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materials letters

Materials Letters xxx (2010) xxx-xxx

Contents lists available at ScienceDirect



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Materials Letters

journal homepage: www.elsevier.com/locate/matlet

Nanoscale color control of TiO₂ films with embedded Au nanoparticles

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A R T I C L E I N F O

Article history: Received 1 July 2010 Accepted 9 August 2010 Available online xxxx

ABSTRACT

We demonstrate an efficient nanoscale control of the optical properties of TiO_2 films by tuning the Surface 15 Plasmon Resonance (SPR) in the embedded Au nanoparticles. The films were grown by reactive magnetron 16 sputtering. SPR tuning was achieved by different annealings, which affected the shape and size of the Au 17 nanoparticles, and also the phase of the dielectric matrix. These changes promoted the variations on the 18 optical properties. As shown by the modeling of the effective dielectric function of the TiO_2/Au in the SPR 19 region, the variation of their optical absorption spectra correlates with morphological changes. 20

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25 1. Introduction

The interest in composite materials containing metal nanoparticles 26(NPs) embedded in dielectric matrices is related to their potential 2728 application in a wide range of technological applications, such as colored coatings [1], solar cells [2], sensors [3,4], antibacterial [5] 29photocatalysis [6-8], and nonlinear optics [2,9,10]. TiO₂ is a 30 transparent semiconductor material with a wide band gap $E_g = 3.2$ -31 3.4 (eV) and high refractive index (n = 2.5-2.9) used for metal-32 dielectric composites designed for obtaining desired optical proper-33 ties in the visible range. 34

The brilliant colors of composites containing noble metal inclu-35 sions are due to SPRs in the metallic phase [11]. Films with well-36 separated embedded metallic NPs with dimensions significantly 37 smaller than the wavelength of the exciting light are characterized 38 by a peak in the visible range of the absorption spectra. The band 39 width, intensity, and position of the absorption maximum depend on 40 41 the surrounding dielectric matrix, the size, distribution and especially on the shape of the NPs. This fact allows to tune the optical properties 42of the composite, (i) by changing the refractive index of the matrix 4344 (n_h) and (ii) by modifying the morphology and distribution of the metallic inclusions changing the aspect ratio of metallic NPs [12]. 45

46 **2. Experimental details**

47 TiO₂/Au composite films were deposited on glass/quartz sub-48 strates, and *in-situ* Au doped by one step reactive magnetron 49 sputtering process, at a constant temperature of 150 °C [13]. The 50 deposited films were annealed in order to promote changes in the

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0167-577X/\$ - see front matter © 2010 Elsevier B.V. All rights reserved. doi:10.1016/j.matlet.2010.08.031

morphology, distribution and structural features of *in-situ* grown gold 51 NPs. The doping leads to an average Au volume fraction, f_{Au} , of about 52 12 at.% as determined by the Rutherford backscattering spectrometry 53 (RBS). 54

The crystalline structure of as-grown and annealed films was 55 investigated by X-ray diffraction (XRD), using a Philips PW 1710 56 diffractometer (Cu-K_{α} radiation) operating in a Bragg-Brentano 57 configuration. The XRD studies allowed to study the film structure 58 concerning both the Au NPs and the TiO₂ matrix. Transmission 59 electron microscopy (TEM) employing a Hitachy 800H apparatus was 60 used to characterize the shape, size and spatial distribution of NPs. 61 Color coordinates and absorption spectra were measured using a 62 commercial MINOLTA CM-2600d and a UV-vis-NIR spectrophotom- 63 eter (UV-3101) respectively [4,13].

3. Results and discussion

According to the RBS data, there were no measurable changes in 66 the Au doping profiles across the 300 nm of the entire films' thickness 67 during the annealing. The SPR-mediated color control has been 68 achieved by means of annealing of TiO₂/Au composite films grown by 69 reactive magnetron sputtering. Fig. 1 shows the color coordinates 70 (L^*, a^*, b^*) and the change after annealing at different temperatures. 71 Concerning the TiO₂ matrix, the XRD studies revealed that it is 72 amorphous in as-grown and at low temperature annealed conditions. 73 At 500 °C the TiO₂ dielectric matrix starts to crystallize in an anatase-74 type structure, transforming to rutile above 700 °C. Simultaneously, 75 the Au atoms are organized in crystalline nanoparticles (revealing a 76 fcc-type structure, with the (111) preferential growth orientation) 77 [4,13].

