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In situ fabrication of polyacrylate-silver nanocomposite through

photoinduced tandem reactions involving eosin dye

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ABSTRACT

A silver-acrylate nanocomposite was prepared using a novel one-pot strategy involving eosin

dye as visible sensitizer and an amine derivative as radicals source. The mechanism

highlighted by steady state photolysis and time resolved absorption spectroscopy lies on the

initial formation of a strong ion pair complex between eosin and Ag+. Upon visible

irradiation, the excited triplet state of the metal complex efficiently abstracts a hydrogen atom

from the amine and produces a α-aminoalkyl radical. In acrylate monomer matrix, such a

strong reactive species initiates a free radical photopolymerization and also provides the

reduction of the silver cation. Through this 'in-situ' fabrication method, the kinetics formation

of the nanocomposite and its detailed structural analysis are characterized by UV-Visible,

real-time FTIR absorption spectroscopy and by transmission electron microscopy.

Keywords: Nanocomposite, Silver nanoparticles, Eosin

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1. Introduction

Progresses in size- and shape-selective synthetic routes to produce metal nanoparticles have become the subject of extensive investigations over past decade. Their size-dependent physical properties[1,2] clearly constitute one of the most attractive aspect which has promoted the emergence of a very large set of applications in optics, [3-6] microelectronics, [7,8] high density data storage, [9,10] catalysis [11,12] and energy conversion.[13,14] Nanocomposite materials, which combine the properties of polymers and silver nanoparticles, are considered to be promising systems for advanced functional applications.[15] For instance, coaxial nanocables[16] with a nanometric silver core and covered by a polymer shell exhibit superior performance as nanocircuits[8] or nanosenors.[7] Such nanowires can be synthesized by either solution-phase methods using capping reactants[17,18] or by templating methods using inorganic porous material,[19] polymer membranes,[20,21] or block copolymer.[22] Another type of nanocomposites consists in a dispersion of metal particles into a polymeric matrix such as poly(vinyl alcohol),[23,24] poly(methyl methacrylate)[25] or poly(vinyl pyrrole) for instance.[26,27] Typically, in-situ and ex-situ methods[28-30] are the two main synthetic routes that lead to silver-polymer nanocomposites. The ex-situ method consists in dispersing dry silver nanoparticles produced beforehand by an appropriated chemical process into a polymer medium to generate highly ordered periodic arrays of nanoparticles.[31] In the in-situ method, metal nanoparticles are generated directly in the matrix through chemical reduction of a cationic precursor previously dissolved in the bulk.[24] Photochemical reduction by irradiating a dye sensitizer in the presence of silver ions also provides a convenient way to produce embedded nanoparticles in polymer.[21,32-35] Moreover, a concomitant functionalization of the nanoparticles with dyes can occur, which is a valuable feature for applications in photochemical solar cells.[14,36] In all cases, the high-performance of nanomaterials depends on the controlled distribution of uniformly shaped and sized particles. Therefore, the development of synthetic strategies to control particles growth and/or agglomeration during nanocomposite fabrication appears quite obviously as a key challenge.

The present work reports a strategy involving the photoinduced formation of homogeneous silver nanoparticles in a polyacrylate polymer stemming from a cross linking photopolymerization of pentaerythritol triacrylate monomer (Scheme 1a). In this one-step approach, a combined system employing eosin dye as visible photosensitizer (Scheme 1b) and

N-methyl diethanolamine as co-initiator is used to photogenerate Ag nanoparticles and to photoinitiate a free radical polymerization at the same time. After a mechanistic insight provided by steady state and laser flash photolysis experiments, the analysis of the elementary steps leading to the metal nanocomposite will be correlated to the structural properties of the materials.

2. Experimental section

2.1. Reagents

Silver nitrate (AgNO₃) with a purity >99% was purchased from Fluka. Eosin Y disodium salt, N-methyl diethanolamine (MDEA), tetra-n-butylamonium hexafluoroborate were purchased from Aldrich. Mixture of pentaerythritol tri- and tetra-acrylate (PETA) monomers stabilized with monomethyl ether hydroquinone (MEHQ - 400 ppm) was purchased from Sartomer. All solvents were Aldrich and Fluka Spectrograde.

