Characteristics of Pd and Pt Nanoparticles Produced by Nanosecond Laser Irradiations of Thin Films Deposited on Topographically-Structured Transparent Conductive Oxides

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Abstract: Pd and Pt nanoparticles on Fluorine-doped tin oxide (FTO) are produced. This outcome is reached by processing nanoscale-thick Pd and Pt films deposited on the FTO surface by nanosecond laser pulse. Such laser processes are demonstrated to initiate a dewetting phenomenon in the deposited metal films and lead to the formation of the nanoparticles. In particular, the effect of the film’s thickness on the mean size of the nanoparticles, when fixed the laser fluence, is studied. Our results indicate that the substrate topography influences the dewetting process of the metal films and, as a consequence, impacts on the nanoparticle characteristics. The results concerning the Pd and Pt nanoparticles’ sizes versus starting films thickness and substrate topography are discussed. In particular, the presented discussion is based on the elucidation of the effect of the substrate topography effect on the dewetting process through the excess of chemical potential. Finally, Raman analysis on the fabricated samples are presented. They show, in particular for the case of the Pd nanoparticles on FTO, a pronounced Raman signal enhancement imputable to plasmonic effects.

Keywords: Pd; Pt; FTO; laser irradiations; dewetting; nanoparticles

1. Introduction

Palladium (Pd) and Platinum (Pt) nanoparticles (NPs) are widely exploited to design and fabricate nanostructured innovative sensing, catalyst, electronic, energy production and storage devices [1–8]. From a general point of view, this type of nano- system captures great interest due to size-dependent structure and properties, high specific surface area and reactivity. From a specific point of view, for example, some notable chemical reactions occurring at surfaces take advantage of the enhanced catalytic activity of Pd and Pt when in the nanoscale form [9–13]. In addition, the physico-chemical properties of Pd and Pt NPs can be largely tuned by the control of their size and shape. This is an important property of the NPs for several technological applications. A critical issue is the formation of NPs with a desired size that should take place directly on a support. Hence, for the exploitation of these nanostructures in real devices, the design of simple, cost-effective, versatile, high-throughput fabrication methods on suitable supporting surfaces, allowing desired NPs size and shape control, is of paramount importance [14]. Obviously, the detailed understanding of the basic microscopic mechanisms governing the process involved is crucial in assuring the desired NPs control.
So, we report here on the exploitation of a laser-based approach to fabricate Pd and Pt NPs supported on a functional surface. The approach is based on the sputtering-deposition of nanoscale-thick Pd or Pt films onto the surface of a Fluorine-doped Tin Oxide (SnO$_2$:F) layer on glass (soda-lime), that is, Fluorine-doped tin oxide (FTO/glass) substrate. FTO is a transparent conductive oxide which use is largely diffused as coating in solar cell devices or for photoelectrochemical reactions on surface [15–19]. Then, nanosecond laser irradiations were carried on the metallic films surface so to exploit the potentialities of the laser-matter interaction for surface nanostructuration [20–30]. In particular, we processed, by the laser pulse, Pd or Pt films of various thickness d [20–37]. The molten-phase dewetting process of the Pd and Pt films is observed to lead to the formation of NPs for which the mean diameter $<D>$ and mean surface density $<N>$ are quantified. Our results show, in particular, the effect of the FTO surface topography on the resulting NPs size through the geometry-dependent chemical potential, an effect which is absent for the dewetting phenomenon of thin metal films on flat surfaces [22–32]. This effect is discussed starting from previous results on metallic film dewetting occurring on intentionally patterned surfaces [33–36]. We show that this effect is crucial in the present case since the used FTO substrate (similarly to standard FTO used in real devices) does not present a flat surface on a microscopic scale. Instead, due to standard deposition processes [17,38,39], the FTO layer on the glass slide surface is obtained as structured in micrometric randomly-arranged FTO pyramids.

Finally, we present, also, Raman analysis on the fabricated samples finding, for the case of the Pd NPs, a pronounced Raman signal enhancement imputable to intense plasmonic effects.

From a general point of view, this work completes our previous studies on the molten phase dewetting of pure (Au) or alloy (AuPd) metals films on transparent conductive oxides [17,40,41], extending these studies to other metals such as Pd and Pt whose nanostructuration by laser pulsed-induced dewetting is rarely studied (Pt) [42,43] or, practically, absent (Pd).

