Research Article

Femtosecond nonlinear absorption and optical limiting action in nanoplatelet CuFe₂O₄-decorated rGO nanocomposites



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Abstract

Using 800 nm infrared, 150 femtosecond (fs), 80 MHz high repetition rate laser pulse excitation, intensity-dependent $(I_0 = 245-735 \text{ MW/cm}^2)$ measurements on the nonlinear absorption of copper ferrite hetero-architecture functionalized with three rGO contents (15, 25 and 40 wt%) were performed. Copper ferrite–rGO nanocomposite demonstrated a peculiar W-pattern [a peak (saturable absorption) with two valleys [reverse saturable absorption (RSA)] at extreme] in the open aperture Z-scan data for the decorated (15 wt%) rGO systems recorded at a low peak intensity ($I_0 = 245 \text{ MW/cm}^2$). The Z-scan traces were dominated by RSA characteristics due to two-photon absorption process at high peak intensities. The variation in intensity-dependent nonlinear absorption coefficient and ground-state absorption at laser excitation wavelength revealed the presence of excited-state absorption (ESA). The maximum two-photon absorption coefficient was recorded for CuFe₂O₄–(40 wt%) rGO at 735 MW/cm² excitation, which is attributed to the combined contribution of rGO and CuFe₂O₄ that is four times higher than pure rGO and CuFe₂O₄. The presence of layer like CuFe₂O₄ nanoplatelets upon the layer-structured rGO offers an added advantage in enhancing the nonlinear absorption coefficient. Near-resonant ESA-based optical limiters comprising CuFe₂O₄–rGO can possibly be used as a versatile optical limiting material for eye safety against intense, high repetition rate fs laser pulses.

Keywords Reduced graphene oxide · Nonlinear absorption · Nonlinear refraction · Optical limiting

1 Introduction

Lasers these days are unavoidable tools in cutting edge technologies such as optics/photonics, and in particular, intense, femtosecond (fs)/picosecond (ps) laser pulses are in high demand in automobile, micro-fabrication and ultrafast measurement areas [1–7]. Such fs lasers, with capacity of delivering high intensities at short intervals of time, are capable of damaging the optical components, and therefore, numerous attempts are being made in preventing these laser-based accidents damaging the human eyes and precious sensors. The desideratum of ideal optical power limiters to delicate optical components (including the human eyes) with a low damage threshold due to

deliberate or unintentional exposure to these high power lasers is one of the ultimate challenges for the optics community [8–11]. Consequently, the development of optical limiting devices has shown tremendous advancement due to its possible utility in photonic application like laser safety devices, optical shutter, pulse shaping and light stabilizing sources [12–14]. However, the development of such materials (transparent at low intensity and translucent at high intensity) in ultrashort (fs) pulse time scale is still a work in progress. Several nonlinear mechanisms such as nonlinear absorption (NLA), nonlinear refraction (NLR) and nonlinear scattering (NLS) have been studied extensively in a variety of materials and have been engaged for achieving superior optical limiters [15, 16].

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Among these diverse phenomena, materials that possess nonlinear absorption can yield effective energy-absorbing optical limiter for various laser pulses. Since the intensitydependent NLA measurements can provide additional information on the nature of nonlinear absorption, Z-scan experiments with different input intensities of excitation were performed to measure the nonlinear transmissivity of the sample. Graphene possesses interesting nonlinear absorption and refraction properties because of the interband transitions that are independent of the broad wavelength. To improvise the nonlinearity with tunability, graphene oxides (cheaper alternates of graphene) are often decorated with noble metal atom, porphyrin, phthalocyanine or covalent functionalization groups [17-20]. The remarkable light absorption and broad band emission of oxide derivatives of graphene (GO-graphene oxide, rGO-reduced graphene oxide) due to extended π -conjugate system have shown significant interest in laser damage protecting units. Here, the sp² hybridization from oxygen-containing groups left out after exfoliation varies which leads to change in NLO properties of rGO. Also, these residual functional groups upon the surface of rGO prevent aggregation and percolate various metal oxides such as Fe₂O₃, ZnFe₂O₄, TiO₂, ZnO and BBO [21–25]. Extensive efforts by various research groups have been extended toward understanding the nonlinear optical (NLO) properties of carbon-based graphene and graphene nanocomposites like graphene/epoxy resin [26], carbon black/epoxy resin [27], functionalized graphene fluoride [28], graphene families [29], graphene oxide [30], cobalt phthalocyanine [31], layered GO [32], polymer–GO [33], GO–Fe₂O₃ [21] and graphene [34]. Most of these materials have demonstrated superior NLO coefficients, which is essential for photonic applications. Particularly, metals incorporated in graphene nanocomposites depict impressive NLO and optical limiting performance owing to the combination of different mechanisms. Among metal oxide loading, ferrite stands high and recent work on ferritedecorated rGO showed remarkable ultrafast nonlinearity characteristics. With intensity-dependent high repetition rate fs laser excitation, CdFe₂O₄-rGO nanocomposite illustrated a shifting of RSA to SA because of the involvement of various molecules with respect to excitation intensity [35]. Impressed by this factor, we have extended our earlier investigations on ferrite-decorated rGO systems. Earlier reports showcase that the combination of rGO with CuFe₂O₄ can yield potential systems with strong nonlinearity [36]. Also ferrites are investigated in different optoelectronic applications as it possess the advantages of (a) owning 3d¹⁰ 4 s¹ configuration in the valence shell, (b) available at low cost and (c) environmentally benign [37-39]. So using the different intensities ($l_0 = 245, 490$ and 735 MW/cm²) of Ti:Sapphire (150 fs, 800 nm and 80 MHz)