2.2. Instrumentation

The absorption measurements were carried out with a Perkin Elmer Lambda 2 spectrometer. A FluoroMax 4 Luminescence Spectrometer was used for the fluorescence and time gated phosphorescence. Low temperature measurements were performed in a glassy matrix of isopropanol using a 5-mm diameter quartz tube inside a Dewar filled with liquid nitrogen. The emission spectra were spectrally corrected in all cases.

Laser flash photolysis at nanosecond time scale was carried out with an Edinburgh Analytical Instruments LP900 equipped with a 450-W pulsed Xe arc lamp, a Czerny-Turner monochromator and a fast photomultiplier. The samples were irradiated with the second harmonic ($\lambda = 532$ nm, ~ 10 ns, 4 mJ per pulse) of a Nd/YAG Powerlite 9010 from Continuum. The sample was purged with argon for 15 min prior to photophysical studies.

The acrylate formulation, typically Eosin (0.1 wt %), MDEA (3 % wt) and AgNO $_3$ (1 wt %) in PETA, was sandwiched between two glass plates with a calibrated thickness spacer setting the optical path length to 30 μ m. Photochemical reactions were carried out at 532 nm with a cw 532 nm Verdi laser from Coherent. The progress of the reaction was monitored via UV-Vis absorption spectra.

The photopolymerization was followed up in situ by real-time Fourier transformed infrared spectroscopy with an AVATAR 360 FTIR spectrometer from Nicolet. The laminated formulation (25 μ m thick), deposited on a BaF₂ pellet, were irradiated at 532 nm with a green laser diode module from Crystalaser. The conversion rates were deduced from the disappearance of the vinyl C=C stretching vibration band at 1630 cm⁻¹.

Transmission electron microscopy (TEM) was used to characterize the size and shape of Ag nanoparticles. The nanocomposites were cut by means of a microtome (LKB model 8800) and placed onto the observation grid to get their TEM images. Transmission electron microscopy measurement is carried out at 200 kV using a Philips CM20 instrument with Lab6 cathode.

The ¹³C NMR spectra were recorded in CD₃CN on a 400 MHz Brucker NMR spectrometer using cyclohexane as the internal standard.

3. Results and discussion

3.1. Formation of a strong ion-pair complex with Ag⁺

The lowest energy absorption band of EO^{2-} is localized in the 450-575 nm region with a maximum at 530 nm in acetonitrile (Figure 1). This band corresponds to the S_0 - S_1 electronic transition with a π - π^* character localized on the xanthene moiety.[37] The geometry optimization by AM1 method reveals that 2-benzoate and xanthene sub-units are twisted with an angle of 77°, which indicates a weak electronic conjugation between the two aromatic moieties. The fluorescence spectrum of EO^{2-} is mirror image of its absorption band with a slight Stockes shift (535 cm⁻¹) suggesting a weak geometrical relaxation in the singlet state. The 0-0 transition energy measured from the intercept of normalized absorption and fluorescence spectra corresponds to 2.31 eV in acetonitrile. The phosphorescence spectrum of EO^{2-} is located in the 600-700 nm range with a maximum emission at 652 nm which leads to a triplet energy of about 1.90 eV (inset figure 1).

As depicted in figure 2, the addition of silver nitrate to a solution of EO^{2-} in acetonitrile leads to a slight red shift of the last absorption band of the chromophore ($\Delta\lambda_{max}=+4$ nm). The presence of two isosbestic points located at 498 nm and 530 nm clearly indicates the occurrence of equilibrium reaction. Moreover, the addition of tetra-n-butylamonium hexafluoroborate (3 x 10^{-2} M) excludes any variation of the ionic strength during the reaction. Therefore we can ascribe these effects to a coordination reaction of Ag^+ by EO^{2-} which leads

to the formation ion-pair complex [EO-Ag] $^-$. According to the stronger acidic character of the hydroxylic group with respect to that of the carboxylic one[38,39], such a cation exchange reaction should mainly involve the chelation of silver cation by the 'phenolate' oxygen. This is also in line with the Pearson's hard-soft acid-base (HSAB) principle[40-43] which presumes a better stabilizing interaction of Ag^+ (Soft Lewis acid) with the phenolate group than with the carboxylate function. However the participation of the carboxylate function to the coordination reaction can not be excluded. In order to determine the reaction constants, the absorption changes as a function of the concentration of $AgNO_3$ were analyzed according to the model proposed by Valeur et al[44] which is extended to the formation of [1:1] and [1:2] ligand-metal complexes[45]. The inset Figure 2 depicts the best-fitting curve to the experimental data using the method of least squares, log $K_{1:1}$ exhibits a high value of ca. 4.7 ± 0.3 whereas log $K_{1:2}$ has a value of ca. 5.0 ± 0.4 . Such a slight difference suggests that the formation of the [1:2] complex can be reasonably neglected with respect to our working condition (i.e. $[Ag^+]/[EO^2^-] < 50$).

