

Plasmonic hot-carrier extraction: mechanisms of electron emission

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Abstract

When plasmonic nanoparticles are coupled with semiconductors, highly energetic hot carriers can be extracted from the metal–semiconductor interface for various applications in light energy conversion. Hot charge–carrier extraction upon plasmon decay using such interface has been argued to occur after the formation of an intermediate electron population with an uniform momentum distribution. The efficiency of the charge separation process is thus discussed to be limited by this spatial homogeneity in certain plasmon-induced applications. Here we demonstrate using visible pump,

near-infrared probe transient absorption spectroscopy, that increases in the contact area between metal and semiconductor leads to an increase in the quantum yield for hot electron injection that are not consistent with a homogeneous energy-momentum distribution of hot-electrons. Instead, further analysis of the experimental data suggests that the highly energetic electrons are emitted across the interface via a surface photoemission mechanism.

Nanoscale heterogeneous structures consisting of metal-semiconductor junctions have great potential in applications related to photonic energy conversion. When plasmonic metal nanostructures are in direct contact with a semiconductor, a Schottky barrier can be formed at the metal-semiconductor interface² [Figure 1(A)]. Hot charge carriers that result from non-radiative plasmon decay can be emitted across this interface, leading to charge-separated states that are useful for various applications, including photodetectors,³ photovoltaics⁴ and photochemistry.⁵ The reported collection efficiency of these electrons is low, typically $< 1\%$.^{3,4,6-9} However, Schottky photodetectors are valuable due to their fast response and low photon-energy detection capability.¹ Furthermore, the operational lifetime of photocatalytic Schottky devices surpasses that of more efficient semiconductors.⁷

It has been postulated that non-radiative plasmon relaxation results in a transient population of energetic electrons with an uniform spatial distribution of momenta.^{10,11} This spatial uniformity limits the efficiency of charge-separation in metal-semiconductor junctions.¹² Typically, the metal-semiconductor interface consists of a single, planar junction^{3,4} [Figure 1(B)] where only those electrons that simultaneously move toward the single interface and have sufficient kinetic energies to overcome potential energy barriers can be emitted (shaded area in the momentum-space sphere shown in Figure 1(B)). Simplified models predict that, for moderate electron energies, only a small fraction of the total hot-electron population resulting from plasmon decay meet these criteria, leading to poor quantum yields for hot-electron extraction.¹³

Other limiting factors for hot-electron extraction stem from properties intrinsic to metals,

