Electrostatic control over optically-pumped hot electrons in optical gap antennas

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Abstract

We investigate the influence of a static electric field on the incoherent nonlinear response of an unloaded electrically-contacted nanoscale optical gap antenna. Upon excitation by a tightly focused near-infrared femtosecond laser beam, a transient elevated temperature of the electronic distribution results in a broadband emission of nonlinear photoluminescence (N-PL). We demonstrate a modulation of the yield at which driving photons are frequency up-converted by means of an external control of the electronic surface charge density. We show that the electron temperature and consequently the N-PL intensity can be enhanced or reduced depending on the command polarity and the strength of the control static field. A modulation depth larger than 100% is observed for activation voltages of a few volts.

Plasmonic metal nanoantennas have been attracting an ever growing attention due to their ability to strongly enhance electromagnetic fields at the nanoscale and consequently improve weak nonlinear optical processes. The development of advanced simulating tools combined with modern nanofabrication facilities undoubtedly pushed the general understanding to a level where nonlinear plasmonics can be deployed as a device technology vehicle. From the microscopic point of view, the nonlinear responses occurring in metal optical antennas, subject to an ultrafast intense electromagnetic pulse, arise from the swift generation of an out-of-equilibrium electron distribution and its associated thermalization dynamics. Externally controlling these parameters is a difficult task. Therefore, nonlinear behaviors are essentially adjusted by engineering the strength of the local electromagnetic field either by varying the geometry during the nanofabrication stage or by optimizing the pumping conditions. While the latter has the unique benefit to dynamically manipulate coherent nonlinear signals at ultrafast time scales, it is certainly not a viable approach for integrating nonlinear plasmonic functional units in a photonic circuit architecture. A complementary alternative strategy is to control the nonlinear activity by an electrical command applied to the system. A few demonstrators were reported in the literature where the generation of second harmonic light was tuned by plasmonic gaps loaded with nonlinear electro-optical materials.

Our work brings an additional stone in this general context. Upon excitation by a tightly focused femtosecond near infrared pulsed laser, Au nanostructures are known to react nonlinearly by producing, among other nonlinear processes, second-harmonic light and a broad up-converted photoluminescence spectrum. The latter was originally attributed to a two-photon interband absorption process because of its near quadratic power dependence. However, alternative explanations were recently proposed including inelastic transitions, higher-order nonlinearities and radiative decay of a transient out-of-equilibrium electron distribution. The pump laser transfers...
energy to the electrons in the metal producing a non-Fermi distribution for a few femtoseconds. Subsequently, electron-electron scattering leads to thermalization of the electron gas which can reach several thousands of Kelvins. Before the lattice and the electron gas reach equilibrium in picoseconds time range, electrons will remain hot and possibly emit a thermal electromagnetic continuum. The presence of a hot electron distribution and its different characteristic relaxation times were clearly identified in the mechanism leading to nonlinear photoluminescence (N-PL) emission. This broadband N-PL and its associated non-equilibrium electron distribution is ubiquitous of metal nanostructures and has been observed for a wide range of geometries ranging from simple resonant nanoparticles, to coupled antennas and extended two-dimensional forms.

We demonstrate in this report that a command voltage can enhance or reduce the nonlinear photoluminescence produced by electrically-contacted Au optical gap antennas without relying on the co-integration of nonlinear dielectrics, in other words, the interstice of the antennas remains unloaded. This is achieved by adjusting the electronic temperature responsible for the N-PL via an electrical control of the surface electron density.

![Diagram of experimental setup](image)

**Figure 1**: (a) Colorized scanning electron microscope (SEM) image of a Au optical 50 nm gap antenna used in the experiment. Au is colorized by a yellow hue. (b) Experimental setup. A Ti : Sapphire laser emits 180 fs pulses at 808 nm. The laser is focused on the optical gap antenna, which is electrically connected to a voltage source. The nonlinear photoluminescence is detected by an avalanche photodiode. (c) Confocal scanned image of the nonlinear photoluminescence emitted by the device. The area approximately covers the image in (a).

Electrically-contacted gap antennas taking the form of two gold tapers separated by a sub-100 nm gap are produced by a two-step lithography. Figure 1(a) shows a scanning electron micrograph of a 50 nm gap antenna produced by electron beam lithography in a first step. In the second lithography step, the microscopic electrodes contacting individual optical antennas are fabricated by photolithography. Details about the fabrication process are presented in Sect.1 of the Supporting Information.

