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Deterministic radiative coupling between plasmonic nanoantennas and semiconducting nanowire quantum dots

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Abstract

We report on the deterministic coupling between single semiconducting nanowire quantum dots emitting in visible and plasmonic Au nanoantennas. Both systems are separately and carefully characterized through micro-photoluminescence and cathodoluminescence. A two-step realignment process using cathodoluminescence allows for electron-beam lithography of Au antennas near individual nanowire quantum dots with a precision of 50 nm. A complete set of optical properties was measured before and after antenna fabrication. They evidence both an increase of the nanowire absorption, and an improvement of the quantum dot emission rate up to a factor of two in presence of the antenna.

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(Some figures may appear in colour only in the online journal)

1. Introduction

Recent progress in semiconductor growth research now allows for the fabrication of nanowire (NW) structures which are of great interest owing to their high crystalline quality, strain-free character and practical geometry. A key step towards the realization of nano-optical circuits and applications relies on the coupling of single NWs with other structures, like photonic crystal cavities \cite{1} or plasmonic nanoantennas (NAs) \cite{2–5}. To better use the advantages and the versatility of NWs, it is now necessary to adapt previous studies on self-assembled or colloidal quantum dots (QDs) to their NW counterparts. While numerous results have been obtained in the coupling of self-assembled \cite{6–10} and colloidal \cite{11, 12} QDs with plasmonic nanostructures, all the existing studies on control of the optical properties of nanowire quantum dots (NWQDs) rely on a photonic approach, using the nanowire itself as an antenna \cite{13–15}. Following previous work in our group \cite{8} on droplet epitaxy QDs, we show how combining cathodoluminescence (CL) with a standard e-beam lithography technique allows us to fabricate, at will, plasmonic NAs in the vicinity of II-VI NWQDs emitting around 620 nm. Both systems are initially fully characterized using CL for the antennas and micro-photoluminescence (μPL) as well as time-resolved spectroscopy for the emitters. Previous studies have made use of morphological criteria to detect nanoemitters, using scanning electron microscopy (SEM) or atomic force microscopy \cite{7}. In contrast, our method has the advantage of allowing localization of embedded structures which are undetectable with the previous methods.

2. Nanowire quantum dot properties

Our nanoemitters are single (Cd,Mn)Te QDs (Mn fraction \(\approx 5\%\)) inserted inside ZnTe/(Zn,Mg)Te core/shell nanowires (figure 1(a)). The NWs are grown by molecular beam epitaxy...
on a (111)B GaAs substrate covered by a 500 nm thick buffer layer of ZnTe [16, 17]. Dewetted gold droplets are used as catalysts, and temperature variations allow us to favor different growth mechanisms resulting in the final heterostructure. The core growth is both longitudinal and lateral, leading to the conical aspect of the nanowire, with a core diameter of 20 nm at the top to ≈200 nm at the base, and a 20 nm thick shell. The QD emission is measured by confocal μPL spectroscopy. It features a main emission peak around 620 nm, blue-shifted by 150 nm compared to the exciton transition in bulk CdTe due to the confinement and the strain induced by the surrounding shell [19]. It is significantly broadened (FWHM 10 nm) by the presence of the magnetic Mn atoms creating a fluctuating magnetic field inside the QD, randomly shifting the exciton line in time by Zeeman effect [20] (figure 1(b)). Amongst all the studied NWs, the central emission wavelength ranges from 605 to 630 nm.

NWs are detached from their growth substrate and dispersed onto a host substrate for optical studies. The host is a Si substrate pre-patterned by optical lithography and dry etching to fabricate coarse localization marks. It is covered by a 100 nm Au layer deposited by e-gun evaporation and a 250 nm thick Al₂O₃ spacing layer by atomic layer deposition. The mirror and the spacer thickness are designed to give constructive interferences between the light directly emitted from the QD and the light reflected by the mirror in order to maximize luminescence collection. NWs are finally dispersed by mechanical contact between the host and growth substrates.

