Enhanced generation and anisotropic Coulomb scattering of hot electrons in an ultra-broadband plasmonic nanopatch metasurface

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The creation of energetic electrons through plasmon excitation of nanostructures before thermalization has been proposed for a wide number of applications in optical energy conversion and ultrafast nanophotonics. However, the use of “nonthermal” electrons is primarily limited by both a low generation efficiency and their ultrafast decay. We report experimental and theoretical results on the use of broadband plasmonic nanopatch metasurfaces comprising a gold substrate coupled to silver nanocubes that produce large concentrations of hot electrons, which we measure using transient absorption spectroscopy. We find evidence for three subpopulations of nonthermal carriers, which we propose arise from anisotropic electron-electron scattering within sp-bands near the Fermi surface. The bimetallic character of the metasurface strongly impacts the physics, with dissipation occurring primarily in the gold, whereas the quantum process of hot electron generation takes place in both components. Our calculations show that the choice of geometry and materials is crucial for producing strong ultrafast nonthermal electron components.
There is wide-ranging motivation to understand hot electrons in optically excited plasmonic systems for a number of diverse applications. Hot electrons have been demonstrated to inject over large interfacial energy barriers, enabling sensitization of plasmonic Schottky photodetectors to sub-bandgap photons, nanoscopy with high spatial and chemical sensitivity, and photocatalyzed reactions, including hydrogen dissociation on plasmonic nanoparticles. The ultrafast optical response from hot electrons has further use in nonlinear optics, ultrafast optical switching, and beam steering. However, what are usually termed “hot” electrons fall into two disparate populations of excited carriers: “nonthermal” electrons with an initial stepwise distribution extending up to the photon energy from the Fermi level and possessing an indefinite temperature, and “thermal” electrons with a quasi-equilibrated Fermi–Dirac distribution and an electronic temperature elevated above that of the surrounding lattice. Due to their relatively low energies, thermal electrons are ineffective for carrier injection and slow to relax. For this reason, nonthermal carriers are of greater relevance for light-harvesting and ultrafast optics applications.

Nonthermal electrons form by plasmon dephasing from its wavelike state to a high-energy charge pair in the metal’s conduction bands. This process occurs through indirect intraband transitions involving large changes in carrier momentum (Fig. 1a), distinct from the momentum-conserving interband (IB) transitions at energies above the optical bandgap. Due to the breaking of momentum-matching conditions, nonthermal electron generation is predominantly a quantum process driven by optical electric field hot spots and surface-assisted scattering (Supplementary Note 1). Energy is then redistributed among the carriers through electron–electron (e–e) scattering (or Coulomb scattering) to form a thermal electron population (Fig. 1b), which subsequently relaxes through electron–phonon (e–p) scattering until thermal equilibrium is reached with the surrounding lattice. We note that the diagrams in Fig. 1b represent an ideal case, where an instantaneous pulse excites only nonthermal carriers. However, in reality, both thermal and nonthermal carriers are initially produced during a finite pump pulse (Supplementary Fig. 1), with the relative amount of nonthermal generation determined by the local electric field and the Drude response of the metal.

Nonthermal electrons present considerable challenges to examine and utilize because they are short lived, generally created with low probability, yield only weak perturbations to the overall permittivity of the metal, and exhibit a broad distribution of scattering rates. The broad distribution of scattering rates is a function of their energy; as nonthermal electrons scatter closer to the Fermi level their relaxation rate slows due to Pauli exclusion effects. This is generally described in the context of Fermi liquid theory (FLT), which predicts an e–e scattering rate proportional to \((E - E_F)^2\). This results in values of ~0.1 fs\(^{-1}\) at 1 eV energies in noble metals such as Ag (Supplementary Note 2) and can be 2–3 orders of magnitude slower near the Fermi energy \((E_F)\). In FLT, the nonthermal e–e scattering is assumed to occur isotropically in a parabolic band structure (Fig. 1a).