The sample is then transferred to an inverted optical microscope (Nikon, Eclipse) equipped with a two-dimensional piezoelectric sample scanner (Mad City Labs, NanoLP-100). A simplified sketched of the setup is pictured in Fig. 1(b). A high numerical aperture (N.A.) objective (×100, N.A.=1.49) focuses on the sample the light of Ti:Sapphire laser (Coherent, Chameleon) emitting 180 fs long pulses at a repetition rate of 80 MHz and tuned at a wavelength of 808 nm. The nonlinear up-converted photoluminescence of the sample is collected by the same objective and is spectrally discriminated from the incident wavelength by a dichroic beamsplitter and a bandpass filter. Except otherwise specified, the second-harmonic signal simultaneously generated along side the N-PL is also rejected by a notch filter centered at 405 nm. The two leads defining the gap of the optical antenna are electrically connected to a function generator delivering a square command voltage \( V(t) \). Considering the resistance of the electrical connections (\( \sim 100 \, \Omega \)), the electrical potential is mostly dropped at the interstice. For an approximately 50 nm gap size formed by two plane-parallel plates, the electric field across the two leads is in excess of \( E = 2 \times 10^7 \, \text{V} \cdot \text{m}^{-1} \) per applied volt. The photoluminescence signal locally emitted by the device is detected confocally by a photon counting avalanche photodiode module (Excilites, SPCM-AQRH) either as a function of the \((x, y)\) position of the sample with respect to a fixed focus spot, or as a function of the applied voltage \( V(t) \) for a given position of the sample.

An example of the nonlinear \((x, y)\) photoluminescence map is illustrated in Fig. 1(c) for a null bias. Here the optical gap antenna is laterally scanned through the focus of the laser and the N-PL photon counts is integrated over the emitted spectrum (low pass filter at a cut-off wavelength of 680 nm). The average incident power is 290 μW and the polarization is aligned with the main axis. As anticipated, the highest N-PL signal occurs at the gap of the antenna where the incident field is enhanced. The edges of the electrodes are also visible as a faint background in the image. As expected, the N-PL emission yield evolves nonlinearly with the incident...
Figure 2: (a) Time traces showing the response of the N-PL to a 12 V square waveform with a period of 2 s. (b) Another N-PL time trace obtained on a different antenna with a 20 V square waveform and a period of 800 ms. The N-PL enhancement is approximately 500 \%.

Laser power as shown in Sect. 2 of the supporting information.

Figure 2(a) shows an example of N-PL voltage-induced modulation when the laser is situated closed to the antenna gap. Here, a 12 V square voltage waveform is applied to the electrical contacts at time $t = 8$ s. The voltage signal is recorded with a limited number of samples due to a limitation of the acquisition electronic when monitoring simultaneously the voltage and the APD counter. At null voltage, the N-PL count rate is constant at 270 kcount·s$^{-1}$ indicating a residual lateral shift from the pixels featuring the highest count rate at the gap center (≈ 520 kcount·s$^{-1}$). When the voltage is applied, the N-PL rate increases to reach approximately 400 kcount·s$^{-1}$, an on-state enhancement of 1.5 fold. The rate goes back to its zero-bias off-state level between two voltage pulses with an apparent delay. Further discussion will address this point later on. The enhancement factor depends on extrinsic parameters such as the applied voltage and the laser power as discussed later, but also on intrinsic figures, essentially linked to the fluctuations of the geometry of the optical gap antenna imparted by the nanofabrication process. As an illustration of the above arguments, Fig. 2(b) shows a significantly greater modulation depth obtained from a 100 nm interstice, a 20 V square waveform with a period of 800 ms and an average laser power of 49 \( \mu \)W. Although there is a small deviation in the N-PL maximum rate over time, the on-state enhancement reaches nearly 500 \% in this experiment despite a large inte-electrode spacing.