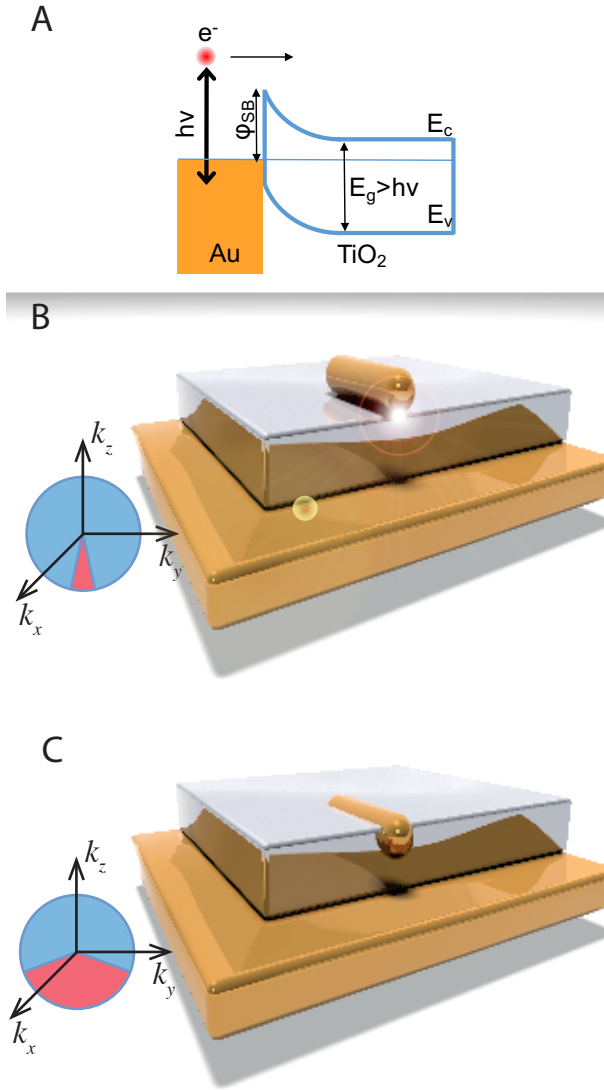


Figure 1: **Partially-embedded Au nanorods.** (A) Band diagram of a metal–semiconductor contact for the extraction of plasmonic hot–charge carriers. E_g , E_c and E_v are the bandgap, conduction band and valence band of the semiconductor material respectively. φ_{SB} is the Schottky barrier.¹ Absorption of a photon of energy $h\nu$ ($<E_g$) and non–radiative relaxation could lead to electron injection from the metal to the semiconductor. (B) Metal–on–semiconductor structure: On momentum space, there is a narrow escape cone for electron injection. (C) When the metal nanostructure is partially embedded, the number of interfaces available for electron injection increases.

including ultra-fast carrier-carrier relaxation pathways¹⁴ and broad energy distributions of hot-carriers, which stem from an almost continuous density of electronic energy states that exist below the Fermi level of the metal.¹⁰ These limitations can be overcome via a direct charge-transfer mechanism¹⁵ which can take place when the metallic nanostructure and the semiconductor material have a strong electronic interaction,¹⁶ a situation that has led to quantum yields for charge separation of $> 20\%$.¹⁵ Another approach to increase the charge-separation quantum yield is to increase the contact area between metal and semiconductor [Figure 1(C)].⁹ This in principle allows for a larger fraction of the excited hot electron population to be extracted into the semiconductor material (shaded area in the momentum-space sphere shown in Figure 1(C)). Whilst this concept was demonstrated in the past by Knight et al.⁹, their approach is of limited scalability due to the use of complex top-down nanofabrication techniques. Furthermore, top-down fabrication approaches set a lower-limit to the minimum size of the metal nanostructures (> 20 nm), which in turn set the optical response of the nanostructures to the NIR region of the spectrum. This implies a low electrochemical potential of the charge-separated state limiting the range of applications in hot-electron photo-redox catalysis.¹⁷ An important point to note is that full coverage of the plasmonic nanostructures by the semiconductor could hinder hot-carrier extraction.

Here we demonstrate that increases in the metal-semiconductor contact area result in increased yields for electron injection that are not consistent with an intermediate spatially-isotropic distribution of electron momenta. We achieve the increased metal-semiconductor contact area by means of a simple, large scale and low cost fabrication process, using nanocrystalline Au rods as the building blocks. The electron injection yields are estimated from ultra-fast pump-probe spectroscopy measurements, which demonstrate a $2.7\times$ increase in the quantum yield for electron injection with contact area. This increase is consistent with a surface photoemission mechanism.

In Figure 2 we show a schematic of the “lift-off” process we have developed to create partially embedded metal-semiconductor structures over macroscopic areas. Firstly (step

