Effect of Ultrafast Broadband Nonlinear Optical Responses by Doping Silver into Ti$_3$C$_2$ Nanosheets at Visible Spectra

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Abstract: Ti$_3$C$_2$ nanosheet is a newly discovered two-dimensional (2D) clan. It turns out to have encouraging applications for electromagnetic shielding and energy storage. Here, Ag@ Ti$_3$C$_2$ hybrids are precisely synthesized by using the one-step solution processing method. Also, their ultrafast broadband nonlinear optical responses in the visible region are studied systematically through nanosecond open-aperture Z-scan and transient absorption techniques. The mechanism of two-photon absorption (TPA) is disclosed in the visible region (409–532 nm). When the laser energy is low and the wavelength is longer than 400 nm, nonlinear absorption cannot happen. Meanwhile, as the laser energy increases, two photons will be absorbed by the electrons in the valence band and the electrons will jump to the conduction band. The process is named as two-photon absorption which will make the specimen show reverse saturable absorption (RSA) properties. What is more, the ultrafast carrier dynamics of the specimen are studied by using the transient absorption. The result shows that the decay contains two phases: the fast and then the slow one. The two phases first come from electron–phonon and then from phonon–phonon interactions, respectively. The electron transfer and charge carrier trapping processes are further verified by the outcomes of similar measurements on Ag@ Ti$_3$C$_2$ hybrids. Besides, the two decay processes increase together with the pump fluence. These results show that Ti$_3$C$_2$ nanosheet has potential applications in broadband optical limiter.

Keywords: 2D metamaterial; MXene; optical nonlinear

1. Introduction

With the discovery of graphene, much attention has been placed on two-dimensional (2D) materials to promote the research of modern nonlinear optics owing to their layers of atomic structure, strong light–matter interaction and huge nonlinear optical effect [1]. They have been found to have numerous applications in photonics and optoelectronics [2]. Due to their remarkable optical properties, many 2D materials such as few-layer bismuthene [3], transition metal dichalcogenides (TMD), black phosphorus (BP) [4] and antimonene [5], have promising prospects for mode-locked laser, optical modulator, ultrafast laser generation, photodetectors, etc. [6]. On the other hand, some inherent flaws of them remain a challenge for further applications. For example, graphene has a direct bandgap and BP exhibits unstable characters in the air [7,8]. As a result, the final application of promising 2D nonlinear optical (NLO) materials still has a long way to go. As the latest branch of 2D materials, MXenes have captured much attention among scholars due to their fine conductivity, large elastic moduli, great electric capacity, tunable bandgap, and more admirable optical transparency [9–13]. So far, more than 30 types of different MXenes have been reported, showing the growing members of the MXene family [14]. Moreover, the abundant nature of chemistries and electronic band structure enrich the application base on MXenes.
from energy storage and biomedicine to nonlinear photonics and nuclear waste management [15,16]. Typically, to achieve the application-oriented demands for MXenes, various modification techniques have been performed to obtain the same desired functionalities as the previous studies on colloidal solutions of nanoparticles (NPs) hybridized by graphene, TMDs, etc. [17]. Furthermore, the breakthrough of hybridizing NPs for NLO application has emerged in the past several years in terms of their defect states, surface control, and excellent plasmonic properties [18–20]. Ti3Ce, initially synthesized by Gogotsi et al. in 2011 [21], is the first member of the MXene family. It has gained widespread use in NLO components [14], optoelectronic devices [22] and electromagnetic applications [23] owing to its unique features of atomic structures. While, in visible light region, silver (Ag) can act as an excellent dopant accompanied by functional advantages of increasing the excited absorption cross-section and promoting absorption in the excited state. Silver nanoparticles, though they oxidize rapidly in the air, are favored for their bactericidal properties. The method of not being oxidized in oxygen is still being investigated. As an intriguing semiconductor noble metal hybrid, the Ag@Ti3C2 nanostructure had been successfully synthesized by Satheeshkumar et al. It shows the encouraging Surface-Enhanced Raman Scattering (SERS) [24] and surface plasmon resonance (SPR) properties [25]. The effect of SPR arising from the Ag nanoparticles plays a vital role in the proposed structure in other works [26–28]. With noble metal inducing SPR, the NLO responses can be enhanced significantly and the outperformance can happen in pulsed lasing [29].

