Femtosecond Field-Driven On-Chip Unidirectional Electronic Currents in Nonadiabatic Tunneling Regime

Liping Shi, Ihar Babushkin,* Anton Husakou, Oliver Melchert, Bettina Frank, Juemin Yi, Gustav Wetzel, Ayhan Demircan, Christoph Lienau, Harald Giessen, Misha Ivanov, Uwe Morgner, and Milutin Kovacev

Recently, asymmetric plasmonic nanojunctions have shown promise as on-chip electronic devices to convert femtosecond optical pulses to current bursts, with a bandwidth of multi-terahertz scale, although yet at low temperatures and pressures. Such nanoscale devices are of great interest for novel ultrafast electronics and opto-electronic applications. Here, the device is operated in air and at room temperature, revealing the mechanisms of photoemission from plasmonic nanojunctions, and the fundamental limitations on the speed of optical-to-electronic conversion. Inter-cycle interference of coherent electronic wavepackets results in a complex energy electron distribution and birth of multiphoton effects. This energy structure, as well as reshaping of the wavepackets during their propagation from one tip to the other, determine the ultrafast dynamics of the current. It is shown that, up to some level of approximation, the electron flight time is well-determined by the mean ponderomotive velocity in the driving field.

1. Introduction

As known from atomic physics,[1–4] with increase of the incident laser intensity, the ionization dynamics undergoes a transition from a relatively slow multiphoton process to fast sub-cycle bursts, referred to as tunneling ionization. The dimensionless Keldysh parameter \( \gamma = 2 \pi \tau_T / T \) characterizes, relative to the optical cycle duration \( T \), a typical time \( \tau_T = \sqrt{2 m \phi / |eE|} \), required for an electron to nearly completely leave the atom with the ionization potential \( \phi \) in an external field of the amplitude \( E \) (here \( m \) and \( e \) are the electron mass and charge). In the multiphoton regime (\( \gamma \gg 1 \)), electrons require many optical cycles to be ionized. It is easier to describe such a process in the frequency-domain as an absorption of \( n = \phi / h \omega \) photons of energy \( h \omega \). But in the tunneling regime (\( \gamma \ll 1 \)) electrons escape from the nucleus during a small fraction of an optical cycle, and thus the description in the time-domain is more convenient. Nevertheless, time- and frequency-domain descriptions represent the same process. In particular, multiphoton dynamics can be also described in time-domain, as an interference among the electronic wavepackets[1,3]
created at different optical cycles. This fact was utilized in the 
Yudin–Ivanov (Y–I) model,[6] where both multiphoton and tun-
neling regimes were described in a single formula. In this unified 
description, the time-domain tunneling picture is used. Yet, the 
electron ionization dynamics differ in these two regimes, demon-
strating sub-cycle features in the tunneling regime and much 
slower dynamics in the multiphoton one. In the intermediate 
regime, \( \gamma \approx 1 \), both fast and slow components appear.

This tunneling picture arose from atomic physics, and has 
also been proven valid for photoemission at surfaces of metallic 
nanopits.[7–12] Although the electronic wavefunctions inside the 
metal are not localized, an approximation of localized wavefunc-
tions still yields good results when considering ionization from 
metallic surfaces.[8,9,11,12] Strong near-field enhancement near 
the nanopits triggers electrons near the Fermi level to tunnel through 
the surface on a sub-cycle timescale,[7,8,12–23] which is of particu-
lar interest in ultrafast time-resolved electron nano-scopy.[12,21–25]

Interference of the wavepackets tunneled at different cycles gives 
rise to pronounced peaks in the electron spectra separated by the 
energy of the pump photons.[8,11,12,21]

Recently, ultrafast electron emission from gold dimer nanoan-
tennas (nanojunctions) with gap sizes down to the few nanome-
ter scale has attracted great interest for on-chip petahertz 
electronics.[25–27] When embedding such a nanojunction into a 
closed circuit, unidirectional electronic optically controllable cur-
rents bursts, which vary on the sub-cycle scale, may arise.[26,27] 

if few-cycle driving pulses with controlled carrier-envelope phase 
(CEP) are used. Especially interesting in this respect is the recent 
proposal to create ultrafast unidirectional currents in asymmet-
ric nanostructures,[28] which enables a DC bias without necessity 
of CEP control of the driving pulse.

Since gold nanoantennas cannot withstand strong electric 
fields due to the limitations of photothermal damage as well as 
with the near-field enhancement due to the metal.[29–32] the quasistatic 
tunneling regime \( (\gamma \ll 1) \) is rather impractical. In contrast, the interme-
diate regime \( (\gamma \approx 1) \) is much more attractive. However, in this 
regime, the nature of the photoemission and the corresponding 
electron dynamics are still poorly understood, despite being 
of crucial importance for investigating the limitations on the speed 
and bandwidth of on-chip ultrafast electronic devices. The pio-
niering work in ref. [28] left several important questions open.
For instance, it suggested that dynamical lowering of the barrier 
(Schottky effect) can significantly reduce the scaling of multi-
photon photoemission with intensity. However, this assumption 
has no analogies in other systems in strong optical fields like 
atoms or molecules. Besides, typical velocities of electrons in the 
nanogap remain unclarified, although they are crucial to under-
stand the bandwidth limits of the plasmonic electronic devices.

