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Experimental Evidence for Non-Thermal Contributions to Plasmon-Enhanced Electrochemical Oxidation Reactions

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Abstract: Photocatalysis based on plasmonic nanoparticles has emerged as a promising approach to facilitate light-driven reactions under far milder conditions than thermal catalysis. Several effects, such as strong local electromagnetic fields, increased electron and lattice temperatures, or the transfer of non-thermal charge carriers could contribute to the reaction rate enhancement. In order to understand plasmon-enhanced catalysis and to enable plasmonic platforms, a distinction between the different underlying effects is required. We investigate the electrochemical model reactions oxidative hydroxide adsorption and glucose oxidation and deconvolve the enhancement processes via their dependence on excitation wavelength. We observe that non-thermal effects contribute significantly to the plasmonic enhancement.

Introduction
Plasmonic nanoparticles (NPs) are currently intensively discussed as an enhancing agent for light-driven heterogeneous catalytic conversions.1–6 The possibility of photo-generated carriers initiating the reaction might lower its operating temperature, which is usually a problem in terms of catalyst stability, high required thermal input and reaction selectivity. Thus, an operating temperature reduction is desirable.7 In a pioneering work, Christopher et al. observed a strong increase of the catalytic reaction rate on plasmonic silver nanoparticles upon illumination with visible light and a scaling of the rate enhancement with the incident optical power.8 The authors proposed a combination of electromagnetic field enhancement and extracted electrons to be responsible for the enhancement. However, their early model did not account for the currently discussed specifics of the plasmon excitation.9–14 Looking at gold nanoparticles (AuNPs) as plasmonic model system, different processes could contribute to the reaction rate enhancement (cf. Figure 1):

1) During the coherent drive of the plasmon, light is focused into strongly enhanced, localized electric fields around the NPs.15 These strong fields could affect the energies of molecular (anti-)bonding states e.g. of adsorbates involved in the reaction, as for example suggested for the dissociation of O2.16

With the subsequent plasmon decay, several processes occur that could also lead to reaction rate enhancements:9,17–20

2) Landau damping of the plasmon results in a non-equilibrium, non-thermal carrier distribution, i.e. electrons and holes are excited to states far from the Fermi level. These could enhance the reaction via a direct21 or indirect22–26 carrier transfer between the NPs and the adsorbate-binding molecular orbitals.2,4,25,27,28
3) Carrier-carrier scattering leads to an equalization of the non-thermal carrier distribution, resulting in a bath of thermalized excited electrons in equilibrium. Within 10-1000 fs these could also populate electronic states in the adsorbate molecule.\textsuperscript{18}

4) Electron-phonon coupling occurs on picosecond timescales and turns the thermalized carrier bath into local heat (phonons)\textsuperscript{19,20}. The elevated NP temperatures could enhance a phonon-driven chemical reaction that occurs at or close to the surface.

![Figure 1: Left: Schemes of the different possible reaction rate enhancing processes. Middle: Expected wavelength-dependencies for the reaction rate enhancement \( r \). Right: Expected wavelength-dependent rate enhancements after normalization to the absorption (Abs).](image)

To optimize plasmon-assisted catalysis towards higher reaction rates, it is important to know which of the above processes is the most relevant. The relative importance is currently a matter of debate\textsuperscript{29}. A dominance of purely thermal effects was proposed, ruling out the injection of non-thermal carriers\textsuperscript{30,31}. Other studies report the influence of both, thermal and electronic enhancements\textsuperscript{32-35}. For example, Zhou et al.\textsuperscript{34} measured the photocatalyst’s surface (local) temperature using an infrared thermometer during ammonia decomposition and concluded that the thermal contribution caused by the determined heating cannot solely explain the observed enhancement in the product formation rate. Consequently, a recent study by Zhan et al.\textsuperscript{36} employs excitation-power and wavelength-dependent measurements to disentangle thermal and non-thermal effects in electrochemical reactions. Christopher et al. proposed a similar methodology for studying the plasmonic reaction enhancement\textsuperscript{8}. An approach by Li et al. employed two different illumination geometries on a metal-semiconductor catalyst for carbon dioxide methanation. The comparison of the indirect excitation (a barrier layer absorbs all light so that the metal
below can only generate a thermal enhancement) with the direct (no barrier layer so that all enhancement mechanisms can occur on the metal side) excitation of the plasmon pointed towards significant non-thermal contributions.\textsuperscript{37}