SN Applied Sciences A SPRINGER NATURE journal laser pulses, NLO activities of a series of CuFe₂O₄-rGO nanocomposites were measured. It is well known that inclusion of copper ferrite can induce excited states near the excitation wavelength, and thus, there is a possibility to demonstrate ESA-based optical limiters. Therefore, the motivation of the work is to demonstrate a near-resonant excited-state absorption-based optical limiter comprising CuFe₂O₄-rGO with improved NLO performance for eye safety against intense, high repetition rate fs laser pulses. Interesting transition of nonlinear absorption mechanism (SA, W-pattern, sequential 2PA) with respect to the composition of samples was observed. Thus, the basic of understanding the influence of elemental composition (Cu) and morphology in altering the nonlinear optical performance of the material is analyzed.

2 Experimental details

2.1 Preparation of solar exfoliated reduced graphene oxide

Reduced graphene oxide (rGO) that resembles the internal structure of graphene is chosen as host element since graphene is expensive to be produced in bulk quantities. Using inexpensive graphite as a starting precursor, graphene oxide was prepared by modified Hummer's method [40]. The process of oxidation was achieved by reacting graphite (1 g, Alfa Aeser) with sodium nitrate (1 g, EMPLURA) in the presence of sulfuric acid (4 ml). The process of oxidation was continued by the addition of potassium permanganate (6 g, EMPLURA), and the reaction being exothermic excess of heat was released. The solution was immersed in ice bath to control the temperature well below 20 °C. Hydrogen peroxide (6 ml) hydrated solution was used to terminate the oxidation process, and the obtained black power was washed (with HCl and H₂O), centrifuged (8000 rpm, 10 min) and dried (60 °C, 12 h). The best way to remove the attached functional groups without destructing the layered structure of graphene is to reduce graphite oxide using solar radiation. The process of exfoliation was activated by focusing solar radiation upon prepared graphene oxide using convex lens of focal length 100 cm. The transition of light brown to dark black accompanied by the sudden increase in the volume ascertains the formation of solar exfoliated reduced graphene oxide [41]. Here the process of de-oxygenation in graphene oxide occurred due to localized thermal excitation (150-200 °C in 1-2 s) arising from the photoacoustic effect originated through focused solar radiation.

2.2 Preparation of CuFe₂O₄-rGO nanocomposite

The decoration of copper ferrite with different load contents upon reduced graphene oxide to form CuFe₂O₄-rGO nanocomposites was achieved by a one-step hydrothermal method. Here, nitrates of copper and ferric (1:2 molar ratio) were used as precursor to form CuFe₂O₄. Upon reduced graphene oxide (rGO) with various contents (15, 25 and 40 wt%), copper ferrite metal oxide was formed. The oxygen functionalities on rGO sheets serve as nucleation sites to accommodate copper and ferrous oxides dissociating from the respective nitrates and thus forming CuFe₂O₄-rGO nanocomposites. The reaction was carried out in an autoclave unit (volume of 150 mL) with rGO dispersed metal nitrate solutions being heated at 180 °C under ambient pressure. Solvable byproducts in the precipitate were removed with water washing and then dried in air atmosphere (60 °C, 24 h). The above procedure was carried out for 0, 15, 25, 40 wt% of rGO to attain pure CuFe₂O₄ nanoparticles and CuFe₂O₄-rGO nanocomposites.