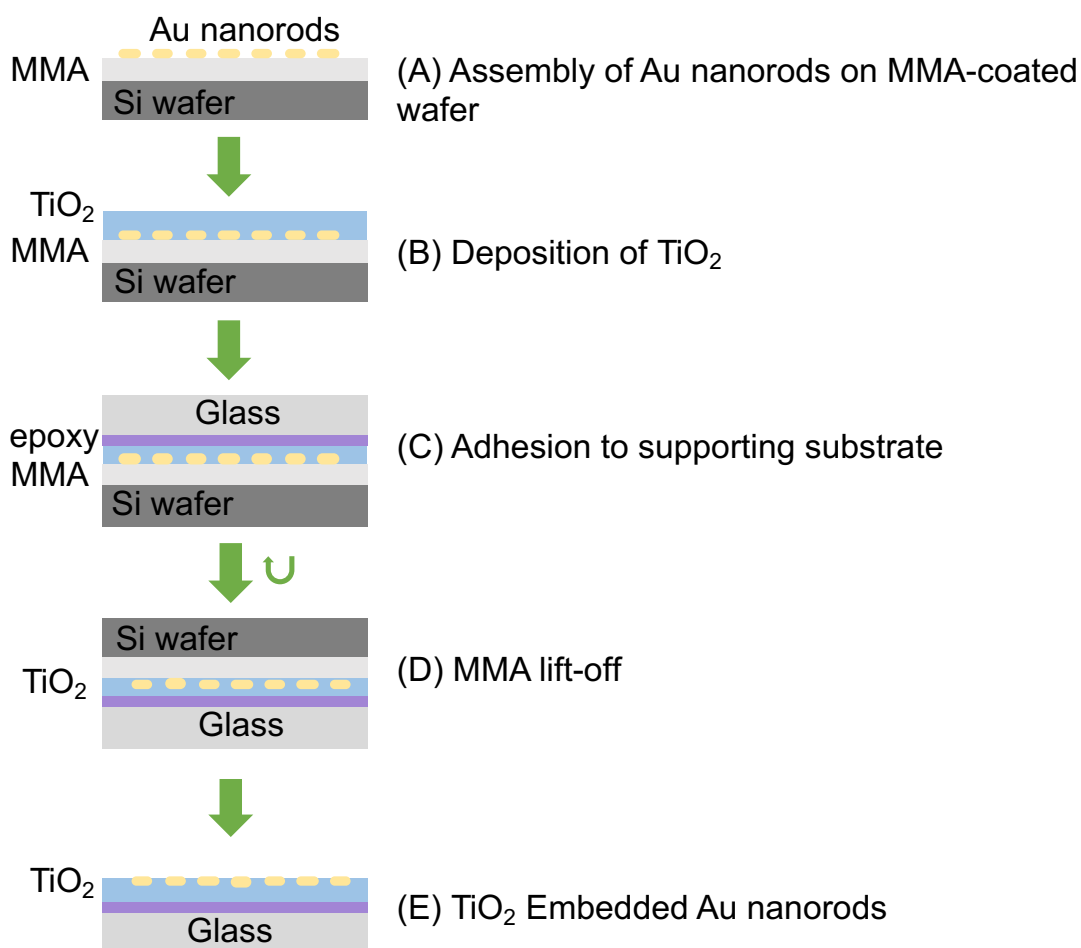


Figure 2: **Nanofabrication.** (A) Methyl methacrylate (MMA) is spin-coated on a Si wafer and Au nanorods are self-assembled on the MMA layer. (B) Deposition of 50 nm TiO₂ via electron beam evaporation. (C) The sample is adhered to a glass substrate via an epoxy layer and subsequently in step (D) it is immersed (upside-down) in acetone to remove the MMA layer and release the structure from Si wafer to form the final (E) partially embedded structure.

A, Figure 2), a thin layer of methyl methacrylate (MMA, an acetone-soluble material) is spin-coated onto a cleaned Si wafer and Au nanorods are self-assembled (by means of electrostatic interactions) on the surface of MMA by immersing the MMA/Si wafer in an aqueous dispersion containing Au nanorod colloid (more details in the methods section). After the Au nanorods are self-assembled onto the MMA/Si wafer, a layer of titanium dioxide (TiO₂, 50 nm) is deposited onto the structure via a standard electron beam evaporation process (step B, Figure 2). The thickness of this layer must exceed the diameter of the Au nanorods (estimated to be ~ 20 nm, supporting information figure S1) in order to ensure almost complete coverage of the nanostructures. At this point, more layers could be added, including a mirror, which would result in the creation of metal-semiconductor-metal structures that can exhibit increased light absorption¹⁸⁻²⁰ (supporting information section S2). An epoxy adhesion layer is then used to bond the complete structure to a glass substrate (step C, Figure 2). Next (step D, Figure 2), the sample is immersed in acetone for 3 hours to dissolve the MMA layer and release the Si wafer, resulting in structures where Au nanorods are partially buried within TiO₂ (step E, Figure 2). For the experiments to be described below, we have also prepared a reference sample consisting of *non-embedded* Au nanorods on TiO₂. In this case, the self-assembly of Au nanorods is performed on the surface of TiO₂ after the acetone lift-off step (step E, Figure 2).

