A TiN@Au-NR Plasmonic Structure with Tunable Surface Plasmon Resonance Depending on TiN to Au Thickness Ratio

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Abstract

TiN nanostructures have been shown to exhibit promising plasmonic properties and are potential candidates for various applications, including energy harvesting. However, these properties also show a strong dependence on the processing conditions which have been reported to affect metallicity of TiN. Herein, we report on layered TiN@Au-nanorods (NRs) nanostructures consisting of 20 nm TiN thin layer that is magnetron sputtered on Au-NRs of variable length, yielding different TiN/Au thickness ratios (R). While a 20 nm TiN layer sputtered on the same substrate on which the Au-NRs are grown shows a weak absorption peak in the near IR region, an intense and broad plasmonic peak that lies red from the transverse plasmonic peak of monolithic Au-NRs layer is observed for TiN@Au-NRs. The red-shift is shown to increase with increasing R, attaining 100 nm for R = 1, together with an intense tail in the IR region. These results are interpreted in terms of a strong coupling between TiN and Au that drastically affects the plasmonic behavior of the structure. The results are contrasted with those on Pd@Au-NRs where only a slight blue-shift of few nanometers from the Au peak is observed. Potential applications of the TiN@Au-NRs are mainly in energy harvesting such as water splitting and photocatalysis using electromagnetic radiation in a broad wavelength range, as well as medical applications. Pd@Au-NRs may be used as electrocatalysts with plasmonic enhancement, e.g., for the hydrogen evolution reaction.

Keywords Composite 1D-nanostructures · TiN layer on Au-nanorods · Pd layer on Au-nanorods · Plasmonic absorption · Tunable plasmonics · FDTD

Introduction

The most widely known and applied plasmonic nanostructures are based on the noble metals Ag and Au that exhibit localized surface plasmon resonance (LSPR) which is a collective dipolar oscillation of conduction electrons in phase with the incident electromagnetic field. Plasmonic nanostructures find a wide range of applications spanning the fields of biosensing and energy harvesting to photocatalysis [1–7]. In the last few years, however, transition metal nitrides showing metallic conduction, prominent among them is TiN, have seen a flurry of research activities as emerging plasmonic nanomaterials with properties purportedly surpassing those of noble metals [8]. Recently a few applications have been demonstrated using TiN as a plasmonic nanomaterial for hot carrier injection in the wide visible range of the optical spectrum. For instance, Ishi et al. [9] report on a TiN/ZnO/TiN structure that shows a higher photocatalytic performance under (simulated) sunlight, with photocurrent responses surpassing those of Au/ZnO/Au [9]. TiN is also CMOS (complementary metal-oxide-semiconductor) compatible and has been recently reported for self-powered photodetectors when used as a backside illuminated TiN layer on p-type silicon [10].

There are undeniable advantages of TiN that understandably accrue from its high hardness (for instance, TiN is widely known as a hard coating for high speed steel tools) and corrosion resistance which should yield robust plasmonic devices. However, the optical and also the electronic properties of TiN very much depend on the processing conditions which control the amount of impurities, mostly oxygen, and particle size. For instance, Yick et al. show that surface chemistry and particle size of TiN nanoparticles (NPs), fabricated by the transferred arc plasma technique, affect the plasmonic peak...
position and its shape [11]. In particular, they demonstrate that enhancing the metallic character of the NPs through high-applied currents results in a blue-shift of the plasmonic peak. TiN films with the “best” plasmonic behavior were claimed by Sugavaneshwar et al. who used pulsed laser deposition on silicon and polymeric substrates [12]. Their analytically calculated scattering and absorption spectra point to plasmonic properties similar to those of Au-NPs.