3.2. Mechanistic aspect of Ag nanoparticles photogeneration in solution

Figure 3 shows the evolution of the transient absorption spectrum of EO²⁻ in acetonitrile. The spectrum mainly consists of a broad band located in the 560-700 nm range. This band was previously assigned[46-48] to the absorption of the triplet of EO²⁻ (³EO²⁻), its transient monitored at 580 nm decays mono-exponentially with a lifetime of 45 μs. The spectrum also exhibits a bleaching in the 520 nm region which corresponds to the depletion of the ground state. The transient absorption spectrum below 490 nm corresponds to the superimposition of the absorption of three species: the triplet of eosin, the semi-oxidized (EO^{*-}) and semi-reduced (EO^{*-}) radical ions. The two latter species are generated through a self-quenching of ³EO²⁻ by ground state molecules:

$${}^{3}EO^{2-} + EO^{2-} \rightarrow EO^{2-} + EO^{2-}$$
 ${}^{3}EO^{2-} + EO^{2-} \rightarrow EO^{*-} + EO^{*3-}$

In acetonitrile, these long-lived radical ions are known[47,48] to have strongly overlapped absorption bands. The maximum absorption wavelength of EO^{*3-} is reported at 430 nm (49) whereas EO^{*-} shows a band centred at 450 nm (39,50).

The addition of 50 eq. of Ag⁺ to a solution of EO²⁻ leads to a quantitative conversion of the dye to its ion-pair complex, [EO-Ag]⁻. As shown in Figure 4, the transient absorption

spectrum remains globally invariant for wavelength above 490 nm. The triplet lifetime measured from the transient at 580 nm drops to a value of 25 µs. Consecutive to the triplet quenching, the development of a long time component is observed in the transient monitored at 450 nm (inset Figure 4) which indicates the growth of a new band in this spectral window. This band can be confidently assigned to the absorption of semi-oxidized eosin as compared with that observed by Lindqvist (47) and Grossweiner (48,50,51). Besides, the transient absorption spectrum exhibits an other band centred at 370 nm. Such a broad band which collapses within the first 2 µs after laser pulse corresponds to the well-established band of Ag⁰ (17,52,53). Hence we have a clear evidence of a photoinduced electron transfer from ³EO²⁻ to Ag⁺. However, probably due to its relative low reaction rate, such a process can not produce efficiently silver nanoparticles. This is confirmed by steady state photolysis experiment. The changes in the spectral features observed upon laser excitation of eosin dye and AgNO₃ (50 eq.) solution in acetonitrile are depicted in Figure 5A. Upon increasing the irradiation time, the absorption band of EO²⁻ is progressively collapsing, however we fail to observe the formation of silver nanoparticles. The same photolysis experiment is performed with a solution of acetonitrile containing eosin dye, AgNO3 and MDEA. In this case, a new band centred around 435 nm and characteristic of the surface plasmon (SP) band of silver nanoparticles is clearly developing (Figure 5B). Obviously, the reduction of Ag⁺ in presence of MDEA is only possible with EO²⁻ and under visible irradiation. This confirms that the formation of silver nanoparticles is promoted by the excited dye and its mediating role with MDEA. The amine is known to undergo a hydrogen abstraction reaction with the triplet state of eosin (3EO2-). The sequential reaction first involves an electron transfer reaction from amine to ³EO²⁻ then a proton transfer within ion pair formed between amine radical cation and eosin radical anion (49,54). Hence, the reaction should initially produce a α-aminoalkyl radical and the conjugated acid of the semi-reduced eosin (EOH*2-). However the latter species which exhibits a transient absorption band around 375 nm is in equilibrium with its conjugate base EO^{*3} whose absorption band is located at 430 nm. The equilibrium has been observed on flash photolysis of EO²⁻ in aqueous media, with a pK_a of 7 associated with the acidic form (47,48,51). The bimolecular quenching rate constant of ³EO²⁻ by MDEA (k₀) can be determined from the dependence of the pseudo-first-order decay of the triplet dye on the concentration of MDEA according to the equation:

$$k_{obs} = k_0 + k_q[MDEA]$$

In this condition, the bimolecular quenching rate exhibits a value of $2.5(\pm\,0.1)$ x $10^6\,M^{-1}$ s⁻¹, in good accordance with literature (54). This low rate constant suggests typical hydrogen atom transfer reaction instead of electron-proton transfer sequence. Parallely to the quenching of triplet eosin, EO*3- is detected at 430 nm. This indirectly confirms the production of the α -aminoalkyl radical which can not be detected in the spectral window provided by laser flash photolysis.