In the following sections, we investigate the influence of the extrinsic control parameters on the modulation of the N-PL rate. The parameters can be controlled with greater precision than intrinsic parameters such as a gap size or electrode morphology, which are plagued by inherent fabrication uncertainties. We start by studying the response to the amplitude of the control voltage $V(t)$. Figures 3(a) and 3(b) show time traces of the N-PL rate when the amplitude of the applied square waveform is varied in an upward and downward sweep, respectively. The data obtained during the sweeps are reproducible indicating that neither the voltage range sampled nor the laser power irreversibly degraded the optical gap antenna. At that stage of the discussion, one can draw preliminary conclusions. The graphs unambiguously show that
the N-PL flux is sensitive to the applied voltage bias, providing thus an external electrical handle to manipulate the nonlinear activity. Since the N-PL sits on relatively large offset at zero bias, the voltage-induced modulation becomes visible only after a certain voltage value. In the present experiment, the modulation remains undetected for $V < 5 \text{ V}$. Figure 3(c) illustrates the voltage dependence of the mean N-PL rate estimated during the on-state of the voltage duty cycle. A quasi linear trend is observed. The proportionality between the amplitude of the bias and the rate of N-PL is a parameter which may prove useful for realizing a commanded device. The dotted line at $222 \text{ kcount s}^{-1}$ is the mean flux of the N-PL calculated in the time sequence when the signal is unperturbed by the voltage.

Crucial to the observation of the N-PL modulation is the precise location of the laser spot with respect to the antenna gap. Typically, for a given bias voltage and gap size, the strength of the modulation strongly depends on the adjustment of the diffraction-limited observation area. Centering the gap in the laser focus does not necessarily render the strongest modulation. Likewise, no modulation is observed when probing the N-PL at the edges of the tapered section. We find that the modulation is generally the strongest when the tips of the tapers forming the bow-tie are selectively illuminated. Figure 4(a) and 4(b) show the time evolution of the N-PL signal when the sample is displaced from the maximum response of the gap by approximately 100 nm on either side. In this experiment, the antenna gap is nominally 80 nm. In Fig. 4(a), the displacement of the sample with respect to the focus favors a N-PL signal partially overlapping the positive electrode and one retrieves the behavior observed previously where the on-state enhances the N-PL signal. In Fig. 4(b), the sample is slightly moved with respect to the laser focus to favor a N-PL response from the grounded electrode. Clearly, the N-PL rate and the amplitude are now out-of-phase. The dotted lines in the graphs indicate the approximate off-state N-PL rate (null voltage). Upon application of the voltage bias, the N-PL is no longer enhanced but suffers from a significant reduction of its rate compared to its steady-state value at $V = 0 \text{ V}$. This behavior suggests that the polarity of the electrode contributing to the photoluminescence signal plays an important role in the control mechanism. We confirmed the above observation by changing the polarity of the amplitude without changing the position of the sample (overlap over the ground electrode). Results are shown in Fig. 4(c). Again, the dashed line at $270 \text{ kcount s}^{-1}$ is the N-PL rate for $V = 0 \text{ V}$. Clearly, an on-state positive bias (here $V = +20 \text{ V}$) quenches the N-PL while a on-state negative bias ($V = -20 \text{ V}$) gives rise to an enhanced nonlinear signal.

Enhancement or quenching of the N-PL rate when the voltage is turned on and off is not immediate. The time traces in Fig. 2, Fig. 3 and Fig. 4 show that the temporal evolution of the N-PL is occurring with a slow dynamics and is not symmetric with respect to the voltage edge. In Fig. 5(a) the average rise time $\tau_r$ and decay time $\tau_d$ measured at 10% and 90% of the modulation are.
\( \tau_r = 150 \pm 30 \text{ ms} \) and \( \tau_d = 390 \pm 80 \text{ ms} \). However, the figures are here indicative. We have observed a large variability from antenna to antenna, but also between positions of the sample with respect to the laser focus. When the on-state quenches the signal, the asymmetry reverses as illustrated in Fig. 5(b). However, the slower time is always associated to the settle down of the N-PL when the voltage goes to 0 V suggesting that a diffusive process is at play. If the kinetics of the N-PL signal would be mainly driven by such a slow time response, electrical signals with a duration smaller than the \( \tau_r \) would not significantly affect the N-PL rate. Figure 5(c) displays an experiment where the voltage bias is a 10 ms pulse. The modulation here remains substantial (\( \sim 90\% \)) suggesting that the kinetics is driven by a combination of a fast and a slow process.