2. Materials and Methods

Fluorine-doped Tin Oxide (SnO$_2$:F, FTO)/glass (soda-lime) slides, from KINTEC factory (Hung Hom, Kowloon, Hong Kong) [44], were the starting substrates. According to our previous characterizations [17], the average transmittance of these slides, corresponding to incident light wavelength from 400 to 1100 nm, is 73.2% and the FTO sheet resistivity is 8.6 ohm/sq. For each metal deposition, some glass/FTO slides were inserted within the vacuum chamber of a RF Emitech K550X sputter coater (Quorum Technologies, East Sussex, UK). In addition, Si slides were introduced with the glass/FTO slides. They served as reference samples for successive check of the effective thickness of the deposited metal films. Pd or Pt films were sputter-deposited onto the FTO surface using 99.999% purity Pd or Pt targets using Ar as sputtering gas. In particular, the control of the effective thickness d of the deposited metal films is reached by the control of the deposition time (in the range 1–240 s) and the emission current (in the range 5–50 mA). Successive Rutherford backscattering spectrometry analysis (using 2-MeV $^4$He$^+$ incident ions and backscattered at 165°) on the metal films deposited on the reference Si slides allowed the association of several couples (deposition time, emission current) to a specific thickness d of the Pd or Pt film. For the analysis presented in this work we chose, in particular, $d_{\text{Pd}} = 3.0, 7.5, 17.6, 27.9$ nm for the Pd films, and $d_{\text{Pt}} = 1.9, 7.5, 12.2, 19.5$ nm for the Pt films, with a measurement error of 5%.

The metal films supported on the FTO surface of the glass/FTO slides were processed by nanosecond laser irradiation (one pulse per film). In order to carry out the irradiations, we used a nanosecond pulsed (12 ns) Nd: yttrium aluminum garnet YAG laser working at 532 nm (Quanta-ray PRO-Series pulsed Nd:YAG laser, Spectra Physics, Santa Clara, CA, USA). In our experimental setup, the laser spot is circular with a diameter of approximately 4 mm. A Gaussian shape characterizes the spatial distribution of the laser intensity. Specifically, the laser intensity decreases from the chosen maximum value to the 97% of this maximum value within a circular area of 600 µm in diameter. For the experiments presented in this work, a laser fluence having as maximum 0.50 J/cm$^2$ (measurement
error of 0.025 J/cm²). So, we consider that within a circular spot with diameter 600 μm centered on the entire laser spot (4 mm in diameter), the metal film was irradiated by one pulse with 12 ns duration and fluence 0.50 J/cm². All the analyses refer, then, to the NPs formed in this region.

Microscopic morphological analyses were carried out by using a scanning electron microscope (SEM, Zeiss FEG-SEM Supra 25 Microscope, Carl Zeiss Microscopy, New York, NY, USA). The Gatan Digital Micrograph software (version 3.0, GATAN Inc., Pleasanton, CA, USA) was used to extract quantitative information by the SEM images. In particular, to extract from the SEM images the mean diameter <D> of the metal NPs, for each image a threshold on the brightness of the image was set, so that the bright regions in the image, with intensity value 1, represent the NPs and the dark regions, with intensity value 0, represent the supporting substrate. The diameter D of a NP is measured as the diameter of the smaller circle inscribing the NP. The mean value <D> of the diameter of the NPs (and the corresponding error arising as the standard deviation) for each sample has been quantified on a statistical population of 400 NPs. In addition, the mean surface density <N> (number of particles per unit area) was evaluated by direct counting and averaging on several SEM images.

Raman spectra on the samples were acquired by using a micro-Raman spectrometer (Confocal Raman–AFM SNOM, WITec ALPHA300RS, Ulm, Germany) equipped with a charge-coupled device (CCD) system. A 532 nm laser line (output power ~32 mW, power on the sample ranging from 0.5 to 10 mW), a grating with 600 grooves mm⁻¹ and a 100× differential interference contrast (DIC) objective (NA 0.9) were used.

3. Results and Discussion

First, we analyzed, by SEM imaging, the topography of the reference FTO surface. So, Figure 1 reports a typical SEM image of the bare FTO surface. It appears that the FTO layer on the glass substrate has a complex non-flat topography. In fact, by the inspection of this plan-view image, the FTO layer results to be formed by micrometric structures having an approximately truncated-pyramidal shape and which, randomly arranged, cover the entire surface of the glass slide. We point out that this is the standard morphology for deposited FTO layers [38,39] and that it is desired in several applications (solar cells, etc.) due to its ability in efficiently scatter light [17,18,39,41]. Then, in Figures 2 and 3, some exemplificative microscopies of the FTO surface covered by the Pd and Pt films are reported. Figure 2 refers to the Pd covered FTO surface with increasing effective thickness of the covering metal film: \( d_{\text{Pd}} = 3.0 \) nm (a) and \( d_{\text{Pd}} = 27.9 \) nm (b). Figure 3 refers to the Pt covered FTO surface with increasing effective thickness of the covering metal film: \( d_{\text{Pt}} = 1.9 \) nm (a) and \( d_{\text{Pt}} = 19.5 \) nm (b). Increasing the amount of sputtered Pd or Pt (i.e., increasing the effective thickness of the deposited metal film), an evolution of the surface morphology can be observed: the pyramidal structuration of the FTO layer is yet recognizable; however, over the pyramids’ surfaces a nano-granular morphology develops (see, in particular, Figures 2b and 3b)

![Figure 1](image-url)  
*Figure 1.* FTO bare surface as imaged by SEM. The surface structuration in pyramidal-like structures is recognizable.
This morphology is the standard one, in the initial stages of growth (i.e., nucleation and growth) for metal films deposited on non-metal substrates and which leading growth process is the Volmer-Weber growth mode [3,17,25,26,31,41,45–47].

