simon, G. Ultrafast Dynamics of Nonequilibrium Electrons in Metals under Femtosecond Laser Irradiation. Phys. Rev. B 2002, 65, 245303.

(17) Gerstner, V.; Knoll, A.; Pfeiffer, W.; Thon, A.; Gerber, G. Femtosecond Laser Assisted Scanning Tunneling Microscopy. J. Appl. Phys. 2000, 88, 4851.

(18) Gerstner, V.; Thon, A.; Pfeiffer, W. Thermal Effects in Pulsed Laser Assisted Scanning Tunneling Microscopy. J. Appl. Phys. 2000, 87, 2574.

(19) Keulhofer, C.; Foreman, S. M.; Gerlich, S.; Kasevich, M. A. Ultrastable Laser-Triggered Emission from Hafnium Carbide Tips. Phys. Rev. B 2012, 86, 035405.

(20) Yanagisawa, H.; Hengsberger, M.; Leuenberger, D.; Klöckner, M.; Hafner, C.; Greber, T.; Osterwalder, J. Energy Distribution Curves of Ultrashort Laser-Induced Field Emission and Their Implications for Electron Dynamics. Phys. Rev. Lett. 2011, 107, 087601.

(21) Yanagisawa, H.; Schnep, S.; Hafner, C.; Hengsberger, M.; Kim, D. E.; Klinger, M. F.; Landsman, A.; Gallmann, L.; Osterwalder, J. Delayed Electron Emission in Strong-Field Driven Tunneling from a Metallic Nanotip in the Multi-Electron Regime. Sci. Rep. 2016, 6, 35877.

(22) Barwick, B.; Corder, C.; Strohaber, J.; Chandler-Smith, N.; Uiterwaal, C.; Batelaan, H. Laser-Induced Ultrafast Electron Emission from a Field Emission Tip. New J. Phys. 2007, 9, 142.

(23) Riffé, D. M.; Wang, X. Y.; Downer, M. C.; Fisher, D. L.; Tajima, T.; Erskine, J. L.; More, R. M. Femtosecond Thermionic Emission from Metals in the Space-Charge-Limited Regime. J. Opt. Soc. Am. B 1993, 10, 1424–1435.
(46) Choi, D.; Kim, C. S.; Naveh, D.; Chung, S.; Warren, A. P.; Nuhfer, N. T.; Toney, M. F.; Coffey, K. R.; Barmak, K. Electron Mean Free Path of Tungsten and the Electrical Resistivity of Epitaxial (110) Tungsten Films. *Phys. Rev. B - Condens. Matter Mater. Phys.* 2012, 86, 045432.

(47) Nathan, V.; Guenther, A. H.; Mitra, S. S. Review of Multiphoton Absorption in Crystalline Solids. *J. Opt. Soc. Am. B* 1985, 2, 294-316.

(48) Lin, Z.; Zhigilei, L. V.; Celli, V. Electron-Phonon Coupling and Electron Heat Capacity of Metals under Conditions of Strong Electron-Phonon Nonequilibrium. *Phys. Rev. B* 2008, 77, 075133.

(49) Fujimoto, J. G.; Liu, J. M.; Ippen, E. P.; Bloembergen, N. Femtosecond Laser Interaction with Metallic Tungsten and Nonequilibrium Electron and Lattice Temperatures. *Phys. Rev. Lett.* 1984, 53, 1837-1840.

(50) Girardeau-Montaut, J. P.; Girardeau-Montaut, C. Theory of Ultrashort Nonlinear Multiphoton Photoelectric Emission from Metals. *Phys. Rev. B* 1995, 51, 13560-13567.

(51) Thomas, S.; Krüger, M.; Förster, M.; Schenk, M.; Hommelhoff, P. Probing of Optical Near-Fields by Electron Rescattering on the 1 Nm Scale. *Nano Lett.* 2013, 13, 4790-4794.

(52) Simmons, J. G. Generalized Formula for the Electric Tunnel Effect between Similar Electrodes Separated by a Thin Insulating Film. *J. Appl. Phys.* 1963, 34, 1793-1803.

(53) Seifert, T.; Jaiswal, S.; Martens, U.; Hannegan, J.; Braun, L.; Maldonado, P.; Freimuth, F.; Kronenberg, A.; Henrizi, J.; Radu, L.; Beaurepaire, E.; Mokrousov, Y.; Oppeneer, P. M.; Jourdan, M.; Jakob, G.; Turchinovich, D.; Hayden, L. M.; Wolf, M.; Münzenberg, M.; Klaua, M.; Kampfrath, T. Efficient Metallic Spintronic Emitters of Ultrabroadband Terahertz Radiation. *Nat. Photonics* 2016, 10, 483-488.