Up-conversion luminescence coupled to plasmonic gold nanorods for light harvesting and hydrogen production

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The conversion of infra-red to visible-light, the so called “up-conversion luminescence”, and its use for H\textsubscript{2} production allowing for a larger fraction of sun light on earth to be used is one of the most important light harvesting aims. We have prepared, characterized and tested a tri-functional material composed of an up-converter (Na\textsubscript{3}YF\textsubscript{5}:Er\textsuperscript{3+},Tm\textsuperscript{3+}), plasmonic gold nanorods (Au NRs), and a CdS semiconductor. Tests with gas phase methanol/O\textsubscript{2} resulted in the production of H\textsubscript{2} and CO\textsubscript{2} upon excitation with a 980 nm IR light. Equally important, these products were observed (although with a third of the value obtained by the ternary system) for the binary system (Na\textsubscript{3}YF\textsubscript{5}:Er\textsuperscript{3+},Tm\textsuperscript{3+}/Au NRs); without the CdS semiconductor. This is the first time that H\textsubscript{2} was made photo-catalytically using plasmonic Au nanoparticles upon initial excitation with infra-red light and is attributed to the up-conversion process.

The low light flux and weak energy density of sunlight on earth (ca. 1 kW/m\textsuperscript{2}), half of which is composed of infra-red light, are two limiting factors affecting the progress of solar-driven processes. While considerable progress has been made in photovoltaic solar cells, their still relatively high cost makes them non-competitive, compared to fossil fuel, for energy intensive systems (such as those used in chemical and transport industries). Photo-catalytic materials are to date far less efficient than photovoltaics and are thus unpractical for energy harvesting. Many limiting factors are behind this lack of progress. Most active semiconductor materials are either unstable in water\textsuperscript{\textsuperscript{1,2,3}} or do not possess the electronic band edge requirements\textsuperscript{4} for the redox reaction needed for water splitting. Actually, because of thermodynamics, most of stable semiconductors have large band gaps (typically > 3 eV), such as TiO\textsubscript{2}, SrTiO\textsubscript{3} and GaN limiting their use to specific small scale applications, which cannot compete with fossil fuel based processes\textsuperscript{5}. Over the last two decades, considerable efforts were undertaken to address these; such as anion doping to decrease the wide band gap of semiconductors\textsuperscript{6,7,8}, multi-phase semiconductors\textsuperscript{9,10,11,12}, plasmonic metals to improve light harvesting\textsuperscript{13,14,15} and specific 3D-arhctecture to increase light harvesting and decrease charge carriers’ recombination rates\textsuperscript{6,17,18,19}, with limited success. One promising alternative to address the wide band gap semiconductors issue is to change the light frequency rather than changing the materials energy gap and thus allow their use, since they are stable in water, for H\textsubscript{2} generation. In that regard, up-conversion (UC) luminescence is attractive. However, the very weak efficiency of this system makes this process so far of marginal importance. A second requirement is thus needed and it is light amplification. A possible approach for that is the use of a plasmonic system poised to enhance the local electric field by many orders of magnitudes\textsuperscript{20} at the interface of the UC material, thus in theory an efficient ternary (UC-plasmonic-catalyst material) system can be made. UC luminescence is the sequential absorption of two or more photons and can take place through a number of pathways, which have been discussed in various reviews and articles\textsuperscript{21-27}. Since Auzel and co-workers\textsuperscript{28} reported their pioneering work on lanthanide-based UC processes most materials studied are based on lanthanide ions, Ln\textsuperscript{3+}, such as Erbium (Er\textsuperscript{3+}), Holium (Ho\textsuperscript{3+}) and Thulium (Tm\textsuperscript{3+}) in an inorganic host containing Ytterbium (Yb\textsuperscript{3+}).

In this work, we have used an inorganic host material, Na\textsubscript{3}YF\textsubscript{5}, in which Yb cations are dispersed: Na\textsubscript{3}YF\textsubscript{5}:Yb. This results in light absorption at 980 nm which when doped with Tm cations an up-converter emission takes place at difference wavelengths. The absorbance at 980 nm and the emissions, need to be tuned with the semiconductor band gap as well as with the plasmon resonance energy, to fulfil the compositional and structural requirements. We have prepared the up-conversion material (Na\textsubscript{3}YF\textsubscript{5}:Yb, Tm) with Tm at. % =1.67 % (with respect to Y and Yb) (figures 1 and S1A). Figures S1B and S1C show the response of the UC material to light frequency. We have prepared a few materials with different Tm%; the one reported here gave the best performance.

Figure 1A presents the NaY(Yb)F\textsubscript{4}:Tm which is composed of individual crystallitles of about 150-250 nm in size. EDX analyses show the presence of F, Na, Yb and Y (the Cu signal comes from the TEM grid); The crystalline nature of the NaY(Yb)F\textsubscript{4} is shown in Figure 1B. Au/CdS/NaY(Yb)F\textsubscript{4}:Tm is presented in Figures 1C-1F. The large particles correspond to NaY(Yb)F\textsubscript{4}, whereas the smaller ones are CdS. In addition, Au nanorods are also present in contact with CdS nanoparticles. There exists an intimate contact between all the phases present. Figures 1C and 1D correspond to two images recorded in TEM and STEM-HAADF modes, respectively, of the same area of the sample.