2.3 Z-scan experiment

Open aperture Z-scan experiments were carried out on $CuFe_2O_4$ -(15, 25 and 40 wt%) rGO nanocomposite to study the nonlinear absorption at various input intensities. An excitation source of Ti:Sapphire laser oscillator (Chameleon, M/s Coherent) was used providing output at a wavelength of 800 nm, pulses with duration of 150 fs at 80 MHz repetition rate. The laser pulses were focused using 100-mm focus lens, and the corresponding beam waist was estimated to be ~ 25.5 µm at 800 nm, and all the experiments were performed with various input peak intensities (245–735 MW/cm²). From the recorded open aperture Z-scan data, the variation in normalized transmittance with respect to light fluence [F(z)] was derived using standard relations [22].

3 Results and discussion

3.1 Preliminary confirmation and structural studies

Recorded XPS patterns of pristine and decorated GO are illustrated in Fig. 1a. For rGO, peaks that represent carbon (285.3 eV) and oxygen (529.7 eV) environment of the molecule are shown in Fig. 1a. C 1s XPS (Fig. 1b) pattern of reduced graphene oxide has peaks at 284.5 eV (C=C), 285.8 eV (C=O) and 288.6 eV (C=O) which represents different functional groups of reduced graphene oxide. The wide-scan XPS spectrum of CuFe₂O₄–(15 wt%) rGO nano-composite obviously depicts the characteristic peaks at 285.01 eV, 532.83 eV, 722. 99 eV, 710.10 eV and 932.80 eV,



Fig. 1 XPS spectra of **a** rGO and CuFe₂O₄–(15 wt%) rGO nanocomposites, **b** C 1s XPS spectra of rGO and **c** Cu 2p, **d** Fe 2p core-level spectrum of CuFe₂O₄–(15 wt%) rGO nanocomposites. The peaks in **b** were fitted with Gaussian curves, and the green lines represent the components of Gaussian fit made on C 1s XPS spectrum

corresponding to the binding energy of C 1s, O 1s, Fe 2p and Cu 2p, respectively. Figure 1c (Fe) and Fig. 1d (Cu) represent the core-level XPS pattern for composite. Peaks at binding energies of 713.10 eV (Fe $2p_{1/2}$), 718.3 eV (Fe³⁺), 726.14 eV (Fe $2p_{1/2}$), 935.11 (Cu $2p_{3/2}$) and 944.25 eV (Cu $2p_{1/2}$) ensure the presence of metals (copper and ferrous) ions in the composite which is found to match well with the literature [42, 43].

X-ray diffraction patterns were recorded to confirm the formation of the support as well as the composite, and data are shown in Fig. 2. In the XRD of GO, the disappearance of graphite diffraction peak ($2\theta = 26^{\circ}$) and the formation of characteristic peak at lower angle $(2\theta = 10.6^{\circ})$ confirm the complete oxidation during Hummer's method [22]. The expected reverse observation (graphene oxide turning themselves as graphene like structure) of no diffraction at lower angle and broad weak XRD peak at 24° that corresponds to (002) plane of rGO was observed in solar exfoliated product. For CuFe₂O₄-decorated rGO hetero-architecture, the presence of (200), (211), (422), (004), (312) and (224) reflection planes was ascribed to the tetragonal structure of CuFe₂O₄ [JCPDS: 34-0425]. In the CuFe₂O₄-rGO hetero-architecture, exfoliation of rGO occurred because of crystal growth of copper ferrite between the rGO sheets. In the recorded FTIR spectrum (Fig. 3), rGO shows broad vibrational band due to the stretching mode of hydroxyl groups (3420 cm⁻¹). Upon decoration, rGO shows a decrease in absorption intensity of 3420 cm⁻¹(O–H), 1725 cm⁻¹,(C=O) and 1120 cm⁻¹



Fig. 2 XRD patterns of pure and decorated $CuFe_2O_4$ -rGO composites

(C–O) which indicate that rGO was further reduced to a great extent during the incorporation of $CuFe_2O_4$. New absorption bands observed at 545 cm⁻¹ (Cu²⁺ stretching in the octahedral site) and 454 cm⁻¹ (Fe³⁺ stretching in the tetrahedral site) belong to $CuFe_2O_4$ in the composite. For $CuFe_2O_4$ –(40 wt%) rGO sample, the FTIR spectrum showed peaks at 3427 cm⁻¹ (O–H), 2928 cm⁻¹ (–CH₂), 1726 cm⁻¹ (C–O), 1620 (C=C) and 1125 cm⁻¹ (C–O), and in the case of (25 wt%) rGO, the peak at 2928 cm⁻¹ (–CH₂) vanished for the nanocomposite. The effective removal of functional groups in $CuFe_2O_4$ –(15 wt%) rGO was observed as many of the functional groups almost vanished in the IR spectra of the composite.