Figure 6 shows the evolution of the transient absorption spectrum of a solution containing EO^{2-} , $AgNO_3$ and MDEA. At very short delay after laser pulse, the absorption band of Ag^0 located at 370 nm reaches its maximum, its transient can be confidently fitted with one exponential leading to a short lifetime of 290 ns. Interestingly, the fast decrease of the metal absorption band reveals the presence of a weak band at 375 nm which is ascribable to the absorption of EOH^{*2-} . As the intensive transient absorption band of EO^{*3-} is concomitantly developing, we have clear indication that the reduction of Ag^+ does not proceed through the oxidation of EO^{*3-} but through an electron transfer from the α -aminoalkyl radical. Carboncentred free radicals with adjacent nitrogen or oxygen are potent reducing agents in solution[55-59]. For instance, it has been reported that the photogenerated alkoxy benzyl radicals[33] or ketyl radicals[60] can convert efficiently Ag^+ to Ag^0 . Hence, we can assume that the main reaction route leading to the generation of Ag nanoparticles can be described as follows:

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[EO-Ag] $^{-}$ + MDEA \rightarrow EOH $^{\bullet 2^{-}}$ + Ag $^{+}$ + MDEA $^{\bullet}$
Ag $^{+}$ + MDEA $^{\bullet}$ \rightarrow Ag 0 + MDEA $^{+}$

The initial coordination of eosin dye with silver cation undoubtedly constitutes an advantage since the photogeneration of the α -aminoalkyl radical and its consecutive oxidation by Ag^+ are occurring in the same reactive cage. In PETA monomer, the ion-pair complex formation appears all the more useful as its presence should strongly counterbalance the effect of a slow diffusion in high viscous medium.

3.3. Reduction of Ag⁺ ions in a PETA matrix

The fabrication of metal nanocomposite is performed using pentaerythritol triacrylate (PETA) as monomer. Surprisingly, the dilution of AgNO₃ in PETA matrix leads to a very slow change in colour. The absorption spectrum of the formulation clearly shows the growth of a band centred at 435 nm which corresponds to the SP band of Ag nanoparticles (inset Figure 7). The

TEM image of the sample indicates a polydisperse distribution of spherical silver nanoparticles (Figure 7). Hence a reduction of silver cations is occurring in the monomer. The involvement of MEHQ (used as monomer stabilizer) in this reaction can be excluded since no reduction was detected upon addition of large amount of this phenol derivative to a solution of the silver salt in acetonitrile. Therefore the pentaerythritol triacrylate induces a slow reduction of Ag⁺. Moreover, the 13 C NMR spectrum of the monomer in acetonitrile- d_3 is notably modified in the presence of Ag⁺. While main part of the NMR spectrum remains invariant in presence of Ag⁺, the signals of vinylic carbons[61] are gradually shifted by increasing the concentration of AgNO₃. For instance, upon addition of 10 eq. of Ag⁺, the signal of methylene carbons ($\delta = 131.51$ ppm) is shifted to low-field region (± 0.2 ppm) whereas the methine one $(\delta : 128.29 \text{ ppm})$ shifts to high-field region (-0.2 ppm). It should be noted that Ag⁺ is wellknown to form complexes with a large set of alkenes[62-64], including methyl acrylate derivatives [65]. Therefore, the formation of a class $T \square \text{ complex}[64]$ between Ag^+ and PETA is proposed. Such a structure implies a trigonal environment around Ag+ with a chelation of the cation by the three vinylic functions. As consequence, the close proximity between the hydroxyl group of PETA and Ag⁺ should promote a redox reaction and lead to the formation of Ag nanoparticles. The oxidation of primary alcohols (R-CH₂OH) by Ag⁺ is also well established, the reaction is slow and requires heating to be accelerated as follows[66-68]:

$$2 \text{ Ag}^{+} + \text{R-CH}_2\text{OH} \rightarrow 2 \text{ Ag}^{0} + 2 \text{ H}^{+} + \text{R-CHO}$$

However such a reaction is a very slow process and constitutes a minor route for Ag⁺ reduction in comparison with the photoinduced generation of Ag nanoparticles described hereafter.