The possible contributing processes have different dependencies on the excitation conditions (cf. Figure 1):

**Enhanced near fields**

The local fields around AuNPs are strongly enhanced when exciting in resonance with the plasmon.\textsuperscript{15} The excitation-wavelength dependence of the near field enhancement does in general not track the absorption, which is known as near-field far-field shift.\textsuperscript{38} In the case of small AuNPs, interband transitions lead to a decrease in local field enhancement at short wavelengths and an increase in absorption.\textsuperscript{15,39} On the other hand, the absorption drops faster towards the near-infrared than the field enhancement, because it scales linearly with the excitation energy (apart from the plasmon resonance), while the field enhancement does not.\textsuperscript{15} Figure 2 displays a model calculation for small AuNPs. The field enhancement normalized to the absorption possesses an excitation-wavelength dependence. In case the population of the reactants’ molecular (anti-)bonding orbitals promoted by the local field would be the main contribution to the reaction rate enhancement, the normalization to the absorbed optical power would lead to no clear resonance, but to a rate increase towards longer wavelengths (Figure 2, inset).

The complicated dependencies of the electronic structure of adsorbed molecules makes the scaling of the reaction rate enhancement with incident power hard to define. However, several studies report surface-enhanced Raman scattering as field driven process to scale super linear with power.\textsuperscript{40–42}

![Figure 2](image-url): Calculated absorption cross section and field enhancement, averaged over the surface of a 12 nm AuNP using Mie theory. The refractive index of the surrounding medium was set to $n = 1.6$ to account for the ITO substrate and the molecules surrounding the gold nanoparticle. An analytic model from Ref.\textsuperscript{43} was used for the dielectric function of gold. Inset: Average near field intensity enhancement, normalized to the calculated absorption cross section.

**Non-thermalized hot carriers**
Landau damping is a momentum-conserving process that allows donating the full photon energy into a single intraband transition, resulting in non-thermal electrons and holes.\textsuperscript{12} In this case, the exact carrier distribution depends on the excitation wavelength.\textsuperscript{14,44} Absorbed low-energy photons result in electrons excited close to the Fermi edge. Photons with higher energies can excite non-thermal electrons far from the Fermi edge.\textsuperscript{14,26,45} Even higher photon energies can excite interband transitions from the d-band into states above the Fermi level.\textsuperscript{13} For the case that non-thermalized hot electrons are the main contribution to the reaction rate enhancement, a spectrum of the reaction rate enhancement normalized to the absorbed number of photons would show a peak in the spectral range where hot electrons are generated most effectively.

**Thermalized hot carriers**
The lifetimes of non-equilibrium carriers is very short (100 fs) and their excess energy is quickly redistributed within the electron gas.\textsuperscript{12,13} As the estimated heat capacity of the electron gas in AuNPs is a linear quantity in the temperature range up to 3000 K and independent of the excitation-mechanism,\textsuperscript{11} the temperature of the thermalized electron gas only depends on the amount of absorbed optical energy. Consequently, for thermalized hot carriers as the main contribution to the reaction rate enhancement, a normalization to the absorption would result in a vanishing wavelength dependence.

Both, non-thermalized and thermalized hot carrier-based enhancements would lead to a linear increase of the reaction rate with absorbed optical power due to the linearity between photon absorption and carrier generation.

**Local heating**
The heat capacity of AuNPs does not differ significantly from bulk Au. The rise in temperature thus depends linearly on the total amount of absorbed optical energy.\textsuperscript{10,11,46} If temperature was the main reaction rate enhancing contribution, the wavelength dependence of the rate enhancement would reproduce the absorption spectrum. Similar to the case of thermalized hot carriers, the reaction rate enhancement, normalized to the absorbed optical power would not show any clear dependence on the excitation.

