Tailoring active color rendering and multiband photodetection in a vanadium-dioxide-based metamaterial absorber

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Metamaterials have demonstrated exotic electromagnetic properties, which offer a good platform for realizing light absorption, photodetection, filtering, and so on. However, broadband multifunctional metamaterial absorbers are restricted in cascaded structures. Here, broadband multifunctional properties were realized by introducing vanadium dioxide into a metamaterial absorber. Through the modified design and highly efficient utilization of multiple resonant modes, both plasmonic tunable color filters and near-infrared photodetectors can be simultaneously achieved by this construction. Meanwhile, active color and a photodetection band in the near-infrared range can become tunable with the insulating–metallic transition of vanadium dioxide. Thus, the variations of rendering colors could correspondingly indicate shifts of the near-infrared photodetection bands. This method theoretically confirms the feasibility of designing multifunctional devices via a vanadium-dioxide-based metamaterial absorber, which holds great promise for future versatile utilization of multiple physical mechanisms to achieve numerous functionalities in a simple nanostructure or device. © 2018 Chinese Laser Press

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1. INTRODUCTION

Metamaterials, artificial composite materials or structures, exhibit salient advantages in terms of controlling and manipulating light–matter interactions at subwavelength scale [1,2]. Therefore, they enable spectral engineering, phase manipulation, polarization control, such as light harvesting [3,4], filtering [5,6], modulation [7], optical angular momentum generation [8,9], holograms [10], polarization conversion [11,12], and many more processes. In general, particular spectral properties in such artificial engineered metamaterials emerge from optical resonances induced by composite nanostructures. Therefore, spectral information strongly relies on the size, shape, and arrangement of metallic or dielectric nanostructures. Most importantly, tailoring all parameters provides an efficient means for spectral management and therefore reaps significant benefits for achieving tunable spectra, especially in the visible range [13–15]. However, it will also be stuck in the fixed morphological nanostructure. With respect to this limitation, integrating or embedding the materials or structures with tunable optical properties could be a better approach to overcome the bottleneck [7,16–18].

So far, most research efforts in the visible range have been exerted on tunable structural colors via microelectromechanical systems (MEMS) [19], photonic crystals [20], mechanical deformation strategies [18,21], optoelectronic or electrochemical methods [22–24], etc. MEMS offer an avenue to form a gap changing Fabry–Perot (FP) cavity, which can produce a continuously varying color in the whole visible range. However, it is unfavorable for optoelectronic integration due to the complex control system. Photonic crystals can also provide a dynamically tunable capability of resonant wavelengths, i.e., colors, with the changes of effective refractive indices in the photonic crystal structures. Mechanical deformation strategies could support the changes of structural periods, which will produce dynamically tunable colors due to the changing of effective refractive index. In addition, optoelectronic methods can also tune the permittivity of materials such as transparent conductive oxides and graphene. Alternatively, electrochemical methods can afford the deposition or dissolution of noble metals, which directly alters morphological nanostructures, such as size and volume. Therefore, it will cause shifts of the resonant wavelengths, which can be regarded as the changing of colors in
the visible frequency. Nevertheless, these strategies for changing colors, possessing both reversible dynamic response and distinct optical behavior, are naturally single function, which means that these structures or devices will work only in the visible spectral range for color filtering, color printing, sensing, and so on. While the rendering colors of the designed devices can represent the working state in the other spectra, like a pH indicator, it will therefore enhance efficiency and cost effectiveness. Phase-change materials exhibit broadband electromagnetic response with tunable properties, such as germanium-antimony-tellurium alloys (Ge$_2$Sb$_2$Te$_5$) [25–28], and vanadium dioxide (VO$_2$) [29], which could be excellent candidates for multiband and multifunctional applications. VO$_2$, an iconic example of a correlated electron material, has received significant attention because of its insulating–metallic transition (IMT) at ∼68°C [30,31]. In addition, the phase change due to a Mott transition in VO$_2$ can directly change its refractive index, which has been proved by applying temperature [32,33], electrical fields [34], or light [35,36] in the visible, infrared, and other spectral regions [29,32,35,36]. However, most of functional devices based on VO$_2$ work in a relatively narrow band, for example, it may work only as an infrared photodetector at 10.6 μm [37].