![Figure 2](image1.png)

**Figure 2.** SEM images of the FTO surface covered by Pd film with effective thickness of: (a) 3 nm; (b) 27.9 nm. The presence of the films can be recognized in more rough pyramids surfaces with respect to bare surfaces.

![Figure 3](image2.png)

**Figure 3.** SEM images of the FTO surface covered by Pt film with effective thickness of: (a) 1.9 nm, (b) 19.5 nm. The presence of the films can be recognized in more rough pyramids surfaces with respect to bare surfaces.

However, in late stages of growth, increasing the amount of deposited atoms, the nucleated small metal islands grow until to contacting each other giving rise to a coalescing process [45–47]. In this stage, a percolative surface morphology of the metal film is obtained in the sense that it is formed by ramified interconnected nano-islands separated by small gaps [45–47]. Finally, continuing to deposit more and more material, holes in the growing metal film are filled to form a continuous rough film [45–47].

On each metal-covered sample, we carried out the laser irradiation. Then, we used the SEM analysis to study the metal surface morphology evolution versus the metal film thickness, see Figures 4 and 5. In general, the pulsed laser-induced dewetting of the metal film is observed, that is, the 0.50 J/cm² laser irradiation for 12 ns causes the melting of the Pd or Pt film followed by the film rupture and retreating processes towards the formation of NPs of circular section (Figures 4 and 5). The spherical (or almost-spherical) shape of the NPs can be easily recognized from the SEM images in Figures 4 and 5 where in several cases NPs are observed from a tilted configuration due to the surface structuration of the FTO surface (i.e., the NPs placed on the lateral surfaces of the FTO pyramids).

In fact, generally, metal films on transparent conductive oxides are highly non-wetting systems [26] so that, after the dewetting process, spherical or almost-spherical NPs are obtained (i.e., maximization of the contact angle).
These perforations are likely to occur where the film is thinner since the fluence required for melting present experiments for which the pulse duration is \( \tau \). then, then split into droplets by the Rayleigh instability \([24–27]\) and minimizing the total surface energy that this dewetting process starts by the film perforation which leads to holes formation in the film. Dewetting \([17,22–27,41–43,48]\) followed by the film's solidification. Shortly, it is widely accepted that this dewetting process starts by a molten-phase dewetting process. In fact, in standard metals (such as Pd and Pt) the thermal equilibrium between hot electrons and phonons is established, typically, within a characteristic time \( t_{eq} \approx 50 \text{ ps} \) \([17,22–27,41–43,48]\). Using nanosecond pulsed laser to generate heat in the film, as in the present experiments for which the pulse duration is \( \tau = 12 \text{ ns} \) so that \( \tau >> t_{eq} \) then the metal film melting dynamics is the main process \([17,22–27,41–43,48]\). Therefore, in the present experiments, the Pd and Pt dewetting process occurs with the films being in the molten state (i.e., spinodal dewetting \([17,22–27,41–43,48]\)) followed by the film’s solidification. Shortly, it is widely accepted that this dewetting process starts by the film perforation which leads to holes formation in the film. These perforations are likely to occur where the film is thinner since the fluence required for melting is lower for thinner films \([24]\). Deposited metal films, in fact, are, typically, not perfectly uniform in thickness presenting, instead, a natural roughness (local variation in thickness). The metal at the hole edges forms the so-called rims which retract drawing away from the center of the hole. The retreat ing molten films in neighboring holes coalesce into metal filaments (some solidified Pd filaments are recognizable, for example, in Figure 4a). These filaments are thermodynamically unstable so that, then, then split into droplets by the Rayleigh instability \([24–27]\) and minimizing the total surface energy of the system. In the final stage, after solidification, an array of metal NPs uniformly covering the laser-irradiated surface is obtained.

Figure 4. SEM images of the FTO surface covered by Pd film after the 0.50 J/cm\(^2\) laser pulse ((a) 3 nm-thick, (b) 27.9 nm-thick). The formation of almost-spherical particles on the FTO surface is recognizable both in (a,b).

Figure 5. SEM images of the FTO surface covered by Pt film after the 0.50 J/cm\(^2\) laser pulse ((a) 1.9 nm-thick; (b) 19.5 nm-thick). In (a) the Pt film is unchanged, while in (b) the formation of almost-spherical particles on the FTO surface is recognizable.
We proceeded, then, to the quantitative analysis of the Pd and Pt NPs characteristics using the SEM images. In particular, for each sample, distributions of the NPs diameter $D$ were constructed as described in the experimental section. As an example of the procedure, Figure 6a reports a part of the SEM image of Figure 4a concerning the NPs obtained by the laser irradiation of the 3 nm-thick Pd film and in Figure 6b how it was treated in luminosity and contrast to identify the NPs (circular white regions). Using the scale marker, for each circular region the diameter can be identified so to construct the NPs diameters distribution. In addition, by direct counting of the circular white regions, the NPs surface density can be derived.