Figure 3(A) and (B) show surface topography images obtained with Atomic Force Microscopy (AFM) for the resulting embedded and non-embedded structures. In Figure 3(A), the Au nanorods are embedded within the TiO₂ and the AFM topography image consists of a random collection of pits denoting the location where the Au are embedded. In fact, the nanorods are shown to be buried ~ 2 nm below the TiO₂ surface (supporting information figure S2). On the contrary, the surface topography of Figure 3(B) clearly reveals the geometrical outline of the Au nanorods that rest on top of the surface of the TiO₂ film. Figures 3(C) and (D) show scanning electron microscopy images of both kinds of structures [3(C):embedded, and 3(D): non-embedded] where the presence of Au nanorods on both sam-

ples is clearly visible. Together, the images of Figure 3 indicate that the process described in Figure 2 leads to structures where Au nanorods are partially buried inside TiO₂.

Figure 3(E) shows the measured optical extinction [defined as $-\ln(T)$, T is the transmission spectrum] for the structures of Figure 3, along with the measured absorption spectrum for the Au nanorods in solution. The spectrum in solution is dominated by a single band that corresponds to the longitudinal localised surface plasmon resonance of the Au nanorods. The extinction spectra of the Au nanorods when placed on top and when partially buried inside a thin film of TiO₂ show redshifts, which are attributed to the presence of the high refractive index of the TiO₂. Numerical full wave simulations were carried out to identify the observed extinction bands, as shown in Figure 3(F). In figure 3(F) we show the calculated absorption spectra for idealised structures consisting of spherically-capped Au nanorods resting on top of a thin film with refractive index 2.4, and for a similar structure embedded to a variable extent in the film. By immersing the Au nanorod in the semiconductor, the localised surface plasmon resonances (both longitudinal and transverse) redshift in proportion to the amount of surface coverage.

Following Landau (non-radiative) damping of surface plasmon resonances, the resulting hot-electrons have a finite probability of injection into the semiconductor material.¹⁰ Once injected, these carriers rapidly relax to the lower states of the conduction band of the semiconductor, and can thus absorb both visible and near-infrared light enabling the detection of these charge-transfer processes through visible pump near-IR probe spectroscopy, as we describe next.^{21,22}

As shown in the diagram of Figure 4(A), we photo-excited the samples using ultra-short light pulses spectrally tuned to the localised surface plasmon resonance of the Au nanorods, which according to the data of Figure 3(E), was 634 nm for the non-embedded structures and 732 nm for the embedded structures. These wavelengths are significantly below the band-gap of TiO₂ and consequently, no direct excitation of electron-hole pairs in this material is possible. After non-radiative relaxation of the localised surface plasmon resonances, injection