3.4. Fabrication of silver nanoparticle embedded polymer

A formulation of the PETA monomer containing EO²⁻ (0.1 wt %) and MDEA (3 % wt) [69] and AgNO₃ (1 wt %) is photopolymerized upon irradiation at 532 nm (2.5 mW cm⁻²). The conversion of the acrylate double bonds was monitored from the decrease of the IR band at 1630 cm⁻¹. In Figure 8 are compared the conversion rate related to this formulation with that of a reference one without AgNO₃. Both samples show a similar rate of polymerization (within our experimental error) with an acrylate final double bond conversion of about 60 %. This final degree of double bond conversion should be ascribed to the acrylate trifunctionality. As the functionality increased, the viscosity of the resin increased, with the

resulting gel-effect and the higher cross linked density, set a limit to the extent of conversion[70]. We can clearly notice that the addition of Ag^+ does not perturb the photopolymerization kinetics. Therefore, the quantum yield of generation of MDEA $^{\bullet}$ is sufficiently high to enable the free radical photoinitiation and the photogeneration of Ag nanoparticles. Moreover after 10 min exposure, the reference sample turns from pink to colourless whereas the sample with Ag^+ turned from pink to yellowish (inset Figure 8).

The absorption spectrum of the sample undergoes important changes during visible irradiation (Figure 9). The absorption band of EO²⁻ decreased progressively whereas the SP developed in the 350-500 nm region. This band exhibited a maximum wavelength at 437 nm with a FWHM of 115 nm. Careful analysis of the absorption spectrum over the illumination period clearly reveals a two-step process:

- (i) The first step corresponds to the photobleaching of EO²⁻ with the concomitant growth of the surface plasmon band as indicated by the linear correlation between the absorbance at 532 nm vs. absorbance at 432 nm (inset of Figure 9).
- (*ii*) In a second step, the SP band continues to develop under irradiation even though the absorption band of EO²⁻ has completely disappeared. Small particles formed in the first step contribute to the further reduction of Ag⁺ through an autocatalytic process where the residual PETA monomers should act as sacrificial donor. Ultimately, the absorbance of SP stabilized to ca twice that measured at the end of EO²⁻ bleaching.

Figure 10 shows the linear correlation between the final absorbance reached by the SP band at its maximum and the initial amount of EO^{2-} ; this correlation suggests that the initial step determines the final concentration of nanoparticles in the polymer. The bright field TEM micrographs of silver-nanocomposites displayed in Figure 11 corroborate the correlation between the density of nanoparticles and the initial amount of EO^{2-} . Throughout the photoreduction process, the symmetrical shape of the SP band denotes a monodisperse size distribution of the silver nanoparticles. Electron microscopy analysis of the polymer indicated the formation of monodisperse spherical metallic particles in nanometer range size. Analysis of a population of ca. hundred particles from a portion of the grid indicated that their average diameter is 5.0 ± 0.9 nm respectively 5.0 ± 0.7 nm. The particles were homogeneous in size and no agglomeration was observed. Such a homogeneous distribution can be accounted by the stabilizing effect of the PETA matrix. Through its acrylate functions, the PETA monomer can easily adsorb on the nanoparticle surface. In this regards, the capping effect of unsaturated long-chain agents was previously observed by Wang *et al.*[71], who correlated the size of silver particles with the number of double bonds in the chain.

4. Conclusion

A strong ion pair complex is formed at the ground state between eosin dianion chromophore and Ag^+ . The triplet state of such a metal complex dye conserves its high hydrogen abstraction propensity toward N-methyl diethanolamine. The α -aminoalkyl radicals that are consequently produced upon visible irradiation lead to a consecutive reduction of the coordinated silver cations. Moreover, in triacrylate monomer matrix, the quantum yield of radicals generation is still high and allows a simultaneous free radical polymerization. Hence, an "in situ" synthesis of silver nanoparticle embedded polymer is clearly demonstrated with the advantage that the particles density in the polymer can be tuned by controlling the initial concentration of dye. Finally, the obtained nanocomposite exhibits spherical shaped Ag nanoparticles whose homogeneous dispersion is ascribable to the capping effect of monomer acrylate branches that are not engaged in the polymer net.