Figure 6. (a) SEM image showing the NPs produced on the FTO surface after the laser irradiation of the 3 nm-thick deposited Pd film (it is a part of the SEM image in Figure 4a). (b) SEM image in (a) suitably treated in luminosity and contrast to identify the NPs (circular white regions) so to extract NPs size and number per unit area.

Figure 7 reports, then, some examples of obtained diameter distributions: (a,b) are the diameters for the NPs obtained by the laser irradiation of the 3 nm-thick ((a)) and 27.9 nm-thick ((b)) Pd films deposited on the FTO surface; (c,d) are the diameter distributions for the NPs obtained by the laser irradiation of the 7.5 nm-thick ((c)) and 19.5 nm-thick ((d)) Pt films deposited on the FTO surface. As easily recognizable, (a,c) shows a monomodal size distribution while (b,d) bimodal sizes distributions.

Each diameter’s distribution was, hence, statistically analyzed to extract the mean diameter $<D>$ (and the corresponding standard deviation $\Delta D$) of the NPs. In addition, we evaluated, also, the mean NPs surface density $<N>$ and the corresponding errors. The results are summarized in the plots in Figure 8. In particular, performing these analyses, in some samples we observed a monomodal size distribution (one gaussian distribution) indicative of an unique NPs population to which was associated an unique mean diameter $<D>$ and mean NPs surface density $<N>$. In other samples, we observed a bimodal size distribution (two gaussian distributions) indicative of two NPs sub-populations to which were associated mean diameters $<D_1>$ and $<D_2>$ and mean NPs surface densities $<N_1>$ and $<N_2>$.

Figure 7. Cont.
Figure 7. Examples of NPs diameter distributions: The diameters distributions for the NPs obtained by the laser irradiation of the (a) 3 nm-thick and (b) 27.9 nm-thick Pd films deposited on the FTO surface; The diameters distributions for the NPs obtained by the laser irradiation of the (c) 7.5 nm-thick and (d) 19.5 nm-thick Pt films deposited on the FTO surface. As easily recognizable, (a,c) shows a monomodal size distribution while (b,d) bimodal sizes distributions.

Figure 8. Mean diameter $<D>$ ((a)) and mean surface density $<N>$ ((b)) of the Pd NPs obtained by the 0.50 J/cm² laser irradiation of the 3 nm, 7.5 nm, 17.6 nm, 27.9 nm thick deposited Pd films. Mean diameter $<D>$ ((c)) and mean surface density $<N>$ ((d)) of the Pt NPs obtained by the 0.50 J/cm² laser irradiation of the 1.9, 7.5, 12.2, 19.5 nm-thick deposited Pt films. The indication of two values for $<D>$ or $<N>$ for the same sample means the presence of two different NPs sub-populations in that sample.

Specific values for $<D_1>$, $<D_2>$, $<N_1>$ and $<N_2>$ are, further, summarized in Table 1 for the various samples.

To discuss these data, first of all we recall some basis of the thin metal film melting process induced by pulsed laser irradiations. In the first approximation, a simple model [24,26] considers the metal film of thickness $d$ as irradiated by one laser pulse of duration $\tau$ with a top-hat temporal profile and establishes (as reliable hypothesis) the total heat (per unit area) generated in the film as $S = I\tau(1-r)[1-\exp(-\alpha d)]$ being $I$ the laser pulse intensity (W/cm²), $r$ the film reflectivity at the laser wavelength, $\alpha^{-1}$ the film optical absorption length at the laser wavelength ($\alpha$ is called the absorption coefficient). Note, however, that for a thin film, $r$ and $\alpha$ depend on $d$. On the basis of this model, the consequent temperature rise in the thin film is $\Delta T = S/Cd$ where $C$ is the film heat capacity per unit
volume. So, the melting (and molten-phase dewetting) characteristics of a thin metal film depend on optical and thermal characteristics of the film \((d, r, \alpha, C, \text{melting temperature})\). In principle, differences in these parameters for Pd and Pt justify the observed differences in \(<D_1>\) and \(<N_1>\) when fixed the film thickness, the laser fluence and pulse duration. This is the case for the 7.5 nm-thick Pd and Pt films for which, on the same substrate, two Pd NPs populations and an unique Pt NPs population are, respectively, obtained. On the other hand, it is easy to understand why the laser irradiation of the 1.9 nm-thick Pt film does not produce a dewetting process (i.e., formation of the NPs). In this case, clearly, \(d<<\alpha^{-1}\) (for metal films, generally, \(\alpha^{-1}\approx10\text{ nm}\)) and/or \(r\) is so low that the Pt film does not absorb energy from the laser pulse: the film temperature does not change and the film is unaffected by the laser irradiation.