Nanostructured Au is a well-established plasmonic material, while TiN is an emerging one with plasmonic properties that depend on the chemistry achieved (see above). Questions arise as to how the plasmonic properties will change in the case of hybrid TiN-Au nanostructures and what are the means to control them. It is the aim of the present work to address these questions. For this purpose, we investigate the plasmonic properties of TiN-NPs@Au-NRs as a function of TiN to Au-NR thickness ratio (R) and show a unique plasmonic absorption behavior that arises from the coupling between TiN-NPs and Au-NRs. Depending on R, the plasmonic absorption red-shifts by up to 100 nm, along with a high background that extends into the NIR region. We contrast these results with those obtained on Pd-NPs coated Au-NR having the same Au/Pd ratio, where rather a blue-shift of few nanometers of the absorption peak was observed. For the purpose of this study, we fabricated Au-NRs of different length using electrodeposition in anodized aluminum oxide (AAO) template films on glass substrates. After exposing the Au-NR structure, TiN and Pd films of 20 nm thickness were sputtered on them via physical vapor deposition.

Materials and Methods

Synthesis and Characterization

The following chemicals were used as purchased: oxalic acid dihydrate 99% (Roth, Karlsruhe, Germany), phosphoric acid 85% (Roth, Karlsruhe, Germany), sodium hydroxide (Roth, Karlsruhe, Germany), and gold(III) chloride trihydrate (Sigma-Aldrich, Steinheim, Germany). Deionized water was used to prepare aqueous solutions.

Aluminum thin films of approximately 500 nm were deposited on a borosilicate glass/Ti (2 nm)/Au (5 nm) heterostructure via electron beam evaporation (PVD75, Lesker, Jefferson Hills, USA, equipped with an electron beam system (Ferrotec GmbH, Unterensingen, Germany)). Anodization was conducted potentiostatically in 0.1 M oxalic acid at 70 V in a two-electrode set-up with a Pt mesh as the counter electrode, using an electrochemical workstation (Keithley 2400 SM, Cleveland, USA). After anodization, barrier layer removal and pore widening were performed in phosphoric acid (5 wt.% in water) for 50 min at 30 °C [13, 14]. Au-NRs were grown into the AAO pores via electrodeposition in 8 mM HAuCl₄ aqueous electrolyte using a three-electrode setup, with the Au/AAO template working electrode, a Pt counter electrode, and an Ag/AgCl reference electrode, under potentiostatic conditions (0.1 V) using an electrochemical workstation (Princeton Potentiostat/Galvanostat Model 263A, Oak Ridge, USA). The Au-NRs length was adjusted via controlling deposition time. After electrodeposition, the 1D nanostructures were exposed by dissolving the AAO film in an aqueous solution of NaOH (5 wt.%) at room temperature. Subsequently, the supported Au-NRs of different lengths were introduced together with a glass/Ti (2 nm)/Au substrate (the same substrate used to grow the Au-NRs) into the sputtering chamber. Twenty-nanometer-thick TiN and Pd films were RF sputtered on the Au-NRs samples and the supporting substrate using pure targets (TiN (Kurt Lesker Company, Jefferson Hills, USA) and Pd (ESG Edelmetall GmbH, Rheinstetten, Germany)) under an approximate argon partial pressure of 10⁻³ mbar. The sputtering power was set to 120 W for Pd at room temperature and 220 W for TiN at 250 °C. The film thickness was monitored by a quartz crystal microbalance. It should be pointed out that the sputtering of 20 nm TiN (or Pd) was conducted simultaneously, under the same conditions for all the samples under study.

The microstructure and morphology of the samples were characterized with a high-resolution scanning electron microscope (SEM Ultra Plus, ZEISS, Oberkochen, Germany) operating in the secondary (SE) and energy selective backscattered (ESB) electron modes. The SEM is also equipped with an energy dispersive X-ray spectroscopy (EDS) package (INCAx-spect, Oxford Instruments, High Wycombe, UK). The structure was characterized by X-ray diffraction (XRD, X’Pert Pro diffractometer PANalytical, Eindhoven, Netherland) in grazing incidence diffraction mode with constant θ = 1° using monochromatic Cu Kα radiation with λ = 1.5418 Å. The device has a full width to half maximum resolution of 0.03°. The absorption spectra were recorded using a UV-Vis-NIR system in transmission mode (Lambda 950 UV–Vis-NIR Spectrometer, Perkin Elmer, Llantrisant, UK). The substrate (glass/Ti (2 nm)/Au (5 nm)) was used as reference. X-ray photoelectron spectra were obtained on a TiN film using Omicron full lab system (Sciencia Omicron GmbH, Taunusstein, Germany) and Al Kα radiation. The films were first cleaned using argon ions sputtering before analysis.