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FIGURES AND SCHEMES CAPTIONS

- Figure 1: Normalized absorption and fluorescence spectra of EO²⁻ in acetonitrile. Inset: Time-gated phosphorescence spectrum of EO²⁻ in glassy matrix of isopropanol (delay: 50 μs)
- **Figure 2**: Evolution of the absorption spectrum of EO^{2-} (2.4 x 10^{-6} M) upon addition of AgNO₃ (solvent : acetonitrile, [MDEA] = 10^{-4} M, [nBu₄N,BF₆] = 3 x 10^{-2} M) Inset : Dependence of the absorbance at 535 nm vs. concentration of silver cations (squares). Best fitting curve (line)
- Figure 3: Time-resolved transient absorption spectra recorded at different delays after 532 nm laser pulse excitation of deaerated acetonitrile solution containing EO^{2-} (5 x 10^{-6} M) and nBu_4N , BF_6 (10^{-2} M)
- Figure 4: Time-resolved transient absorption spectra recorded at different delays after 532 nm laser pulse excitation of deaerated acetonitrile solution containing EO^{2-} (5 x 10^{-6} M), AgNO₃ (2.5 x 10^{-4} M) and nBu₄N,BF₆ (10^{-2} M) Inset: Absorption-time profile recorded at 450 nm
- Figure 5 : Evolution of the absorption spectra of two irradiated mixtures in acetonitrile $(\lambda_{irr}=534 \text{ nm}). \text{ Solution } \textbf{A}: EO^{2\text{-}}(10^{\text{-}5} \text{ M}), \text{AgNO}_3 (5 \text{ x } 10^{\text{-}4} \text{ M}) \text{ and } \text{Bu}_4\text{N}, \text{BF}_6 \\ (10^{\text{-}2} \text{ M}); \text{ Solution } \textbf{B}: EO^{2\text{-}}(10^{\text{-}5} \text{ M}), \text{AgNO}_3 (5 \text{ x } 10^{\text{-}4} \text{ M}), \text{MDEA} (10^{\text{-}2} \text{ M}) \text{ and } \\ \text{nBu}_4\text{N}, \text{BF}_6 \ (10^{\text{-}2} \text{ M})$
- Figure 6: Time-resolved transient absorption spectra recorded at different delays after 532 nm laser pulse excitation of deaerated acetonitrile solution containing EO²⁻ (5 x 10⁻⁶ M), AgNO₃ (2.5 x 10⁻⁴ M), MDEA (5 x 10⁻³ M) and nBu₄N,BF₆ (10⁻² M)

Figure 7: TEM image of silver nanoparticles embedded in pentaerythritol triacrylate (AgNO₃, 1 wt %).

Inset: Growth of the absorption spectrum of Ag nanoparticles (optical path of the cuvette: 1 cm)

Figure 8 : Real-time FTIR kinetics conversion curves for visible-curing at 532 nm of resins with (1) EO 2 / MDEA, 0.1 wt % / 3 wt % (2) EO 2 / MDEA / AgNO₃, 0.1 wt % / 3 wt % / 1 wt %

Inset: View of 30µm thick samples (1) and (2) after curing

Figure 9: Time evolution of the absorption spectrum of the resin during photopolymerization EO 2 -/ MDEA / AgNO $_3$, 0.1 wt % / 3 wt % / 1 wt % (optical path of the cuvette : 30 μ m)

Inset: Linear correlation between absorption at 532 nm and 432 nm

Figure 10: (A) Kinetics of absorption at 432 nm for various samples with increasing amounts of EO^{2-} (MDEA / AgNO₃, 3 wt % / 1 wt %). (B) Correlation between the final absorbance of plasmon band with the initial amount of EO^{2-}

Figure 11: TEM image of silver nanoparticles embedded in a polyacrylate matrix, with their respective size distribution. $EO^{2-}/MDEA/AgNO_3$: (a) 0.06 wt % / 3 wt % / 1 wt % and (b) 0.28 wt % / 3 wt % / 1 wt %

Scheme 1: Structures of pentaerythritol triacrylate (PETA) and of Eosin Y disodium salt