Here, to acquire the functionalization of Ti3C2 for NLO applications, we have built a hybridized system of Ti3C2 and Ag NPs with the sole step without an external reducing agent or surfactant as illustrated in the abstract above which is environmentally benign [24]. Ultrafast carrier dynamic and nonlinear absorption of Ag@Ti3C2 hybrids in the visible spectral region are researched. It is significant to reveal the nonlinear absorption properties of that at a broadband wavelength range from 409 to 532 nm by nanosecond Z scan measurement. Besides, the character of optical limiter is studied in detail. The carrier dynamics of this hybrid are investigated with femtosecond transient absorption experiments which can be used to study the decaying channels at various energy levels with multiple probe wavelengths.

2. Specimen and Experiments

The Ti3C2 used in this work is manufactured by selectively etching Al from the MAX precursors Ti3AlC2Tx. Synthesizing details can be found elsewhere [30]. The Ag@Ti3C2 hybrids are made ready by mingling the Ti3C2 nanosheet dispersion and AgNO3 solution. They are shown in Figure 1. First, we re-dispersed 3 mL of original Ti3C2 nanosheet colloidal solution (1 mg/mL) in 30 mL of AgNO3 solution (1 mg/mL) and sonicated the mixture for 30 min. Then, we centrifuged the obtained colloidal solution of hybrid nanocomposites of Ag@Ti3C2 hybrids at 12,000 rpm for 20 min and re-dispersed in 30 mL deionized water. By scanning electron microscopy (SEM, ZEISS ULTRA 55, Oberkochen, Germany), we observed the Microstructure features of Ag@Ti3C2 hybrids and Ti3C2. Furthermore, the monolayered structures are displayed by transmission electron microscopy (TEM, (FEI Tecnai G2 F20, Hillsboro, OR, USA), and the element mapping was captured by STEM-EDX mapping. At room temperature, we surveyed the absorbance spectra of Ti3C2 nanosheet with a UV-vis spectrometer ((TU-1901, Persee, Auburn, CA, USA). With 6 ns Q-switched Nd: YAG nanosecond pulse laser ((Surelite II, Continuum, San Jose, CA, USA), we applied the approaches of open-aperture Z-scan to detect the NLO absorption features of Ag@Ti3C2 hybrids solutions. To evade the thermally induced nonlinear scattering, the laser repetition rate was set at 5 Hz. To obtain the lasers (410–700 nm), an optical parametric oscillator (OPO) was employed [31]. The laser beam (ω0 = 200 µm) was focused through a lens (f = 20 cm) onto a quartz cuvette (2 mm) full of Ag@Ti3C2. While the specimen changes its position on the translation stage (TSA200, Zolix, Beijing, China), the transmittance signals are detected precisely, with the Z-scan signals gathered by an energy detector (J-10MB-LE, Coherent, Santa Clara, CA, USA) and recorded through an energy
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The ultrafast carrier dynamics were revealed by femtosecond transient absorption spectroscopy measurements. The laser pulse (~35 fs, 800 nm and 1 kHz) is originally created through Ti: sapphire regenerative amplifier ((Astrella, Coherent, Santa Clara, CA, USA). What is more, a BBO crystal is employed to double the photon energy to 400 nm. A sapphire plate is adopted to get white-light continuum (420 to 750 nm) as a probe beam which is concentrated through a quartz cuvette (2 mm) filled with Ag@ Ti3C2 aqueous solution. The intensity of both transmitted probe and reference beams were detected by two spectrometers (Avantes-950F, Avantes, Apeldoorn, the Netherlands), respectively, after the specimen was delayed relatively by a delay line (TSA200, Zolix, Beijing, China). All the signals were input to a computer synchronously. Through a standard specimen, pump and probe beams are overlapped spatially. By using optical Kerr signal from the SiO2 substrate, we affirmed the group-velocity dispersion of the white-light continuum [32].

3. Result and Discussion

The morphologic structures of Ag@ Ti3C2 and Ti3C2 are shown in Figure 2. The TEM images and HRTEM images EDX analysis pictures of Ti3C2 are presented in Figure 2a,b, ~1 μm monolayered Ti3C2 nanosheet can be observed and Ti, C, and O elements are displayed in Figure 2a. Aside from the inherent Ti–C bond, the Ti3C2 nanosheets are predominantly oxygen-terminated. A finer structure is shown in Figure 2b. Through applying a mixture of AgNO3 and Ti3C2 solution during synthesis of the Ag@ Ti3C2 rather than pure Ag, situ Ag doping of the Ti3C2 structure is obtained without compromising the accordion-like Ti3C2 structure. Plenty of Ag nanoparticles was dispersed uniformly in Ti3C2, which can be observed in Figure 2c. Meanwhile, the observed structure, which can be seen in the X-ray powder diffraction (XRD, Seifert-FPM, Freiberg, Germany) pattern in Figure 3a, is consistent with morphological results. In order to further illustrate the microscopic results of the material, the selected area of the yellow square in Figure 2a,e is enlarged to obtain Figure 2b,d. The cross-section of layered Ti3C2 structure in Figure 2d and the distributed Ag on Ti3C2 flakes are shown. The SPR effect dominates by the particle size, gap distance among adjacent AgNPs [26], resulting in the hybridization of surface plasmon resonance [33] and gap plasmon resonance [27]. In this work, the average particle size of AgNPs was ~6–10 nm. Figure 2e presents the TEM micrographs, respectively. The experiment results proved that Ag nanoparticles (the bright particles) are inlaid into Ti3C2 nanosheet with one accord. Furthermore, in order to determine the composition of the obtained Ag@ Ti3C2, Ti, C, O, and Ag elements are shown in EDX analysis in Figure 2f.
In brief, all the results presented below show that the doping Ag is uniformly distributed throughout the entire Ti$_3$C$_2$ structure.