Finite-Difference-Time-Domain (FDTD) Calculations

The FDTD method implemented in MEEP code was used to calculate the absorption and the field enhancement of Au-NR, Pd@Au-NR, and TiN@Au-NR. The cylindrical symmetry was used to model the structures, and the dielectric functions were described by Lorentz-Drude model. The dielectric function parameters of the different materials were obtained from selected sources: those of TiN were fitted by Patsalas et al.
[15, 16]; those of Au were reported by Hao et al. [17] and those of Pd by Rakici et al. [18]. The Au-NR diameter was set to 80 nm, and its length was varied between 20 and 200 nm. The (Pd-) TiN layer was modeled as a disk of 80 nm diameter and 20 nm thickness (see Fig. S1 (supplementary material) for a schematic representation of the model).

### Results and Discussion

#### Structure and Optical Properties

Table 1 lists the different TiN/Au and Pd/Au thickness ratios investigated. The results are compared with those of a 20 nm TiN and Pd thin films on the same substrate heterostructure, deposited under the same conditions.

| R = TiN/Au (Pd/Au) | Au-NRs length (nm) | TiN/Pd thickness (nm) |
|-------------------|--------------------|----------------------|
| 0.10              | 200                | 20                   |
| 0.20              | 100                | 20                   |
| 0.40              | 50                 | 20                   |
| 1.00              | 20                 | 20                   |

Table 1: The investigated TiN/Au thickness ratios (R)

The absorption spectra of the Au-NRs before and after coating with the TiN and Pd layers are illustrated in Fig. 3. The transverse plasmonic peak of the Au-NRs lies at its usual value of 518 nm for the 200 nm long NR (Fig. 3a). Decreasing the Au-NR length (decreasing the aspect ratio), e.g., to 20 nm, leads to a slight red-shift of approximately 40 nm of the maximum together with a noticeable broadening of the peak and diminishing intensity. It should be mentioned that the longitudinal plasmonic peak was not observed due to the experimental set-up which allows measurement only parallel to the NR long axis. The 20 nm TiN layer shows a weak absorption peak at approximately 868 nm which is somewhat higher than the experimental values reported by Yick et al. for TiN particles processed under different conditions and suspended in water [11]. Both our value and those of Yick et al. [11] are, however, largely beyond the calculated values of Sugavaneshwar et al. for their “best” TiN films [12]. The Pd-NP layer shows only an intense background in the whole wavelength range investigated (Fig. S3, supplementary material). At R = 0.1, TiN@Au-NR shows a double peak that upon comparison with that of pure Au-NR of the same length is a clear indication of a strong coupling between the Au-NRs and the TiN layer. Deconvolution of the double peak yields a slightly red-shifted Au peak at 524 nm; if we attribute the second peak to TiN, a substantial blue-shift of its plasmonic peak, which now lies at 563 nm, can be suggested. Further, we notice a damping of the plasmonic absorption of the Au-NRs together with a supposedly strong increase in the absorption peak intensity of TiN. For R = 0.2, a broad absorption peak is obtained that appears to be composed of two smeared maxima (see inset of Fig. 3d), but the peak positions are rather similar. For R > 0.2, the broad absorption peaks observed show a substantial red-shift; e.g., for R = 1, the peak position lies at 640 nm, and an intense tail that extends into the near IR region can be noted (compare the dashed-line curves to the continuous line ones). This absorption in the near infrared region makes these nanomaterials particularly interesting for biomedical devices requiring wavelengths in the range of 1000–1200 nm where the human skin is transparent [19, 20]. The behavior above completely differs from that of the Pd@Au-NR nanostructures of similar R, where barely any difference in the absorption wavelength with increasing R can be noticed, lest for an overall damping of the absorption peak intensity. The Pd@Au-NR results are in agreement with those published by Xiang et al. on Au-NR -core-Pd- shell structures [21], where Pd has been shown to blue-shift the plasmonic absorption peaks of Au-NRs and dampen their intensity, depending on Pd thickness.