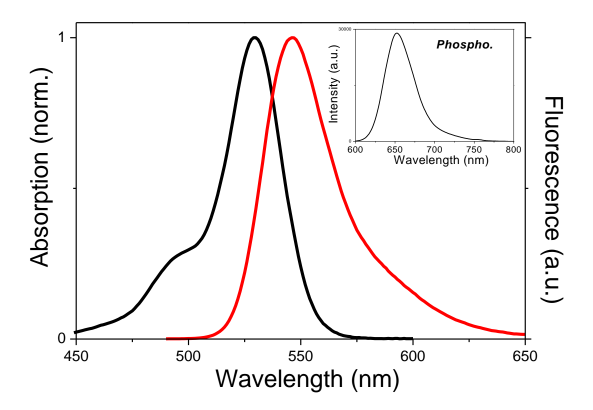


Figure 1

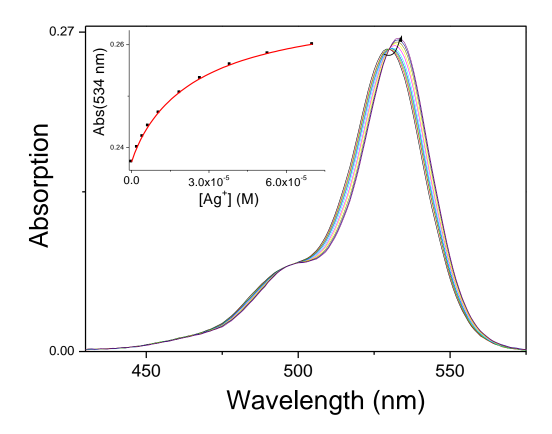


Figure 2

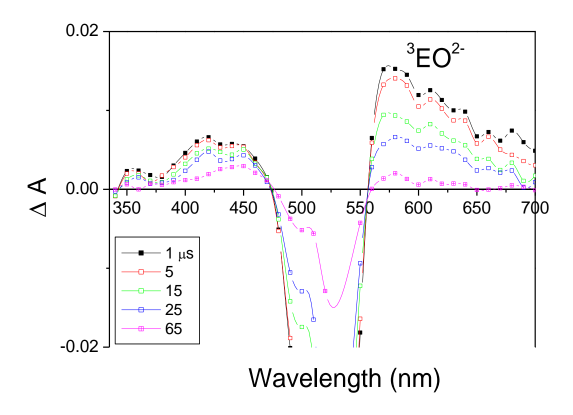


Figure 3

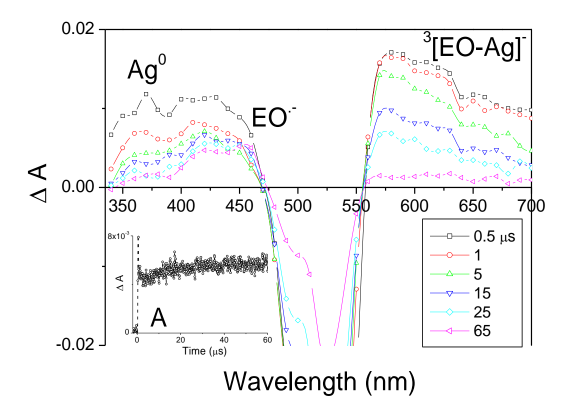


Figure 4

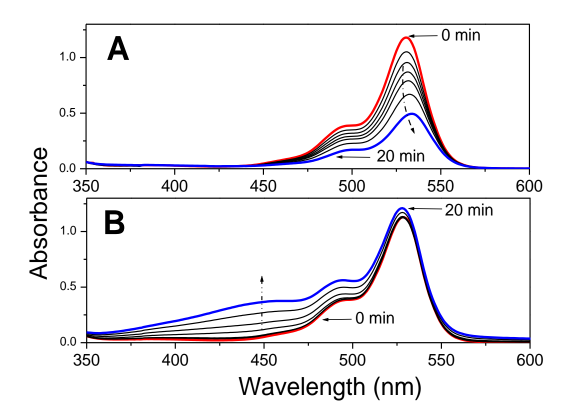


Figure 5

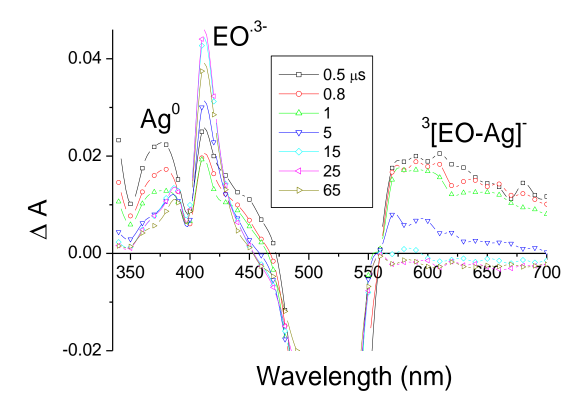


Figure 6