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A good contact exists between NaY(Yb)F$_4$ and Au-CdS. A HRTEM image corresponding to the square depicted in Figure 1C is shown in Figure 1E. The crystallinity and good contact between the different phases is clear. The bottom right inset in Figure 1F shows another STEM-HAADF image exhibiting CdS particles and Au nanorods. The upper part of the image in the white square is shown in HRTEM mode in Figure 1F. The bottom left of Figure 1F presents an FT image with spots at 3.2 Å, corresponding to the (101) crystallographic planes of the CdS support particle while those at 2.4 and 2.0 Å correspond to the (111) and (200) crystallographic planes of the metallic Au nanorod, which is oriented along the [110] crystallographic direction. To summarize, the Au-CdS-NaY(Yb)F$_4$ sample is constituted by NaY(Yb)F$_4$ crystallites of 150-250 nm surrounded by CdS nanoparticles of 40-80 nm. Some of the CdS nanoparticles are decorated by Au metal nanorods. There is a good contact between the different phases present in the sample and no agglomeration or segregation of a single phase occurs.

The system was excited with a femto second laser (Libra F-HE, Coherent) which has a maximum power of 4W/cm$^2$ in the IR region. As seen in figure S1 the up-conversion emissions of light at 800nm and 477nm are produced upon excitation with 980 nm light. In order to test this we have prepared a CdS semiconductor (that absorbs light at around 500nm) on top of which we have put Au nanorods (d=10nm and L=41nm, 0.25 wt. %, the remaining is CdS; see methodology section). The CdS was prepared and characterized following a previous work. We have measured the Au nano-rods for their particular plasmon resonance response in the IR and Visible (figure 2) light range. This has resulted in extending the Au plasmon energy into the IR region due to both the transversal and longitudinal plasmons. This wide spectral range, most likely due to light scattering, was in our favor, as it coincided with the absorbance edge of the up-converter (980 nm); and this is needed to enhance light adsorption which in turn is poised to enhance light emission. Figure 2 shows that the presence of Au nanorods on top of CdS has resulted in light absorption above 800nm, extending up to 1000nm; thus covering the absorption edge of the UC material.

NaYF$_4$ (Yb):Tm crystals have a gagarinite group structure with five bands of 2A$_g$ + 3E$_{Ig}$, Figure 3. The most enhanced phonons modes were at 253.4, 303.5, 366.2, 495.4 and 632.6 cm$^{-1}$, these phonons modes are similar to un-doped NaYF$_4$ samples. The contribution of Tm was not observed due to its low concentration.

The electron phonon coupling in these lattices takes place at relatively low energy phonon modes. Due to this property, NaYF$_4$ is considered the best host for up conversion with different rare earth ions. The Raman spectrum of Au-CdS shows two phonon vibrational modes, first and second order LO modes at 301.6 and 601.5 cm$^{-1}$. A shift of 5.8 cm$^{-1}$ in the case of Au-CdS/NaYF$_4$/Yb:Tm is noticed when compared to Au-CdS. We have excited the complete UC catalyst with 980 nm light (the absorbance edge of the UC) in the presence of gas phase methanol/air. Then we have monitored both H$_2$ and CO$_2$ production. CdS can only work as a semiconductor photo-catalyst upon excitation with visible light range and is therefore inactive when excited with 980nm light. The needed wavelength is provided by the up-converter material since a fraction of the 980nm light is converted to 802 nm and 477 nm which excite both the Au nano-rod and the CdS, respectively. The results were positive, in other words H$_2$ and CO$_2$ were detected and monitored (figure 4).