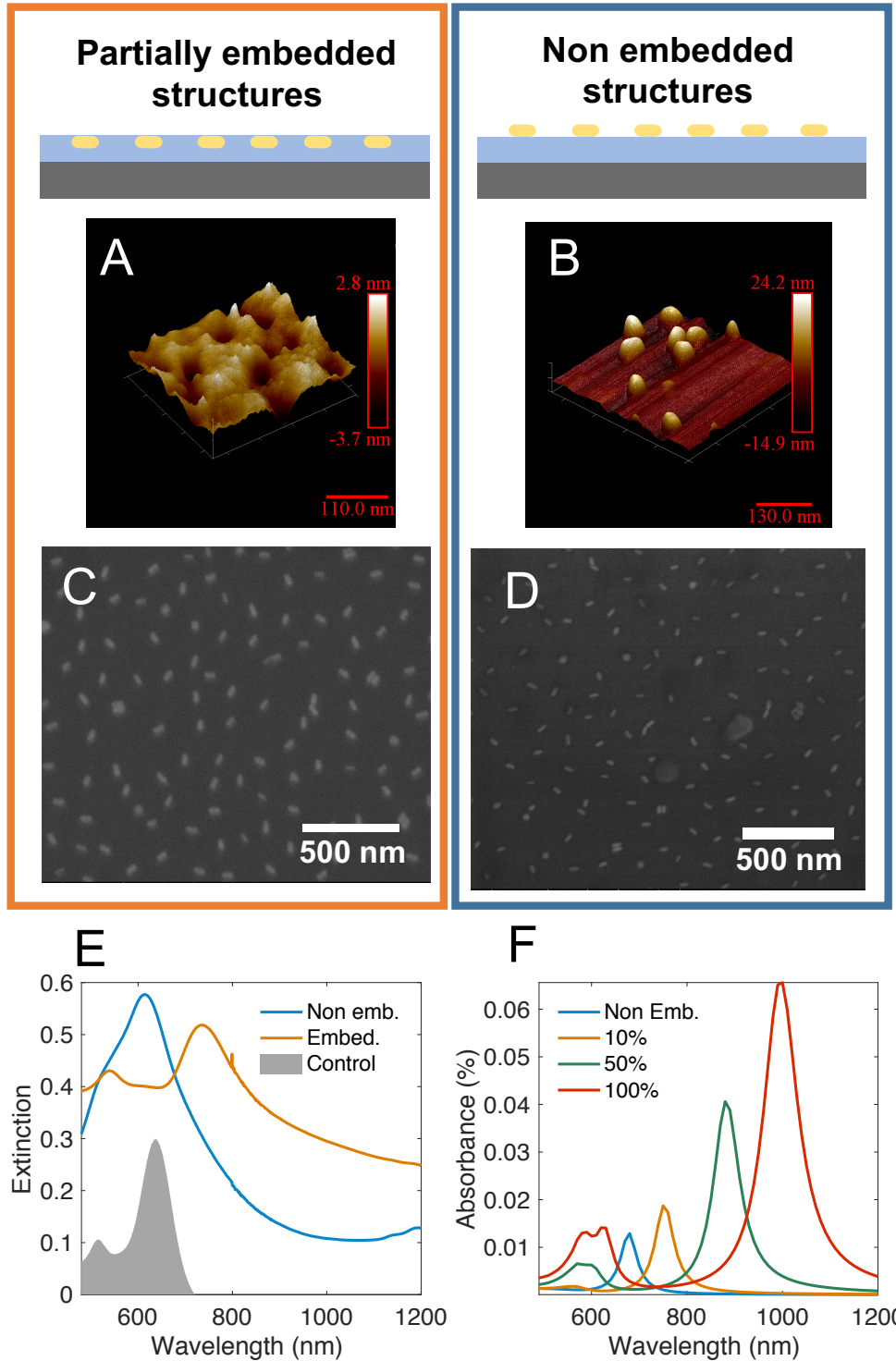


Figure 3: **Structural characterisation and Optical properties.** (A) and (B) are atomic force microscopy topography images of the embedded and non-embedded structures respectively. (C) and (D) are the corresponding scanning electron images. (E) Measured extinction $[-\ln(T)]$ spectra of the embedded, non-embedded structures along with the absorption spectrum of the Au nanorods in solution. (F) Calculated absorption spectra using a full-wave numerical solution of Maxwell equations.