Table 1. Summary of the values of \(<D_1>, <D_2>, <N_1>, <N_2>\) for the NPs in the samples.

| Thickness of the Film-Material | \(<D_1>(\text{nm})\) | \(<N_1>(\times10^9\text{ cm}^{-2})\) | \(<D_2>(\text{nm})\) | \(<N_2>(\times10^9\text{ cm}^{-2})\) |
|-------------------------------|------------------------|---------------------------------|------------------------|---------------------------------|
| 3 nm-Pd                       | -50                    | -4.5                            | 0                      | 0                               |
| 7.5 nm-Pd                     | -75                    | -2                              | -100                   | -0.5                            |
| 17.6 nm-Pd                    | -70                    | -1.5                            | -150                   | -1.5                            |
| 27.9 nm-Pd                    | -100                   | -2                              | -220                   | -2                              |
| 1.9 nm-Pt                     | 0                      | 0                               | 0                      | 0                               |
| 7.5 nm-Pt                     | -70                    | -3.2                            | 0                      | 0                               |
| 12.2 nm-Pt                    | -75                    | -0.6                            | -220                   | -0.6                            |
| 19.5 nm-Pt                    | -70                    | -1.6                            | -180                   | -2.7                            |

Shortly, we also discuss the laser irradiation effect on the underlaying FTO substrate. In fact, the melting temperatures for Pt and Pd are, about, 2040 and 1828 K, respectively, while the melting temperature for the SnO\(_2\) is around 1900 K. So, in the laser irradiation processes we could expect, also, a partial melting of the FTO substrate due to the high temperatures achieved for melting, at least, the Pt films. To discuss this point, for example, we recall the results obtained by Font et al. [49] regarding the heat transfer mechanisms involved in thin metal films melting on a SiO\(_2\) layer and exposed to laser irradiation. In this work, the authors present a model describing the heat transfer and phase change of the metal/substrate system during nanosecond laser irradiation with parameters similar to those presented in our work. In particular, Font et al. [49] considered a Cu film/10 nm-thick SiO\(_2\) layer/Si substrate system irradiated by a pulsed laser characterized by a Gaussian beam, and they studied the effect of the laser parameters, including energy density and pulse duration, on possible melting of the SiO\(_2\) layer. Interestingly, they found that maximum thickness of the melted SiO\(_2\) region can be of the same order of magnitude as the thickness of the metal film. The calculations carried out by the authors are based on some assumptions. In particular, for example, they supposed that the substrate is completely transparent to the laser radiation, so that the metal film absorbs the energy of the laser and transfers it to the substrate by conduction. This condition is fulfilled, also, by the FTO/glass substrate used for the experiments presented by us. To describe the heating and possible melting of the substrate, they considered, obviously, heat conduction in each layer of the system. In the baying equations, so, the thermal conductivity of the SiO\(_2\) layer (in solid and liquid phases) is involved. This value, however, is very different from that characterizing the FTO layer. In fact, for example, in the solid phase, for SiO\(_2\) typical value for the thermal conductivity is 1.4 W/m·K [49]. Instead, SnO\(_2\) is characterized by a much higher thermal conductivity, of the order of 40 W/m·K [50].

Clearly, the temperature increase in the film depends on how it dissipates the laser-generated heat [24]: concerning a metal film deposited on a low thermal conductive substrate, the laser-generated heat will remain, mostly, confined in the metal film resulting in a higher increase of temperature of the system. This means that for the metal films deposited on the FTO substrate the rapid dissipation of the laser-generated heat through the FTO substrate results in a lower increase of the system temperature compared to the case for which the same metal films are deposited on a lower thermal conductivity
substrate (as SiO₂). In fact, comparing the SEM images in Figures 4 and 5 (Pd and Pt films on FTO after the laser irradiations) to the SEM image in Figure 1 (bare and untreated FTO), it seems clear that the surface morphology of the FTO substrate is unchanged after the laser irradiations. This is an indication of the fact that even if the laser-generated heat determines the temperature increase of the metal films above the film material melting temperature, the relatively high thermal conductivity of the FTO probably results in a temperature increase of the FTO layer below its melting temperature. This is the situation for the 0.50 J/cm² laser pulse. This situation is strongly changed, for example, using a 0.75 J/cm² laser pulse for which the SEM images (not shown) clearly showed a dramatic change of the FTO surface morphology: in this case, the FTO pyramids are no more recognizable being the FTO surface more similar to a flat surface. This is, probably, consequence of the higher temperature reached by the FTO layer, higher than the FTO melting temperature. In this case, so, also the FTO layer melts during the laser irradiations and, also, an FTO morphology change is involved in the overall process. This is a further reason for which we limit the data presented here to the 0.50 J/cm² laser pulse, so involving only the Pd and Pt molten-phase dewetting process.