The photocatalytic methanol reaction in the presence of O$_2$ and/or H$_2$O can be summarized by three equations.
The observed ratio of the reaction rates between H₂ and CO₂ in figure 4A is above the stoichiometric ratio of 2 (equation 2) and this is due to the presence of gas phase water. Methanol is an excellent hole-scavenger allowing for excited electrons to reduce H⁺ (of surface -OH) to H₂[56]. After e injections into the valence band (VB) methanol gives formaldehyde which in turn is oxidized (with molecular or dissociated oxygen anions radicals) to give CO₂ upon further e injections into the VB[57]. In the presence of water, the reaction gives more H₂ (equation 3) and the energy needed to drive its reaction (endothermic) is provided from light, in this case. The traces of methane might be due to farther reaction of CO₂ with hydrogen atoms[58]. Because Au NRs are capped with an organic compound we have then repeated the experiments in the absence of methanol (blank), to test their possible contribution into the reaction and the results were negative (figure S2). This indicates that H₂ and CO₂ do not originate from the ligand capping the NRs. Then we have tested the UC material without the semiconductor CdS but with Au nano-rods. Previous reports have indicated that Au nanoparticles alone can reduce H⁺ to H₂ once excited with visible light[59,60,61]. Indeed the visible light (and probably the 800nm IR light) both generated by the UC made the reaction possible (with ca. 0.25-0.30 of the activity obtained when CdS was present). This indicates that H₂ production may in part originate from the direct catalytic reaction of the Au NRs upon excitation. The ratio of the production rate of H₂ to CO₂ in the absence of CdS is found to be less than 1. The absence of the semiconductor (absence of energy levels for e transfer from methanol to the VB of CdS) is probably the reason where in this case only hot e from Au NRs are involved in the reaction which in turn makes oxygen anions radicals routes more efficient, generating more CO₂ by direct oxidation of methanol. The system does not work in the absence of a metal (since it is needed to trap excited electrons) so one cannot, separately test the extent of enhancement due to Au NRs alone without the addition of another metal such Pd NP, but that is the subject of a separate study. Quantitative quantum yield of the reaction is not possible in our experimental set up, and is most likely very weak. Yet the mere observation of catalytic H₂ production indicates that converting IR light to visible light, needed to excite both a visible light driven semiconductor as well as the two plasmonic resonance of Au nanorods (transverse and longitudinal modes at ca. 500 and 800nm, respectively, as seen in figure 2), is possible and can reduce hydrogen ions to molecular hydrogen.

**Materials and methodology**

0.538 g of Yttrium (III) nitrate hexahydrate (Sigma Aldrich 99+), 0.260 g of Ytterbium (III) nitrate pentahydrate (Sigma Aldrich 99.9 %) and 0.015 g of Thulium (III) nitrate pentahydrate (Sigma Aldrich 99.9 %) were dissolved in 75 mL de-ionized water. 5.777 g of citric acid was dissolved into the pre-mentioned mixture to obtain a concentration of 0.4 M and citric acid to rare earth metal ratio of 4. In a separate flask, 3.78 g of NaF were dissolved in 75 mL of de-ionized water to obtain a 1.2 M concentration. The two mixtures were left under stirring for 1 hour after which, the NaF solution was added to the rare earth metal solution dropwise. After mixing the two solutions, the resulted mixture was left stirring for half an hour then transferred into a Teflon-lined autoclave (where only ¾ of the autoclave was filled with solution). The solution was then treated hydrothermally at 180°C for 24 hours. After completion, the product was washed three times with de-ionized water and once with ethanol.

![Figure 4](Image)

**Figure 4**

Gas phase photoreaction products of methanol under 980 nm excitation over A. 0.25 wt. % Au/CdS-(NaYF₄-Yb-Tm) with a 1 to 1 ratio. B. 0.25 wt. % Au/(NaYF₄-Yb-Tm). Ambient-air + gas phase methanol. Humidity is ca. 50% at 20°C, one atm. (ca.2 kPa). Methanol vapor pressure is ca. 10 kPa. Rates for each product are indicated inside the figure.

**Au/CdS/Upconverter Preparation**

Gold nanorods colloidal suspension is acquired from Sigma Aldrich with a 10 nm diameter and 41 nm in length. Au concentration is estimated to be greater than 30 μg/mL in H₂O. The amount of cetyl trimethylammonium bromide, C₁₅H₃₂NBr (CTAB) ligand on the metal (used to stabilize the nanorods) is estimated to be < 0.1 wt. %. CdS is prepared by precipitation of Na₂S and Cd(NO₃)₂ followed by calcination under inert atmosphere at 600°C for four hours. 0.25 wt. % Au/CdS is made by mixing 120 mg of CdS with 10 mL of gold colloidal suspension and drying at 90°C overnight under stirring.
Photoreaction at 980 nm Excitation
15 mg of 0.25 wt. % Au nanorods/Cds are mixed with 15 mg of (NaYF₄ – 28mol% Yb – 0.75mol% Tm) and sonicated in ethanol for several minutes. The mixture is then deposited on glass and the solvent dried at 70°C. Inside a 6 mL reactor, one drop of methanol (ca. 0.05 mL to ensure surface saturation) is added along with the coated slide and the reactor is sealed. The catalyst is then excited with ~0.15W at 980 nm (spot size is close to 3 mm) light provided from a 100 fs pulsed laser (Coherent), filters were provided from Concept Design Production (CDP) Systems Corp (http://www.cdp-sys.com).

Raman analysis were performed using a Thermo Scientific™ DXR™ Raman equipment. Catalyst samples were analysed from 50 to 3400 cm⁻¹ with multiple scans (16scans) with exposure time of 0.5 seconds, the wavelength of the laser used for the analysis was 532 nm, with the power of 8mW and the spot size was 2.1μm. The aperture opening was 50μm and grating had 900 lines per mm. High-Resolution Transmission Electron Microscopy (HRTEM) was carried out at 200 kV with a JEOL JEM 2100 instrument equipped with a LaB₆ source. The point-to-point resolution of the microscope was 0.19 nm. Samples were deposited on holey-carbon-coated Cu grids from alcohol suspensions.

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Notes and references

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