Optical spectra, together with the corresponding morphological 79 data, are depicted in Fig. 2. As it can be seen the SPR related absorption 80 of light occurs only in the films annealed at 400 °C or higher 81

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Fig. 1. Color coordinates (L^*, a^*, b^*) and the appearance of TiO₂/Au composite films annealed at different temperatures.

temperatures. The intensity of the resonance absorption increases with the annealing temperature and so does the NP size, as revealed by the TEM images of the same figure. At higher annealing temperatures the SPR band shifts to longer wavelengths and changes its shape. From the theoretical result obtained for a perfectly spherical metallic particle, often referred to as the Mie theory [14] the SPR condition is:

$$_{s}+2\varepsilon_{h}=0 \tag{1}$$

90 where $\varepsilon_h = n_h^2$ is the dielectric constant of the host (matrix) and ε_s 91 denotes the metal-dielectric function. In the simple Drude model,

$$\varepsilon_{s}(\omega) = \varepsilon_{\infty} \left[1 - \frac{\omega_{p}^{2}}{\omega \left(\omega + i\Gamma_{p}\right)} \right]$$
(2)

92 where ε_{∞} is a constant, ω_p is the plasma frequency and Γ_p is a damping 94 parameter. Assuming $\Gamma_p << \omega_p$, it follows from Eqs. (1) and (2) that the 95 SPR frequency for a sphere is given by

$$\omega_{SPR} = \frac{\omega_p}{\sqrt{1 + 2\varepsilon_h / \varepsilon_\infty}}.$$
(3)

700 600 500 A=-ln(T) (a.u.) 400 2 300 1 200 As-Dep n 400 800 900 500 600 700 Wavelength (nm)

Fig. 2. Optical transmittance spectra (plotted in logarithmic scale) of a set of TiO_2/Au composite films annealed at different temperatures. TEM images of four representative films are also shown.

The width of a single SPR is determined mostly by the plasmon 98 damping which depends on the NP size. For spherical Au NPs the 99 following expression has been proposed [15].

$$\hbar\Gamma_p(R) = \hbar\Gamma_p(\text{bulk}) + g_s v_F / R = 0.0244 + 0.922g_s / R \quad [\text{eV}], \qquad (4)$$

where v_F is the Fermi velocity, g_s is a geometrical factor of the order of 102 unity and the NP radius R is in nanometers (For the geometrical factor, 103 a value of $g_s \approx 0.7$ has been suggested) [16]. The intensity of the 104 resonance is inversely proportional to Γ_p and directly proportional to 105 the particle volume (R^3). Thus, the theory predicts that the intensity of 106 the SPR band decreases and its broadening increases with the 107 decrease of the NP size. It is experimentally demonstrated when 108 samples annealed at 300 °C and 400 °C are compared (Fig. 2). 109 However, for higher temperatures the absorption band becomes 110 broader and its shape changes suggesting that perhaps there are more 111 than one resonance involved.

The single SPR defined by Eqs. (1)-(3) is split if the particles are 113 not spherical [17]. Axial-symmetric nanorods are supposed to 114 produce two SPRs, 115

$$\omega_{SPR}^{'} = \frac{\omega_{p}}{\sqrt{1 + (\eta_{\parallel}^{-1} - 1)\varepsilon_{h}^{'}/\varepsilon_{\infty}}}; \ \omega_{SPR}^{''} = \frac{\omega_{p}}{\sqrt{1 + (\eta_{\perp}^{-1} - 1)\varepsilon_{h}^{'}/\varepsilon_{\infty}}}$$
(5)

where η_{\parallel} and η_{\perp} are so called depolarization coefficients [18], which 116 are some geometrical factors, non-negative numbers obeying the 118 relation $\eta_{\parallel} + 2\eta_{\perp} = 1$. The first resonance takes place for electromagnetic wave polarized along the nanorod axis while the second one 120 corresponds to the perpendicular polarization. If nanorods are 121 embedded in a matrix in a random-orientation fashion, then one 122 should expect to observe both SPRs. If we approximate nanorods by 123 elongated (prolate) spheroids with excentricity *e*, the depolarization 124 coefficients are given by [18]: 125

$$\eta_{\parallel} = \frac{1 - e^2}{e^3} \left(\frac{1}{2} \log \frac{1 + e}{1 - e} - e \right) \le \frac{1}{3}; \ \eta_{\perp} = \frac{1 - \eta_{\parallel}}{2} \ge \frac{1}{3}.$$
(6)