Let us review the different mechanisms that can explain a voltage-sensitive N-PL response. Possible contributors include electrostriction, field-induced morphological reorganization, third-order \( \chi^{(3)} \) nonlinearity, and various hot electron phenomena. Electrostriction is a mechanical deformation of any material upon applying an electrical stress. In the gap geometry considered here, the phenomenon would manifest itself as a variation of gap width due to an elastic deformation of the Au terminals, the supporting SiO\(_2\) substrate, or a combination thereof. The electromagnetic response of the gap is essentially dictated by the distance separating the two Au electrodes. Within the range of the gap sizes considered here (tens of nanometers), the local electromagnetic field in the gap is strongly enhanced when near-field coupling between the two metal sides is favored. Because of the intrinsic nonlinear nature of the detected luminescence, the signal is a probe of the local electromagnetic field built up in the gap region. Therefore compressive or tensile in-plane stresses induced by the applied voltage would simultaneously increase or reduce the N-PL rate. The electrostrictive strain in the material features a quadratic dependence with the electric field, therefore its sign does not change direction when the field is reversed and cannot explain the polarity-dependent on-state enhancement or quenching of the N-PL rates observed experimentally in Fig. 4. We confirmed the absence of an electromechanical deformation of the gap by simultaneously monitoring second-harmonic photons (SHG) produced along side the nonlinear photoluminescence.

The SHG signal was previously reported to be sensitive to the size of the gap in coupled geometries and to subtle variations of the shape of a nanostructure. With the optical antennas studied here, the SHG response is generally weaker than the emitted N-PL. Figure 6(a) shows the times traces of the N-PL and SHG signals collected from a 20 nm gap (nominal) biased with 7 V square waveform. The N-PL follows the voltage drive while the modulation of the SHG remains very weak upon applying a bias in this unloaded optical gap antenna. Figure 6(b) shows the Fourier transform of the varying signals around the modulation frequency (here 0.5 Hz). The graph hints to the presence of a residual response of the SHG to the time-varying applied field, but remains many orders of magnitudes smaller than the N-PL response.

Another phenomenon that could contribute to a change in the N-PL rate is a modification of the morphology of the optical gap antenna either im-

![Figure 5](attachment:image_url)
Figure 6: (a) Time traces of voltage $V$, N-PL and SHG responses obtained from a 20 nm wide gap antenna. The N-PL rate is modified upon application of 7 V bias while the SHG remains weakly affected. (b) Fourier transform of the modulated signals. The N-PL peak present at the modulation frequency (0.5 Hz) is significantly stronger than the SHG.

posed by the laser$^{22}$ or by a field-induced reorganization of the metal interface.$^{35,36}$ Since the modulation amplitude does not significantly change in the time traces presented above, we conclude that the structure under illumination does not undergo an irreversible reshaping and the intensity of the laser remains below the damage threshold.$^{27}$ Like second-harmonic generation arising in small Au structures, N-PL is believed to be a surface effect.$^{5,37}$ Hence, a reversible electric field-induced reorganization of the metal outer boundaries would necessarily affect the strength of both the N-PL and SHG signals. Reordering of the Au outmost atomic layers was found to occur for local field magnitudes in excess of $25 \times 10^9$ V·m$^{-1}$,$^{35}$ a value two orders of magnitude higher than typical fields applied in our experiment. Even considering an increased mobility of Au adatoms due to a local laser-induced rise of the temperature, it unlikely explains the difference between on-state enhancement and on-state quenching observed for the N-PL. Although noisy, the SHG signal in Fig. 6 confirms that there is no drastic reorganization of the antenna’s surface during the experiment.

An additional possible explanation for the voltage modulation of the N-PL is the activation of a Kerr-type nonlinear susceptibility induced by the strong electrostatic field developing in the gap region. The field displaces charges by polarizing the two sides of the gap thereby creating a disturbance in the number density of the electrons. This sheet of polarization, confined to the first atomic layers, can be taken into account by introducing a correction to the complex dielectric function of Au.$^{38}$ However, like in electrostriction, the corrective term depends quadratically on the electric field. The presence of $\chi^{(3)}$ nonlinearity is thus not able to account for the N-PL sensitivity to the sign of the voltage and the linear-like trend shown in Fig. 3(c).