The phase transitions of hybridization researched by XRD are displayed in Figure 3a. Two peaks of Ag diffraction and Ti$_3$C$_2$ that are shifted to a much lower angle compared with that of the specimen revealed the expansion of the interlayer distance. In Figure 3b, the Ag@Ti$_3$C$_2$ shows an absorption peak at ~436 nm, which is due to the formation of Ag nanoparticles [24]. As shown in Figure 3b, two absorption peaks are observed at 245 and 325 nm, respectively. These results are under expectation.

As shown in Figure 3b, we find the high absorption in the UV region ranging from 225 to 375 nm as a result of the band-gap energy of the oxidized MXene, which is also predicted by previous theoretical calculations [34,35].
Open aperture Z-scan data of Ti$_3$C$_2$ dispersions are shown in Figure 4a–d. It is obtained at four laser wavelengths of 409, 436, 500, and 532 nm. Specially, in Figure 4b, at the resonant wavelength of 436 nm, the measurement is applied to study the optical nonlinearity of the Ag@Ti$_3$C$_2$ nanosheet under the laser pulse energy of 0.11, 0.36 and 0.64 GW/cm$^2$. When the specimen reaches the focal plane ($z = 0$), we notice the formation of a shallower valley, which indicates that reverse saturable absorption (RSA) arises. A greater enhancement of optical nonlinearity than that of the Ti$_3$C$_2$ nanosheet is observed in our previous work [30]. The experiment results under the other three wavelengths also exhibit RSA properties. In Figure 4a, when incident pulsed laser energy is 0.11 and 0.36 GW/cm$^2$, the normalized transmittances of specimens remain unchanged and no signals are observed. When the pulsed laser energy reaches 0.64 GW/cm$^2$, the transmittances decrease correspondingly. As shown in Figure 4b,c, when the energy is from 0.11 to 0.64 GW/cm$^2$, a deeper valley appears. In Figure 4d, as RSA is shown under the energy of 0.36–0.64 GW/cm$^2$, no experimental signal is observed under 0.11 GW/cm$^2$. The findings of the experiments in Figure 4a,d disclose that the result of RSA is not clear at fewer inputs of energy. Accordingly, we raised the input energy to carry out the following experiments. In short, broad-band RSA appears on Ag@Ti$_3$C$_2$ nanosheet, which proves that it is potentially applicable to optical limiting devices.

The absorbance spectra in Figure 3b show that the one-photon absorption effect only takes place when the wavelength amounts to no more than 400 nm. Thus, the physical process of RSA can be concluded as follows. When laser energy is low, nonlinear absorption cannot happen. However, with the laser energy increased, two photons will be absorbed by the electrons in the valence band and the latter will jump to the conduction band. To put it another way, two-photon absorption would occur and make the specimen display RSA [36].

Electron transfer between Ti$_3$C$_2$ and Ag atoms will arise when Ag nanoparticles are doped into it. From Figure 5b, we should notice TA spectra of the Ag@Ti$_3$C$_2$ and the occurrence of the excited state absorption. The strengthened absorption, compared with pure Ti$_3$C$_2$, may arise from the absorption of photo-induced surface plasmon in Ag nanoparticles [31].
Based on the above analysis, we combine the SA and the RSA to achieve the total absorption coefficient as follows [37]:

$$a(I) = \frac{\alpha_0}{1 + (I/I_s)} + \beta I$$  \hspace{1cm} (1)$$

Here in Equation (1), $\alpha_0$ signifies the linear absorption coefficient, $I$ the pulse laser intensity, and $I_s$ the saturable intensity. To suppress interference of the solvents in the nonlinear effect, we apply the Z-scan measurement to the solvents within the same parameters. The normalized transmission of the open aperture Z-scan is illustrated as below [38]:

$$T = \sum_{m=0}^{\infty} \left[ -q_0(z) \right]^m (m + 1)^{\frac{3}{2}} \approx 1 - \frac{\beta I_0 L_{\text{eff}}}{2 \sqrt{2}(1 + z^2)}$$  \hspace{1cm} (2)$$