The results above, both in terms of peak shift and intensity, unambiguously suggest that there is a strong coupling between the plasmonic absorptions of Au and TiN, essentially resulting in a behavior completely different from that of monolithic Au-NRs. While a plethora of work has been published on hybrid nanostructures, including Ag/Au particles and Au nanorods as well as Au (Ag)/dielectrics in solution (e.g., see the review article by Cortie et al. [22]), nothing has been reported so far on the coupling between noble metal nanostructures and transition metal nitrides. We might treat the present experimental findings assuming, first, the existence of two plasmonic peaks of TiN@Au-NR. Deconvoluting the double/broad absorption peaks, we may then obtain one peak corresponding to that of Au-NR that is...
slightly red-shifted, most probably because of the difference in the dielectric permittivity of both materials; the second peak might be attributed to TiN with rather a large blue-shift which might be rationalized in terms of increased metallicity of the TiN layer owing to carrier transfer from Au. Indeed, depending on its chemical composition, TiN is reported with a carrier concentration in the range of 10^22/cm^3. According to Sugavaneshwar et al. [12], increasing metallicity of TiN boosts carrier concentration to approximately 8 × 10^22/cm^3, and this would lead to plasmonic absorption at wavelengths in the 600 nm range. Metallic TiN has reportedly a higher Fermi level and lower work function than Au [23, 24]. This should result in electron transfer from TiN to Au, as depicted in Fig. 4a, that is compensated by hole formation in TiN. However, the material presently dealt with has most probably oxygen defects, as corroborated by its plasmonic peak that lies red-shifted by about 250 nm from that of a metallic TiN. Further support of the presence of oxygen defects is provided by the XPS spectra of the TiN layer surface, which show oxygen defects (Fig. S4, supplementary material). Consequently, the presence of oxygen defects could change the material’s behavior to a more semi-conducting one, and
the band diagram in Fig. 4a no more holds. For instance, Lui et al. show that NTiO behaves as a semi-conductor with a band gap that depends on the amount of oxygen defects [25]. Upon excitation, hot electron injection from Au into the conduction band of the Ti(N, O)x structure is expected to increase its metallic character (Fig. 4b) that should explain the strong blue-shift of the plasmonic peak of TiN. When R is increased (this is done by maintaining the TiN thickness at 20 nm while varying the of the Au-NR length downwards to 20 nm), meaning that the nanostructure tends towards Au-core-TiN-shell, the blue-shift of TiN is still substantial, but the peak intensity decreases, while the near-infrared absorption increases over a wide wavelength range. However, it is important to note that, overall, the absorption intensity of all the structures with $R \geq 0.2$ is higher than that of monolithic Au-NRs (compare Fig. 3a and d) which further confirms the strong coupling between Au and TiN.

In another reasoning, we could attribute the plasmonic absorption behavior of the coupled TiN/Au-NRs structures to the difference in permittivity of both materials that red-shifts the Au plasmonic peak and this to higher wave lengths, depending on TiN/Au-NR -thickness ratio. In this case, we assume that the plasmonic absorption of TiN is overridden by the high background. However, because the TiN layer thickness is maintained constant and the interface between the Au-nanostructures and TiN should not change with decreasing Au-NR length and taking into account the plasmonic absorption behavior of pristine Au-NRs, the strong red-shift increase with increasing R (approximately 100 nm for R = 1) and the higher absorbance observed for the structures with R $\geq 0.2$ remain to be explained.

Either way, the experimental results above impressively demonstrate the tunability of the plasmonic absorption of TiN@Au-NR structure in a rather large wave length range. The structure is endowed with the additional advantages of chemical and mechanical stability, thanks to the TiN layer, as mentioned in the introduction above.