In this study, we introduce VO$_2$ into the metal–insulator–metal (MIM) metamaterial absorber to create dynamic color display and near-infrared multiband photodetection. The property of freely designed working wavelength in the metamaterial absorber can realize near-infrared photodetection due to the hot-electron and photocarriers generated in the plasmonic absorber can realize near-infrared photodetection due to the finite-difference time-domain (FDTD) method. Filling factor FF is the duty ratio of the nanoparticles. The thickness $h$ of the silver nanoparticles is globally set as 70 nm to make further fabrication easier. $t_{\text{insulator}}$ and $t_{\text{VO2}}$ are the thickness of the insulator and the VO$_2$ film, respectively. Moreover, the thickness of the silver film on the bottom is more than 100 nm to act as a reflective mirror. The refractive index of the insulator is 2.0. Silver is used as the metal material with its dielectric constant from the data of Palik [39]. The optical constants of VO$_2$ at the insulating state and the metallic state are extracted from Ref. [32].

### 3. RESULTS AND DISCUSSION

Figure 2(a) shows the calculated spectra with the phase transitions of VO$_2$ at $t_{\text{VO2}} = 20$ nm, $t_{\text{insulator}} = 5$ nm, period $p = 470$ nm, and filling factor FF = 0.3. It can be clearly seen that the spectrum has changed significantly. The resonant wavelength shifts from 1467 to 1928 nm in the near-infrared range. When the period decreases from 470 to 440 nm and the identical period of a unit functional metamaterial structure along the $x$ and $y$ directions. Filling factor FF is the duty ratio of the nanoparticles. The thickness $h$ of the silver nanoparticles is globally set as 70 nm to make further fabrication easier. $t_{\text{insulator}}$ and $t_{\text{VO2}}$ are the thickness of the insulator and the VO$_2$ film, respectively. Moreover, the thickness of the silver film on the bottom is more than 100 nm to act as a reflective mirror. The refractive index of the insulator is 2.0. Silver is used as the metal material with its dielectric constant from the data of Palik [39]. The optical constants of VO$_2$ at the insulating state and the metallic state are extracted from Ref. [32].

![Fig. 1. Schematic of the VO$_2$-based metamaterial photodetector with active color rendering.](image-url)

![Fig. 2. Calculated reflection spectra of the VO$_2$-based metamaterial absorber with phase transitions from the insulating state to the metallic state at $t_{\text{VO2}} = 20$ nm, $t_{\text{insulator}} = 5$ nm, (a) $p = 470$ nm, FF = 0.3, (b) $p = 440$ nm, FF = 0.5, and (c) $p = 260$ nm, FF = 0.6. The solid line represents the reflective spectrum at the insulating state, while the dashed line stands for the spectrum at the metallic state. (d) The calculated CIE 1931 chromaticity diagram of the corresponding spectra showing in (a)–(c). The red triangle formed by the red squares as the triangle vertices shows the color gamut when VO$_2$ is at the insulating state, while the blue triangle formed by the blue circles represents the color space at the metallic state.](image-url)
filling factor becomes 0.5, as shown in Fig. 2(b), the resonant wavelength in the near-infrared shifts from 2224 to 3016 nm. When the period \( p \) and filling factor FF are, respectively, 260 nm and 0.6, the resonant wavelength in the near-infrared also varies from 1686 to 2213 nm with phase transition, as observed in Fig. 2(c). These results confirm the initial resonant wavelength shift toward the longer wavelength with phase transition of \( \text{VO}_2 \), where the shifting of resonant wavelengths could support multiband photodetection. Moreover, the changes of the reflection at the initial resonant wavelength provide the switching state. Meanwhile, differences of the spectra between the insulating state and the metallic state have also occurred in the visible regime. To further investigate the differences of the spectra between the two states in the visible frequency range, the chromaticity is calculated by interpreting the reflective spectrum with respect to the color-matching functions of the human eye as defined by the International Commission on Illumination (Commission Internationale de l’Eclairage, CIE) [40,41]. The data in the CIE chromaticity diagram can intuitively represent the variations of the structural colors. As shown in Fig. 2(d), the red squares stand for the corresponding colors of the reflective spectra in Figs. 2(a)–2(c) at the insulating state of \( \text{VO}_2 \), while the blue circular dots represent those at the metallic state. Thus, the square dots can be regarded as the vertices of a triangle that shows the color gamut when \( \text{VO}_2 \) is at the insulating state, while the blue triangle with the blue circular dots represents the color space at the metallic state. It can be clearly seen that color has changed dramatically with phase transition; for example, the color shifts from green to aquamarine at period \( p = 440 \) nm and filling factor \( FF = 0.5 \). In addition, the associated rendering colors of the spectra under D65 illumination (North daylight illuminant) are depicted in the inset. It, therefore, confirms the color changes with phase transition. As a consequence, when the phase transition occurs, the resonant wavelength at the near-infrared range and the spectrum in the visible frequency (or color) range will simultaneously change, which can be identified as an appreciable synergetic relationship between the photodetection band in the near-infrared and the actively rendering color. Moreover, the slight variations of the colors can directly indicate significant shifts of the photodetection band in the near-infrared range. Also, the initial and subsequent bands of photodetection with the insulating–metallic transition jointly contribute to broadband photodetection in the near-infrared range.