Here in Equation (2), $\beta$ signifies the nonlinear absorption coefficient; $I_0$ the peak intensity at the focus; $L$ the distance that the light passes through the specimen; $L_{\text{eff}}$ the effective interaction length which could be elaborated as $L_{\text{eff}} = (1 - e^{-a_0})/a_0 z$; $z$ the displacement of the specimen along the stage from the focal point ($z = 0$); and $z_0$ the Rayleigh diffraction length.

In Figure 4, we use Equations (1) and (2) to achieve the theoretical fit and the nonlinear absorption coefficient $\beta$, the former is drawn in solid lines and latter listed in Table 1. The conclusion is that the TPA coefficients are obtained along with each wavelength increase, 

**Figure 4.** Normalized transmission of Ag@Ti$_3$C$_2$ dispersion for open aperture Z-scan at the wavelength of (a) 409 nm, (c) 500 nm, and (d) 532 nm. (b) Ag@Ti$_3$C$_2$ and Ti$_3$C$_2$ nanosheet dispersion open aperture Z-scan data for comparison.
and laser irradiance increases synchronously. These findings are advantageous when they are used to limit the optical intensity to protect human eyes as well as optical devices.

Table 1. Nonlinear optical parameters of the Ti$_3$C$_2$ nanosheet.

| λ (nm) | $I_0$ (10$^{-2}$ GW/cm$^2$) | β (10$^{-9}$ cm/mW) |
|--------|----------------|------------------|
| 409    | 0.74            | -                |
|        | 1.10            | -                |
|        | 1.40            | 1.80 ± 0.11      |
| 500    | 0.74            | 0.91 ± 0.06      |
|        | 1.10            | 1.12 ± 0.09      |
|        | 1.40            | 1.81 ± 0.12      |
| 436    | 0.74            | 1.03 ± 0.08      |
|        | 1.10            | 1.32 ± 0.10      |
|        | 1.40            | 1.93 ± 0.13      |
| 532    | 0.74            | -                |
|        | 1.10            | 0.41 ± 0.05      |
|        | 1.40            | 0.78 ± 0.06      |

We take the measure of broadband transient absorption to study the ultrafast carrier dynamics of Ag@Ti$_3$C$_2$ and Ti$_3$C$_2$ nanosheet. To find the different carrier dynamics between Ag@Ti$_3$C$_2$ and Ti$_3$C$_2$ nanosheet, the broadband TA under the 400 nm pump is set at the constant pump fluence of $6.4 \times 10^3$ mW/cm$^2$. In Figure 6a,c, the TA spectra with multiple probe beams (450 to 600 nm) of Ag@Ti$_3$C$_2$ and Ti$_3$C$_2$ nanosheet are demonstrated by a 2D map comprising TA signals in a temporal and spectral sense. Obviously, a brighter zone is given in Figure 5a than that in Figure 5b, which implies that enhanced signals emerged according to Ag nanoparticles in the Ti$_3$C$_2$ nanosheet. The plasmonic effects, that of SPR is arising from the Ag nanoparticles [27]. As shown in Figure 5b,d, five differential horizontal cuts through the 2D map illustrate the absorption spectra that vary with distinct delay time. In Figure 5b, a positive absorption in Ag@Ti$_3$C$_2$ proves that the excited state absorption (ESA) occurs in the whole spectral region. Furthermore, the amplitude of the Transient Absorption (TA) spectrum descends while the delay time ascends, which could be ascribed to the carrier relaxation. We adopt the curve of 0 ps delay time (black) as a reference signal before the Ti$_3$C$_2$ nanosheet being excited. The transition process between occupied and unoccupied states results in photo-induced absorption that leads to the peak of Ag@Ti$_3$C$_2$ at the wavelength of 495 nm, which is the mutual proof of the observations of RSA in Z-scan experiment [39–41]. It costs the TA signal ~50 ps to relax from the peak to zero in the full waveband. In Figure 5d, the semblable behavior is shown in Ti$_3$C$_2$ nanosheet, and in ~25 ps, the TA signal relaxes to zero. It is quite interesting that Ag@Ti$_3$C$_2$ exhibits enhanced nonlinear optical in contrast with that in pure Ti$_3$C$_2$ nanosheet.