Here we employ asymmetric plasmonic nanojunctions to pro-
duce ultrafast unidirectional currents in a way similar to used in 
ref. [28]. In contrast to ref. [28], we operate the device at room 
temperature and in ambient conditions. Furthermore, we develop a 
modification of Y–I nonadiabatic tunneling model adapted for 
gold nanostructures. It allows us to clarify the origin of the 
photoemission currents. We investigate ultrafast dynamics of elec-
tronic wavepackets in the nanojunctions, and reconsider funda-
mental limitations in speed and bandwidth of such optical field-
driven electronic devices.

2. Experimental Section

Femtosecond pulses from a Ti:sapphire oscillator with a repeti-
tion rate of \( r = 100 \) MHz were tightly focused onto an array of 
asymmetric plasmonic nanojunctions. The plasmonic nano-
junctions were fabricated by focused ion beam milling of a 100 
nm thick Au film on a sapphire substrate. Figure 1a depicts an 
overview scanning electron microscopy (SEM) image of the 
nanostructure. The laser spectrum spanned from 650 to 1000 nm 
with a central wavelength of 840 nm. Two broadband double-
chirped mirrors and a pair of fused silica wedges were employed 
to control the dispersion of the pulses. The pulses were charac-
terized by dispersion-scan method[13] and the shortest duration 
was retrieved to be \( \approx 7.6 \) fs. The laser beam diameter on the 
sample was estimated to be \( \approx 7 \) \( \mu \)m, which corresponded to a si-
multaneous illumination of about \( N \approx 100 \) unit cells of the nano-
junctions. Therefore, for the highest incident pulse energy of 1 
nJ, the peak electric field of the pump laser was estimated to be 
\( E_0 \approx 1.6 \) \( \text{V nm}^{-1} \). Figure 1b displays a representative unit cell 
of the nanojunctions, which consists of a tip-to-tip triangular Au 
needles with a gap of 50 nm. However, the upper tip was made 
much sharper than the bottom one, leading to an asymmetric dis-
tribution of the near-field enhancement, as shown in Figure 1d,e. 
The sharper tip exhibited much higher near-field strength. As a 
result, an effective negative bias arose from the upper tip to the 
bottom one, breaking the symmetry of electronic transport. The 
upper Au needle acted as an emitter electrode and the bottom 
one functioned as a collector. Therefore, a nonzero net photoe-
mission current was expected to be observed when integrating 
over the entire pulse width.

The time-integrated photoemission current \( J \) was measured by a 
low-noise amplifier. Meanwhile, the plasmonic-enhanced third 
harmonic generation[34,35] was employed to optimize the laser 
dispersion and the spatial overlap between laser focus plane and 
the plasmonic nanojunctions. Figure 1c shows the measured 
photoemission current (blue curve) and third harmonic signal 
(red curve) at various thickness of the inserted silica wedges. The 
sensitive dependence of the current on the pulse duration con-
firmed that the electron emission was induced by nonthermal 
processes, because the thermal effects do not depend on the pulse 
duration for femtosecond laser. At the shortest pulse duration, 
the integrated current read \( J = 0.6 \) nA, which translated to 40 
electrons per pulse in average. As shown in Figure 1f, the nan-
odevice withstands a long-term exposure.

In Figure 2a a log–log dependence of the photoemission cur-
rent on the incident laser electric field strength \( J = E \) curve is 
shown, with an observed slope of \( \approx 2 \). The integrated current 
versus the polarization of the incident laser is also studied(\( J \)–\( \theta \) 
curve, cf. Figure 2b). Here \( \theta \) was defined as the crossing an-
gle between electric field direction and tip-to-tip orientation of 
the nanojunction. The current followed a fourth-order power of 
\( \cos \theta \), corresponding to a \( J–E \) curve with a coefficient \( n = 2 \), that 
is, \( J \propto (E')^2 = I' \), consistent with directly measured \( J–E \) curve in 
Figure 2a. Considering that the electric near-field enhance-
ment factor at a nanopit is inverse proportional to its radius of 
(curvature[37] as shown by the SEM in Figure 1b, the ratio of pho-
onemission rate from the emitter with respect to the collector was 
estimated to be \( (R/r)^4 \approx 40 \). Therefore, the photoemission from
the collector is ignored in the below theoretical sections. It should be pointed out that these results also agree with those of ref. [28]. However, the photoemission process was investigated using the time-domain rather than frequency-domain approach, and a conclusion opposite to that of ref. [28] was drawn, as discussed later.

3. Theoretical Description

3.1. Yudin–Ivanov Approach

For the near-field enhancement in our structure the Keldysh parameter is estimated to be in the intermediate range (c.f., gray area in the inset of Figure 2a). In the framework of atomic physics, a model named Y–I formula[6] works well in the range from multiphoton to tunneling regime (red curve in the inset of Figure 2a) and keeps correct inter-and intra-cycle ionization dynamics. Assuming the driving field in the form of \( E(t) = E(t) \cos \omega t \), where \( \cos \omega t \) denotes the fast oscillating component, and \( E(t) \) the slow-varying envelope, the cycle-resolved ionization rate \( \Gamma \) is given by (in atomic units, that is, frequency \( \omega \), time \( t \), ionization potential \( \phi \), and field \( E \) are measured in the corresponding Hartree units \( \omega_a = 0.26 \text{ rad as}^{-1}, t_a = 24.2 \text{ as}, \phi_a = 27.21 \text{ eV}, \) and \( E_a = 514.2 \text{ V nm}^{-1} \):}

\[
\Gamma(t) = \frac{\pi}{\tau_r} \exp \left( -\sigma_0 \frac{E(t)^2}{\omega^2} \right) \left[ \frac{2\kappa^2}{E(t)} \right]^{22/\kappa} \exp \left[ -\frac{E(t)^2}{2\omega^2} \sigma_1 \sin^2(\omega t) \right] \tag{1}
\]