When assuming the rate of a chemical reaction scaling in an Arrhenius type with the temperature, the reaction rate would increase exponentially with increasing incident optical power for sole thermally enhanced processes.

To quantify the excitation-wavelength dependence experimentally, a reaction with a directly observable rate is required. When regarding the charge flow between the plasmonic metal and its surrounding as an electrical current that occurs due to illumination under defined conditions, one can directly relate this current to the number of carriers contributing to the reaction’s enhancement. This suggests an electrochemical approach in which the plasmonic metal represents the working electrode under investigation. The charges extracted from/into the adjacent phase (electrolyte) are compensated at a counter electrode delivering an externally measurable current. This approach enables to screen an arbitrary reaction (cathodic, i.e. extraction of electrons from the plasmonic metal working electrode, or anodic, the extraction from or compensation of holes in the metal) by applying the respective necessary electrochemical potential between this working electrode and an inert reference electrode (i.e. at which no charge transfer reactions occur). In this setup we used AuNPs immobilized on an optically passive and electrochemically inert electrode (In-doped SnO\textsubscript{2}, ITO) to compare the reaction current under dark with that under defined illumination conditions. The difference of both over time is averaged giving the photocurrent \(I_{\text{ph}}\), a quantitative measure of the steady-state photo-enhancement of the reaction rate. As model reactions with broad relevance we screened the alkaline adsorption of hydroxide on the Au surface and – upon addition of glucose to the electrolyte – the oxidation of glucose.\textsuperscript{23,47-50} We varied power and wavelength of the optical excitation to disentangle thermal and non-thermal influences.
We observed linear dependencies of the reaction rates on the incident optical power, whereas its effectivity (slope of $I_{ph}$ vs. absorbed power) strongly varies with the excitation wavelength, showing a maximum at the plasmon resonance. The normalization to the absorbed optical power allows us to conclude that the main contribution to the reaction rate enhancement are non-thermal carriers and not thermalized electrons, elevated NP temperatures or strong local fields.

**Results & Discussion**

AuNPs were synthesized with a mean diameter of 12.3 ± 0.8 nm (see Supporting Information Figure S1) and immobilized on an ITO-covered glass slide to prepare AuNP/ITO electrodes. As apparent in the scanning electron microscopy (SEM) images (Figures 3a-b), AuNPs cover the ITO surface with negligible agglomerate formation. SEM investigations (see Supporting Information Figure S2) reveal no relevant AuNP detachment from the substrate after the electrochemical experiments.

![Figure 3: (a,b) SEM images of the AuNP-ITO electrode surface. (c) Cyclic voltammogram in glucose-containing (500 mM) and glucose-free base electrolyte (500 mM KOH). The current was normalized to the AuNP electrochemically active surface area (ECSA, see Supporting Information Figure S3) to give the current density $j$. Scan rate: 50 mV/s. Arrows indicate the scan directions.](image)

The electrodes were used for the glucose oxidation in alkaline environment ($pH = 14$).

In alkaline environment, alcohols (as which we consider glucose) deprotonate to alkoxydes which are then oxidized on a surface decorated with oxide species.\textsuperscript{47,51,52} The latter result from hydroxide adsorption according to:

$$\text{Au} + \text{OH}^-_{\text{aq}} \rightarrow \text{Au} - \text{OH}_{\text{ads}} + e^- \quad (1)$$

which occurs at potentials similar to the oxidation potential of glucose. Glucose oxidation to gluconic acid then occurs via a one-electron transfer between the metal and the electrolyte:\textsuperscript{48}

$$\text{C}_6\text{H}_{12}\text{O}_6 + \text{Au} - \text{OH}_{\text{ads}} \rightarrow \text{Au} - \text{C}_6\text{H}_{12}\text{O}_7_{\text{ads}} + \text{H}^+ + e^- \quad (2)$$

Recently, Wang et al. demonstrated the photo-enhancement of alkaline glucose oxidation on AuNPs.\textsuperscript{23} A broad range of parameters that affect the photo enhancement of the reaction were reported, i.e. the incident light wavelength, the power of the light source and the $pH$ of the electrolyte solution.

