Strong-field ionization of clusters using two-cycle pulses at 1.8 μm

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The interaction of intense laser pulses with nanoscale particles leads to the production of high-energy electrons, ions, neutral atoms, neutrons and photons. Up to now, investigations have focused on near-infrared to X-ray laser pulses consisting of many optical cycles. Here we study strong-field ionization of rare-gas clusters (10⁴ to 10⁵ atoms) using two-cycle 1.8 μm laser pulses to access a new interaction regime in the limit where the electron dynamics are dominated by the laser field and the cluster atoms do not have time to move significantly. The emission of fast electrons with kinetic energies exceeding 3 keV is observed using laser pulses with a wavelength of 1.8 μm and an intensity of 1 × 10¹⁵ W/cm², whereas only electrons below 500 eV are observed at 800 nm using a similar intensity and pulse duration. Fast electrons are preferentially emitted along the laser polarization direction, showing that they are driven out from the cluster by the laser field. In addition to direct electron emission, an electron rescattering plateau is observed. Scaling to even longer wavelengths is expected to result in a highly directional current of energetic electrons on a few-femtosecond timescale.

In the past few years, ultrashort infrared (IR) laser pulses with wavelengths >1 μm have become an important tool for the investigation of strong-field physics in atoms, molecules and solids¹–⁴. The use of these laser pulses is often motivated by the large ponderomotive energies that electrons can acquire, which scale as the square of the wavelength. As a consequence, electrons can be efficiently accelerated up to high kinetic energies, which has been exploited for the production of high-harmonic generation (HHG) pulses in the keV range⁵ and for the development of bright hard x-ray sources⁶. Furthermore, the use of strong IR laser fields has led to an advancement of HHG schemes in solids⁷–⁹ and to the development of time-resolved photoelectron holography⁸.

Sub-wavelength nanoscale particles represent a distinct class of targets, and were found to exhibit a fundamentally different behaviour in comparison to atoms, molecules and solids, when exposed to strong laser fields. While the particles themselves have a near-solid density, the distances between the particles are large, meaning that energy cannot be dissipated into the environment, as is the case for solids. The ionization of nanoscale clusters by intense laser pulses has been extensively studied in different wavelength regimes during the past two decades. Up to now, the largest number of experiments has been performed using laser pulses <1 μm⁹–²¹. Particularly intriguing results in this wavelength range were the emission of bright X-ray radiation⁴⁰, highly charged ions¹¹, fast ions¹² and neutrals¹³ with MeV kinetic energies, as well as the observation of correlated electronic decay²⁰. The efficient absorption of near-infrared (NIR, which in this manuscript only refers to the wavelength regime around 800 nm) laser energy in clusters creates extreme conditions, which even resulted in the observation of nuclear fusion¹⁴. With the advent of free-electron laser and high-flux HHG sources, intense laser-cluster experiments have been extended to shorter wavelengths in the extreme-ultraviolet (XUV)²²–²⁹ and X-ray regimes³⁰–³².

Ionization of clusters by intense IR laser pulses >1 μm allows one to exploit both the high ponderomotive energies that electrons can acquire in the laser field and the efficient absorption of laser energy by nanoscale particles. Potential applications include sources of high-energy electrons, ions and photons. Up to now, IR strong-field ionization dynamics of clusters using laser wavelengths >1 μm have been largely unexplored, in contrast to the wavelength regimes from the NIR to the X-ray range. In ref. 33, XUV fluorescence from molecular clusters was studied following ionization by 1.45 μm pulses. The emission of fluorescence light is a signature of Rydberg atom and ion formation by electron–ion recombination processes during cluster expansion, which radiatively decay on a nanosecond timescale³⁰,³¹. In contrast, the investigation of fast electrons emitted from clusters interacting with intense NIR laser fields provided insight into the dynamics taking place on a timescale of 7 to 1000 fs.³²,³³,³⁷. By using laser pulse durations in the impulsive limit, i.e. so short that the atoms/ions do not move significantly

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During the laser pulse, it becomes possible to isolate the electron dynamics taking place early during the first few cycles of the laser pulse. This regime has led to the surprising observation that the ion emission was peaked in the direction perpendicular to the laser polarization\(^{16}\).