Here \( Z \) is the effective atomic charge, \( \kappa = \sqrt{2\phi}, \sigma_0 = \frac{1}{2}(r^2 + \frac{1}{2}) \ln C - \frac{1}{2}r \sqrt{1 + r^2}, \ C = 1 + 2r \sqrt{1 + r^2} + 2r^2, \) and \( \sigma_1 = \ln C - 2r \sqrt{1 + r^2} \). The averaged photoemission rate over a single optical cycle of Equation (1) reads (in atomic units):

\[
\Gamma_{av} = \frac{\pi}{\tau_r} \exp \left( -\sigma_0 \frac{E(t)^2}{\omega^2} \right) \left[ \frac{2\kappa^2}{E(t)} \right]^{22/\kappa} \left[ \frac{2\omega^3}{\pi E(t)} \sigma_1 \right]^{1/2} \tag{2}
\]

To proceed further, it is important to understand the basic idea behind the approach leading to Equation (1) and Equation (2). The population in continuum \( W \) is represented as an integral over all partial amplitudes \( a_p \) of the ionized electron with the
momentum $p$: $W = \int_p |a_p|^2d^3p$. The corresponding amplitude $a_p$, under reasonable approximations can be obtained as

$$a_p \approx \int_{-\infty}^{t'} e^{-iS(t,t')/\hbar}dt'$$

(3)

where $S(t,t')$ is the action:

$$S(t,t') = (\phi + \frac{p^2}{2m})(t-t') + \frac{1}{2m} \int_{\tau}^{t'} (p + eA(t''))^2 d\tau''$$

(4)

where $A(t)$ is the vector potential corresponding to the driving electric field $E(t)$. Following Equation (3), every amplitude $a_p$ is the result of summation over all partial amplitudes, each having the phase $S(t,t')$ defined by Equation (4). The first term in Equation (4) corresponds to phase shift gained by overcoming the barrier, and the second term, so-called Volkov phase, corresponds to the phase electron gains in the electric field. Action $S(t,t')$ changes quickly for all points except the stationary ones, that is, obeying $\partial S/\partial t' = 0$. As a result, integration over the fast oscillating argument yields zero everywhere except at the stationary points (this fact constitutes the essence of so-called stationary phase, or saddle-point, method). The stationary condition results in a complex value of $t'$ equal to $t'$, which can be found analytically. The ionization rate can be, up to the insignificant prefactor, calculated as $\Gamma(t) \approx \exp(-2im[S(t,t')]/\hbar)$. Under reasonable approximations, this expression can be calculated analytically and gives rise to Equation (1).

The Y–I model, as mentioned above, is written for a rather general case, without the details of the potential. The only parameters referring to a particular system are the effective charge $Z$ and the ionization potential $\phi$. For gold nanostructures, we should take into account the ability of electron density on the surface to redistribute, on a femtosecond time scale, to “screen” the ionized electron. This screening is in a good approximation described by the mirror-charge model,\[18,39\] In this model, if we consider an ideal metal, the outgoing electron “feels” a charge at the position mirrored relative to the metal surface, but with the charge sign inverted. The distance from electron to this effective “ion,” represented by mirroring charge, is twice the distance to the surface. Thus the attraction force becomes four times smaller than for the case of an electron and a single atom. This attraction force reduction can be taken into account by introducing the effective charge $1/4$ instead of $1$ in the Y–I formula. Furthermore, in a non-ideal metal, the effective charge value is modified by a factor $[1 - \epsilon]/(1 + \epsilon)$, where $\epsilon$ is the complex susceptibility of the metal. As a result, the effective charge in Equation (1) is governed by the equation:

$$Z = \frac{1}{4} \left[ 1 - \frac{1 - \epsilon}{1 + \epsilon} \right]$$

(5)

We fit the experimental data in Figure 2a by the modified Y–I model and obtain the field enhancement factor to be $g \approx 14.2$. This is a very reasonable value which is in a good agreement with the numerical simulation (c.f. Figure 1d,e). Accordingly, the Keldysh parameter $\gamma$ in our experiments is evaluated to be in the range from 0.8 to 1.7. The fitting was made without considering the spatial profile of the pulse. It is however easy to see that in the present case, taking into account the Gaussian profile leads to only a constant pre-factor $1/4$. Indeed, since we have, with a good precision $J \approx E_0^2$, spatial integrating of the current $I(r)$ (were $r$ is the radial coordinate) taking into the Gaussian profile of the field $E_0 \approx \exp(-r^2/2\sigma^2)$ ($\sigma$ is the pulse width) will give the constant factor $1/4$ in comparison with the same integration, performed over the constant-field distribution with the same area.