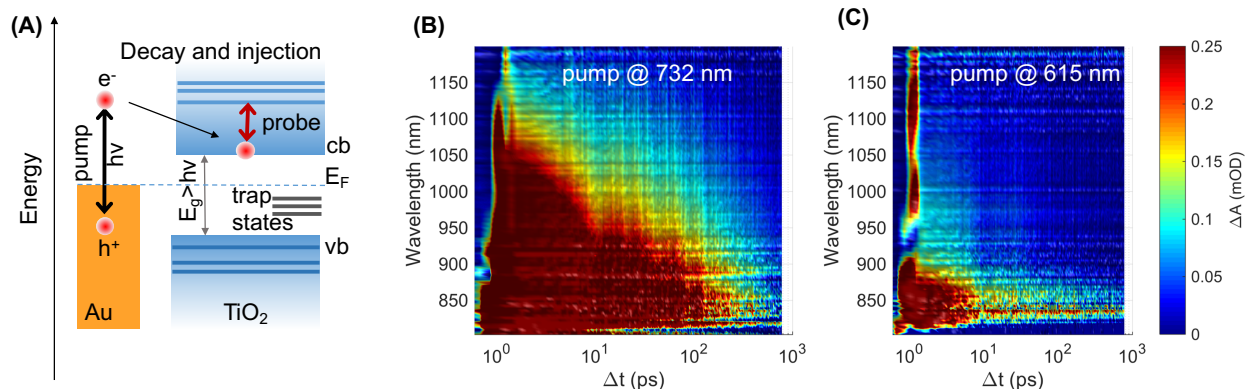


Figure 4: **Ultra-fast pump probe spectroscopy.** (A) Pump pulses excite localised surface plasmon resonances in the Au nanorods which, after non-radiative decay, can result in the injection of hot-electrons into the conduction band (cb) of TiO_2 . Intraband excitation of these electrons is possible with NIR probe pulses. Also shown are the relevant energy levels of the system: E_F Fermi level of the metal, vb: valence band and E_g : semiconductor bandgap. (B) and (C) show the transient absorption ΔA spectra as a function of probe wavelength and pump-probe delay time (Δt) for the (B) partially and the (C) non-embedded structures. The pump wavelengths are indicated in the figures.

of electrons into the TiO_2 can take place, leading to a transient population of electrons at the bottom of the conduction band of the semiconductor [Figure 4(A)]. These electrons can absorb near-infrared light (via intraband electronic transitions) and are thus detectable through the changes in the absorption of time-delayed near-IR light probe pulses.²²

Figure 4 (B) and (C) show the results of these measurements as a function of probe wavelength and pump-probe time delay. In general, the measured transient absorption (ΔA) spectra indicate excited state absorption ($\Delta A > 0$), where the amplitude of the measured signals are larger for the samples where the Au nanorods are partially embedded in TiO_2 . In the time domain, the detected signals consist of a fast rise (sub-picosecond time-scale) and a rapid decay ($\Delta t < 5$ ps) which is followed by a slower, multi-exponential decay ($\Delta t > 5$ ps). It has been documented that the electron population in TiO_2 decreases with time due to an interplay of processes such as: recombination with the positive holes present in Au,²³ surface electron trapping ($\Delta t < 200$ fs, Figure 4 shows a representation of such states)²⁴ and charge transfer to ambient oxygen molecules ($10^{-6} - 10^{-3}$ s),²⁵ processes with marked differences in time scales which can account for the multi-rate decay. The transient absorption

changes measured for partially embedded Au nanorods in metal–semiconductor–nanorod structures (Figure S4), exhibit damped oscillations with a strong Fourier component at ~ 20 GHz (supporting information sections S2 and S3). We assign this oscillation frequency to mechanical (breathing mode) oscillations of the nanorods. These oscillations are part of the non–radiative decay channels available to localised surface plasmon resonances in the Au nanorods,¹⁴ and occur in competition with electron injection across the Au–TiO₂ interface.