Let us go back to the origin of the nonlinear photoluminescence and the presence of a hot electron distribution. It has been debated that the up-converted emission and its associated nonlinearity may not arise from a multiple absorption events followed by an interband recombination of electron-hole pairs. Instead, it is hypothesized that the nonlinear behavior stems from the emission process. The out-of-equilibrium electronic distribution generated by the femtosecond light pulses absorbed by the metal radiates.$^{22}$ It is interesting to note that this scenario bears strong similarity with the anomalous light emission observed in electron-fed antennas, where an overbias emission, in other words an upconverted signal, was also understood from the framework of a heated electron gas.$^{39,40}$

Figure 7 shows typical N-PL spectra emitted by the gap antennas for two biasing conditions $V = 0$ V and $V = 10$ V. In this example, the bias leads to an on-state enhancement of the N-PL. Both spectra feature a monotonic tail decaying in the visible part of the spectrum and show no evidence of a surface plasmon response.$^{27}$ Following the approach consisting at treating the nonlinear photoluminescence as the glowing fate of hot electrons, we tentatively deduce an effective electron temperature using Planck’s law of radiation:

$$I_{\lambda} \propto \frac{2\pi \hbar c^2}{\lambda^5} \frac{1}{e^{\frac{\hbar c}{k_B T_e}} - 1}$$

where $I_{\lambda}$ is the spectral irradiance, $h$ and $k_B$ are Planck’s constant and Boltzmann’s constant, and $T_e$ is the electron temperature. The solid and dotted lines are fit to the data using Eq. 1. The two fitted curves follow relatively well the spectral de-
dependence of the N-PL considering a temperature of the electron gas at 1542 K when no bias is applied and a raise of \( T_e = 1589 \) K under a 10 V voltage bias. Both temperatures are in the same order of magnitude as previously reported values for electronic temperatures reached after pulsed laser excitation.\(^{26,41}\)

\[
F_0 = \frac{\pi^2}{4} N_e k_B \sqrt{\frac{\pi T_F T_L h \tau_p}{m_e}}
\]  

(3)

\( \tau_p \) is the pulse duration and \( m_e \) is the electron mass. Equation 2 shows that if the electron temperature remains below \( T_s \), it no longer depends on \( 1/N_e \) but on \( 1/\sqrt{N_e} \). A similar dependence was also found by Agranat and co-workers.\(^{13}\)

To check the validity of our reasoning, we estimate what should the laser fluence \( F \) to reach a maximum electron temperature of about 1500 K. Taking a typical experimental laser power at \( 49 \times 10^{-6} \) W and the repetition rate of the laser at \( 80 \times 10^6 \) Hz, the laser pulse energy is \( 0.61 \times 10^{-12} \) J. We find that in order to reach the desired \( T_e \), the illumination area should have a diameter of approximately 450 nm. This value is in very good agreement with the point spread function of the high N.A. objective used in this study.

In Eq. 2, the only parameter that may be changed upon applying a bias is the local density of electrons present at the surface of the two capacitively coupled electrodes. The influence of carrier concentration on the intensity of linear photoluminescence was already pointed out for organic polymers.\(^{23}\) The static electric field dropped in the gap introduces polarization charges present within the Thomas-Fermi screening length of the metal leading to an accumulation of electrons on the ground side and a depletion of electrons on the positive lead. Using Eq. 2, we find that \( \frac{\Delta T_e}{T_s} = \frac{1}{2} \frac{\Delta N_e}{N_e} \). Assuming an unperturbed electron density at \( N_e = 5.9 \times 10^{28} \) m\(^{-3}\) for \( V_b = 0 \) V, we evaluate the required change of \( N_e \) to account for an increased electron temperature \( \Delta T_e = 1598 - 1542 = 47 \) K when switching the bias from \( V_b = 0 \) V to \( V_b = 10 \) V. We find that \( \Delta T_e \) can be explained by \( \Delta N_e = 3.59 \) nm\(^{-3}\). To obtain a surface charge density, we make the assumption that most of the density change is contained within the Thomas-Fermi screening length \( L_{TF} \), which is about 60 pm for Au.\(^{15}\) Correspondingly, the change of the electron surface density is then \( \Delta N_e \times L_{TF} = 3.4 \times 10^{-2} \) C.m\(^{-2}\).