Regarding the standard dewetting process of metal films on flat surfaces [22–31], a unique NPs population is expected with a mean size, which increases by increasing the starting thickness of the deposited film. The observation of the formation of two different NP populations in some samples is, thus, indicative of the effect of the substrate non-flat surface topography on the film dewetting process, influencing, then, the size of the NPs. To describe this effect we, briefly, discuss some of the results due to Gierman and Thompson [33]. They intentionally patterned, by using electron beam lithography, a SiO₂ surface in spatially ordered arrays of inverted pyramid shaped pits. However, different patterns were designed by changing the spatial period of the pits (175 or 377 nm) and the pit-to-mesa width ratio (1.5, 1.9, 5.3). Onto the so-patterned SiO₂ surface, Au films of different thickness (16, 21, 32 nm) were evaporated. The Au films solid-state dewetting process was activated by 800 °C annealing process. The microscopic analysis revealed for the resulting Au NPs originated in the dewetting process, a spatial arrangement on the substrate surface and size distribution dependent on the geometric characteristics of the surface pattern (spatial period of the pits and pit-to-mesa width ratio) in combination with the Au film thickness. On the basis of the combination of these parameters: in some cases the preferential formation of a single NP per pits with no NPs on the mesa was observed; in other cases, the formation of one NP per pit with some other NPs on the mesa was achieved; in further cases, the generation of multiple NPs per pit with some other NPs on the mesa resulted; finally, some other combinations lead to the formation of multiple NPs per pit with some other NPs on the mesa. These results clearly demonstrated the crucial effect of the surface topography on the deposited metal film dewetting also giving the possibility to study, quantitatively, this effect taking into account the excess of local chemical potential as modulated by the finite local curvatures on the substrate surface [33]. In fact, considering the dewetting process of a thin metal film on a flat surface, it is clear that the driving force for the process is, solely, the minimization of the total surface and interface energy of the system. For a flat surface, the local radius of curvature is, in each surface point, $R \rightarrow \infty$. The corresponding local curvature is $\kappa = 1/R$ to which is related, by the Gibbs–Thomson relation, the local excess of chemical potential $\Delta \mu = \kappa \gamma \Omega \rightarrow 0$ (being $\gamma$ the film surface energy and $\Omega$ film atomic volume. Consequently, the dewetting process of a thin film on a flat surface is not affected by the surface topography. The situation changes if the surface is not flat, that is, if it presents finite local curvatures (as in the case of the patterned surfaces used by Giermann and Thompson). In this case, due to the finite values of $R$, the non-zero values of $\Delta \mu = \kappa \gamma \Omega$ introduce an additional driving force for the film dewetting process determining, in particular, a preferential material diffusion from the positions with $\kappa > 0$ (peaks or ridges) to the positions with $\kappa < 0$ (valleys), see the representative scheme in Figure 9. It is clear, then, the modulation of $\Delta \mu$ through the surface geometry impacts on the dewetting process and on the resulting characteristics (spatial arrangement and size) of the NPs originating from this process. Referring to the specific case reported by Giermann and Thompson [33], which is particularly useful for the interpretation of our data, the edge
of a pit is characterized by $\Delta \mu \propto 1/R_A > 0$ while the apex of an inverted pyramid is characterized by $\Delta \mu \propto -1/R_B < 0$, see Figure 9. It follows that, to minimize the total surface and interface energy of the system and to establish a global condition for which $\Delta \mu = 0$, the film material will diffuse away from the pit edge toward the pit apex and here, preferentially, will form the NPs. However, since the local curvature at the pit edge and the apex decreases with increasing film thickness, the driving force for flow from the edge to the apex also decreases with increasing film thickness. So, overall, the effect becomes dependent on the film thickness.

![Figure 9](image_url) Representation of a thin film conformally covering a patterned surface. The curvature at the pit edge, $R_A$, and at the inverted apex, $R_B$ are shown. Reprinted with permission from [33]. Copyright 2005 AIP Publishing.