Using these relations one can see that $\omega'_{SPR} < \omega_{SPR} < \omega''_{SPR}$. The 128 two resonances corresponding to Eq. (4) merge into a single one 129 determined by Eq. (3), when $\eta_{||} = \eta_{\perp} = 1/3$ (spherical NPs). 130

Another effect clearly seen in Fig. 2 is the red shift of the absorption 131 band. This is related to a change in the phase composition of the 132 matrix. As already demonstrated, the transition temperature of the 133 anatase-rutile TiO₂ has been reported on a wide range of tempera-134 tures, depending on the structure and morphology of the film growth 135 process [19,20], and occurs typically between 700 and 800 °C for the 136 studied samples, as it is reported in previous publications [13]. The 137 two phases have different values of the refractive index, $n_{anatase}$ = 2.5 138 and n_{rutile} = 2.9 at λ = 550 nm proposed [21]. As the annealing 139 temperature was increased, the matrix became more crystalline, 140 increasing the refractive index, but some amorphous TiO₂ phase 141 remains. Anatase is the main crystalline phase until 700 °C, and at 142 higher annealing temperatures the TiO₂ matrix is richer in rutile and 143 the value of ε_h is even higher [4,13].

A model of the optical properties of the TiO₂/Au composite films was 145 performed in order to quantify the arguments. The effective dielectric 146 function of the composite material was calculated using the previously 147 developed modified Maxwell-Garnett (MMG) formalism, which takes 148 into account the dipole–dipole interaction between polarized particles 149 [22,23]. As in the classical Maxwell-Garnett approach, the (complex) 150 effective dielectric function (ε) of the composite is related to the 151 particle's polarizability (α) through the equation: 152

$$\frac{\varepsilon^* - \varepsilon_h}{\varepsilon^* + 2\varepsilon_h} = \frac{4\pi}{3} N\alpha \tag{7}$$

Please cite this article as: Torrell M, et al, Nanoscale color control of TiO₂ films with embedded Au nanoparticles, Mater Lett (2010), doi:10.1016/j.matlet.2010.08.031

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Fig. 3. Calculated and experimental optical absorption spectra of two TiO_2/Au films annealed at 500 and 800 °C, respectively (Au NPs 12 at %).

where *N* is the particle number per unit volume. However, in the 154 MMG the polarizability is renormalized due to the dipole-dipole 155interactions between the particles and therefore this approach is valid 156for higher volume fractions of inclusions (see Refs [4,23] for further 157158details). For the modeling purposes, the complex dielectric function of gold is used, where two extra contributions representing inter-band 159transitions have been included in addition to the Drude term (2) as 160 proposed in the bibliography [24]. Dispersion of the refractive index of 161 the TiO₂ matrix was taken into account according to the semi-162163 empirical relations of Ref [25]. The value of $n_h(\lambda = 550 \text{ nm})$ was fitted by assuming a certain percentage of the amorphous and crystalline 164TiO₂ phases and voids. 165

166 Fig. 3 shows the modeling results of two representative spectra 167 demonstrating the effects of the refractive index of the matrix (SPR band 168position) and NP's shape (band splitting). It is clearly seen that there are two separate resonances already for moderately elongated NPs (e = 0.6169corresponds to the aspect ratio of approximately 1.25). These changes in 170the position and shape of the absorption band result in the variation of 171 the film color coordinates (Fig. 1). The experimental spectra of Fig. 3 also 172show a long absorption tail extending to the near-infrared, not 173 reproduced by the modelling. This absorption is attributed to more 174 complex shapes gold inclusions. This assumption is supported by the 175TEM micrographs of Fig. 2 and it is known that metallic fractal clusters 176 indeed can produce broad absorption spectra [26]. 177

178 4. Conclusions

In conclusion, we have shown that the tuning of the Surface Plasmon
Resonance is possible through the nanoscale control of the nanoparticle
shape and the refractive index of the matrix, both being achieved by
means of annealing treatments at appropriate temperatures.

183 Acknowledgements

This research is sponsored by FEDER funds through the program
 COMPETE-Programa Operacional Factores de Competitividade and by
 249

national funds through FCT-Fundação para a Ciência e a Tecnologia, 186 under the project PTDC/CTM/70037/2006. 187

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