In Figure 5(A), we show a direct comparison of the transient absorption changes for Au nanorods that are partially embedded, non–embedded and a control sample where the nanorods are deposited directly on a glass substrate. For the rods on a glass substrate, no transient signal is observed, which indicates that no electron injection events take place, a result consistent with the expectation that little or no electron–accepting states exist in glass. On the contrary, for the other two cases there is a non–negligible transient signal, which is largest (in magnitude) for the partially embedded samples. We argue that this difference in signal magnitude is largely due to a higher probability of electron injection afforded by the larger number–density of Au–TiO₂ interfaces in the partially embedded structures. We now estimate the changes in hot–electron injection quantum yield using the data derived from the transient absorption spectroscopy. In the small signal limit, the magnitude of the transient absorption signal (that originates from the electrons residing in the semiconductor) is proportional to the number of photons absorbed by the metal nanostructure.^{8,15} The proportionality constant is given by a product of the absorption cross–section and the photon–to–injected–electron quantum yield (supporting information section S4). As shown in the supporting information section (Figure S8), the magnitude of the measured transient absorption increases linearly with pump power, indicating that the experiments were carried out in the small signal limit. We can therefore quantify the increase in hot–electron injection quantum yield R_{QY} , by comparing the magnitude of the measured transient absorption signals for the embedded and non–embedded samples (taking

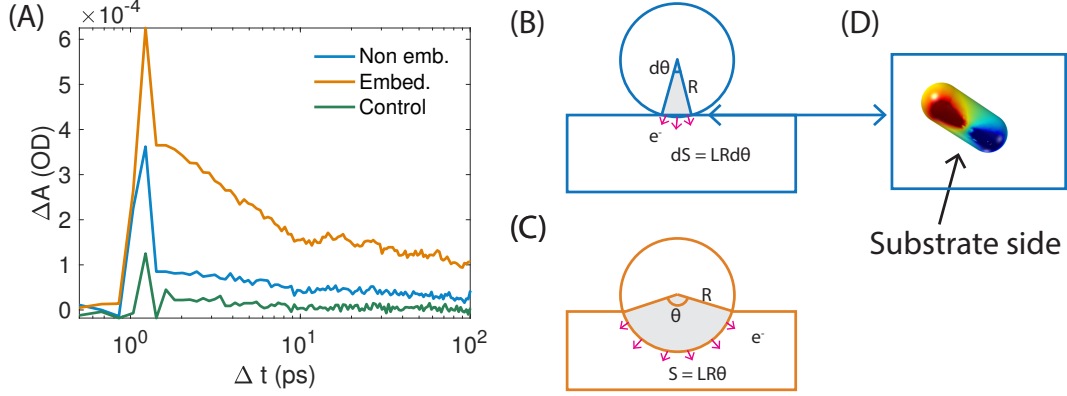


Figure 5: **Amplitude of transient absorption signal.** (A) Time evolution of the transient absorption ΔA signal measured at 1000 nm for samples consisting of Au nanorods on glass, on TiO_2 and partially-embedded in TiO_2 . Also shown are diagrams of the idealised cross-section of cylindrical Au nanorods of radius R and length L (B) on TiO_2 and (C) partially embedded in it. The arrows indicate interface normal vectors. In (B), the Au– TiO_2 contact area is dS , whereas this area increases to S in the case shown in (C). (D) Map of the electric field for a nanorod placed on top of a TiO_2 substrate. The colours indicate the relative sign of the field.

into account the existing differences in the amount of absorbed pump photons):^{8,15}

$$R_{QY} = \frac{\left(\frac{\Delta A_{emb,732nm}(1000nm)}{(1-10^{-OD})j(732nm)} \right)}{\left(\frac{\Delta A_{non-emb,615nm}(1000nm)}{(1-10^{-OD})j(615nm)} \right)}, \quad (1)$$

where $\Delta A_{x,pump}(probe)$ is the measured peak amplitude of the transient change in absorbance at the *probe* wavelength, for sample x , irradiated with *pump* wavelength (these values were averaged between 0.2 and 0.4 ps). j is the incident photon flux and OD the measured optical density for absorption of the pump pulses. Using this equation with the data of Figure 4, we obtain a value of $R_{QY} = 2.7$ indicating that the method outlined in Figure 2 leads to structures with increased hot-electron-injection.