In our case, we observed that increasing the thickness of the Pd film from 3.5 to 27.9 nm and of the Pt film from 7.5 to 19.5 nm, the NPs population evolves from monomodal to bimodal. This indicates the existence of a critical film thickness $d_c$ identifying a variation in the dewetting characteristics of the film, as pictured by the combination of Figures 10 and 11. For $d < d_c$ (generally named “low-thickness” condition) the substrate topography does not affect the film dewetting process (Figure 10). In this condition, the film does not interact with the topographic features of the substrate and, then, the substrate behaves as a flat substrate with respect to the film dewetting. Hence, the film dewets as on a planar substrate originating NPs uniformly arranged over the surface of the substrate. In addition, the size of the dewetted NPs, being the starting thickness of the deposited film very low with respect to local surface curvatures (defined by the characteristic sizes of the FTO pyramids, i.e., height, width, spacing), is not influenced by the substrate topography. For $d > d_c$ (generally named “high-thickness” condition) the geometrical features of the substrate affect the film dewetting process and, hence, the resulting mean size of the formed NPs (Figure 11). In this case the thickness of the deposited film is, at least, comparable to the local surface curvatures. Then, the film dewetting is driven, in addition to the surface energy minimization, by the surface topography through the local excess of chemical potential. This results in the preferential formation of the NPs in-between or over the pyramids. It is clear, hence, that the characteristic sizes of the FTO pyramids affects the final NPs mean size. The overall effect is determined by an interplay between these characteristic sizes (establishing the local surface curvatures) and the thickness $d$ of the film. Therefore, observing Figure 8, in that samples where an unique population of NPs is obtained, the situation $d < d_c$ is realized (dewetting as on a flat surface), whereas in that samples where a two NPs sub-populations are obtained, the situation $d > d_c$ is realized (substrate topography-driven dewetting). Then, we can conclude, also, that $3 \text{ nm} < d_c < 7.5 \text{ nm}$ for the Pd film and that $7.5 \text{ nm} < d_c < 12.2 \text{ nm}$ for the Pt film. However, for sake of completeness, we also mention that, according to previous experimental and theoretical studies on dewetting of films on patterned surfaces [51–54], a critical thickness $d_c$ for the film separating different effects of the substrate topography on the dewetting process is related to additional parameters than the local excess of chemical potential $\Delta \mu$. In particular, it is known that a fixed substrate pattern can influence the dewetting pathways of supported film depending on the film thickness by acting on the
length-scale of the processes (holes nucleation, film retreating, Rayleigh instability, etc.) involved in the dewetting phenomenon. Dewetting on a flat substrate progresses with the formation and growth of randomly placed holes with a certain mean length-scale. Eventually, coalescence of holes produces an isotropic collection of droplets. The average diameters of the dewetted structures depend on the initial film thickness. On the other hand, specific spatial periodicity of patterns on the substrate can act on a deposited film by imposing conditions on the length-scale of the processes involved in the dewetting phenomenon (in addition to the effect imposed by local excess of chemical potential on the material film diffusive processes). An a-priori theoretical determination of the critical thickness $d_c$ should take into account, also, such effects but this is out of the scope of the present experiments.

**Figure 10.** Picture of the result of the “low-thickness”-film dewetting process on the FTO topographically structured surface. Small NPs are uniformly distributed over the entire FTO surface (as on a flat surface).

**Figure 11.** Picture of the result of the “high-thickness”-film dewetting process on the FTO topographically structured surface. Large NPs are preferentially formed in-between or over the FTO pyramids.

To conclude the analyses of the characteristics of the Pd and Pt NPs produced on the FTO substrate, we performed, also, Raman analysis. Figure 12 displays, first of all, the Raman spectrum of the bare FTO/glass substrate, in black. There are two fundamental Raman scattering peaks which are those characteristic of rutile SnO$_2$ single crystal [55]. For pure SnO$_2$, the characteristic and intense band at 625 cm$^{-1}$ is due to the $A_{1g}$ vibration mode of SnO$_2$. The weak band around 478 cm$^{-1}$ is due to the $E_g$ vibration modes of SnO$_2$ [55]. Figure 12 reports, also, as examples of the general behaviour, Raman spectra of the FTO/glass substrate covered by Pd or Pt NPs originating from the laser heating of the deposited films: in red the spectrum of the substrate covered by Pd NPs obtained by the laser irradiations of the 27.9 nm-thick Pd film; in blue the spectrum of the substrate covered by Pd NPs obtained by the laser irradiations of the 17.6 nm-thick Pd film, in green the spectrum of the substrate
covered by Pt NPs obtained by the laser irradiations of the 19.5 nm-thick Pt film. The general effect is that, compared with bare FTO, the intensity of Pd NPs/FTO samples become stronger.

![Raman spectra](image)

**Figure 12.** Raman spectra corresponding to the bare FTO substrate (black), FTO covered by Pd NPs obtained by the laser irradiation of the 27.9 nm-thick Pd film (red) and of the 17.6 nm-thick Pd film (blue), FTO covered by Pt NPs obtained by the laser irradiation of the 19.5 nm-thick Pt film. SERS effect can be particularly recognized in the blue spectrum.

On the contrary, the presence of the Pt NPs on the FTO surface, does not influence the FTO peak intensity, indicating no Pt NPs-FTO interaction. In particular, the Pd NPs SERS (Surface Enhancement Raman Scattering) enhancement may be due to Pd NPs surface plasmon resonance [5,56,57]. However, surface plasmon resonance effects also for Pt NPs are expected [5]. The reasons for which these effects lead to intense SERS in the Pd NPs/FTO samples while no SERS effects in the Pt NPs/FTO samples is actually under investigation as a perspective work, requiring a crossing of experimental and simulation-based results. Here, we limit to consider that the SERS enhancement should arise from the effective resonance coupling between the localized surface plasmon of the NPs and the characteristic parameters of the energy band structure of the FTO. A match should be realized in the case of the Pt NPs and FTO, contrary to the case of the Pd NPs and FTO. Besides, several other factors, such as the Ohmic loss, non-radiative Förster energy transfer, lower surface plasmon radiative efficiency, may be responsible for the absence of SERS enhancement in the Pt NPs-FTO systems and these effects have to be considered in a future complete description. However, we mention, for example, that the optical loss of NPs material can be used as a measure to predict the effect in SERS, whereas the optical loss is intended as the imaginary part ($\varepsilon_2$) of the dielectric constant $\varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega)$ [58]. In fact, the imaginary part of the dielectric constant causes Ohmic damping of the electrons oscillations in the metal. Our SERS data seem in agreement with the consideration that several experimental and theoretical works found higher values of $\varepsilon_2$ for bulk, thin films, NPs materials of Pt than of Pd in the UV range [59–61].