In this work, we specifically address the issue of enhancing nonthermal electron generation with the aid of a plasmonic nanocube metasurface geometry that creates strong electromagnetic hot spots. The resulting high concentrations of nonthermal electrons allow us to study their dynamics in great detail. Such a configuration was previously demonstrated in angle-insensitive perfect absorbers, ultrafast emission sources, single photon emitters, and chemical sensors. The use of colloidal silver nanocubes is a simple means of fabricating a conformal metasurface over large areas without the need for nanopatterning. The random orientation of the nanocubes provides an additional advantage in that the optical response is independent of the polarization of incident light. Here we fabricate and optically characterize the steady-state and ultrafast transient response of metasurfaces and experimentally determine the kinetic and spectral response of nonthermal carriers, even when their lifetime is well under 100 fs. Importantly, we propose that the nonthermal carrier decay is anisotropic within the band structure, evidenced by three distinct signatures of the nonthermal carrier decay independent of the polarization of incident light.

**Results**

**Optical characterization of the metasurface.** The metasurfaces employ silver nanocubes with 150 nm edge lengths that are separated from an underlying gold film by a thin
polyvinylpyrrolidone (PVP) shell, a poly(allylamine hydrochloride) (PAH) adhesion layer, and an atomic layer deposition (ALD) grown Al₂O₃ spacer layer, creating an ensemble of nanopatch antennas (Fig. 2a–c and Supplementary Fig. 2). Samples were fabricated using a range of Al₂O₃ spacer thicknesses from 1 to 25 nm. A total of four absorption features are supported in the metasurfaces. These include a substrate-coupled gap plasmon mode in the near-infrared (NIR), a substrate-coupled quadrupolar plasmon mode at ~600 nm, a multipolar plasmon mode in the ultraviolet (UV) to NIR regions of the spectrum. We find excellent agreement between experiments and simulations (Fig. 3b, c), and show an inverse dependence of the quadrupolar and gap plasmon resonance wavelengths with the spacer thickness as a result of coupling to the underlying Ag film as shown in Supplementary Fig. 4a, b.

As can be seen in the steady-state spectra, the full-width-half-maximum of the metasurface gap plasmon resonance is ~200 nm (Fig. 3b, c). This is roughly double the 90 ± 17 nm full-width-half-maximum measured through brightfield absorbance microscopy (see “Methods” section) of individual particles on the metasurface (examples shown in Fig. 3b, inset), indicating some inhomogeneous broadening is present in the ensemble. The simulated absorbance data is also shown in the inset, showing excellent agreement of the calculated linewidth with the experimental data. We attribute this broadening to a finite distribution in particle sizes (Supplementary Fig. 2) that extends the coverage of the gap resonance in the NIR.

**Hot electron production.** Many time-resolved experiments have been performed on various nanoparticle geometries17, 19, 29–31, however in each case, their transient response was dominated by the decay of thermal electrons on the picosecond timescale. Only in a recent gap mode structure has the ultrafast growth and decay of nonthermal carriers been significant32. To confirm the improved nonthermal carrier generation in the nanopatch metasurface geometry, we first experimentally compare transient absorption kinetics at the gap mode in samples with an 8 nm spacer to a thicker 25 nm spacer and a film of bare Ag nanocubes on an SiO₂ substrate, as shown in Fig. 4a, b. (A detailed discussion of our transient absorption measurements will be addressed in the following section). As can be seen, the ultrafast (<300 fs) decay attributed to nonthermal carriers is strongest for the 8 nm spacer, reduced for the 25 nm spacer, and nonexistent for nanocubes on SiO₂ without the metasurface geometry.