The current between the AuNP/ITO electrode and the counter electrode serves as a metric to quantify the extent of potential and illumination-dependent charge transfer. The required potential range for glucose oxidation was identified by cyclic voltammetry (CV) measurements under dark conditions (Figure 3c). The voltammogram (red curve) shows a broad oxidation peak at approximately +300 mV upon anodic potential ($E$) increase, which is attributed to the oxidation of glucose to gluconic acid...
according to Eq. (2). Upon further potential increase, the peak slowly decays due to a saturation of the surface with adsorbing gluconic acid species (Au – C₆H₁₂O₇,ads) and further increases at potential ranges where higher oxides form according to:

\[ \text{Au} - \text{OH}_{\text{ads}} + \text{OH}^{-}_{\text{aq}} \rightarrow \text{Au} - \text{O}_{\text{ads}} + \text{H}_2\text{O} + e^- \]  \hspace{1cm} (3)

and the oxidation of the initially non-covered Au sites:

\[ 2\text{Au} + 6\text{OH}^-_{\text{aq}} \rightarrow \text{Au}_2\text{O}_3 + 3\text{H}_2\text{O} + 6e^- \]  \hspace{1cm} (4)

In the cathodic scan, i.e. after scan reversal, a second oxidation peak at approximately +100 mV is observed, which can also be attributed to the adsorption-controlled glucose oxidation onset in coincidence with the onset of stripping of gluconic acid and/or oxygen species from the oxide-covered Au (at +200 mV, according to the reverse of Eq.s (1), (3) and (4)). The observed behavior is equal to that observed earlier for AuNP/ITO electrodes. This hydroxide adsorption was screened in a separate experiment using a glucose-free electrolyte. The corresponding curve in Figure 3c shows the hydroxide adsorption onset during the anodic scan at approximately 0 mV and occurring throughout the entire anodic potential range, originating from the oxide formation-stripping behavior of Au in alkaline conditions. The subsequent scan reversal leads to a slight oxide stripping peak at +50 mV followed by the reduction of dissolved oxygen residuals.

To assess the plasmon-based reaction enhancement, we carried out chronoamperometry measurements (i.e. a potential step from the open-circuit potential to a potential in the glucose oxidation and simultaneous Au surface oxidation regime as identified by CV under dark conditions; see Supporting Information Figures S4A-B for the corresponding current transients under dark and illumination). All transients show a current decrease in the first seconds of the experiments, which we attribute to the transport of electroactive species to the electrode and capacitive charging upon switch-on. At later times of the experiment (after 30 s) the currents were stable. To obtain the illumination-related current transient (photocurrent \( I_{\text{ph}}(t) \)), we subtracted the dark current from that of the illuminated electrode. \( I_{\text{ph}}(t) \) was averaged within the time interval [30; 60 s] to obtain the steady state photocurrent \( I_{\text{ph}} \) (cf. Supporting Information Figures S4C-D). Figure 4 displays \( I_{\text{ph}} \) for different applied potentials \( E \) and in different electrolyte solutions. We observe a small photocurrent over the entire \( E \) range in the (glucose-free) base electrolyte whereas the photocurrent steadily increases when glucose is present. Relative to the dark current, the current enhancement in the base electrolyte is large at lower potentials, i.e. where the hydroxide monolayer adsorption sets on (due to the normalization to small dark currents, the relative enhancement is apparently high at potentials before this onset although photocurrents are in fact low) and decreases with increasing potential. In contrast, the current enhancement has a clear maximum at \( E = +300 \text{ mV} \) when glucose is present (coinciding with the CV peak potential, Figure 3c). We conclude from this similarity of the CV peak with the relative \( I_{\text{ph}} \) peak that the photo-enhancement of the glucose oxidation rate is proportional to its dark oxidation rate. Although an electrode potential shift of a few mV due to illumination has been reported, the effect is not expected to lead to the observed large current increase and is furthermore immediately compensated by the potentiotstat. Therefore, it can be neglected. We thus focused on +300 mV as the working potential. We assume that in case of the glucose-containing electrolyte at this potential both reactions (hydroxide adsorption by Eq. (1) and glucose oxidation according to Eq. (2)) occur whereas only the first proceeds in the glucose-free electrolyte.
Figure 4: Photoresponse of AuNP/ITO electrodes glucose oxidation and hydroxide adsorption. Absolute (left ordinate, dashed lines) and relative (ratio of $I_{ph}$ to dark current $I_{dark}$, right ordinate, straight lines) photocurrents vs. applied potential $E$, excited at 454 nm with 35.7 mW in the glucose-containing (red) and the glucose-free base electrolyte (black). Error bars represent the standard deviation within the steady-state duration of the experiment.