As shown in the inset of Figure 2a, the Y–I formula approaches the multiphoton limit at low intensities, and approximates the tunneling limit as given by the F–N equation at high intensities. We remark that there is another analytical expression, so-called Keldysh formula\[1,40\] which works in the tunneling, multiphoton and intermediate regimes as well. It is obtained using the same stationary phase method as the Y–I formula, but the corresponding integrals are taken in the frequency space. However, the advantages of the Y–I formula are the applicability to very short (yet single-color) pulses and a possibility to describe the dynamics inside the laser cycle (c.f. the inset in Figure 2b), whereas
the Keldysh equation was written for the quasi-monochromatic wave and gives only cycle-averaged ionization rate. On the other hand, we note the recent works[41,42] where the Keldysh formula was extended to ultrashort pulses of some particular shapes. For the particular parameters presented here there is no possibility to reliably differentiate between Y–I and Keldysh formulas. This differentiation can be done, for instance, for even shorter pulses in the case if the CEP control is implemented. In this situation the change of the CEP leads to change of the result in the case of Y–I formula but in the framework of the Keldysh formula the result should be CEP-independent.

As depicted in Figure 2, the current J is nearly proportional to \( P^2 \). According to the Y–I model, we identify this \( P^2 \)-law as an indication that we are in the intermediate regime, and thus invoking the Schottky effect is not necessary to explain this behavior as it is done in refs. [28, 40]. In the following paragraphs, we return to this point in details.

### 3.2. Schrödinger Equation

To get deeper insight into the dynamics of the photoemission and subsequent electron propagation between the tips, we simulate the emission by numerical solution of the following 1D time-dependent Schrödinger equation (TDSE) in the Coulomb gauge (\( \partial_x A = 0 \)):

\[
\frac{i\hbar}{2m} \frac{\partial \psi(x, t)}{\partial t} = \left( p + eA(x, t) \right)^2 + V(x) \psi(x, t)
\]

where \( \psi(x, t) \) is the electronic wavefunction, \( p = -i\hbar \frac{\partial}{\partial x} \), \( A(x, t) \) is the vector potential which takes into account spatial field inhomogeneity as a pre-factor \( e^{-\delta x} \), \( \delta = 1.0 \text{ ns}^{-1} \), \( V(x) \) is a rectangular asymmetric potential

\[
V(x) = \begin{cases} 
-V_0, & \text{if } |x| < a; \\
0, & \text{if } x > a; \\
\infty, & \text{if } x < -a,
\end{cases}
\]

where \( a = 0.106 \text{ nm}, \ V_0 = 16.94 \text{ eV} \) are selected in such a way that i) the potential has exactly one bound state and ii) the ionization potential of this bound state equals to the work function of gold (5.1 eV). We note that this potential assumes that the wavefunction inside the metal is localized. It was shown that this assumption does not significantly influence the ionization rate.[9] Moreover, the experimentally observed electron spectra[8,10] and even electron dynamics[43] are good described by this rather simple approach. Therefore, it is widely used to model the ionization.[8,10,12-21,44-46] In the potential defined by Equation (7), ionization can occur only in the positive direction of \( x \). The electrons leaving the emitter are accelerated by the field and propagate toward the collector. These electrons are considered to be fully absorbed, that is, reflection on the potential of the collector is neglected. We model this by adding to the potential the soft absorbing boundary \( V(x) \rightarrow V(x) - i \alpha (x) \) with \( \alpha = a_0 (1 + erf((x - s) / \delta)) / 2 \), where \( \delta = 2.65 \text{ pm}, \ a_0 = 1 \text{ au}, \) and \( s = 50 \text{ nm} \) is the distance between the emitter and collector. The simulation was made by a split-step method, with separate evaluation of the terms \( \approx p^2, pA + Ap, A^2, \) and \( V \); the action of \( p \) was calculated using the fast Fourier transform.

The resulting dynamics of the electronic wavefunction \( \psi(x, t) \) is shown in Figure 3a for the peak driving field amplitude \( gE_0 = 12.3 \text{ V nm}^{-1} \) (corresponding to \( \gamma = 1.68 \), clearly in the transient regime) and pulse duration of \( 7 \text{ fs} \) according to the experiment. One can see from Figure 3a, that at every positive-field subcycle of the driving electric field, an ionization event takes place: noticeable part of the electron is released close to extremum of the electric field. As one can see from Figure 3c, the free electron density increases starting from the maximum of the electric field, achieves its maximum one quarter of the cycle later, and then decreases because some part of electrons returns back. After being ionized, the part of electron, which does not return, propagates toward the collector. Interestingly, upon the propagation the electron wavepacket is separated into distinct well-visible “beams” marked by \( n = 1, 2, ..., \) every of them propagating with velocity clearly different from the others. The field inhomogeneity plays only relatively minor role in this dynamics. If it is removed (\( \delta = 0 \), see dashed line in Figure 3b) the beam structure remains the same, only most of the electrons move faster to the collector.

Ionized electrons reach, after some propagation, the collector, producing the current \( J(t) \) given as

\[
J(t) = \frac{\hbar \left| \psi \frac{\partial \psi}{\partial x} - \psi^* \frac{\partial \psi^*}{\partial x} \right|}{2im}
\]

Here, \( \psi \) is taken at the surface of the collector. The above mentioned “beams,” which are visible in Figure 3a, manifest themselves as short spikes of \( J(t) \) as can be seen in Figure 3b.