To investigate the physical mechanism of the multifunctional \( \text{VO}_2 \)-based metamaterial absorber, the reflection characteristics of the proposed structure are scrutinized with various periods, and the filling factor FF, \( t_{\text{VO}_2} \), and \( t_{\text{insulator}} \) are fixed at 0.5, 20 nm, and 5 nm, respectively. Figure 3(a) shows the calculated reflection spectra at both the insulating and metallic states with the period ranging from 220 to 700 nm in increments of 40 nm. The optical responses manifest a substantially lower reflection at particular wavelengths according to the periods of the nanoparticles, following four trends indicated by the dashed lines marked A, B, C, and D from the shorter wavelengths to the longer ones. These dashed lines can be clearly seen in the contour maps of the reflection spectra with more detailed data, which are normalized to the area of the unit cell at the insulating state and the metallic state of the \( \text{VO}_2 \) film, respectively, as shown in Figs. 3(b) and 3(c). Lines A (cyan dashed line), B (white dashed line), and C (magenta dashed line) collectively dominate the spectra in the visible range, i.e., rendering colors, while line D (yellow dashed line) in the near-infrared range possesses low reflection, which means high absorption in the proposed metamaterial structures may support high efficiency photodetection. Moreover, compared to line D in Figs. 3(b) and 3(c), photodetection bands have significant changes. Therefore, owing to the insulating–metallic transition,
multiband photodetection can be achieved in the VO$_2$-based metamaterial absorbers with simultaneous color rendering.

To further explore the origins of lines A–D, we analyze the impact of geometric transformation using a straightforward and physically intuitive procedure with the insulating state of VO$_2$. In these simulations, periods in both the $x$ and $y$ directions are fixed to 500 nm and FF yields to be 0.5 in either the $x$ or $y$ direction. Moreover, the polarization of the illumination light is along the $x$ direction, and the rest of the parameters are the same as those of the insulating state in Fig. 3(a). As a comparison, the structure with no silver nanoparticles has also been calculated as the blank control group. As shown in Fig. 4(a), FF in the $y$ direction is fixed at 0.5, while it varies from 0.3 to 0.7 in the $x$ direction. However, the FFs in the $x$ and $y$ directions presented in Fig. 4(b) are opposite to those in Fig. 4(a). The first dip bands are all around 435 nm in both Figs. 4(a) and 4(b).

Compared to the reflective spectrum with no nanoparticles shown in the dotted line, the entire resonant wavelengths of this mode have slight blueshifts owing to the coupling of the transverse propagation surface plasmon polariton (SPP) modes. Therefore, this mode, line A in Fig. 3(a), may be attributed to the longitudinal FP cavity modes supported by the air/insulator/VO$_2$/Ag cavity. The second dips around 500 nm in both Figs. 4(a) and 4(b) do not change when the structure changes, with the resonant wavelength equal to the period. In addition, these dips shift from 380 to 700 nm in a linear manner, shown as line B in Fig. 3(a), with the period varying from 380 to 700 nm. This is accounted for by Wood’s anomaly for the Ag–air interface, as in the case of diffraction gratings. The wavelength can be dictated by the equation

$$
\lambda = p \cdot \sqrt{\varepsilon_d / \sqrt{i^2 + j^2}},
$$

where $\lambda$ is the wavelength, $p$ is the period, $\varepsilon_d$ is the permittivity of the dielectric at the interface, and $i$ and $j$ are integers signifying the order of resonance. Compared to the first dips around 435 nm, there are significant differences at the third dips [denoted as line C in Fig. 3(a)] between the increasing filling factors in the $x$ and $y$ directions in Figs. 4(a) and 4(b). As shown in Fig. 4(a), the third dips shift from 557 to 713 nm with FF ranging from 0.3 to 0.7. However, it varies only from 647 to 683 nm in Fig. 4(b). That means the shifting of the resonant wavelength is mainly associated with the geometry of the nanoparticles along the polarization direction. Therefore, we regard this mode as the transverse FP cavity modes [42]. In addition, as demonstrated in the insets of Figs. 4(a) and 4(b), the fourth resonant wavelengths [line D in Fig. 3(a)] in the near-infrared range have behavior similar to that of the third ones, either at the insulating state or the metallic state. The longitudinal FP cavity mode (Ag/insulator/VO$_2$/Ag) and localized SPP mode in the MIM metamaterial structure dominate this so-called metamaterial gap plasmon phenomenon [43].