In the following, we use a finite element method software (COMSOL) to obtain the charge distribution in the region of the gap and to compare it to the above figure. The model considers an ideal capacitor with perfectly conductive electrodes, smooth interfaces and a bow-tie shape mimicking the real device. The boundary conditions

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure7.png}
\caption{Spectra of the nonlinear photoluminescence acquired for two applied voltages \( V = 0 \) V and \( V = 10 \) V, respectively. The solid lines are fits using Planck’s law leading to two different electron temperatures.}
\end{figure}

The daunting question to answer is how can a static bias influence the temperature of an out-of-equilibrium electron gas generated by the absorption of a light pulse. In a 1998 paper, A. P. Kanavin and co-workers derived an analytical expression for \( T_e \) when metals are irradiated with femto-second light pulses.\(^{12}\) They found that for times \( t \) longer than the equilibration of the electron distribution, the electron temperature becomes inversely proportional to the electron density \( N_e \) if \( T_e \) exceeds a characteristic temperature \( T_s = \sqrt{\varepsilon_f T_L/k_B} \) where \( \varepsilon_f \) is the Fermi energy and \( T_L \) is the phonon temperature. Considering \( T_L = 300 \) K and \( \varepsilon_f = 5.53 \) eV for Au, \( T_s \sim 4 \times 10^5 \) K. For the laser power used in the present experiment, the electron temperature after the pulse is expected to be smaller than \( T_s \). We thus derive in the Sect. 3–5 of the Supporting Information an analytical expression for \( T_e \) when \( T_e < T_s \). We find

\[
T_e = \sqrt{\frac{F}{F_0}} T_F T_L
\]  

(2)

\( F \) is the laser fluence, \( T_F \) is the Fermi temperature, and \( F_0 \) is a characteristic fluence and writes
are as follows. Two gold terminals with a potential difference of 10 V are separated by a 50 nm gap. The gold structures are resting on a silica glass substrate modeled by a cuboid of 1.5 µm×1.5 µm×100 nm. The superstrate is air. Figure 8(a) shows the electrical potential around the gap region. The upper electrode is here held at 10 V. The magnitude of the calculated electric field is about 2.7×10^8 V·m⁻¹ in Fig. 8(b). The bow-tie geometry leads to a slightly enhanced electric field compared to a capacitor formed by two parallel plates facing each other. Figure 8(c) displays the corresponding surface charge density \( \sigma \). As expected, the largest charge accumulation occurs at the foremost region where the electric field is the strongest. Under a 10 V bias, the value of \( \sigma \) at the interface reaches \( \pm 10^{-2} \text{C}·\text{m}^{-2} \). The simulated number is in comforting agreement with the modification of the electron density estimated above. These consistencies support us in our scenario: the voltage modulation of the N-PL originates from an electrical control of the electron temperature via a modification of the surface charge density. This also explains why modulation of the N-PL is not observed at the edges of the tapered section of the bow-tie since, under a biased condition, the surface charge density preferentially concentrates at the tip. Our finding favors the school of thought explaining the nonlinearity of the photoluminescence emission from the radiative decay of a heated electron gas colliding with the interface. Note that the elevated electronic temperature may still result from a multi-photon absorption process of the laser pulse.

The relationship between the N-PL thermal emission is a nonlinear function of \( T_e \) (Eq. 1) and \( T_e \) is itself inversely proportional to \( \sqrt{N_e} \). We claim that the range of temperatures explored by the voltage activation is small enough so that the trend can be linearized. To support this hypothesis, we investigated the voltage-dependent N-PL emission spectrum of another gap antenna as shown in Fig. 9(a). The data are fitted again with Planck’s law to extract the corresponding electronic temperature. In this set of curves, the spectra (open circles) feature a small resonance in addition to the black-body like tail. Hence, a plasmon contribution in the form of a Gaussian shape is phenomenologically added to Eq. 1 to take into account for the shoulder present at around 600 nm. The solid lines are the resulting fits obtained for the different bias contributions. The fits are satisfactorily following the experimental data points. In Fig. 9(b) we plot the N-PL intensity, integrated over the entire spectral range, as a function of deduced electronic temperature \( T_e \) for all the control voltage values utilized. The evolution features a quasi linear trend in this relatively small excursion of the electronic temperature. The apparent linearity confirms our hypothesis that the explored temperature range is too small to observe the nonlinearity expected from Eq. 1 and explains the linear N-PL evolution observed as function of voltage in Fig. 3(c).