The nature of this dynamics becomes apparent if we consider the picture in the energy space, see Figure 3c, where the energy spectrum of the electrons in dependence on time is shown, as well as in Figure 3d, where energy spectra at specific times \( t_i, \ i = 0, ..., 3 \) are presented. One can see (cf. for instance \( t_0, t_1 \)) that in the beginning of the pulse there is no visible structure in the energy distribution of electrons. Electrons are born with a broad energy spectrum of more than 10 eV width. The energy structure sets up gradually during the next few cycles (see the time events \( t_2, t_3 \)), and, finally close to the end of the pulse (the time event \( t_3 \)) itsettles to be peaked around the multiples of the photon energy \( nh\omega \). This very clearly shows that multiphoton effects (absorption of \( n \) photons) appears only on the inter-cycle scale, as an interference between newly-born parts of the electron wavepacket and the parts which are already present in the continuum.

In even more clear form this inter-cycle interference is schematically shown in Figure 4, which illustrates the physics behind the results in Figure 3c. As it is presented in Figure 4, the newborn electrons have a broad spectrum (black-white gradient) and different phases (color gradient—in the figure we deliberately set the time origin so that the first phase is constant in energy). The interference between parts of the electronic wavepacket created at different cycles leads to clearly visible channels \( n = 0, n = 1, ... \) after the end of the pulse. Note that in Figure 3c these dynamics start to set up already in the middle of the pulse since, unless the wavepacket is not the first one (\( t_0 \) in Figure 3c), the newborn parts of the wavepacket start to interfere with the previously born ones (\( t_1, t_2 \) in Figure 3c).
The condition of constructive interference can be easily obtained as follows: the phase shift $\Phi$ of the electron in the continuum (in the presence of the field) over the optical period $T$ is $\Phi = T(p^2/2m + U_p)/\hbar$, where $U_p = e^2\varepsilon^2/4ma^2$ is the ponderomotive energy. At the same time, the electrons in the metal (near the Fermi energy) will experience the relative phase shift $\Phi_F = -T\Phi/\hbar$. Thus, the relative phase of two partial wavepackets born in the continuum at the two time instants separated by $T$ is $\Phi + \Phi_F$. The corresponding constructive interference condition $\Phi + \Phi_F = 2\pi n$ gives: $p^2/2m + U_p + I_p = n\hbar\omega$, that is, the subsequent peaks in electron energy $p^2/2m$ are separated by the photon energy $\hbar\omega$. This demonstrates that the multiphoton peaks corresponding to the absorption of $n$ photons appear from the interference between the electronic wavepackets created by different optical cycles. Inside the cycle, no such multiphoton effects can be identified. One can explain this also in the terms of the Heisenberg uncertainty relation $\Delta p \Delta x \geq \hbar/2$: in every ionization event, the electrons are born in the very small region surrounding the surface, of the order of $\Delta x \approx 0.1 \text{ nm}$, which means uncertainty in momentum $\Delta p$, corresponding to the kinetic energy in the range from zero to several eV. This uncertainty in momentum is partially “regularized” on the longer, intercycle scale as described above, giving rise to the multiphoton energy structure.

From Figure 4 one can clearly see how to distinguish the impacts from different ionization channels (accomplished by absorption of different number of photons), and to estimate their relative importance: if a certain channel is present in the ionization process, there must be a corresponding peak in the electron energy distribution at the end of this pulse. The “intensity” of every particular peak allows to estimate the relative impact of different channels. To distinguish different channels is important in view of refs. [28, 40] where so called Schottky effect is proposed to describe the current-versus-intensity behavior. The Schottky-effect-based explanation suggests that the potential barrier is lowered by the external field (see Figure 4, red dashed line) so that two instead of four photons are sufficient for ionization.

Figure 3. Dynamics of ionization at the emitter, and current at the collector, according to numerical simulations of the single-electron problem Equation (6) for $gE_0 = 12.3 \text{ V nm}^{-1}$ and 7 fs pulse duration. a) The modulus of the wavefunction of the ionized electron $|\psi(x, t)|$ in dependence on $x$ and $t$. White line shows the driving electric field; as the electronic wavepacket arises in continuum and propagates, it is separated into the well-visible electronic “beams” labeled by $n = 0, 1, \ldots$. Each beam [see also (c)] corresponds to an electron absorbed approximately $n$ photons from the driving field and thus having energy $\approx n\hbar\omega$. b) The electron current $J(t)$ created on the collector surface in dependence on time $t$. Dashed red line shows the current created by a homogeneous field ($\tilde{\alpha} = 0$). The upper inset shows $J(t)$ on a larger times scale, and the lower inset depicts the spectrum of $J(t)$. The vertical lines show the energies corresponding to $n\hbar\omega$. c) Temporal evolution of the free electron energy. The white curve shows again the driving electric field, whereas the horizontal white lines show the energies corresponding to $n\hbar\omega$. d) Energy spectra at the (somewhat arbitrarily selected) times $t_i$ [marked in (c)]. In (c) and (d) one clearly sees multiphoton peaks $n = 1, 2, \ldots$ as they arise on the inter-cycle time scale due to interference, and become more and more pronounced toward the end of the pulse.
This channel corresponds to \( n = 2 \) in Figure 4 and in Figure 3, in contrast to “direct” tunneling which corresponds to \( n = 0 \) (green line in Figure 4).