$I_{ph}$ versus illumination wavelengths $\lambda$ from 454 – 802 nm is displayed in Figure 5a.

Figure 5: (a) Photocurrent $I_{ph}$ vs. excitation wavelength $\lambda$ for both electrolytes at $E = +300$ mV. The measured values are normalized to the incident optical power (cf. Supporting Information Figure S5 and Figure 1, central column). (b) $I_{ph}$ vs. incident optical power $P$, at different excitation wavelengths. The data correspond to the situation displayed in the central column of Figure 1. As expected from the preceding discussion, we observe a strong dependence of $I_{ph}$ on $\lambda$ with a pronounced maximum around the plasmon resonance in the glucose oxidation case. $I_{ph}$ for increasing incident optical power at different excitation wavelengths is displayed in Figure 5b. $I_{ph}$ scales linearly with the power in all cases. These two observations alone would apply to all potential reaction rate-enhancing processes.
$I_{\text{ph}}$ scales linearly with the reaction rate enhancement via the number of electrons excited via absorption. The normalization of $I_{\text{ph}}$ to the optical absorption of the electrode then corresponds to the situations displayed in the right column of Figure 1, as discussed above. To measure the absorption, micro-absorption spectroscopy was applied (see dashed line in Figure 6a). The experiment accounts for the reflected and transmitted light (see Supporting Information Figure S6). The spectrum of $I_{\text{ph}}$ normalized with the absorbed power ($P_{\text{abs}} = P \cdot \text{Absorption}$) (Figure 6a) clearly displays a non-vanishing dependence on the excitation wavelength, for both electrolytes showing a maximum around $\lambda = 590$ nm. We find higher rate-enhancements in the glucose-containing solution than in the base electrolyte by a factor of 1.7. According to Eq. (2), glucose oxidation first demands for an adsorbed OH species (i.e. one charge to be transferred). The oxidation of glucose then demands for another charge. We can deduce that the enhancement refers to both reactions (1) and (2), while deconvolving both appears rather speculative due to different adsorbate coverages in both cases. The retaining wavelength dependence of the normalized $I_{\text{ph}}$ (as a measure for the reaction rate enhancement) provides a strong indication that non-thermal effects contribute significantly to the enhancement of both oxidation reactions (apparently, the photoexcitation enhancement of the glucose oxidation occurs via a similar mechanism as the hydroxide adsorption, whereas the enhancement is stronger for glucose oxidation). As discussed earlier, thermal effects would vanish after normalization to the absorption and local fields would show a more linear dependence in the observed wavelength regime. $I_{\text{ph}}$ is independent from the duration of illumination (see Supporting Information Figures S4C-D), i.e. from any heat uptake. The slope of the power dependence of $I_{\text{ph}}$ (Figure 6b) normalized to the absorbed optical power $P_{\text{abs}}$ represents the effectivity of the reaction rate-enhancing process. It is a measure for the reaction events per absorbed photon energy. Thermal processes would show a non-linear behavior with increasing power (see discussion above). This is observed in none of the regarded cases. For non-thermal carriers, the slope is expected to be linear and larger for wavelengths where non-thermal carriers are excited effectively. Following the present understanding of the plasmon decay, this is what we observe: Landau damping of the plasmon is effectively a plasmon–electron scattering process, where the energy of a
single plasmon quantum generates an electron–hole pair.\textsuperscript{13} Plasmon excitation
thus effectively generate non-thermal carriers, where the excess energy scales with the photon energy.\textsuperscript{22,56,57} Both of these two important and exemplary reactions seem to be enhanced by the injection of non-thermal carriers resulting from the plasmon decay, but each with different sensitivity. Therein, the plasmonic enhancement of hydroxide adsorption, being a prerequisite to many alkaline oxidation reactions, further emphasizes the potential of hot carrier injection for e.g. oxygen evolution or green fuel (e.g. methanol) formation reactions. In order to achieve an even more efficient enhancement by hot charge carriers, further studies addressing the generation of non-equilibrium carrier distributions, as well as their lifetimes and diffusion are important.\textsuperscript{12,58} Furthermore, the observed difference in reaction enhancement between the oxidation and the hydroxide adsorption points towards complex and convolved dependencies.