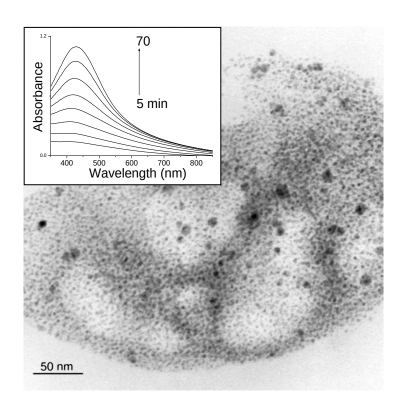


Figure 7

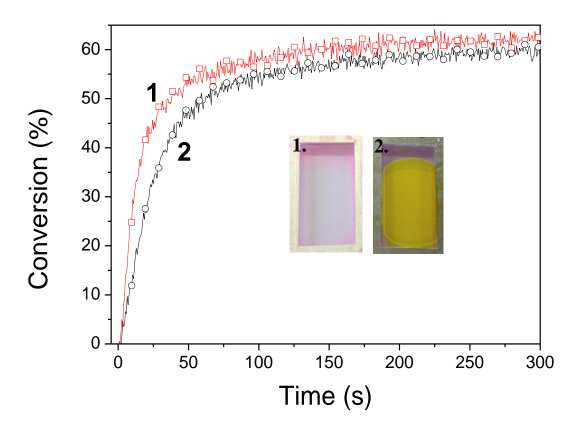


Figure 8

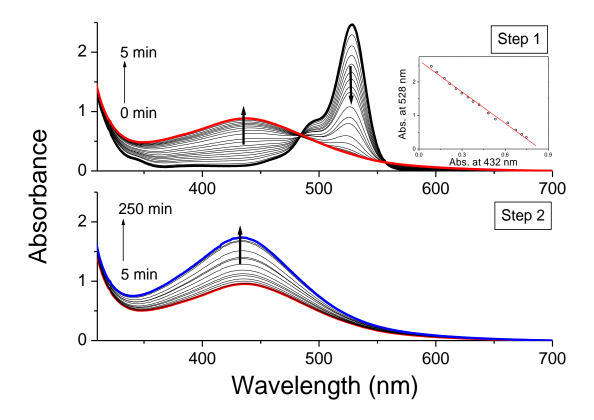


Figure 9

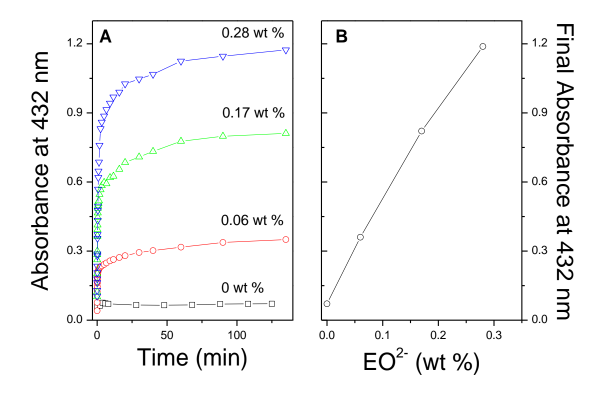
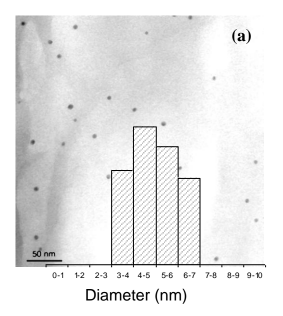


Figure 10



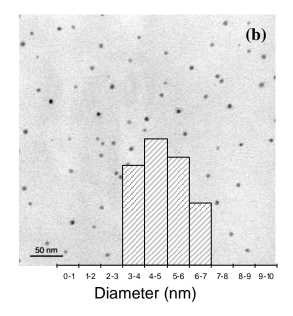


Figure 11

Scheme 1