As one can see from Figures 3c and 4, the peak at \( 2\hbar \omega \) (\( n = 2 \)) could which could correspond to Schottky effect indeed appears, but it is also clearly not a dominating one. The peak corresponding to “direct” tunneling \( n = 0 \) has even higher amplitude. The “anomalous” dependence of the current \( J \) on intensity \( I (f \approx f^2) \) is thus not explained by the Schottky effect but by the fact that we are in the transition region from tunnel to multiphoton ionization, and the impacts from different channels \( n \) have close amplitudes and add up to give the observed scaling.

Interestingly, the emergence of the multiphoton energy structure of the electronic wavepacket allows to understand, what happens with an electron as it propagates between the tips (Figure 3a) and thus to interpret the resulting current \( J(t) \) in Figure 3b. Indeed, different electronic “beams” in Figure 3a and different peaks in \( J(t) \) in Figure 3b correspond to different peaks in the electron energy in Figure 3c. The slowest beam \( n = 0 \) corresponds to the lowest-energy electrons, which absorbed just enough energy to get through the barrier; \( n = 1 \) corresponds to the electrons which absorbed one photon more, and so on. Every beam corresponds to electrons which have a mean velocity around \( v = \sqrt{2\hbar \omega /m} \). Because of diffractive spreading of the wavepacket, its width grows as \( \sqrt{t} \). The slowest wavepackets spend longer time to overcome the nanogap, and thus spread stronger. The resulting structure of the current \( J(t) \) in Figure 3b is thus the collection of peaks, every of them having increasing width due to increasing diffractive spreading. Although the peak with \( n = 0 \) carries the largest part of the whole electron probability, it is also broadened at strongest, so that as it arrives to the collector, it has a relatively small amplitude (see inset to Figure 3b where this peak is visualized). The same is true for other, not too high values of \( n \): the diffractive spreading significantly decreases the corresponding amplitude in \( J(t) \). On the other hand, the amount of the electrons contained in the subsequent \( n \) decreases as \( n \) increases. The global maximum of \( J(t) \) is thus the result of this interplay between the diffractive spreading and energy balance of individual beams. In our configuration, the most intense peaks appear at around \( n = 7 \sim 11 \).

As follows from Figure 3b, although the shape of the current bunch occurring at the collector is rather complicated, one can define some quantities characterizing it, in particular i) the time delay in respect to the center of the pump pulse, and ii) the duration of the bunch. Here we define the above mentioned delay via the position (in time) of the highest peak of \( J(t) \), and the width is defined by full-width half-maximum. Such defined delay and duration are shown in Figure 4 for different sizes \( s \) of the nanogap. For the particular case of Figure 3b with \( s = 50 \) nm, the current \( J(t) \) peaks at around 40 fs (that is, delayed by around 30 fs from the pulse center) and the duration of the current bunch is of the order of 15 fs.

One can see from Figure 5 that the delay of the current spike does not, generally speaking, grow linearly with the gap size. This is explained by the reshaping of the electronic wavepacket described above. This reshaping is, as already mentioned, a result of interplay between the diffractive spreading and electron energy balance. Nevertheless, if we compare the delay of \( J(t) \) with the naive estimation for the flight time \( t_f = s/v_p \), where \( s \) is the size of the nanogap, and \( v_p \) is the mean velocity corresponding to the ponderomotive energy \( U_p = e^2\epsilon_0^2/4m\omega^2 \) (red dotted line in Figure 4), we observe quite a good correspondence. This indicates that, although the electron wave package has a complicated shape, its propagation can be described with a velocity which corresponds to the mean ponderomotive energy of the electron in the field. The speed of the devices increases thus with decrease of the nanogap. Recently, the gaps in the order of nm were achieved.[47]
4. Conclusion

In conclusion, we utilized an array of spatially asymmetric nanojunctions to break the symmetry of the ionization process and to generate an ultrafast optically switchable on-chip electronic current at room temperature and under the standard conditions. Generation of currents up to 0.6 nA by a few-cycle driver pulse with a random carrier-envelope phase, without using a DC bias, is possible in this way. Introducing CEP control to this scheme could additionally increase the efficiency, but for relatively long pulse durations we used here such increase should be only minor. We have extended the Y–I model, which works well for the tunnelling, multiphoton and intermediate regimes of atoms, to gold nanostructures. The excellent agreement of the Y–I model allowed to establish the leading current formation mechanism: the observed data can be well explained by assuming nonadiabatic tunnelling through the barrier. This conclusion is supported by direct simulations of electron ionization dynamics using the time-dependent Schrödinger equation, which demonstrated, that any effects manifesting as an absorption of several photons occur by the inter-cycle interference of electronic wavepackets, and are undefined on the sub-cycle time scale. No signatures of prevailing influence of the Schottky barrier lowering was found. In contrast, the unusual current scaling \( J \propto I^2 \) is explained via the joint influence of all multiphoton channels taking place in the non-adiabatic tunnelling regime, that is, in the transition region between tunnelling and multiphoton ionization. Our detailed view of the electron dynamics allowed to determine the limits on the speed of such devices. We observe that the shape of the electron wavepacket is rather complicated: the flying electrons are separated into “beams,” each of them having the velocity corresponding to certain number of absorbed photons. Nevertheless, in average, the flight time of electrons in the nanogap is determined, to a good precision, by the ponderomotive velocity of electrons in the driving field. This suggests that the primary way to increase the speed could be not only to decrease the gap but also to increase the ponderomotive energy, which does not automatically mean increasing the peak field: multicolor driving fields could also help at this.\[^{[48]}\]