In the TA experiment, we used a 400 nm (~3.1 eV) pump light laser whose energy is stronger than the energy bandgap of Ti$_3$C$_2$ nanosheets (~2.04 eV). In general, after the specimen is pumped by incident pulses, electrons in the valence band (VB) will be excited and jumped onto the conduction band (CB) through the Franck–Condon transition in several femtoseconds. Meanwhile, the holes will remain in the valence band [42]. Afterwards, carriers that have been excited by photons will be immediately transformed into hot carriers with Fermi–Dirac distribution. Consequently, the hot carriers will follow different relaxation processes through electron–electron and electron–phonon scatterings, which produce the electrons to the conduction band minimum in several picoseconds. In the end, the electrons will cool down, and in tens of picoseconds, it will relax back to VB and reunite with the remaining holes.
To examine the carrier dynamics at multiple wavelengths, we chart Figure 6a,b to illustrate the normalized dynamics curves at the wavelength of 470, 485, 500 and 525 nm, among which Figure 6a explains the optical transmission responses of Ag@Ti3C2 that include a fast decay component and a slow one. We attribute the two components to their corresponding decay processes. After excitation, the fast decay component comes first and Coulomb-induced hot carriers will be trapped from the core state to the surface state. Their spare energy will be released via optical phonon scattering (~4.5 ps). Then comes the slow decay component and the cooled carriers will pass through nonradiative transition to the ground state within ~43 ps [20,43]. The two different decay components are demonstrated by Equation (3), the biexponential decay function as shown below [44]:

$$\frac{\Delta T}{T} = A_1 \exp\left(-\frac{t}{\tau_1}\right) + A_2 \exp\left(-\frac{t}{\tau_2}\right)$$  \hspace{1cm} (3)

where $A_1$, $A_2$ are the amplitudes of each decay components. $\tau_1$ and $\tau_2$ refer to two parameters, describing fast and slow decay lifetimes.
4. Conclusions

The NLO absorption properties of Ti$_3$C$_2$ nanosheet is systematically researched through an OA z-scan technique to reach the conclusion that at broadband visible range (409–532 nm), Ti$_3$C$_2$ nanosheet exhibits RSA properties which are resulting mainly from TPA. What is more, we adopt femtosecond transient absorption spectroscopy to study ultrafast dynamics of the specimen to find that its relaxation contains a fast decay component (~4 ps), resulting from electron–phonon interaction, and a slow one (~12 ps), from Ag nanoparticles, the excited carriers from the CB of Ti$_3$C$_2$ will first move to the sp band of the particles and then turn back to VB of Ag@Ti$_3$C$_2$. This process will happen in a longer lifetime (50 ps) than the direct relaxation in pure Ti$_3$C$_2$ nanosheet, which leads to the elevation of RSA, a similar phenomenon found in graphene [47].

Table 2. Carrier dynamics parameters of the Ti$_3$C$_2$ nanosheet.

| Samples       | $\lambda$ (nm) | $\tau_1$ (ps) | $\tau_2$ (ps) |
|--------------|----------------|---------------|---------------|
| Ag@Ti$_3$C$_2$ | 470            | 4.5           | 33.9          |
|              | 485            | 4.6           | 36.5          |
|              | 500            | 4.2           | 43.1          |
|              | 520            | 3.9           | 45.8          |
| Ti$_3$C$_2$ nanosheet | 470            | 3.8           | 18.0          |
|              | 485            | 3.2           | 27.4          |
|              | 500            | 4.6           | 22.5          |
|              | 520            | 4.6           | 19.9          |

It is argued that electrons on the lower energy states tend to be probed more readily than those on higher energy states. An analogous phenomenon was observed in graphite [45]. As Ag has tremendous states density, the excited electrons in the CB of Ti$_3$C$_2$ will jump to the d band of Ag atoms before the bleaching effect of VB vanishes [46]. Thus, the process prolongs the lifetime of excited electrons which will lead to a stronger RSA response. Afterwards, the electrons in the d band of Ag atoms will jump to the corresponding excited state of functional teams which will absorb the laser energy and jump to the higher energy level. Hence, the enhanced RSA is induced. Ornamented with Ag nanoparticles, the excited carriers from the CB of Ti$_3$C$_2$ will first move to the sp band of the particles and then turn back to VB of Ag@Ti$_3$C$_2$. This process will happen in a longer lifetime (50 ps) than the direct relaxation in pure Ti$_3$C$_2$ nanosheet, which leads to the elevation of RSA, a similar phenomenon found in graphene [47].
phonon–phonon interaction. Besides, the two decay times increase with pump fluence. The experiment confirms that Ti$_3$C$_2$ nanosheets can be used in ultrafast optoelectronics and optical limiters.

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