A first selection of NWs is performed using CL at 5 K [8, 16]. The luminescence from the QD is spectrally filtered and detected by an avalanche photodiode (APD). CL imaging of the emission profile at the QD luminescence wavelength is obtained for a set of NWs. To avoid spurious emission from excitons trapped in defects at the base of the NWs, we conserve only structures presenting a well-localized emission in the top half of the conical NW and having the typical spectral structure of figure 1(b). The selected QDs are then fully characterized by μPL spectroscopy at 5 K. The NWs are excited at 447 nm by a frequency doubled, ps-pulsed Ti: Sapphire laser. Time-resolved measurements are performed at very low pumping power, well below saturation of the QD, with a spectral integration bandwidth of 2 nm to ensure that we detect only the excitonic transition, resulting in a monoexponential time trace. They show decay times ranging from 0.2 to 1.18 ns (figure 1(c)). The average lifetime (\(\tau = 0.68\) ns) is about five times longer than for self-assembled CdTe/ZnTe QDs [21]. In addition, the emission from the QD is linearly polarized in a direction parallel to the NW axis in 70% of cases, and orthogonal in all the others. The average degree of linear polarization is 0.7 ± 0.2. These optical properties can be explained by the very small energy difference between the valence bands of CdTe and ZnTe [20]. The resulting nature of the hole ground state strongly depends on additional energy shifts induced by strain or QD aspect ratio [19, 22]. In our case, the orbital hole wavefunction is probably poorly confined inside the QD, resulting in low electron–hole wavefunction overlap and thus long radiative decay times. We note that the measured decay rate 1/\(\tau = \gamma_r + \gamma_n\), where \(\gamma_r\) (resp. \(\gamma_n\)) is the radiative (resp. non-radiative) decay rate, might be limited by non-radiative processes. Furthermore, a large variability from dot to dot strongly mixes the light- and heavy-hole bands, explaining the polarization results. This variability is not detrimental to our study as we compare a complete set of optical properties on the one and same NW before and after antenna fabrication.

### 3. Nanoantenna characterization

Au nanoantennas are fabricated on the host substrate, in a region empty of NWs. They consist in single rectangles of 70 nm fixed width, as sketched in figure 2(a). Their length \(L\) varies from 85 to 140 nm. They are fabricated by electron-beam lithography on a 200 nm thick Poly(methyl methacrylate) (PMMA) bi-layer resist, using PMMA with molecular weights of 50 kDa and 950 kDa to create a mechanical mask with the resist. 35 nm of Au is deposited on the sample by e-gun evaporation. N-methyl-2-pyrrolidone (NMP) lift-off with 80 °C heating is finally performed to remove the resist.

The antennas are characterized using CL at room temperature [23, 24]. The CL spectrum of each antenna is obtained by raster-scanning the electron beam over its surface. It presents a large peak at a fixed wavelength around 580 nm and a second resonance at lower energy. CL spectra for all the antennas are presented in figure 2(b), revealing that the second resonance red-shifts with increasing antenna length. We note that the optimization of the Al₂O₃ spacer thickness in order to increase the collected signal at 620 nm results in destructive interferences for the collected light in the 700 to 800 nm wavelength range, decreasing the detection contrast in this spectral region. Further information is obtained by imaging the plasmon local density of states (LDOS) [23]. It is

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**Figure 1.** (a) Nanowire morphology: an Au droplet serves as a catalyst to grow a CdTe quantum dot inserted in a ZnTe conical nanowire. A lateral regrowth of ZnMgTe creates a shell around this core. (b) Typical μPL spectrum of a nanowire QD and (c) corresponding decay rate measurement, integrating a bandwidth of 10 nm around 622 nm. A monoexponential fit (red line) gives a lifetime \(\tau = 1.18 \pm 0.01\) ns.
Figure 2. (a) Sketch of the NW–NA geometry (left: side view, right: top view), to scale. (b) Cathodoluminescence spectra of the rectangular nanoantennas, with Gaussian fits of the main component (red dashes) and the dipolar mode blue dashed-dotted line discussed in the text. The green line corresponds to (e). The blue line with squares indicates the fitted central wavelength of the dipolar resonance. The blue and brown shaded areas correspond to the integrated bandwidth in (c) (blue) and (d) (brown) respectively. (c) and (d) LDOS imaging of the 111 nm long antenna around 700 nm and 600 nm respectively, with 100 nm spectral integration bandwidth. The blue and brown rectangles correspond to the antenna boundary. (e) Cut profile of the CL intensity along $\lambda_{QD} = 620$ nm (green solid line in (b)) showing an increase of the measured LDOS for an antenna length of 97 nm due to the onset of the dipolar mode.