Acknowledgements

L.S. and I.B. contributed equally to this work. The authors acknowledge support from Deutsche Forschungsgemeinschaft (DFG) (KO 3798/4-1, BA 4156/4-2, MO 850-19/2, MO 850-23/1) and from German Research Foundation under Germany’s Excellence Strategy EXC-2123 and Germany’s Excellence Strategy within the Cluster of Excellence PhoenixD (EXC 2122, Project ID 390833453), Lower Saxony through ‘Quanten und Nanometrologie’ (QUANOMET, Project Nanophotonik). H.G. and B.F. are supported by National Natural Science Foundation of China (No. 12004314), the open project program of Nanometrologie’ (QUANOMET, Project Nanophotonik). H. G. and B. F. gratefully acknowledge funding from Deutsche Forschungsgemeinschaft (DFG) (KO 3798/4-1, MO 850-19/2, MO 850-23/1) and from German Research Foundation under Grant No. Z1A040010. M.I acknowledges support by the DFG priority program QUTIF under grant agreement IV 152/6-2 and the funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement No 899794.

Open access funding enabled and organized by Projekt DEAL.

Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

Research data are not shared.

Keywords

ionization, nanostructures, optoelectronics

Received: October 28, 2020
Revised: April 9, 2021
Published online:

[1] L. Keldysh, Sov. Phys. J. Exp. Theor. Phys. 1965, 20, 1307.
[2] E. Mevel, P. Breger, R. Trainham, G. Petit, P. Agostini, A. Migus, J.-P. Chambaret, A. Antonetti, Phys. Rev. Lett. 1993, 70, 406.
[3] F. Lindner, M. G. Schäftel, H. Walter, A. Baltuska, E. Goulielmakis, F. Krausz, D. Milošević, D. Bauer, W. Becker, G. G. Paulus, Phys. Rev. Lett. 2005, 95, 040401.
[4] I. Babushkin, C. Brée, C. M. Dietrich, A. Demircan, U. Morgner, A. Hussakou, J. Mod. Opt. 2017, 64, 1078.
[5] H. Zimmermann, S. Patchkovskii, M. Ivanov, U. Eichmann, Phys. Rev. Lett. 2017, 118, 013003.
[6] G. L. Yulin, M. Y. Ivanov, Phys. Rev. A 2001, 64, 013409.
[7] R. Bormann, M. Geidel, A. Weismann, S. Yalunin, C. Ropers, Phys. Rev. Lett. 2010, 105, 146701.
[8] M. Krüger, M. Schenk, P. Hommelhoff, Nature 2011, 475, 78.
[9] S. Y. Yalunin, M. Geidel, C. Ropers, Phys. Rev. B 2011, 84, 195426.
[10] M. Krüger, M. Schenk, M. Förster, P. Hommelhoff, J. Phys. B: At., Mol. Opt. Phys. 2012, 45, 074006.
[11] M. Krüger, M. Schenk, P. Hommelhoff, G. Wachtler, L. Lemel, J. Burgdörfer, New J. Phys. 2012, 14, 0805109.
[12] M. Krüger, C. Lemell, G. Wachtler, J. Burgdörfer, P. Hommelhoff, J. Phys. B: At., Mol. Opt. Phys. 2018, 51, 172001.
[13] P. Dombi, A. Horl, P. Rätz, I. Márón, A. Tügler, J. R. Krenn, U. Hohenester, Nano Lett. 2013, 13, 674.
[14] G. Herink, D. Solli, M. Geidel, C. Ropers, Nature 2012, 483, 190.
[15] B. Piglosiewicz, S. Schmidt, D. J. Park, J. Vogelsang, P. Groß, C. Manzoni, P. Farinello, G. Cerullo, C. Lienau, Nat. Photonics 2014, 8, 37.
[16] P. Racz, Z. Pápá, I. Márton, J. Budai, P. Wrobel, T. Stefaniuk, C. Prietl, J. R. Krenn, P. Dombi, Nano Lett. 2017, 17, 1181.
[17] D. J. Park, B. Piglosiewicz, S. Schmidt, H. Kollmann, M. Mascheck, C. Lienau, Phys. Rev. Lett. 2012, 109, 244803.
[18] J. Vogelsang, J. Robin, B. J. Nagy, P. Dombi, D. Rosenkranz, M. Schiek, P. Groß, C. Lienau, Nano Lett. 2015, 15, 4685.
[19] F. Schertz, M. Schmelzeisen, M. Kreiter, H.-J. Elmers, G. Schönhense, Phys. Rev. Lett. 2012, 108, 237602.
[20] J. Robin, J. Vogelsang, B. J. Nagy, P. Dombi, P. Groß, C. Lienau, in Conf. on Lasers and Electro-Optics, OSA Publishing, Washington, DC 2016, p. FTh4B.
[21] P. Dombi, Z. Pápá, J. Vogelsang, S. V. Yalunin, M. Sivis, G. Herink, S. Schäfer, P. Groß, C. Ropers, C. Lienau, Rev. Mod. Phys. 2020, 92, 025003.
[22] K. E. Priebe, C. Rathje, S. V. Yalunin, T. Hohage, A. Feist, S. Schäfer, C. Ropers, Nat. Photonics. 2017, 11, 793.
[23] A. Feist, N. Bach, N. R. da Silva, T. Danz, M. Möller, K. E. Priebe, T. Domröse, J. G. Gatzmann, S. Rost, J. Schauss, S. Strauch, R. Bormann, M. Sivis, Schäfer, C. Ropers, Ultramicroscopy 2017, 176, 63.
[24] S. Zhou, K. Chen, M. T. Cole, Z. Li, J. Chen, C. Li, Q. Dai, Adv. Mater. 2019, 1805845.