The observation of the unusually strong ultrafast response in the kinetics in our samples can be explained qualitatively by calculating the rates of nonthermal carrier generation. A simple quantum formalism describing the rate of optical generation of nonthermal carriers involves integration over the surface area of a nanoparticle:

\[
\text{Rate}_{\text{nonthermal}} = \sum_{i=Au,Ag} \frac{2e^2 E_{fi}^2}{\hbar^2 \omega \left(\omega_0 \right)^2} \int_{S_{nc}} \left| E_n(\theta, \phi) \right|^2 dS,
\]

where \(E_{fi}\) is the normal electric field inside a nanocrystal near the surface, \(E_n\) is the Fermi energy of the metal, and the integral in Eq. 1 is taken over the surface of a nanocrystal \(S_{nc}\)32. Such calculations (Fig. 4c) clearly indicate that the generation of hot carriers becomes strongly amplified at the plasmonic wavelength in samples with small gaps due to the formation of plasmonic hot spots. Moreover, we can estimate an averaged number of energetic electrons (i.e., electrons in the interval \(E_F < E < E_F + h\omega_0\)) during a short excitation pulse (80 fs is assumed here to match the experiment) as:

\[
N_{\text{Nonthermal,avg}} = \sum_{i=Au,Ag} \text{Rate}_{\text{nonthermal},i} \cdot \tau_{e-e,i} \left(1 - \frac{1 - e^{-\Delta t/\tau_{e-e,i}}}{\Delta t/\tau_{e-e,i}}\right),
\]

where \(\tau_{e-e,i}\) is the characteristic e–e scattering lifetime of an energetic electron in the corresponding metal \((i = Au, Ag)\) and \(\Delta t\) is the pulse duration. The Coulomb scattering time used above is defined as \(\Delta t = 4\pi e_0 \left(\omega_0 \right)^2 \left| E_{fi} \right|^2 /\hbar\omega_0\), where the material constant \(\tau_{0,i}\) is given by Supplementary Eq. (5). The above equations were written for a steady-state regime under CW illumination since the
optical pulse duration is longer than the plasmonic relaxation time. For calculations, we use a flux of $2.5 \times 10^8 \text{W cm}^{-2}$ that is typical for our experiments. As can be seen in Fig. 4d, it is the quantity $N_{\text{Nonthermal}, \text{avg}}$ that directly creates the ultrafast nonlinear response, which is highly enhanced by the gap plasmons in the NIR range. In other words, the NIR hot spots govern the population of energetic nonthermal electrons with energies in the interval $E_0 < E < E_0 + \hbar \omega$. The key parameter here is the plasmonic energy that is reduced for the red-shifted gap plasmon modes. There are two physical mechanisms responsible for the favorable features of hot electrons for the gap plasmon: (1) the quantum amplitude $(\hbar \omega)^{-3}$ in Eq. 1, and (2) electrons in the NIR gap plasmons have lower energies and therefore are much longer lived since $\tau_{e-e} \propto (\hbar \omega)^{-2}$ (Eq. 2 and supplementary Eq. (1b)). When energetic electrons become longer lived, the population (Eq. 2) strongly increases. We also note that the gap plasmon also has increased absorption cross sections (theory data in Supplementary Note 1 and Supplementary Fig. 5). Basically, we can see that certain aspects of the gap plasmon optical response are amplified in the NIR range.

To further highlight the benefits of the nanopatch metasurface geometry, we benchmark its nonthermal electron generation rate against other nanopatch configurations through electrodynamic simulations (Fig. 5 and Supplementary Fig. 6). For these calculations, we evaluate the peak generation rate when excited at the gap plasmon resonance (and other resonances when multiple modes are supported in the structure). In all cases, the nanoparticle volume was fixed at $(150 \text{ nm})^3$ to remain consistent with our nanopatch metasurface samples (Fig. 2a and Supplementary Fig. 2). We find up to $\sim 10 \times$ higher production in the metasurface geometry (configurations 5 and 7–10) compared to bare nanospheres, nanorods, or nanocubes (configurations 1–4 and 6). The highest rates of production in the metasurface geometry arise from the electromagnetic hot spot generated within the film-nanocube gap. We also observe a $\sim 20\%$ higher nonthermal electron generation rate when employing Ag instead of Au nanocubes in the same geometry (configurations 9 and 10). This can be attributed to less damping and a longer momentum relaxation time in the Ag nanocubes. Additionally, metasurfaces with thinner spacer layers exhibit higher nonthermal electron generation rates owing in part to an increased optical field strength (Supplementary Note 1 and Supplementary Fig. 7). These results are consistent with the previous literature, where nonthermal carrier generation was shown to be a surface effect enhanced by strong optical fields and hot spots. For example, quantum efficiencies as high as 90% have been predicted for nonthermal electron generation in dimer nanostructures with small gaps.