There is one part of the data set shown above that is not completely explained by the proposed mechanism. As shown in Fig. 5, the modulation is characterized by a combination of a slow and a fast process. Clearly, the charge and the discharge of
the gap is dictated by the value of the capacitance, which is here calculated at $C = 6.5 \times 10^{-18}$ F. Considering a 50 $\Omega$ load, the gap is polarized within a fraction of a femtosecond. Inevitably, the presence of a millisecond dynamics hints the presence of additional mechanisms not taken into account so far.

The electric fields generated in the gap are fairly strong: the calculation for a 10 V applied voltage shown in Fig. [3(b)] indicates an electric field approaching $3 \times 10^8$ V-m$^{-1}$. Such a large field imposes an electrical stress on the SiO$_2$ substrate, which lead to a wear out of the oxide and eventually to its dielectric breakdown, when a percolation path forms between the two electrodes. The alteration of the insulator layer is associated to the presence of native and stress-induced electron traps in the oxide.

Among the many different contributors leading to the creation of charge trapping defects and states located at the Au/SiO$_2$ interfaces, a relevant mechanism for the discussion involves the electrical injection of hot holes and hot electrons in the oxide by Fowler-Nordheim tunneling through the triangular potential barrier. In the present experiment, we may discriminate two populations of hot carriers. A first population gains its energy by the lateral field applied across the gap. These carriers injected in the dielectric typically may be trapped by native defects, and may as well generate new trapping sites by releasing hydrogen species that are diffusing toward the counter electrode and producing interface states and electron traps. These traps can be charged upon application of the bias by tunneling of electrons, and discharged on removal of the voltage pulse. The detrapping lifetime depends on the nature of the centers, their location within the dielectric, the strength and the duration of the electrical stress and a wide range of values spanning $10^{-12}$ s to $10^3$ s are reported in the literature.

A second population of hot electrons, at the origin of the N-PL signal, is created following the absorption of the laser pulse in the metal electrodes and swiftly decays within about a picosecond. This photo-induced out-of-equilibrium electronic distribution drifts with the static electric field whenever $V$ is applied. We hypothesize that the slow part of the nonlinear photoluminescence kinetics is dictated by charging and discharging of field-induced electron traps in the dielectric barrier. In an attempt to verify this claim, we identify the formation of defects in the oxide by monitor-

![Figure 10: Time traces showing the voltage, the N-PL signal and the current. The N-PL responds to the bias with an on-state enhancement. At $t \sim 11$ s, an onset of leakage current is measured resulting from the degradation of the electrically stressed dielectric medium in the gap.](image)
Other type of amorphous dielectrics, such as HfO$_2$, will also feature electrical-induced wear-out. We discarded crystalline quartz substrate because of its piezoelectric and triboluminescence responses.

We can draw two sets of conclusions from the series of experiments presented here. The first set has a technological character. We have experimentally demonstrated that the nonlinear response of electrically connected optical gap antennas can be externally amplified or attenuated by a control voltage. The modulation occurs without the addition of nonlinear polymers and relies on the intrinsic metal nonlinearity. This type of electrical control, demonstrated here with Au nonlinear photoluminescence (and to a lesser extend with the second-harmonic generation), opens new avenues for actuating and reconfiguring more complex device architectures such as plasmonic logic gates for instance. The second set of conclusions addresses the physical origin of the voltage-controlled nonlinear photoluminescence yield. We propose a scenario by which the local electron temperature at the metal interface is modified by a sheet of polarization charges created by the static electric field. By changing the surface density of electrons to about $10^{-2}$ C m$^{-2}$, the temperature of the electronic distribution can be adjusted by $\sim$ 50 K. This approach can be combined with designs optimized for generating non-uniform electron temperature in order to bring additional agility. The present finding focuses on the response of a bow-tie shape, combining a large N-PL yield overlapping spatially with the region where the electrical potential is dropped. These conditions may be found in other geometries.

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Supporting Information Available: fabrication protocol, power dependence of the nonlinear emission, simplified model of metal heating by a short laser pulse, perturbation theory, peak electron temperature at the contact surface. This material is available free of charge via the Internet at http://pubs.acs.org.

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