[25] J. Schoetz, Z. Wang, E. Pisanty, M. Lewenstein, M. F. Kling, M. Ciappina, ACS Photonics 2019, 6, 3057.

[26] T. Rybka, M. Ludwig, M. F. Schmalz, V. Knittel, D. Brida, A. Leitenstorfer, Nat. Photonics 2016, 10, 667.

[27] M. Ludwig, G. Aguirregabiria, F. Ritzkowsky, T. Rybka, D. C. Marinica, J. Aizpurua, A. G. Borisov, A. Leitenstorfer, D. Brida, Nat. Phys. 2019, 16, 341.

[28] C. Karnetzky, P. Zimmermann, C. Trummer, C. D. Sierra, M. Wörle, R. Kienberger, A. Holleitner, Nat. Commun. 2018, 9, 2471.

[29] N. Pfüllmann, C. Waltermann, M. Noack, S. Rausch, T. Nagy, C. Reinhardt, M. Kovačev, V. Knittel, R. Bratschitsch, D. Akemeier, A. Hütten, A. Leitenstorfer, M. Morgner, New J. Phys. 2013, 15, 093027.

[30] W. P. Putnam, R. G. Hobbs, P. D. Keathley, K. K. Berggren, F. X. Kärtner, Nat. Phys. 2017, 13, 335.

[31] L. Shi, B. Iwan, Q. Ripault, J. R. Andrade, S. Han, H. Kim, W. Boutu, D. Franz, R. Nicolas, T. Heidenblut, C. Reinhardt, B. Bastiaens, T. Nagy, I. Babushkin, U. Morgner, S.-W. Kim, G. Steinmeyer, H. Merdji, M. Kovacev, Phys. Rev. Appl. 2018, 9, 024001.

[32] L. Shi, R. Nicolas, J. R. Andrade, W. Boutu, D. Franz, T. Heidenblut, C. Reinhardt, U. Morgner, H. Merdji, M. Kovacev, ACS Photonics 2018, 5, 1208.

[33] L. Shi, J. R. Andrade, A. Tajalli, J. Geng, J.-M. Yi, T. Heidenblut, F. Seegerink, I. Babushkin, M. Kholidotsova, H. Merdji, B. Bastiaens, U. Morgner, Milutin Kovacev, Nano Lett. 2019, 19, 4779.

[34] M. Hentschel, T. Utikal, H. Giessen, M. Lippitz, Nano Lett. 2012, 12, 3778.

[35] L. Shi, B. Iwan, R. Nicolas, Q. Ripault, J. R. Andrade, S. Han, H. Kim, W. Boutu, D. Franz, T. Heidenblut, C. Reinhardt, H. M. J. Bastiaens, G. T. Nagy, I. Babushkin, U. Morgner, S.-W. Kim, G. Steinmeyer, H. Merdji, M. Kovacev, Optica 2017, 4, 1038.

[36] A. Plech, V. Kotaidis, M. Lorenc, J. Boneberg, Nat. Phys. 2006, 2, 44.

[37] L. Novotny, B. Hecht, Principles of Nano-Optics, Cambridge University Press, Cambridge 2012.

[38] J. D. Jackson, Classical Electrodynamics, Wiley, New York 1962.

[39] V. Sahni, K.-P. Bohnen, Phys. Rev. B 1985, 31, 7651.

[40] P. Zimmermann, A. Hötger, N. Fernandez, A. Nolinder, K. Müller, J. J. Finley, A. W. Holleitner, Nano Lett. 2019, 19, 1172.

[41] L. V. Keldysh, Phys.-Usp. 2017, 60, 1187.

[42] S. Kim, T. Schmude, G. Burkard, A. S. Moskalenko, arXiv:2011.15125 2020.

[43] Ph. Dienstbier, personal communication, 2021.

[44] P. Hommelhoff, C. Kealhofer, M. A. Kasevich, in IEEE International Frequency Control Symp. and Exposition, IEEE, Piscataway, NJ 2006, pp. 470–474.

[45] S. V. Yalunin, G. Herink, D. R. Solli, M. Krüger, P. Hommelhoff, M. Diehn, A. Munk, C. Ropers, Ann. Phys. 2013, 525, L12.

[46] P. D. Keathley, A. Sell, W. P. Putnam, S. Guerrera, L. Velásquez-Garcia, F. X. Kärtner, Ann. Phys. 2013, 525, 144.

[47] M. Garg, K. Kern, Science 2020, 367, 411.

[48] P. G. d. A. Martínez, I. Babushkin, L. Bergé, S. Skupin, E. Cabrera-Granado, C. Köhler, U. Morgner, A. Husakou, J. Herrmann, Phys. Rev. Lett. 2015, 114, 183901.