By exploring the optical response with variable filling factors at insulating state with period $p = 500$ nm, we preliminarily

![Fig. 4](image-url)
elucidate the core physical mechanisms underlying the selective spectra dips in reflection. To scrutinize the relative modes or phenomena, we plot the magnetic field distribution at the resonant wavelengths of lines A, C, and D with period \( p = 500 \text{ nm} \) and filling factor \( FF = 0.5 \) in both the \( x \) and \( y \) directions. The other parameters are the same as the insulating ones in Fig. 3(a). As shown in Fig. 4(c), the field distribution at wavelength of 440 nm exhibits a legible feature of a longitudinal FP cavity mode as the higher field intensity located at the VO₂-film–Ag mirror interface. Meanwhile, the SPPs excited at the sidewalls (Ag–air interface) of the nanoparticles exhibit a weak coupling with the FP cavity mode. Thus, the longitudinal FP cavity mode and the SPP mode act collectively to inhibit the reflection around specific wavelengths, marked as line A. Alternatively, the SPPs excited at the backside of the nanoparticles (Ag–insulator interface) carry the coupling electromagnetic power and propagate transversely, forming a transverse FP resonator, as illustrated in Fig. 4(d). The field distribution at wavelength of 650 nm (denoted as line C) locates mainly in the dielectric layers between the upper and lower metal layers where the field maximum tends to be concentrated in the center area of the slits, which is coincident with the transverse FP mode. As presented in Fig. 4(e), the field profile at resonant wavelength at 2500 nm (denoted as line D) shows a standard FP mode in the MIM metamaterial structure. The expected metamaterial absorber can be regarded as a combination of a microcavity and surface plasmonic structures, where both the surface plasmon resonances near field enhancement and light recycling contribute to the high light absorption shown in Fig. 3. Thus, hot-electron photodetection with the plasmonic structures and photodetection with the VO₂-based cavity photodetector jointly contribute to the total photodetection of the VO₂-based metamaterial absorber.

Having analyzed the physical mechanisms in the proposed metamaterial absorber, we build a database of reflection spectra in the visible and near-infrared ranges with different periods and filling factors at the insulating state and the metallic state. Although the CIE chromaticity could present the spectra, it cannot demonstrate the real colors as we see in the natural world. To get a better understanding of the actual colors, the corresponding colors of the spectra under D65 illumination are depicted in Fig. 5. Responses have been achieved in nearly the entire visible range with variable geometric parameters at both the insulating and metallic states when the period varies from 260 to 580 nm, and the filling factor changes from 0.3 to 0.6. In addition, colors with the same geometric parameters have changed with the phase transition, so are the photodetection bands in the near-infrared range. For example, the color shifts from green to aquamarine and the photodetection band in the near-infrared range changes from 2224 to 3016 nm with the phase transition at period \( p = 440 \text{ nm} \) and filling factor \( FF = 0.5 \). Consequently, slight changes of color can predict significant changes of photodetection bands. Thus, we can roughly deduce the photodetection band from the rendering color rather than from an additional spectrometer and thermometer.

4. CONCLUSIONS

In summary, a unique strategy for multifunctional devices is proposed to achieve active color rendering and multiband photodetection in a VO₂-based metamaterial absorber. Through the modified design and highly efficient utilization of multiple resonant modes in the VO₂-based metamaterial absorber, both plasmonic tunable color filters and near-infrared photodetectors can be simultaneously achieved, and the filtering colors and photodetection bands will change with the phase transition. Moreover, the slight variations of rendering colors can correspondingly indicate remarkable shifts of bands of photodetection in the near-infrared range. The appreciable synergistic relationship between the photodetection bands in the near-infrared and active rendering colors reveals the feasibility of utilizing VO₂-based metamaterial absorbers, rather than the cascaded structures, for designing multifunctional devices, which may shed some light on expanding the utilization of multiple physical mechanisms to achieve versatile functionalities in a simple nanostructure or device.

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![Fig. 5. Calculated colors evolve with different periods and filling factors at the insulating state and the metallic state under D65 illuminations. The period varies from 260 to 580 nm, and the filling factor changes from 0.3 to 0.6.](image_url)