We observe, finally, in Figure 12, a shift (toward lower wavenumbers) of the $E_g$ peaks in the Pd NPs/FTO samples with respect to the bare FTO substrate, while this shift is absent in the Pt NPs/FTO samples. It is absent, also, before the laser treatment so that it cannot related to substrate modifications due to the sputtering deposition processes. From a general point of view, in Raman spectra, shifting of peaks towards lower or higher wavenumber is related to several factors [62] such as change in the chemical bond length within the sample (the shorter bond length causes to shift higher wavenumber or vice versa), changes in temperature and stress within the sample, and so on. More specifically, a shift in
Raman peaks could be due to the addition of oxygen in the sample. In this regard, the fact that this shift is present only in the case of the Pd NPs could suggest its relation to the Pd NPs surface oxidation [63]. It is widely recognized that Pd NPs are subjected to the interaction with environmental oxygen to form a PdO surface shell [64], while Pt NPs are more stable against this reaction [65]. We can support the conclusion that the Raman peak shift for the sample with Pd NPs could be related to surface oxidation of the NPs, by considering that previous works reported wavelength shift of plasmonic bands for Pd NPs [66] and Ag NPs [67,68] after exposure to air or oxygen with consequent formation of surface oxide (PdO or Ag₂O) for the NPs. To conclude, we recall that the Pd and Pt NPs produced have a quasi spherical shape and we comment on this fact in view of the Raman measurements results: Raman scattering, which is based on inelastic light scattering upon vibrational excitation of molecules and materials is an extremely inefficient process with about one photon out of 10⁷ being inelastically scattered [69]. This drawback can be overcome when the molecules are located near a rough metal surface or metal NPs, which results in a boost of Raman scattering, in the process known as SERS. Upon resonant excitation of strong localized surface plasmon resonances, a greatly enhanced local electromagnetic field (near-field) is generated at the NP surface. This near-field in turn couples to the incident field and re-reemits radiation with the same wavelength (resonant scattering). As the scattering cross section scales with the square of the polarizability (the degree on how easily the electron cloud can be displaced) also the near-field strongly increases for anisotropic NPs due to higher polarizability, as compared to spheres. So, elongated NPs or those with edges and corners, such as nanorods, nanocubes, and nanotriangles, thus generate strong near-fields concentrated at vertices. The presence of the plasmonic near field at the NP surface upon plasmonic excitation increases the polarizability of an adsorbed molecule (compared to a free molecule) due to mutual excitation between induced dipole in the molecule and induced dipole in the NP and leading to enhanced Raman scattering. On the basis of these considerations, a further perspective on the present work lies in the establishment of laser process parameters to controllably produce complex-morphology Pd and Pt NPs to reach higher SERS enhancements. This could be reached, for example, by using femtosecond pulsed laser irradiations [20,21].

4. Conclusions

Pd and Pt NPs onto non-flat FTO substrate were fabricated by exploiting the molten-phase dewetting process of deposited films. The Pd and Pt films were sputter-deposited on the FTO surface, this surface being constituted this surface by FTO microscopic pyramids randomly distributed over a glass slide. The dewetting process of the liquid metal films was initiated by nanosecond laser irradiations at 0.5 J/cm², resulting, after solidification, in the formation of Pd or Pt NPs onto the laser-irradiated FTO surface. However, analyzing the mean diameter <D> of the formed NPs versus the thickness d of the deposited films, an effect of the substrate topography was found. This effect was discussed by the description of the substrate topography influence on the excess of chemical potential driving the dewetting process. We observe that increasing the thickness of the Pd film from 3.5 to 27.9 nm and of the Pt film from 7.5 to 19.5 nm, the NPs population evolves from monomodal to bimodal. This indicates the existence of a critical film thickness d_c identifying a change in the dewetting characteristics of the film: for d < d_c the substrate topography does not influence the film dewetting process, while for d > d_c the topography of the substrate is crucial in determining the dewetting process and, so, the final mean size of the produced NPs. In that samples where an unique population of NPs is obtained, the situation d < d_c is realized (dewetting as on a flat surface), whereas in that samples where a two NPs sub-populations are obtained, the situation d > d_c is realized (substrate topography-driven dewetting). By analyzing the experimental results, we concluded that 3 nm < d_c < 7.5 nm for the Pd film and that 7.5 nm < d_c < 12.2 nm for the Pt film. Finally, the analysis of the Raman spectra for the bare FTO/glass sample and for the Pd NPs/FTO/glass samples and Pt NPs/FTO/glass samples indicated a SERS effect in the Pd NPs/FTO/glass samples due to the effective coupling between the
localized surface plasmon of the Pd NPs and the FTO susbstrate while this effect is absent in the Pt NPs/FTO/glass samples.

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