**Conclusions**

In this work, we investigated the plasmon-enhanced electrochemical glucose oxidation under hydroxide adsorption on AuNPs that are attached to an ITO electrode in an alkaline electrolyte. The photocurrent, i.e. the difference between illuminated and dark current in the steady state, is suitable to quantify a steady state reaction enhancement under minimization of capacitive effects. We estimated the wavelength-dependent photocurrent to separate possible reaction rate enhancing processes. When normalized to the absorbed power, the reaction rate enhancement shows a distinct wavelength dependence with a maximum coinciding with the plasmon resonance. This is not expected neither for thermal nor for processes supported by local fields. Thus, non-thermal carriers seem to be most relevant for the plasmon-assisted conversion enhancement.
Materials and Methods.

Chemicals. Tetrachloroauroic(III) acid (≥99.9% trace metal basis), trisodium citrate dihydrate (≥99.0%), 11-mercaptoundecanoic acid (95%) were purchased from Sigma-Aldrich; ethanol absolute (100.0%) was obtained from VWR. Ethylenediaminetetraacetic acid tetrasodium salt hydrate (EDTA) was obtained from Merck. Glucose was purchased from Sigma and used as received. KCl was purchased from Nanjing Chemical Reagent Co., Ltd. (Nanjing, China). ITO-covered glass (surface resistivity 8-12 Ω/cm² was purchased from Sigma-Aldrich. All aqueous solutions were prepared with Millipore water (resistivity of 18.2 MΩ · cm). HClO₄ and KOH were purchased from Carl Roth, Germany.

AuNP synthesis. Citrate-stabilized AuNPs were synthesized and characterized as published earlier. α-Methoxypoly(ethylene glycol)-κ-(11-mercaptoundecanoate) (PEGMUA, molecular mass: 2 kDa) was synthesized as described previously. AuNPs were functionalized by straightforward ligand exchange of citrate-stabilized AuNPs using a mixture of PEGMUA and MUA in aqueous solution as reported. A PEGMUA/MUA mixture was prepared by mixing different volumes of an aqueous solution of PEGMUA (1 mM) and an ethanolic solution of MUA (1 mM), yielding a mixture with the molar PEGMUA/MUA ratios of 1:3. A solution of PEGMUA/MUA-AuNPs with a final concentration of AuNP of 10 nM was prepared.

AuNPs characterization. Transmission Electron Microscopy (TEM) measurements were performed using a Jeol JEM-1011 instrument operating at 100 kV. For TEM sample preparation, 10 μL of sample solution was drop-casted onto a carbon-coated copper grid, which was placed on a glass slide, and left to dry for at least 24 h. Quantitative analyses of AuNP size distributions based on TEM measurements were conducted using ImageJ.

AuNPs immobilization on ITO substrate. AuNPs were modified on an ITO slide for cyclic voltammetry measurements. Before modification, the ITO substrate with a surface area of 1.44 cm² (1.2 cm x 1.2 cm) was cleaned under ultrasonic agitation subsequently in acetone, isopropanol and water for 15 min each. The slide was thoroughly dried with nitrogen gas, and treated with a PSD-UV4 O₂ plasma cleaner system (Novascan 203 Technologies) for 30 s to further remove organic contaminants from the surface and make the glass hydrophilic. The cleaned slide was then immersed into a freshly prepared diluted acidic solution (pH = 2) of AuNPs for 72 h. The obtained AuNPs/ITO electrode was rinsed with water to clean from non-immobilized NPs, and dried with nitrogen gas. The electrochemically active surface area of this electrode was determined via the capacitance ratio method with 6.69 cm² (see Supporting Information Figure S3, this area corresponds to a geometrical electrode area of 1.44 cm²) and served to normalize the current observed CV experiments.