reconstructed by slowly scanning the electron beam over the antenna and collecting the CL emission as a function of the antenna length around 95 nm due to the contribution of the dipolar mode. It also predicts an enhancement for lengths above 140 nm. This enhancement comes from the onset of a higher order mode theoretically predicted to appear at this length [25, 26]. We note however that for the target coupled NW–NA system, the presence of the very high refractive index nanowire ($n_{ZnTe} \approx 3$) in the near-field of the antenna significantly modifies its plasmonic properties. Hence a red-shift in the plasmon dispersion relation curve is expected, and the enhancement peaks observed in figure 2(e) will occur for smaller antenna lengths.

4. Hybrid NW–NA structures

4.1. Fabrication

Two examples of the target structure are shown in figures 3(d)–(e). Fine alignment marks are first fabricated around the chosen NWs using electron-beam lithography, metal evaporation and lift-off (figure 3(a)). We then record a SEM image (figure 3(b)) together with the CL intensity image at $\lambda_{QD} = 620$ nm (figure 3(c)). Both images are acquired after aligning the microscope beam using the fine alignment marks. Acquisition times are limited to a few seconds to limit mechanical and electrostatic drifts. We fit the QD emission peak with a bi-dimensional Gaussian profile and superimpose it to the corresponding SEM image. It allows us to precisely localize the QD inside the NW in the frame defined by the fine alignment marks. NAs are then fabricated with the same process as the previously characterized antennas. Before insulation of the resist, a final alignment step is performed onto the fine marks under the resist. The antennas have a fixed width $w = 70$ nm and a variable length $L$ from 50 to 140 nm. This range is based on the previous CL results and takes into account the shift due to the presence of the NW. The antenna sides of length $L$ are parallel to the measured polarization direction of the QD emission. We aim at having an antenna to NW gap equal to zero. SEM images reveal an average gap of 12 nm ± 50 nm, thus some antennas are on top of the NW. The final error of 50 nm has different sources. Alignment of the lithography setup has a typical error of 20 nm but is degraded in our case due to the poor contrast in the images of the fine alignment marks under the resist. We evaluate the error coming from the thermal, mechanical and electrostatic drifts during CL at low temperature to be 30 nm. We note that the NW core and shell are already thick (≈50 nm at the QD position). The resulting QD to antenna distance ranges from 50 to 110 nm. This distance is always large enough to prevent luminescence quenching, and the variations due to the positioning error only have a moderate effect on the NA to QD coupling [28].
4.2. Photoluminescence observations

The coupled QD–NA systems are characterized using μPL spectroscopy with a pulsed excitation laser at a wavelength of 447 nm. Figure 4(a) compares the integrated intensity emitted from a representative QD as a function of the exciting laser power before and after antenna fabrication for $L = 63$ nm. The PL intensity first increases with the excitation power and then saturates because of the complete occupation of the discrete excitonic state in the QD. For a same excitation power one clearly sees a much higher collected signal in presence of the NA. Similar measurements on other QDs (present on the same substrate) which have experienced the same process except the final antenna fabrication shows no change in their properties. Previous studies revealed modification of the luminescence collection by redirection of light by the antenna [5, 6, 8]. We have carried out Fourier plane microscopy by imaging the back focal plane of the microscope objective in the μPL setup (see supplementary material) to observe the radiation diagram of the NWQDs [29], and we did not observe a significant change in the radiation pattern after antenna fabrication. Clearly, the higher PL intensity we measure is not due to the redirection of light by the antennas. Additional information is provided by measurements of the exciton lifetime before ($\tau_0$) and after ($\tau_{NA}$) antenna fabrication under the same excitation conditions. We observe no significant change, with an average ratio $\tau_{NA}/\tau_0 = 1.1 \pm 0.2$ (see supplementary materials).