Here we report on the first strong-field ionization experiment in clusters using two-cycle laser pulses at 1.8 \(\mu\)m, meaning that ion motion is effectively frozen. We demonstrate highly efficient electron acceleration, resulting in the observation of keV electrons at much lower laser intensities than in previous NIR experiments\(^{12,35,36}\). A narrow angular distribution shows that electrons are driven out from the cluster by the laser field, and we find signatures of direct electron emission as well as a rescattering plateau. Our results are expected to be relevant to other solid-density nanoscale systems interacting with intense few-cycle IR pulses, in which dense plasmas are generated impulsively, including nanostructures, solids and large (bio-)molecules.

**Results**

**Electron emission angular distributions.** In Fig. 1(a) TOF traces are shown that were recorded from Xe clusters ((\(N = 1 \times 10^5\) atoms) ionized by 1.8 \(\mu\)m pulses with a duration of 12 fs and an intensity of \(1 \times 10^{15}\) W/cm\(^2\) (see Methods for details). In accordance with previous strong-field ionization experiments on clusters in the NIR regime\(^{12,35,36}\), two peaks are observed. While the first peak at 1.5 ns is only influenced by fluctuations of the experimental parameters, the height of the second peak decreases for increasing retarding voltages. We attribute the first peak to the emission of photons in the ultraviolet and XUV range, in accordance with the experiments performed in refs 35 and 36. These photons are emitted as a consequence of Rydberg atom and ion formation in the expanding cluster and their subsequent decay via fluorescence on a timescale of 1 ns\(^{35}\). On this long timescale, any information about the laser polarization direction is lost, since during the cluster expansion many collisions between electrons, atoms and ions have taken place. Therefore, the photon emission is isotropic, and the height of this peak does not change for different laser polarizations. The second peak is attributed to electron emission, in agreement with refs 12, 35 and 36. At a retarding voltage of 3 keV (green curve), a fraction of the electron signal remains visible at 12 ns, showing that electrons with energies in excess of 3 keV are emitted from Xe clusters. This is only slightly lower than the maximum electron kinetic energies of 5 keV\(^{35}\) and 6 keV\(^{36}\) measured in the NIR regime at much higher intensities of \(8 \times 10^{15}\) W/cm\(^2\) and \(6 \times 10^{15}\) W/cm\(^2\), respectively. In contrast to the studies in refs 35 and 36, however, ion motion does not play any significant role, since our pulse duration of 12 fs is too short. We attribute the efficient acceleration of electrons as observed in Fig. 1(a) to the higher ponderomotive potential of the laser field, which is the expected behaviour from studies in atoms reported in ref. 1. This higher ponderomotive potential partially compensates for the lower laser intensity used in our experiment.

Angular electron emission distributions from clusters shown in Fig. 1(b), and were obtained by rotating the linear laser polarization using an achromatic \(\lambda/2\) waveplate. These distributions are peaked along the laser polarization direction. For Xe clusters, the full width at half maximum (FWHM) of the angular distribution is 40° (black curve in Fig. 1(b)), while it is only about 30° for Kr clusters (red curve in Fig. 1(b)). This difference...
could be explained by the larger average cluster size for Xe, which increases the probability of collisions between the escaping electrons and other particles, thus altering the emission direction. Another possible explanation is a different rescattering behaviour in Xe, which is known to be different for different atomic and molecular species.

In comparison, the angular emission distribution of Kr clusters ionized by few-cycle 800 nm pulses is about 55° (blue curve in Fig. 1(b)). The narrower angular width of the electron emission distribution at 1.8 μm compared to 800 nm is attributed to the larger ponderomotive energy and the more efficient acceleration of electrons along the laser polarization direction. Figure 1(c) visualizes the large contrast in the electron emission in the directions parallel and perpendicular to the laser polarization, whereas photon emission is isotropic. In a strong-field ionization experiment using Ne, which does not form clusters at room temperature, we recorded a FWHM of only 12°. This is consistent with earlier measurements on atomic systems, in which a FWHM of about 10° was reported for Ar and Xe at a wavelength of 2 μm.

Electron kinetic energy spectra. In Fig. 2(a), electron kinetic energy spectra for clusters consisting of different atomic species and with different cluster sizes are displayed. While the electron yields obtained from Xe, Kr and Ar clusters are similar for kinetic energies up to about 500 eV (as indicated by the green line), they strongly differ at higher kinetic energies (see black, red and blue lines). The fastest electrons are observed from Xe clusters, followed by Kr and Ar clusters. For comparison, we have recorded electron kinetic energy spectra at 800 nm using a pulse duration of 13 fs and an intensity of 0.8 × 10^{15} W/cm² (U_p ≈ 50 eV) are shown. Here the lines indicate a different behaviour for electrons <100 eV and >100 eV. Note that the absolute values for the data in (a) and (b) are not comparable due to the different experimental conditions. In the inset, the electron spectra from Xe clusters at 1800 nm and 800 nm are directly compared. The data were matched at the first data points.

Figure 2. Electron kinetic energy spectra from clusters. (a) Electron kinetic energy spectra obtained from ionization of Xe, Kr and Ar clusters (with different average sizes as indicated) by 1.8 μm pulses at an intensity of 1 × 10^{15} W/cm² (U_p ≈ 300 eV) and a pulse duration of 12 fs. The spectra were obtained by measuring electron yields at different retarding voltages and differentiating the results. The lines indicate a different behaviour for electrons <500 eV and >500 eV. The gray curve is a TDSE calculation for Ar atoms using the same laser parameters as in the experiment. We chose to match the experimental and theoretical data at the first experimental point at 50 eV. (b) For comparison, electron kinetic energy spectra after ionization of Xe, Kr and Ar clusters by 800 nm pulses at an intensity of 0.8 × 10^{15} W/cm² (U_p ≈ 50 eV) are shown. Here the lines indicate a different behaviour for electrons <100 eV and >100 eV. Note that the absolute values for the data in (a) and (b) are not comparable due to the different experimental conditions. In the inset, the electron spectra from Xe clusters at 1800 nm and 800 nm are directly compared. The data were matched at the first data points.

An electron spectrum from Ar atoms was calculated by solving the time-dependent Schrödinger equation (TDSE) for the same parameters as in the experiment (see Methods for details). The result is shown in Fig. 2(a) (gray curve), where direct electron emission is visible up to 2 U_p, which takes place by quasistatic ionization. Under quasistatic conditions, the yield of direct electrons is given approximately by (atomic units)^41

\[ Y(E_{\text{kin}}) \propto e^{-aE_{\text{kin}}}, \]

where \( E_{\text{kin}} \) is the photoelectron energy. The slope \( a \) is given by \( a = (2\omega^2(2I_p)^{3/2})/(3E_p^2) \), where...
Figure 3. Comparison of different clusters and cluster sizes. Electron yields for Kr clusters with an average size of $\langle N \rangle = 23000$ atoms and $\langle N \rangle = 4500$ atoms as well as for Ar clusters with $\langle N \rangle = 4500$ atoms recorded at different retarding voltages. The yields for the different clusters and cluster sizes may be influenced by a different number of clusters present in the interaction zone for the different conditions. The inset shows a zoom of the high-energy electrons in a linear scale.

$I_p$ is the ionization potential, $F_0$ is the peak electric field of the laser, and $\omega$ is the angular frequency. For the Ar curve in Fig. 2(a), a quasistatic slope of $\approx 0.1$ Hartree$^{-1}$ is expected. The TDSE direct-electron slope is $\approx 0.5$, indicating that ionization is essentially complete by the time the electric field reaches 60% of the nominal peak value. This observation is consistent with the calculated electron dynamics. Interestingly, the experimentally observed slope of $\approx 0.4$ for direct electrons is substantially lower, which is consistent with near-field enhancement that is present in highly ionized clusters$^{42-44}$. Enhanced ionization happens earlier within the laser cycle, meaning that the vector potential at the time of ionization is larger than in the case of a bare atom.

In addition, the calculation shows a rescattering plateau at higher kinetic energies, which originates from electrons that gain energy by a laser-driven interaction with ions$^{45-47}$. Note that our calculation is expected to slightly overestimate the yield of high energy recollision electrons (see Supplementary Information for more detailed information). This plateau is also observed for clusters, where electrons can be rescattered both by individual ions and by the cluster potential$^{48}$. It is striking that the relative yield of rescattered electrons from clusters is more than two orders of magnitude higher than for atoms. This observation is consistent with the large number of ions in the cluster. We note that the electron scattering mean free path in solid Xe is similar to the cluster radius for electron kinetic energies between 1 and 9 eV$^{49}$, and it is expected to increase for larger electron energies. Therefore, electrons that rescatter within the bulk of the cluster may leave the cluster without undergoing additional scattering processes. The rescattering plateau is also visible for 800 nm pulses, but is less pronounced (Fig. 2(b)), in accordance with results obtained in atoms$^1$. Also at 800 nm, the signal from clusters drops more slowly towards higher kinetic energies than for atoms, in particular in the region between 2 $I_p$ and 4 $I_p$.$^{49}$ The inset shown in Fig. 2(b) displays a comparison between electron spectra recorded for 800 nm and 1800 nm. Similar to atoms$^1$, direct electron emission exhibits a different slope for the two different wavelengths, whereas the rescattering plateau shows a similar slope in both cases. Rescattering takes place approximately three quarters of a cycle after ionization$^{50}$. While ionization can occur early during the laser pulse, the intensity has to be high during the following optical cycle so that the electrons can gain high kinetic energies in rescattering. This limits the emission of fast electrons to a few half cycles$^{44,50}$. In the Supplementary Information, a calculated momentum map is presented, showing that rescattered electrons are emitted during at least three half cycles. We note that in addition to direct electron emission and rescattering, thermal electron emission may occur in clusters$^{51}$. It is furthermore noted that laser wakefield acceleration can also lead to the generation of fast electrons, but typically requires much higher laser intensities$^{52}$.

Comparison of different clusters and cluster sizes. In Fig. 3, the electron yield is shown as a function of the retarding voltage for two different average Kr cluster sizes, and for Ar and Kr clusters with the same average size. Note that in comparison to the differentiated data shown in Fig. 2(a), the distribution of total yields at different retarding voltages as presented in Fig. 3 is less noisy, but does not represent a kinetic energy spectrum. We find an increased yield of fast electrons for increasing cluster size. At a retarding voltage of 1.5 kV, the electron yield from larger clusters (black squares) is increased by almost a factor 4 in comparison to small clusters (red circles). This clearly indicates a cluster effect, since one would expect that the shape of the electron kinetic energy spectrum remains unchanged, if only the atomic density was varied. The quantitative differences in the cluster-size dependent spectra can be explained by different widths of the scattering potential that are expected for clusters with different sizes.$^{43}$ The different behaviour of Ar and Kr clusters with the same average size, as observed in Fig. 3, shows that also the atomic structure has an influence on the rescattering plateau. Future time-resolved experiments could give insights into the contributions of the atomic and the cluster potential on rescattering processes.
Discussion
The high relative yield of fast electrons from clusters presented in this work may be due to near-field enhancement effects that were previously studied for clusters and nanoparticles. Model calculations could give insights into the underlying physical mechanisms. However, such calculations for our experimental parameters that would have to take into account both rescattering processes at the potential of single ions and at the potential of the cluster as a whole are at the limits of what is currently feasible. We hope that our work will stimulate the theoretical advances required to allow realistic modelling of such experiments. The understanding of the involved processes could further be improved by performing time-resolved experiments with attosecond resolution.

In summary, we have observed highly efficient acceleration of electrons to multi-keV levels from clusters in strong two-cycle laser fields at 1.8 \( \mu \)m, with an anisotropic electron emission distribution peaked along the laser polarization direction. We found clear signatures of direct and rescattering processes with a much higher probability compared to atoms. This scheme may therefore lead to efficient HHG and even attosecond pulse generation from clusters in the water window. Scaling the ionization wavelength to the multi-terahertz regime is expected to result in a highly directional current of energetic electrons within a few femtoseconds. The presented results are pertinent to laser driven electron dynamics, whenever an intense few-cycle laser pulse interacts with condensed phase matter.

Methods
Experiment. For the generation of 1.8 \( \mu \)m pulses, linearly polarized laser pulses at 800 nm with an energy of 8 mJ and a duration of 30 fs were derived from a Ti:sapphire amplifier system (1 kHz repetition rate), and coupled into an optical parametric amplifier. The linearly polarized output idler pulses at 1.8 \( \mu \)m were spectrally broadened by focusing them into a differentially pumped hollow-core fiber filled with Ar. Temporal compression of these pulses was achieved by a combination of a 1 mm BK7 window and an achromatic \( \lambda/2 \) waveplate consisting of a 1.5 mm fused silica plate and a 1.2 mm MgF\(_2\) plate. In addition, the waveplate was used to rotate the linear polarization. The pulse duration was measured before and after each experimental run making use of spatially-encoded arrangement filter-based spectral phase interferometry for direct electric field reconstruction (SEA-F-SPIDER). The laser path from the output of the fiber to the experimental chamber through air and through the vacuum window slightly stretched the pulse, resulting in a typical pulse duration of 12 fs FWHM. After transmission and reflection losses, a typical pulse energy of 0.45 mJ was applied in the experiment. We used a second hollow-core fiber filled with Ar in order to spectrally broaden the 800 nm pulses. Temporal compression of these pulses was achieved by chirped mirrors and an achromatic waveplate (that was also used for polarization control), giving a pulse duration of 13 fs.

The compressed laser pulses were coupled into a vacuum chamber through a 1 mm thick CaF\(_2\) window, and focused into the interaction zone by a spherical silver mirror with a focal length of 15 cm at near-normal incidence. In the interaction zone, the laser beam and a cluster beam that was generated by a piezoelectric valve intersected at right angles. The average cluster size was controlled by varying the backing pressure between 1 and 7 bar, and estimated according to the Hagena scaling law. Under these conditions clustering is known to be highly efficient and to lead to clusters in the size range from 10\(^3\) to 10\(^5\) atoms, corresponding to average cluster radii between about 3 and 15 nm. We confirmed efficient cluster generation in an independent experiment, where ions with substantial kinetic energies were observed as a result of cluster explosions. A molecular beam skimmer with an orifice diameter of 0.5 mm was used to select the central part of the cluster beam and to maintain a low pressure in the experimental chamber.

A TOF spectrometer with an acceptance angle <3° was used for the simultaneous detection of electrons and photons. A retarding voltage could be applied to the flight tube such that only electrons with kinetic energies above the retarding potential were detected. Kinetic energy spectra were obtained by measuring the electron yields at different retarding voltages and differentiating the results, similarly to previous studies of fast electrons and photons. A retarding voltage could be applied to the flight tube such that only electrons with kinetic energies above the retarding potential were detected. Kinetic energy spectra were obtained by measuring the electron yields at different retarding voltages and differentiating the results, similarly to previous studies of fast electrons and photons.

Calculations. We have calculated a photoelectron spectrum for Ar atoms using TDSE for the same laser parameters as in the experiment (gray curve in Fig. 2(a)). One-electron TDSE calculations used an effective potential designed to reproduce the energies of the low-lying states of an argon atom. TDSE was solved in a velocity-gauge dipole approximation on a spherical grid. Angular momenta up to \( L = 500 \) were included. A uniform radial grid (grid spacing 0.1 Bohr) extended to 450 Bohr from the origin, with a 33-Bohr wide absorbing boundary starting at 417 Bohr. A truncated Gaussian laser pulse with a carrier wavelength of 1800 nm and a FWHM of 12 fs was centered at 36.3 fs. The envelope was smoothly switched to zero between 24 and 36 fs from the envelope peak. A sine carrier envelope phase was used, yielding six half-cycles (three on each side of the envelope peak), where the instantaneous field intensity exceeds 30% of the nominal peak intensity. The electric field of the laser pulse is shown in Fig. 1 of the Supplementary Information. The laser field was polarized along the Y direction. The simulation continued up to 72.6 fs, using time steps of \( \approx 4.84 \times 10^7 \) fs, corresponding to a total of 1.5 \( \times 10^7 \) time steps. The atomic 3p function (\( m = 0 \) = along the laser field polarization direction) was used as the initial state for solving the TDSE. More details about the calculations can be found in the Supplementary Information.

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Author Contributions

B.S. and P.Y. carried out the experiments, and B.S. analysed the experimental data. S.P. performed the theoretical calculations. D.R.A., C.B., P.Y. and C.S. took care of laser alignment, compression and frequency conversion. J.P.M. supervised the project. All authors discussed the results. B.S. and J.P.M. wrote the manuscript, to which the other authors contributed.

Additional Information

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