**FDTD Calculations**

The experimental results depicted above were compared with FDTD calculations using the layered model (Fig. S1, supplementary material). We opted for the dielectric function parameters fitted by Patsalas et al. for TiN because they yield a
behavior rather similar to that of Au [15, 16] and, in this respect, reflect a TiN material with high metallicity. This should allow us to study the behavior of the coupled structure. Light absorption of a single pristine Au-NR was calculated, assuming cylindrical symmetry of the NR. The diameter was kept constant at 80 nm, and the nanorod length was changed. This cylindrical symmetry exhibits two plasmon modes: the longitudinal mode where the oscillations of the electrons are along the cylinder axis and a transverse mode for transverse electrons oscillations. Figure S5 (supplementary material) shows the calculated absorption of differently long Au-NR. For the 200 nm NR, double peaks are observed representing the transversal (509 nm) and longitudinal (780 nm) modes. For the 100 nm NR, the splitting of the plasmonic peaks is not observed due to the low aspect ratio of 1.25; hence, the transversal and longitudinal modes lie in close wavelength range. The calculated absorption peaks are in broad agreement to the experimental results and with the literature [15]. The absorption of a pure TiN thin disk of 20 nm thickness and 80 nm diameter was also calculated. The results depicted in Fig. S7 (supplementary material) show a high background over the visible range with an absorption peak in the UV that is similar to the measured Pd absorption in the literature [26].

Concerning the layered Au-NR/TiN(Pd) structures, it appeared that the FDTD calculations did not render the behavior experimentally observed. In particular, the coupling between TiN and Au-NRs seemed to be underestimated. The reasons are to be found in the simple layered nanorod model together with the dielectric functions selected from the literature. For instance, comparing the dielectric functions of Au [17] and TiN [15, 16] reveals the reason behind the nonsignificant difference between the calculated absorptions of Au-NR and layered structures. The real and imaginary parts of the dielectric functions of TiN and Au behave in a similar way with the only difference residing in a higher imaginary part ($\varepsilon_2$) and a lower quality factor for TiN (broader LSPR peak).

Because the model used is based on the propagation of the electromagnetic wave from one material to the other, with their respective dielectric functions, and does not take into account charge transfer at the interface, the
failed calculation results can be explained by the close dielectric functions used for TiN and Au. Further to the limitations of our model, possible coupling between the TiN nanoparticles in the film, e.g., as nanodimers, which can change the optical behavior of the layered nanostructure, is neglected in the calculations.

In order to circumvent the drawback of the model, the dielectric function of the TiN layer was changed to that of a mixed dielectric function of an Au:TiN alloy (1:1) to account for TiN metallicity [27]. The results obtained for TiN@Au for R = 0.2 and 1 are displayed in Fig. 5; they show a good agreement with the experimental results shown in Fig. 3 and would support our claim of a strong coupling between TiN and Au nanostructures.

**Conclusions**

In the present paper, TiN@Au-NRs and Pd@Au-NRs are processed using a combination of electrodeposition of Au-NR in porous anodic aluminum template films on glass substrates and physical vapor deposition of 20 nm thick TiN and Pd layers. The TiN (Pd) to Au-NRs thickness ratios (R) are tailored by maintaining the TiN (Pd) film thickness of 20 nm and varying the Au-NRs length. A monolithic, nanostructured TiN film sputtered on the same substrate used to grow Au-NRs shows a broad and weak plasmonic peak in the NIR region, while TiN@Au-NRs is characterized by an intense plasmonic peak that lies red from the plasmonic peak of Au-NRs. With increasing R, the peak substantially red-shifts, e.g.,
by 100 nm for R = 1, together with the appearance of a tail in the IR region. This behavior contrasts with that of Pd@Au-NRs which shows only a marginal blue-shift of the Au peak. We surmise that the particular plasmonic properties of TiN@Au-NRs are the result of a strong coupling between TiN and Au-NRs, probably involving carrier transfer from Au to TiN. FDTD calculations using a simple model and dielectric functions from the literature failed to accurately predict the experimental results. However, assuming for TiN, a mixed dielectric function reported for a TiN:Au alloy seems to describe the experimental results. However, assuming for TiN, a mixed dielectric function reported for a TiN:Au alloy seems to improve the agreement between experiment and calculations. Applications of the TiN@Au-NRs structures may encompass mechanically robust plasmonic devices with tunable plasmonic absorption, light harvesting, and photocatalytic nanostructures. Pd@Au-NRs may be particularly useful for photoelectrocatalytic applications, e.g., for the electrocatalytic oxidation of formic acid and for the hydrogen evolution reaction using light enhancement.

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Compliance with Ethical Standards

Conflict of Interest The authors declare that they have no conflict of interest.

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