Having estimated the increase in injection quantum yield, we now discuss if the observed value of R_{QY} is consistent with an increase in the metal–semiconductor contact area resulting from partially embedding the nanorods. R_{QY} can increase with contact area if the population of hot-electrons resulting from plasmon relaxation possesses an isotropic energy–momentum distribution, for in this case the probability for electron injection is partly determined by

the flux of these charge carriers through the metal–semiconductor contact.^{26,27} In Figure 5(B)-(C), we show idealised cross–sections of a cylindrical nanorod (of radius R and length L) placed either on (B) top or (C) embedded in a semiconductor material. Within this idealised geometry, the ratio of contact areas R_A is given by $R_A = S/dS = \theta/d\theta$, where θ is an angle describing the fraction of the cylinder making contact with the semiconductor. Given that $d\theta \ll \theta$, R_A can take values that can significantly exceed the estimated ratio of injection quantum yields R_{QY} . Possible reasons for the discrepancy include: (1) the organic molecules that decorate the Au nanorods hinder the charge transfer events,²⁸ (2) poor crystallinity of the TiO_2 , leading to trap states²⁹ [Figure 4(A)] which quickly trap the injected carriers, (3) non–conformal coating of the surface of the Au nanorods with TiO_2 during physical vapour deposition, or (4) the assumption of spatial invariance of the energy–momentum distribution of hot–electrons is not valid, and this distribution is instead non–uniform.^{9,30,31} Points (1)-(3) highlight some of the limitations of our nanofabrication approach. In particular, it is known that an intense annealing process is required to improve the material properties of the semiconductor,³² but this process will likely melt the Au nanorods. A possible explanation for point (4) is that electron injection into TiO_2 occurs via a *surface mechanism*:³³ a directional emission process wherein an electron moving towards the metal–semiconductor interface absorbs the energy of a photon and thus overcomes the potential energy barrier that exists at the interface.

In the surface photoemission mechanism, the rate of electron injection into the semiconductor is proportional to the (square of the absolute value of) electric field normal to the metal–semiconductor interface (supplementary information section S6). Spatial maps of these fields, shown in Figure 5(D), clearly demonstrate their spatially inhomogeneous nature, and in particular, the strong confinement of electromagnetic fields that exists at the metal–semiconductor interface. This strong localisation, according to the surface photoemission mechanism, implies that increases in the surface coverage of the nanostructure would lead to enhancements in the quantum yield of hot–electron injection that are smaller than

the corresponding increase in surface contact area. In fact, our calculations show that for the geometry of our nanostructures, the largest enhancement is expected to be in the order of $3.26\times$, which is of a similar order of magnitude to our experimental results. This puts forward evidence indicating that the non-radiative plasmon decay and subsequent electron injection into the semiconductor does not occur via the formation of an intermediate hot electron gas with an isotropic momentum distribution, but instead, the energy of the coherent oscillation of charge density (localised surface plasmon resonance) can be transferred to a charge carrier moving to the interface, a directional process.

It is important to note that an estimate of the injection quantum yield for each structure could be made by comparing the measured ΔA signals to those obtained by directly populating the TiO_2 conduction band via transient absorption measurements using pump pulses with energies above the bandgap of the semiconductor³⁴ ($E_g \approx 3.2$ eV for TiO_2 ³⁵).¹⁵ Our instrument is not capable of producing these pump pulses.

In summary, we have shown that the increase in metal-semiconductor contact area lead to a higher efficiency of hot-electron injection into the semiconductor. Detailed analysis of our data suggests that surface photoemission is likely to be the dominant mechanism for hot-electron injection. In order to increase the metal-semiconductor contact area, we have developed a fabrication process that only requires physical vapour deposition techniques and does not involve any complex processes such as electron beam lithography or reactive ion etching, making it highly attractive for large scale fabrication. In particular, we envisage that contrary to other approaches,^{36,37} our method can enable the construction of near-perfect absorbers of light that incorporate partially-buried metal nanostructures.¹⁸ We also envisage that these substrates can be used as active elements in novel sensors,³⁸ photo-detectors³⁹ or in plasmonic photochemistry.^{5,35,40-45}

Supporting Information Available

Experimental section This material is available free of charge via the Internet at <http://pubs.acs.org/>.

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Graphical TOC Entry