AuNPs/ITO electrode characterization. AuNPs/ITO characterization was performed by Scanning Electron Microscopy (SEM) using a Leo Gemini 1530 microscope.

Micro-absorption measurements. The optical setup for micro-absorption measurements was described recently. Briefly, a supercontinuum laser (Fianium, SC-400-4, 450–2400 nm) was used as a light source. The light was guided through a linear polarizer and a beam splitter (Thorlabs, BSW26R) into an inverted microscope (Olympus, IX71). A Leica HCX PL Fluotar 100× objective with a numerical aperture (NA) of 0.9 was used to focus the light onto the sample with a laser power <100 μW. The AuNPs/ITO sample was mounted to a motorized xy-translation stage. The position of the laser spot on
the sample was visible through a microscope camera. The transmitted light intensity \( T \) was collected with an Olympus Mplan FL N BD 100x objective with an NA of 0.9 and guided by a fiber (Ocean Optics, QP600-2-UV-BX for 450–950 nm and BIF600-VIS-NIR for 700–1200 nm) to a spectrometer (Avantes, Avaspec 3648). The reflected light intensity \( R \) was separated from the incoming light with a second beam splitter (Thorlabs, BSW26R) and detected by a spectrometer (same fibers and spectrometer as for the transmitted light). The absorbed light intensity \( A \) was determined by subtracting \( T \) and \( R \) contributions from the source light intensity.

**Electrochemical and photoelectrochemical measurements.** Electrochemical and photoelectrochemical measurements were performed using a ZenniumX potentiostat and a photoelectrochemical cell by Zahner, Germany, equipped with a Pt wire counter electrode. AuNP/ITO electrodes were contacted with Au wire using Ag paste (Plano, Germany) and nail polish for electrical insulation. ECSA of all samples was determined by using the capacitance ratio method\(^6^3\) i.e. performing cyclic voltammetry (CV) cycles under varied scan rates of 2, 5 and 10 mVs\(^{-1}\) in a potential range of ±50 mV around the open-circuit potential (OCP) of the sample in 1 M HClO\(_4\) vs. a Ag/AgCl reference electrode (Figure S2-A). The capacitance (obtained from the averaged slope of the anodic and cathodic charging currents at OCP vs. scan rate) was scaled with the area-specific capacitance (40 μF cm\(^{-2}\))\(^6^3,6^4\) to obtain the ECSA (Figure S2-B). The current in CV (Figure 3c) was normalized to the AuNP ECSA determined with 6.69 cm\(^2\) to give the current density \( j \). For (photo-)electrochemical tests the cell (reference electrode was changed to a Ag wire due to the change in pH from acidic to alkaline) was first filled with freshly degassed 500 mM glucose solution + 500 mM KOH aqueous solution. CV experiments were performed under dark conditions for three successive cycles at 50 mV s\(^{-1}\) scan rate. Subsequent current transients were recorded for 60 s under dark and illuminated conditions in the following order: 1) for varied potentials under 454 nm illumination (15.3 mW, supplied by a tunable light source TLS03 by Zahner, Germany, illuminating the full sample area), 2) for varied intensities at an externally applied potential \( E = +300 \text{ mV vs. Ag at the same wavelength and 3) for varied wavelengths and intensities according to the maximum available emission spectrum of the TLS03 (see Figure S5) at } E = +300 \text{ mV. After this procedure, the electrolyte was removed, the electrode cleaned thoroughly with water and steps 1) - 3) were repeated under freshly degassed 500 mM KOH solution. In addition, in repeated experiments with fresh substrates, no deviations of the experiment results were observed. In control experiments, the ITO substrate neither contributes to the observed dark nor to the photocurrent transients (see Supporting Information Figure S7) and was therefore regarded as passive.

**References**


Graphical Abstract: