Sub-surface laser nanostructuring in stratified metal/dielectric media: a versatile platform towards flexible, durable and large-scale plasmonic writing


a Department of Materials Science and Engineering, University of Ioannina, Ioannina, GR-45110, Greece.
b School of Science and Technology, Nottingham Trent University, Nottingham, NG11 8NS, United Kingdom.
c Department of Physics, Aristotle University of Thessaloniki, Thessaloniki, GR-54124, Greece.
d Faculty of Arts, Computing, Engineering and Sciences at Sheffield Hallam University, Sheffield, S1 1WB, United Kingdom.

*Author to whom correspondence should be addressed. email: nikolaos.kalfagiannis@ntu.ac.uk

Laser nanostructuring of pure ultrathin metal layers or ceramic/metal composite thin films has emerged as a promising route for the fabrication of plasmonic patterns with applications in information storage, cryptography, and security tagging. However, the environmental sensitivity of pure Ag layers and the complexity of ceramic/metal composite film growth hinder the implementation of this technology to large-scale production, as well as its combination with flexible substrates. In the present work we investigate an alternative pathway, namely, starting from non-plasmonic multilayer metal/dielectric layers, whose growth is compatible with large scale production such as in-line sputtering and roll-to-roll deposition, which are then transformed into plasmonic templates by single-shot UV-laser annealing (LA). This entirely cold, large-scale process leads to a subsurface nanoconstruction involving plasmonic Ag nanoparticles embedded in a hard and inert dielectric matrix on top of both rigid and flexible substrates. The subsurface encapsulation of Ag nanoparticles provides durability and long-term stability, while the cold character of LA suits the use of sensitive flexible substrates. The morphology of the final composite film depends primarily on the
nanocrystalline character of the dielectric host and its thermal conductivity. We demonstrate the emergence of a localised surface plasmon resonance, and its tunability depending on the applied fluence and environmental pressure. The results are well explained by theoretical photothermal modeling. Overall, our findings qualify the proposed process as an excellent candidate for versatile, large-scale optical encoding applications.

1. Introduction

Plasmonic materials and devices aim to exploit the unique optical properties of metallic nanostructures to enable routing and manipulation of light at the nanoscale. Lately this field has enabled exciting applications in the areas of chemical and biomedical sensing [1-3], information and communication technologies [4-6], solar energy harvesting [7-12], lighting [13,14], cancer treatment [15,16], optical encoding [17-19] and surface decorations [20] to name a few. A critical parameter in delivering the aforementioned devices is the materials’ preparation methods, which should allow for the production of nanostructures with tunable plasmonic properties. So far, these efforts have been dominated by various techniques such as lithographic processes [21-24], ion beam nanofabrication [25], atomic layer deposition [26], cold welding [27], flash thermal annealing [17,28,29], pattern transfer [30] and template stripping [31,32]. Two issues, however, emerge: Firstly, as far as specific applications are concerned, such as plasmonic writing [17], the protection of these structures from environmental poisoning is of paramount importance; any production process should be able to form plasmonic nanoparticles (NPs) below the dielectric surface. In that context, laser annealing (LA) of AlN:Ag nanocomposites was proposed as a promising route for the production of buried plasmonic NPs for optical encoding [16-18]. LA is a simple patterning tool providing freedom of design, fast processing, compatibility with large-scale manufacturing [18] and allows for the use of inexpensive flexible substrates [33]. Secondly, the scale of the production process would dictate the maximum size of the envisaged devices and their application range in everyday life; processes such as the previously implemented Pulsed Laser Deposition [17] and Confocal Sputtering [16,18] cannot be applied in large-scale due to their
physical limitations and inherent complexity. Therefore, a large-scale and simultaneously cold process is still required for the effective production of buried plasmonic NPs. The deposition of ceramic/metal multilayers and their laser-driven reconstruction would address both the aforementioned limitations, as it is capable of producing metals buried into dielectric ceramics with tunable optical response [32,33], while still compatible with a variety of large-scale deposition configurations such as in-line [34], roll-to-roll [35] or close-field [36] sputtering, which are becoming nowadays the manufacturing routes of choice.

In the present work, we present our entirely-cold route by combining the deposition of ceramic/metal (in particular AlN/Ag and Y$_2$O$_3$/Ag) multilayers with one ultra-short UV LA step. We demonstrate that this LA step is capable of driving the subsurface modification of the metal/dielectric multilayers, and delivers tunable LSPR behavior from Ag NPs that are formed and dispersed in a depth of several nm away from the free surface. This large-scale cold process is very unlike previous attempts dealing with the treatment of thin mono-layered films on top of a rigid substrate [37]. We perform an extensive investigation of the laser-matter interactions in metal/ceramic multilayers, considering the morphological (thicknesses of the individual layers) and microstructural (crystallinity of ceramic layers) features of the multilayers, their thermal conductivity, as well as the fluence of the laser and the environmental pressure during LA. The experimental results are complemented by detailed photo-thermal calculations, which are used to identify the fundamental light-matter interactions and heat diffusion mechanisms in the multilayers, and obtain insights on the basic mechanisms of morphology changes upon LA.

2. Experimental and theoretical methodology

20 Multilayers consisting of alternate thin layers of AlN and Ag were fabricated in an in-house built high vacuum system (base pressure 2×10$^{-6}$ Pa) employing the Dual Cathode Reactive Magnetron Sputtering technique.
Table 1. Summary of the growth and structural features of the AlN/Ag studied samples.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Al Power (Watt)</th>
<th>Ag power (Watt)</th>
<th>AlN crystal structure</th>
<th>Ag layer deposition time (sec)</th>
<th>AlN layer deposition time (sec)</th>
<th>Ag grain size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ML#1)</td>
<td>100</td>
<td>10</td>
<td>wurtzite</td>
<td>50</td>
<td>42</td>
<td>4.7</td>
</tr>
<tr>
<td>(ML#2)</td>
<td>100</td>
<td>10</td>
<td>wurtzite</td>
<td>30</td>
<td>51</td>
<td>4.7</td>
</tr>
<tr>
<td>(ML#3)</td>
<td>100</td>
<td>10</td>
<td>wurtzite</td>
<td>10</td>
<td>60</td>
<td>2.7</td>
</tr>
<tr>
<td>(ML#4)</td>
<td>15</td>
<td>10</td>
<td>amorphous</td>
<td>50</td>
<td>857</td>
<td>4.3</td>
</tr>
<tr>
<td>(ML#5)</td>
<td>15</td>
<td>10</td>
<td>amorphous</td>
<td>30</td>
<td>1028</td>
<td>4.4</td>
</tr>
<tr>
<td>(ML#6)</td>
<td>15</td>
<td>10</td>
<td>amorphous</td>
<td>10</td>
<td>1200</td>
<td>2.5</td>
</tr>
</tbody>
</table>

Additionally, Y₂O₃/Ag multilayers (with individual layer thicknesses equal to the case of ML#5) were grown in a vacuum system (base pressure 5×10⁻⁴ Pa) by the RF magnetron sputtering technique. Both vacuum systems consisted of two 2" unbalanced magnetrons in confocal configuration. All studied samples were deposited on two types of substrates:

i) Czochralski-grown n-type c-Si [100] prime wafers of 1-10 Ωcm resistivity and,

ii) Flexible Poly-EthyleneTerephalate(PET) sheets.

The c-Si substrates were chemically cleaned in an ultrasonic bath, rinsed with de-ionized water and purged with dry N₂ gas. AlN and Ag were deposited from pure Al and Ag targets (2" × 1/4” cylindrical sputter target of purity 99.99%). The formation of the multilayer structure was achieved by the alternate exposure of the substrate to the metal targets in a mixed N₂-Ar plasma conditions. Ar flow rate inside the chamber was set at 12 sccm (leading to a partial pressure of 1.2 Pa) and 8 sccm for N₂ (partial pressure 0.7 Pa). Both gases were 99.999% pure.

Table 1 summarizes the growth and structural features of the AlN/Ag samples studied in this work. Each sample always started and ended with AlN, ensuring the insulation of the metal phase by ambient contamination. The Ag/AlN bilayer was repeated 21 times. The same architecture was followed for the case of Y₂O₃/Ag samples. Two sets of AlN/Ag films were prepared by applying different power values to the Al target, 100 W and 15 W.
Subsequent to growth, LA was performed. The laser source used was an Excimer laser (LAMBDA PHYSIK LPX 305i), which provided un-polarized light pulses of 25 ns in duration and 400-700 mJ/cm²/pulse at 193 nm (ArF). More details on the beam delivery path can be found elsewhere [18,19].

Transmission electron microscopy (TEM) and high resolution TEM (HRTEM) observations were performed using a 200 kV JEOL (2011) LaB6 microscope (point resolution of 0.19 nm and Cs = 0.5 mm). Specimens in cross-section (XTEM) geometry were prepared by the sandwich technique. Mechanical thinning was followed by focused Ar ion milling in a Gatan PIPS and performed in wedge shape using tripod polishing in order to obtain ultrathin samples and minimize the Ar exposure time. To correlate the results with the optical performance, optical reflectance spectroscopy (ORS) measurements were acquired at normal incidence in the spectral range 350–800 nm (1.55–3.54 eV) using deuterium and halogen light sources. The spectroscopic data were normalised to the specular reflectivity of a smooth sputtered Ag film deposited on c-Si(100). X-ray Diffraction (XRD) experiments for the appreciation of the films’ structural quality were carried out in a BRUKER-D8 instrument, equipped with parallel beam optics and a linear strip solid state detector, using the Cu Kα line.

In order to elucidate the heating dynamics involved in our multilayer structures during UV LA, we performed optical and heat transport calculations. In particular, we first solve for the UV light propagation (Maxwell’s equations, solved by the Finite-difference Time Domain (FDTD) method [38-40]) and obtain a detailed map of the absorption profile in the multilayer and the substrate. Our model multilayer system consists of 21 repetitions of a 5nmAg/10nmAlN bilayer caped by 10nm AlN. The index of refraction values at λ=193 nm used in the study are shown in Table 2. The spatial absorption profile is convoluted with the temporal pulse profile (Fig 3(b), inset) and solved in the 1D heat transport equation [41,42]:

\[
c(z)\rho(z)\frac{\partial}{\partial t}T(z,t) = \frac{\partial}{\partial z}[k(z)\frac{\partial}{\partial z}T(z,t)] + Q(z,t)
\]  

where \( \frac{\partial}{\partial t} \), \( \frac{\partial}{\partial z} \) denote partial derivatives with respect to \( t \) and \( z \) respectively, and \( Q(z,t) = a(z) \cdot \phi(t) \) (where \( t \), \( z \) are the time and distance from the top surface respectively) is the laser heating source term defined by the
product of the absorption spatial profile \( a(z) \) and the laser pulse temporal profile \( \phi(t) \), with the total laser fluence given by:

\[
f = \int_{-\infty}^{\infty} \phi(t) dt
\]  

The rest of the terms in the equation are the specific heat capacity \( c \), the mass density \( \rho \) and the thermal conductivity \( k \), all three having a spatial dependence due to the interchange of different materials. Eq. 1 is time-integrated numerically (by a 4th order Runge-Kutta scheme) to obtain the explicit temperature transient. In doing so, we assume that (i) electronic and lattice temperatures are in equilibrium (valid for slow heating times in the ns regime \([42]\)), (ii) radiation and convection losses are insignificant, (iii) material properties (absorption, heat capacity, mass density and thermal conductivity) remain constant during heating. We thus first present our results normalized to the total laser fluence \( f \), i.e. in degrees (K) per mJ/cm\(^2\) and gain valuable physical insight of the thermal transients and thermal gradients that can be developed in the films during LA. Afterwards, we relax assumption (iii) and consider the changes introduced due to a temperature dependent thermal conductivity.

**Table 2.** Values used in the thermal calculations.

<table>
<thead>
<tr>
<th>Material</th>
<th>( n )</th>
<th>( \kappa )</th>
<th>Mass Density (g/cm(^3))</th>
<th>Heat Capacity (J/KgK)</th>
<th>Thermal Conductivity (W/mK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AlN</td>
<td>1.57</td>
<td>0.19</td>
<td>3.26</td>
<td>740</td>
<td>10 - 285</td>
</tr>
<tr>
<td>Ag</td>
<td>1.03</td>
<td>1.18</td>
<td>10.49</td>
<td>235</td>
<td>429</td>
</tr>
<tr>
<td>Si</td>
<td>0.88</td>
<td>2.76</td>
<td>2.33</td>
<td>710</td>
<td>149</td>
</tr>
</tbody>
</table>

**3. Results and Discussion**

As previously mentioned, the AlN crystal structure was varied in the two sets of AlN/Ag multilayers by applying two different power values to the Al target, 100 W and 15 W. This variation leads to different categories of sputtered AlN [19] as indicated by XRD measurements (Fig. 1(a), 1(b)): at 100 W, AlN exhibits XRD peaks corresponding to the (002) and (101) crystal planes of wurtzite AlN (w-AlN) [45]. On the flipside, at 15 W the samples demonstrate a reduction in crystallinity, evident by the absence of any
characteristic peaks for AlN. The Ag(111) peaks of all samples were fitted with Voigt curves and the corresponding Lorentzian broadenings were used to determine the grain size (shown in Table 1) from Scherrer’s formula as described elsewhere [46]. Optical reflectance spectra (ORS) for both “families” of sputtered materials are shown in Fig. 1(c), 1(d). The spectra are in general featureless, resembling the optical response of a bulk metal, except for the two cases of very thin Ag layers (ML#3 and ML#6), were a continuous Ag film has not been achieved and multiple reflections begin to appear accompanied by plasmonic peaks in the lower wavelengths (peaks around 420 and 450 nm for ML#3 and ML#6 respectively).

The produced samples were subjected to LA with an excimer ArF source (193 nm, pulse duration of 20 ns). We found that the optimum seed samples, in terms of their plasmonic response after LA, were ML#3 and ML#5 for w-AlN and a-AlN, respectively. In particular, in the case of w-AlN, the sturdiness of the film and its nanocrystalline character promoted the outdiffusion of Ag when the latter was in high concentration, and therefore only the sample with the lower Ag content (ML#3) developed a plasmonic character after LA. Conversely, in the case of a-AlN, higher Ag concentration (ML#4) made the sample too reflective, whereas
the lower Ag concentration (ML#6) showed poor plasmonic response after LA. Thus, we base our discussion in the best-case scenario for \(w\)-AlN and \(a\)-AlN multilayer structures (samples ML#3 and ML#5, respectively). The effect of a single-pulse LA step onto the structural and optical properties was investigated as a function of fluence (400 – 700 mJ/cm\(^2\)) in an ambient atmosphere or under high pressure (10\(^6\) Pa). Figure 2 illustrates cross sectional TEM images showing the morphology of samples before and after LA at 600 mJ/cm\(^2\).

TEM observations from the as-deposited samples (Fig. 2(a), 2(d)) verified their multilayer character. In both samples Ag was arranged in 21 layers. HRTEM imaging (not shown) identified the \(w\)-AlN presence in sample ML#3 and confirmed the amorphous character of AlN in the case of ML#5. Nonetheless, traces of crystalline \(w\)-AlN were detected by XRD in sample ML#5 after LA (Fig. 1(b), inset), confirming the crystallization of AlN under LA in the vicinity of metal NPs, in agreement with previous HRTEM studies [18]. In both cases, the upper part of the multilayer film appears highly affected by LA. The multilayer morphology is destroyed and the enlarged NPs are rather homogeneously distributed inside the AlN matrix, retaining a high density for the case of sample ML#5; on the contrary, the laser treated part of sample ML#3 is mostly depleted of Ag indicating a higher diffusivity of Ag in \(w\)-AlN compared to \(a\)-AlN, in accordance to what was observed in sputtered AlN:Ag nanocomposite films [19].

Considering LA with a single pulse of 600 mJ/cm\(^2\) at ambient pressure, the \(w\)-AlN/Ag (ML#3) presents a destruction of periodicity affecting only the top six Ag layers, while two more layers are partially destroyed. The sample thickness that retains a stratified arrangement is indicated by a dashed line in the inset of Fig. 2(b), which depicts the contrast intensity profile along the film thickness. In the case of \(a\)-AlN/Ag (ML#5) this structural re-arrangement is more extensive as shown in Fig. 2(e), and it affects the top ten Ag layers, while again two additional layers are partially affected. XRD profile analysis of the Ag(111) peak of the laser treated films confirmed the existence of a bimodal Ag size distribution corresponding to Ag grains located in the higher treated part and the lower untreated part of the film, confirming the TEM results in larger scale.
Figure 2. TEM cross sectional images of: a) ML#3 (w-AlN/Ag): as deposited, b) ML#3: Ambient LA, c) ML#3: High pressure LA, d) ML#5 (a-AlN/Ag): as deposited, e) ML#5: Ambient LA, f) ML#5: High pressure LA. The insets in all images depict contrast intensity profiles (in arbitrary units) obtained along the thickness of the samples. In the LA treated samples the sample thicknesses that retain a stratified arrangement of NPs are indicated by dashed lines. LA fluence for all relevant samples in this figure was set at 600mJ/cm².

Similar results were obtained for LA experiments performed under high pressure for the case of a-AlN (Fig. 2(c)). Typically in LA, a high pressure environment is employed in order to suppress or even eliminate any ablation of the sample. Evidently, a high applied pressure leads to a significant extent of the periodicity destruction depth in the case of w-AlN (Fig. 2(f)). The variation in the NP size distribution and of the total depth of LA process in the films can be attributed to the different thermal conductivities, $k_{\text{AlN}}$, of AlN, which depending on its microstructure may span more than an order of magnitude between crystalline and poorly textured AlN [47,48]. This majorly affects the thermal energy dissipation in the surrounding medium and thus the behaviour of Ag upon LA.

The above considerations make AlN's thermal conductivity a crucial parameter in the LA process. $k_{\text{AlN}}$ is
sensitive to the deposition method and conditions, and can show large variations ranging from about 10 W/mK for amorphous AlN [47] to 285 W/mK for single crystal [48]. To explore its effect, we plot the temperature transient (at the top of the multilayer) for several different values of the AlN thermal conductivity in Fig. 3(b). All other parameters are taken at their crystalline value (see Table 2). A significant difference depending on $k_{\text{AlN}}$ is found for the peak temperature rise when the laser pulse is on, but it quickly diminishes afterwards. The most important effect of $k_{\text{AlN}}$ is shown in the spatial distribution of the peak temperature rise plotted in Fig. 3(c). In particular, for high values of $k_{\text{AlN}}$ the temperature profile is almost constant across the multilayer, even though most of the laser fluence is absorbed within the first 100nm, as shown in Fig. 3(a), where we plot the spatial absorption profile within the multilayer. A combinatory look on Fig. 2 and Fig. 3(a) suggests that layers below the 8th layer of Ag do not efficiently absorb the UV laser light.

This provides a recipe for future practical applications in order to avoid complex structures of 21 repetitions; it seems that 3-4 bilayers would be enough for an effective LA reconstruction featuring a plasmonic response. By the same token, however, we should also note that for fewer bilayers, the cooling through the substrate would be more effective and higher LA fluence could eventually be required. In any case, thinner films would in general result into relatively smaller temperature gradients and thus into more uniform reconstructions even for low $k_{\text{AlN}}$ values.

Going back to our thick, 21-bilayer film, we note that for low values of $k_{\text{AlN}}$, a significant temperature gradient is developed across the multilayer. Higher temperatures close to the surface and lower close to the substrate will result into different annealed structures, explaining our experimental results. We also simulated a multilayer consisting of 10nmAg/5nmAlN (not shown here). As expected, due to the higher percentage of Ag the overall thermal conductivity of the multilayer film is much higher, and the temperature gradients developed are much smaller, even for low values of $k_{\text{AlN}}$. In this case, a more uniform annealing profile in the film is expected.

We stress here that up to now we have only studied the linear heat transport regime to get insight into the temperature transients and gradients that can be developed, and have not considered the thermodynamics of phase changes, e.g. melting and resolidification. Regarding radiation and convection losses from the top
surface, a simple estimate suffices to show that they are of no consequence. For example, in the extreme case of Fig. 3(b) and for a laser fluence of 1000 mJ/cm², a peak temperature of about 6000 K is achieved. If we further assume the extreme cases of unit emissivity, surface-to-air heat transfer coefficient 200 W/m²K (appropriate for forced air cooling[49]) and that this heat exchange with the environment occurs for 30 ns, we get a total energy loss of about 0.1 mJ/cm², which is 4 orders of magnitude smaller that the input fluence, and thus will not have any effect on the temperature transients and profiles.

![Figure 3](image)

Figure 3. (a) UV laser (193 nm) absorption profile in our model structure (21.5nmAg/10nmAlN bilayers on Si), (b) Normalised temperature transient at the top of the multilayer for various values of $k_{AlN}$. The inset shows the pulse profile of the excimer laser, (c) The normalized peak-temperature spatial distribution for the corresponding cases of (b). Large gradients across the film are obtained for low values of $k_{AlN}$, (d) Peak transient temperature distribution as a function of distance from the film's top surface, at different incident laser fluence assuming a nonlinear thermal conductivity for the multilayer structure ($w$-AlN and Ag). Solid lines are for the linear case (zero fluence limit) for $w$-AlN ($k=285$ W/mK) and for an ultra-low conductance $a$-AlN ($k=10$ W/mK), both for a linear Ag thermal conductivity of 429 W/mK. Broken lines are for nonlinear $w$-AlN and Ag for different laser fluence. In the inset we plot the experimental temperature dependent thermal conductivities of Ag [50] and $w$-AlN [48] assumed in this work.

Our critical oversimplifying assumption up to now has to do with neglecting the fact that the thermal parameters of materials are actually temperature dependent. The most striking nonlinearity is found in the thermal conductivity of $w$-AlN, whose relative change with temperature is shown in the inset of Fig. 3(d).
Other parameters such as thermal capacity and mass density also have a thermal dependence, but of a smaller magnitude. Since the highest temperatures are going to be reached inside the film, and to keep our discussion simple, we ignore here nonlinearities associated with the Si substrate. We thus examine the peak transient temperature distribution in the multilayer structure assuming a nonlinear temperature dependence only of the Ag and AlN thermal conductivities. In Fig. 3(d) we plot the peak transient temperature distribution as a function of distance from the film surface for different laser fluence. For comparison we plot the linear response of w-AlN (solid black line) and of an a-AlN with \( k=10 \) W/mK (solid red line). The nonlinear multilayer lines span the space between the two linear curves as the incident fluence increases. Thus the nonlinearity in thermal conductivity increases the temperature gradients developed in the film. Interestingly, the gradients are generally sharper in the nonlinear w-AlN case (e.g. compare the 800 mJ case with the linear a-AlN \( k=10 \) W/mK case, where the peak surface temperatures are the same), meaning that a composite with a matrix of high thermal conductivity (e.g. w-AlN) will get restructured only close to the film’s top surface, while a composite with a matrix of low thermal conductivity (e.g. a-AlN) will get restructured within a wider length inside film. This perfectly explains the behavior observed in our experiments, as seen in the TEM images of Fig. 2(b, e) and 2(c, f).

As expected, the size and distribution of Ag NPs within the amorphous or nanocrystalline AlN matrix determine their LSPR. In Fig. 4(a) and 4(b) we present the ORS data from samples ML#5 and ML#3, respectively. In addition, the insets of Fig. 4(aii) demonstrate an example of a multilayer sample deposited on PET subjected to LA at 700 mJ/cm\(^2\) under high pressure. The digital photos, in white light reflection and transmission mode, clearly present the plasmonic response of the treated areas of the film. The optical response of these samples is in agreement with the above mentioned experimental results and the outcomes of the photo-thermal modeling. Indeed, in the case of ML#5 the low \( k_{AlN} \) values of the a-AlN, allows for high temperatures to be developed inside the film's volume, and thus to NP enlargement with a broad plasmonic response and a monotonic increase in reflection intensity. At the same time the redshift of the LSPR spectral position with respect to the fluence can be attributed either to the formation of bigger Ag NPs and/or to the formation of a more dense environment, possibly by changing the refractive index of the host. Both
assertions seem to be correct, in this case, taking into account the TEM and XRD observations, presented above.

![Graphs showing reflectance vs. wavelength and fluence for different conditions.](image)

Figure 4. (a) ORS of LA processed α-AIN/Ag multilayer under ambient environment (i) and under high pressure (ii), (b) ORS of LA processed w-5 AlN/Ag multilayer under ambient environment (i) and under high pressure (ii). The corresponding spectra from the as grown samples are also shown for comparison. Inset digital photos in (a), in white light reflection and transmission mode, depict sample ML#5 grown on PET subjected to LA at 700 mJ/cm². (c) Maximum reflectance % at the LSPR position (i) and spectral position of the LSPR (ii).

On the contrary, for the case of w-AIN the narrow reconstruction length, in relation to its nanocrystalline character, leads to a non-monotonic behavior of the LSPR characteristics with respect to the fluence. These variations can also be attributed to the different diffusivities of Ag in α-AIN and w-AIN, which lead to a reduction in the Ag content within the film, as mentioned before [19]. Clearly, higher reflectance values are observed at high pressure LA (Fig. 4(c,i)), probably as a result of the suppression of Ag outdiffusion. At the same time no significant shift of the LSPR spectral position is detected in ML#3 (Fig. 4(c,ii)).

So far, we have established that the $k$ of the host determines the fate of the metallic particles in its volume upon LA. This becomes an extra parameter for the manipulation of the LSPR of metal/dielectric media depending on the application. To further explore this argument we tested a multilayer sample consisting of Y₂O₃ and Ag (with similar individual layer thicknesses with sample ML#3). Y₂O₃ was chosen due to its lower thermal conductivity [51] compared to that of AlN [47,48]. Indeed, for the Y₂O₃ host we observe (Fig.
5(a)) that significantly less fluence (45 mJ/cm²) is required in order to observe a plasmonic response. Additionally, substantially lower fluence (150 mJ/cm²) compared to that required for the AlN/Ag multilayers is enough to initiate the deterioration of the plasmon effect.

The aforementioned observations concern the case of Y₂O₃/Ag multilayers grown on Si substrates. If we further grow the same multilayer sample onto PET (which has a much lower thermal conductivity compared to Si) the results vary even more. The deviations in the plasmonic response, due to the substrate effect are presented in Fig. 5(a). Solid lines correspond to the optical response of the samples grown on PET whereas the dashed lines correspond to the samples grown on Si. Due to the reduced cooling offered by PET substrate, we see in Fig. 5(a) that significantly lower laser fluence is required to observe a plasmonic response compared to a Si substrate.

Finally, we present a demonstration of “plasmonic optical reading” on an Y₂O₃/Ag thin film on a PET substrate. The digital photo of Fig. 5(b) represents the sample’s appearance in transmission mode under white light illumination (its appearance in reflection mode is also presented in the inset of Fig. 5(b)). The spot size is 1 mm² and the optical responses of the “NTU” logo, the surrounding matrix and of the as grown sample are shown in Fig. 5(a); specifically, “NTU” spots were fabricated by delivering 25 pulses of 15 mJ/cm² while the surrounding matrix was fabricated by delivering a single pulse of 45 mJ/cm². The minimum feature size of this type of "plasmonic writing" is only limited by the LA wavelength and instrumentation, and could thus easily reach the sum-micron regime.
Figure 5. (a) ORS of sputtered Y$_2$O$_3$/Ag multilayer (as grown and LA) with similar Ag content to ML#3. Dashed lines correspond to the optical response of the sample grown on Si substrate and solid lines to the same sample grown on PET. (b) Digital photo of a flexible plasmonic demonstrator capable of optical reading. Photo captured under white light illumination in transmission and reflection (inset) mode. The spots reading “NTU” were fabricated by LA of the sample delivering 25 pulses at 15 mJ/cm$^2$ whereas the surrounding matrix spots were annealed at 1 pulse of 45 mJ/cm$^2$.

4. Conclusions

In conclusion, multilayers of alternating ceramic (AlN or Y$_2$O$_3$) and Ag layers were subjected to a single shot UV (193 nm) laser annealing, leading to the subsurface formation of Ag NPs with a localized surface plasmon resonance response. An important feature for potential large-scale applications is the simplicity and high capacity of the process employing magnetron sputtering and single-pulse laser annealing. The effect of LA on the structural modification and the resulting optical properties of the films was investigated as a function of laser fluence under ambient atmosphere or under a high Ar pressure. We found that a parameter that majorly affects the annealing process and determines the resulting reconstruction is the thermal conductivity of the dielectric medium, which provides an interesting route for tailoring the optical response of such structures. Detailed photothermal modeling was performed to investigate the laser-matter interactions and heat diffusion in the metal/ceramic multilayers upon LA. The theoretical results cover an extensive range of $k$ values in order to present a generic study for numerous dielectrics. They revealed the paramount role of the thermal conductivity of the dielectric as well as of the relative content of the metal/dielectric components within the multilayer. In particular, low $k$ values ($\alpha$-AlN and Y$_2$O$_3$) lead to a significant temperature gradient across the multilayer structure resulting into broadband size distributions for the Ag particles. On the contrary, high $k$ values (such as in the case of $\omega$-AlN) lead to a narrower size distribution of Ag particles within a shallower depth from the film surface. The developed temperature gradient is also affected by the individual layer thickness and in particular the thickness ratio between Ag and the dielectric in one bilayer: due to the high thermal conductivity of Ag, the higher the Ag content is, the smaller the temperature gradients are within the film, even for low values of $k_{AlN}$.

Finally, we demonstrated the cold character of LA by creating plasmonic templates on flexible polymeric
substrates. From the point of view of applications on optical encoding of information, these examples signify the simplicity, flexibility and versatility of the proposed engineering approach. A additional feature of the produced encapsulated plasmonic NP patterns is their capability of providing two views of the same image with complementary colours in reflection and transmission (overt in white light).

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