To better analyze the effect of the antennas, we evaluate the saturation intensity $I_s$ at high excitation power, and the slope at the origin $g$ of the power-dependent μPL curve. The
ratio of $I_s$ before and after antenna fabrication is plotted as a function of antenna length in figure 4(b), showing a net increase after NA fabrication. As shown in figure 4(c), we also observe a greater increase in the slope $g$ after NA fabrication.

4.3. Discussion

Assuming that we only detect the excitonic transition and considering that the nanowire excitation is done with a pulsed laser, the saturation intensity $I_s$ is equal to $f \times \gamma_{v}/(\gamma_{a} + \gamma_{nr}) \times \alpha_{coll}$, where $f$ is the laser repetition rate, $Y = \gamma_{a}/(\gamma_{a} + \gamma_{nr})$ is the quantum yield and $\alpha_{coll}$ is the fraction of the emitted power collected by the microscope objective. As we measure no change in the radiation diagram after NA fabrication, it is reasonable to assume that $\alpha_{coll}$ is unaffected.

As a consequence the enhancement observed in figure 4(b) is directly related to an increase in the quantum yield $Y$. This enhancement is found to be moderate for antennas with a length greater than 60 nm but stronger for the smallest antennas, up to a factor of 2.5. As expected from section 3, this occurs for antennas that are shorter than the determined resonant length from figure 2(e). Moreover since the QD response at saturation is independent of the excitation power, the change of $I_s$ after antenna fabrication is not due to an increase of the absorption inside the NW.

Nevertheless, CL spectra of the NAs at $\lambda_{exc} = 447$ nm show that there is still a measurable LDOS at this energy that can modify the absorption of the excitation laser inside the NW (see supplementary material). The main contribution to this LDOS comes from the fixed resonance observed in figure 2 and does not depend on the antenna length. The change of the absorption can be retrieved by analyzing the slope at the origin $g$. In the limit of low excitation regime, $g$ is expected to be proportional to $Y \times \alpha_{abs} \times \alpha_{coll}$, where $\alpha_{abs}$ is the fraction of power absorbed in the NW. As $\alpha_{coll}$ remains unchanged after NA fabrication, the ratio $(g/I_{0}/\lambda_{0})$ directly gives the change in $\alpha_{abs}$. The comparison of figures 4(b) and (c) shows that the absorption in the NW is enhanced by a factor of $2.2 \pm 1.1$ over the whole antenna length range. Further illustration of the respective changes in $I_s$ and $g$ are presented in the supplementary material with two other power-dependent PL curves. Our main conclusion is that, for the shorter antenna lengths, enhanced NW absorption and improvement of the quantum yield contribute equally to the four-fold increase of $g$ that we observe.

Finally, lifetime measurements show that there is no change of the total decay rate $\gamma_{a} + \gamma_{nr}$. This is an indication that the total decay rate is dominated by the non-radiative term $\gamma_{nr}$ and is moderately affected by a change in $\gamma_{a}$. Therefore, the observed increase in $Y$ is essentially due to an increase of the QD radiative rate $\gamma_{a}$ because of coupling to the NA.

5. Conclusion

In conclusion, we demonstrated deterministic coupling between single semiconducting nanowire quantum dots and plasmonic nanoantennas using CL and electron-beam lithography with a precision of 50 nm. Our method has the advantage of relying only on the luminescence of the emitters for their precise localization. It also grants full characterization of individual nanoemitters and antennas. CL spectroscopy and LDOS imaging of individual NAs is demonstrated as a powerful technique to experimentally determine antenna parameters for the fabrication of coupled plasmonic-semiconductor emitters. Furthermore, we demonstrate two effects of the NAs on the QDs: an absorption enhancement by a factor of 2, and a light emission enhancement due to radiative coupling to the antenna up to a factor of 2.5 in a spectral region of high plasmonic loss, extending the control of light emission from semiconducting nanostructures towards the visible spectral region. The effect could be greatly increased for a smaller QD-antenna distance, i.e. using a thinner NW shell. Implementation of the method is a crucial step towards fabricating more complex and versatile coupled structures. It can be applied to all kinds of nanoemitters, aiming at controlling their optical properties such as their polarization response [4, 10] or emission diagram [6, 9].

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