Transient absorption spectroscopy. Femtosecond transient absorption measurements were performed to measure the ultrafast response of the plasmonic metasurface (Fig. 6a). In all cases, samples were excited by the NIR pump at the gap plasmon resonance and probed with a continuum pulse spanning the UV-NIR (Fig. 2a). For clarity, we begin by focusing our discussion on samples with an intermediate (8 nm) thickness and later extend our analysis to other spacer thicknesses. For the 8 nm sample, the 1100 nm pump wavelength (1.13 eV) only excites intraband transitions in the conduction bands of both the Au film and Ag nanocubes, owing to their much higher IB transition energies of ~2.5 and ~4.0 eV, respectively. We observe four differential absorption features well matched to the steady-state absorption transitions in the conduction bands of both the Au film and Ag nanocubes, indicating a surface plasmon resonance (SPR) in the Au nanocubes. Figure 6c, d shows cross-sections of the field profiles and induced surface charge distributions at each corresponding peak. In the time domain, the sample exhibits kinetics ranging from ultrafast pulsewidth-limited decays with time constants $<300 \text{ fs}$ to long-lived (~ns) coherent acoustic phonon modes (Supplementary Note 7).
The 8, 25 nm, and SiO2 samples were, respectively, pumped/probed at 1100/1120, 900/920, and 500/365 nm with incident fluences of 40, 40, and 500 μJ/cm² (absorbed fluence kept constant). The ~100 fs response arises from the relaxation of highly energetic nonthermal carriers. Comparing between nonthermal carrier generation rate (c) to estimates of the peak nonthermal carrier density (d) as a function of geometry and pump wavelength. A much larger contribution to the signal is expected for excitations in the NIR.

Separation of scattering processes. To separate out the kinetics and spectral signatures of the individual scattering processes (e–e, e–ph, and phonon–phonon (ph–ph)) from our transient absorption data, we performed a lifetime density analysis (LDA) of the 2D maps (Fig. 7). The LDA method, while computationally intensive, provides a model-independent determination of the constituent lifetimes present in the temporal response along with their amplitudes (Supplementary Note 3). Furthermore, it allows the separation of pulse-width-limited responses from the data when the instrument response is well characterized, as is the case here (Supplementary Fig. 8).

Figure 7a shows the corresponding lifetime density map (LDM) obtained from the transient absorption data in Fig. 6a using LDA. Regardless of probe wavelength, oscillations are present in the amplitude of the LDM, indicating peaks in the lifetime distribution. Vertical dashed lines designate the cutoff wavelengths between absorption features (plasmon modes or IB transition), where the signal reaches a minimum and exhibits isosbestic points. The wavelength range for each was analyzed independently to extract the corresponding spectral (Fig. 7b, c) and kinetic (Figs. 7d–g) components. As is generally reported, we observe a peak with a ~1 ps lifetime corresponding to thermal e–e scattering, a peak with a ~10 ps lifetime corresponding to ph–ph scattering into coherent acoustic modes, and a long decay of ~100 ps corresponding to a spectral shift due to lattice heating.

Surprisingly, we also observe three distinct ultrafast kinetic components corresponding to nonthermal e–e scattering, which we label in Fig. 7d–g according to their respective rates (fast, intermediate, and slow). Across the three plasmon resonances, we observe consistent peaks in each lifetime distribution at 25 ± 5, 105 ± 20, and 246 ± 60 fs, which we find to be independent of pump fluence (Supplementary Fig. 10a–c). This further confirms their nonthermal nature, as the lifetime of thermal electrons is fluence-dependent and proportional to the electron temperature rise relative to the lattice (Supplementary Note 2). Additionally, we find the “fast” component cannot be attributed to the initial plasmon dephasing rate. Since the nanopatch antennas primarily exhibit nonradiative damping of the gap plasmons, the single nanocube homogeneous linewidths of 103 ± 21 meV are a direct measure of the dephasing rate (Fig. 3b, inset). This translates to a 13 ± 2.4 fs dephasing time, consistent with the known intraband nonradiative damping of gold nanorods, which is half the lifetime of the fast carriers.

Although the metasurface comprises both Au and Ag, several factors indicate the kinetics of the three plasmon resonances correspond to carriers predominantly residing in the Ag.
nanocubes. First, the optical fields at the multipolar mode are uncoupled from the gold film and localized on the nanocubes (Figs. 3a and 6c, d; and Supplementary Fig. 3). This ensures any spectral shifts of the multipolar mode are predominantly sensitive to the Ag permittivity. Second, the slowest e–e scattering rate closely matches the previously estimated 220 fs thermalization time for Ag films in the weak perturbation regime. Third, while the Ag nonthermal electrons are confined within the nanocube volume, those in Au are free to diffuse upon generation. The ballistic transport velocity of optically-excited nonthermal carriers in Au has been previously measured to be at least 1 nm fs$^{-1}$, indicating they should rapidly deplete from the gap region of the nanopatch antennas and reduce their local concentration after excitation.

Figure 7b, c further shows the decay-associated spectra (DAS) for the multipolar and gap plasmon resonances. The DAS describe the change in the transient absorption signal for each scattering process. Importantly, we observe distinct DAS for all three nonthermal carrier subpopulations, as well as for the thermal carriers (e–ph). Just like the kinetics, we find the DAS e–e lineshapes are fluence-independent (Supplementary Fig. 10d–j), strongly supporting their assignment to nonthermal carriers. However, the magnitude of the contribution to the overall signal from each subpopulation varies with pump power (Supplementary Fig. 11). With increasing fluence, the fast and intermediate nonthermal carriers have higher relative contributions to the signal than the slower nonthermal carriers. This yields an apparent acceleration of the overall e–e scattering rate of nonthermal carriers (thermalization time) and an inversion of the differential absorbance spectrum at early times in the gap mode between low (20 μJ cm$^{-2}$) and high fluence (130 μJ cm$^{-2}$). Previous studies have characterized this as a transition from a “weak to strong-perturbation regime”$^{17}$; however, until now the origin of this behavior has remained elusive.

Signatures of anisotropic behavior. We propose that the three e–e decay components can be assigned to nonthermal carriers localized near the X, L, and K symmetry points in the band structure, as illustrated schematically in Fig. 8a, b. The concept of nonthermal carriers being distinct in different portions of the band structure, i.e., anisotropic, represents a departure from the bulk treatment of noble metals as isotropic and has recently begun to gain traction in theoretical work (Supplementary Note 2)$^{13, 38}$. First-principles calculations of nonthermal carrier generation and scattering have predicted a range of scattering rates within the Brillouin zone arising from variations in the band curvature (effective electron mass or intraband plasma frequency)$^{13, 38}$, which is consistent with the ~10$^x$ variation in e–e scattering rates measured here. In the case of plasmonic intraband excitation, the breaking of momentum matching constraints enables the excitation of nonthermal carriers with large wavevectors along any band crossing of the Fermi surface. For noble metals such as Ag and Au, this implies a joint excitation of carriers at the X, K, and L symmetry points.

Until now, we have focused our analysis on the three plasmon resonances of the metasurface, which indiscriminately couple to intraband transitions in the Ag nanocubes (blue arrows in Fig. 8b). As a control experiment in support of our observations for Ag nanocubes, we now consider the response of the gold IB (Figs. 3a and 6c, d; and Supplementary Fig. 10d–j). Unlike the purely intraband transitions probed by the plasmon modes (blue arrows in Fig. 8b), the gold IB response selectively probes carrier dynamics at specific points of the band structure (blue arrows in Fig. 8a). In our measurement window, IB transitions near the Fermi edge of the X and L-points are excited at peak energies of 2.4 eV (517 nm) and 2.8 eV (443 nm), respectively$^{39}$. For thermal carriers (e–ph) and slow nonthermal carriers (e–e$\text{slow}$), we observe a strong transient bleach or absorption feature, respectively, at both 436 and 517 nm, closely matching the literature values for IB transitions at the X and L-points. However, the intermediate nonthermal carriers in Au only exhibit a bleach feature near 517 nm, indicating their response is localized near the X-point. Finally, the fast nonthermal carriers exhibit bleach components near the X and L-points, however, the
response near the X-point is shifted to lower energy and is indicative of a carrier population below the Fermi level. We expect that anisotropic carrier scattering also occurs near the K-point, but this transition lies beyond our measurement window.

Compared to the plasmon resonances in the silver nanocubes, the gold IB transitions exhibit much slower rates of e–e scattering overall (Fig. 7f). We observe fluence-independent peaks at lifetimes of 46 ± 13, 172 ± 4, and 338 ± 71 fs. The reduced scattering rate arises from a stronger charge screening by bound d-band electrons in the gold and is roughly proportional to $1/\sqrt{\varepsilon_{\infty}}$ at lower carrier energies, where $\varepsilon_{\infty}$ is the static permittivity of the metal. We find the slow nonthermal e–e scattering time to be ~37% longer in gold, consistent with the predicted value of 35%[17]. Thus, the combination of observing 3 distinct population decays in Au that are slower than Ag by 2 for each $\omega_p$-band (reduced plasma frequency) at higher energies (Supplementary Note 2). To confirm that these trends are solely pump energy dependent and not arising from the antenna geometry, we also excited the 5 nm Al$_2$O$_3$ sample at multiple energies about the gap resonance and observe the same result. The different trends in lifetime as a function of energy support our conclusion that anisotropic electron scattering creates subpopulations at different points in the band structure.

Discussion

A combination of factors likely masked the detection of these nonthermal subpopulations in earlier literature. In uncoupled nanoparticles and metal films, for example, a lack of field hot spots significantly reduces nonthermal carrier generation rates as compared to the nanopatch metasurface. Additionally the analysis of kinetics at single wavelengths (particularly near the IB transitions) biases the response, and at many wavelengths contributions from two of the nonthermal carrier populations can cancel out (Supplementary Note 6 and Supplementary Fig. 11). This can incorrectly give the impression of wavelength-dependent e–e scattering rates when, in fact, a global analysis reveals that different wavelengths measure different percentages of contributions from nonthermal carriers. Finally, the extraction of rate distributions from complex data sets requires the kinds of global analysis methods such as the LDA employed in this work.

In summary, we have measured the optical response of both nonthermal and thermal hot electrons in a plasmonic metasurface using ultrafast transient absorption spectroscopy. This was enabled by the high generation efficiency of nonthermal carriers in the nanopatch geometry, which we showed is a result of the hot spot generated within the gap. Our results show that the choice of geometry and materials are crucial for producing strong ultrafast
Methods

Sample fabrication. Silicon substrates with a 50 nm gold film were coated with ALD-deposited alumina (Al₂O₃) spacers of varying thickness (1–25 nm) by VaporPulse Technologies. PVP-coated silver nanocubes with ~150 nm edge length (nanoComposix, Inc.) were deposited by a previously reported method. Briefly, a single PAH layer was deposited on the Al₂O₃ surface to promote nanocube adhesion. The PAH with an average molecular weight of 58,000 was purchased from Sigma-Aldrich. A solution of 150 nm nanocubes was then prepared by concentrating a 1 mg mL⁻¹ solution of nanocubes in ethanol and re-suspending in 18 MΩ DI water, yielding a final concentration of 4 mg mL⁻¹. An 18 mm round coverslip was placed on top of 15 µL of solution, and the nanocubes were allowed to settle onto the surface for 30 min. The coverslips and excess solution were then removed with DI water and the samples were dried with nitrogen gas.

Nanocube characterization. Samples were imaged under high vacuum (<1×10⁻⁴ Torr) with a JEOL 7500 scanning electron microscope. To measure nanocube dimensions, the area, and min and max Feret’s diameters were extracted using the particle analysis macro in ImageJ. From these values, the edge lengths and radius of curvature for each particle were fit assuming a rounded rectangle shape (Supplementary Fig. 2). Over 440 individual particles were analyzed to obtain reliable statistics.

Reflectivity measurements. Absolute ground-state reflectivities were measured using the integrating sphere of a Perkin Elmer Lambda 950 spectrometer coupled to a P66 photodetector and PMT (Fig. 3b, c and Supplementary Fig. 4). Single nanocube absorption spectra (Fig. 3b inset) were captured using an inverted Nanospec (Acton SP2300 and NIRvana 640, respectively). Unpolarized light from a halogen lamp was both focused and collected through a 100x objective (Olympus LMPlan IR, NA = 0.8) and passed through a 75 µm slit prior to entering the spectrometer to isolate individual particles. Spectra of the specular reflected light from each nanocube were integrated for 200 ms, averaged over 100 exposures to avoid detector saturation, and were normalized to spectra of the bare substrate between particles. Single particle spectra were then fitted to a Lorentzian function to extract the homogeneous linewidths.

Transient absorption spectroscopy. Femtosecond transient absorption spectroscopy was performed using a Ti:sapphire laser with a regenerative 800 nm output.

Fig. 7 Lifetime density analysis of the ultrafast response. a Lifetime density map (LDM) fitted to the differential absorbance data in Fig. 6 and displayed on a log-log scale. Multiple distinct peaks can be observed spanning the entire range of lifetimes at each resonance. b, c Comparison of the decay-associated spectra (DAS) corresponding to fast, intermediate (int), and slow nonthermal e–e scattering and thermal e–ph scattering at the multipoles and gap plasmon resonances. The small fluctuations at 1100 nm are artifacts from residual pump scatter. d–g Lifetime traces of the LDM (blue circles) at the same wavelengths as the kinetics in Fig. 6e. Contributions from nonthermal electron-electron scattering and thermal electron-phonon scattering are indicated (shaded regions) along with the total fitted response (red line). The gray regions are contributions from phonon-photon scattering (~10 ps) and semi-infinite decay due to lattice heating (~100 ps).

nonthermal electron contributions that can be measured through transient absorption. We present experimental evidence of three subpopulations of nonthermal carriers with distinct e–e scattering rates and spectra. We propose that these subpopulations are localized near band crossings of the Fermi surface and decay anisotropically with different time constants. This work supports recent first-principles calculations, predicting variations in nonthermal carrier scattering near the Fermi surface of noble metals. The nonthermal carrier response is shown to extend from the UV to NIR, spanning multiple surface plasmon resonances of the nanoparticle antennas and IB transitions of the underlying gold film. We find an order of magnitude variation in e–e peak scattering times of the three nonthermal subpopulations spanning ~25–250 fs in the Ag nanocubes and ~45–340 fs in the Au film. Harnessing this ultrafast carrier response, in particular at intraband transitions, could enable low-power optical switching well into THz frequencies. Furthermore, one may selectively excite transitions in regions of the band structure with longer lifetimes to leverage the wavevector-dependence of nonthermal carrier scattering, which could improve the efficiency of hot carrier injection. The new insights provided here can impact a diverse range of fields, including telecommunications, nonlinear optics, metamaterials, photocatalysis, photodetectors, and the study of systems far from thermal equilibrium.
Slow nonthermal carriers yield perturbations to transitions near both X and L points. IB transitions at the K point occur at much higher energies, and thus a bleach at the L transition. Intermediate nonthermal carriers appear to be localized solely at the X point where they induce a bleach of the absorbance.

Offset for clarity. The fast nonthermal carriers redshift the X transition, indicative of carriers residing below the Fermi level near the X point, and also exhibit two e-e components supports their assignment to distinct nonthermal carriers. The different dependence of peak lifetime vs. pump energy of the thickness are varied. An opposite trend is observed for the intermediate nonthermal carriers redshift the X transition, indicative of carriers residing below the Fermi level near the X point, and also exhibit a bleach at the L transition. Intermediate nonthermal carriers appear to be localized solely at the X point where they induce a bleach of the absorbance. Slow nonthermal carriers yield perturbations to transitions near both X and L points. IB transitions at the K point occur at much higher energies, and thus are not resolved in the measurement range.
set of normal distributions to extract the lifetime distribution and corresponding DAS for each scattering process. Further details on the LDA and global fitting procedures can be found in Supplementary Notes 3 and 4.

Classical electromagnetic simulations. Classical simulations of the nanopatch structures were performed using classical electrodynamics with COMSOL Multi-physics, assuming standard boundary conditions and using literature values for the dielectric functions of the gold film and silver nanocubes. Calculations assumed a normally incident beam and periodic boundary conditions with the lateral period taken as a large number (500 nm) to avoid interparticle coupling. Silver nanocubes with an edge length of 154 nm and a corner radius of 10 nm (taken from measurements) were modeled with a 3 nm PVP coating, and the surface of the Al2O3 spacer was assumed to be covered with a 2 nm PAH layer. The refractive indices for PVP, PAH, and Al2O3 were taken to be 1.52, 1.4, and 1.77, respectively. Using these conditions, we achieved excellent agreement with experiment for the positions of the plasmon resonances and the homogeneous linewidths (Fig. 3b, c and Supplementary Fig. 4b).

The local dissipation spectra and maps in the text were computed from the standard equation for the local rate of losses:

$$Q_{abs,local} = \left( j \cdot E \right) = \left( m_{\text{metal}} \right) \frac{1}{8 \pi} E_{\text{in}} \cdot E_{\text{out}},$$

where $$E_{\text{in}}$$ and $$E_{\text{metal}}$$ are the complex field amplitude and dielectric constant of the corresponding metal, respectively. Another parameter illustrating characters of the plasmonic modes in our sample is the surface charge shown in Figs. 3a and 6c. It was calculated in the following manner:

$$\Delta E_{\text{surf}} = E_{\text{in}} - E_{\text{out}}$$

where $$E_{\text{in}}$$ and $$E_{\text{out}}$$ are the normal fields near the surface inside and outside the metal, respectively.

Data availability. The data that support the plots within this paper and other findings of this study are available from the corresponding author upon reasonable request.

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M.E.S., G.M.A., M.H.M., A.O.G. and G.P.W.: Conceived and planned the work. J.W.S., G.M.A., A.B.F.M., and D.R.: Fabricated the samples. M.E.S., D.J.G. and G.P.W.: Performed the optical spectroscopy. M.E.S., A.O.G. and G.P.W.: Performed the data analysis. X.-T.K., Z.W. and A.O.G.: Performed quantum and classical electrodynamics calculations. M.H.M., A.O.G. and G.P.W.: Supervised the project. M.E.S., M.H.M., A.O.G. and G.P.W.: Wrote the manuscript with contributions from all authors.

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