Raman spectroscopy is an essential experimental technique for identifying the defects states formed during functionalization of graphene. In recorded Raman spectra with 532 nm laser excitation (Fig. 4) of copper ferrite functionalized upon three rGO contents (15, 25 and 40 wt%), G band (1598 cm⁻¹, 1598 cm⁻¹ and 1590 cm⁻¹) and D band (1300 cm⁻¹, 1327 cm⁻¹ and 1348 cm⁻¹) of graphene were observed. Because of the reduction, I_D/I_G reduces to 0.78 (rGO) which is smaller compared with 0.99 (GO). Also, the CuFe₂O₄–rGO nanocomposite exhibited the characteristic CuFe₂O₄ Raman modes at 216, 281, 392 (E_{2g} and T_{2g}) and 597, 676 cm⁻¹ (A_{1g}). And I_D/I_G ratio of

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Fig. 3 Recorded FTIR of rGO, CuFe $_2O_4$ -(15 wt%) rGO, CuFe $_2O_4$ -(25 wt%) rGO, CuFe $_2O_4$ -(40 wt%) rGO and CuFe $_2O_4$



Fig. 4 Raman spectra of rGO, CuFe₂O₄, CuFe₂O₄-(15 wt%) rGO, CuFe₂O₄-(25 wt%) rGO and CuFe₂O₄-(40 wt%) rGO

nanocomposite was estimated to be 5.06, 1.07 and 0.91, which emphasis the dominance of structural defects in the composite that can result in variation in optical properties. The perfect graphene domain (La) which is inversely

proportional to I_D/I_G ratio was evaluated using the excitation energy (eV) by standard relation [44]. The crystallite sizes of CuFe₂O₄–(15, 25 and 40 wt%) rGO were calculated to be 1.38 nm, 6.53 nm and 7.76 nm, respectively. In fact, the change in I_D/I_G ratio along with La value emphasis the removal of oxygen element organic groups and the incorporation of bond between CuFe₂O₄ and rGO sheets during hydrothermal reaction. Altogether, chemical state (XPS), crystal structure (XRD), molecular structural (FTIR) and defect state (Raman) analysis confirms the successful synthesis of CuFe₂O₄–rGO nanocomposite through the hydrothermal treatment.

3.2 Textural, magnetic and linear optical properties

SEM and TEM techniques were used to analyze the structural morphology of rGO and $CuFe_2O_4$ -decorated rGO hetero-architectures, which inveterate the versatile morphology of $CuFe_2O_4$ -rGO nanostructures. It is manifesting to be observed that in rGO, exfoliated GO appears as transparent, ultrathin and distinguished sheets. Recorded SEM image (Fig. 5) showcases the existence of different morphologies for pure $CuFe_2O_4$ (clusters of platelets) and $CuFe_2O_4$ decorated upon 15 wt% rGO (spindle), 25 wt% rGO (aggregate of platelets and spindle) and 40 wt% rGO (well-distributed thin platelets). Here as graphene content increases, the nucleation formed on the surface of sheets preferred the formation of platelets like structures of CuFe₂O₄. TEM image (Fig. 6) of reduced graphene oxide illustrates curvy and wrinkled sheets of graphene attributed to solar exfoliation process. For pure CuFe₂O₄, the formation of nanoplatelets with 23 nm width was observed. A uniform distribution of nanoplatelets and nanospindles of copper ferrite functionalized upon rGO contents (15, 25 and 40 wt%) with size 9 nm, 17 nm and (67 × 28 nm), respectively, was witnessed. The increase in graphene content leads to the increase in the number of nucleation sites of CuFe₂O₄ and provides a transformation in the morphology from platelets to spindle aided from protrusions expanding with growth. The magnetic property of $CuFe_2O_4$ nanoparticles and $CuFe_2O_4$ -rGO nanocomposite using a vibrating sample magnetometer (VSM) was investigated. Figure 7 illustrates the S-type curve (hysteresis loop) of room temperature magnetized (with an applied magnetic field) nanocomposite. Copper ferrite being the magnetization contributing entity, its size and morphology play a significant role in deciding the ordering and magnetization of the sample. The magnetic remanences of CuFe₂O₄ nanoparticles and CuFe₂O₄-rGO composites were found to be in the range 10^{-6} emu/g which is nearly zero, suggesting its superparamagnetic behavior. Higher saturation magnetization and lower coercivity were observed



Fig. 5 SEM images of **a** rGO (layered structure), **b** CuFe₂O₄ (nanoplatelets), **c** CuFe₂O₄–(40 wt%) rGO (nanoplatelets of copper ferrite upon rGO layers), **d** CuFe₂O₄–(25 wt%) rGO (nanoplatelets of cop-

per ferrite upon rGO layers), ${\bf e}$ CuFe $_2O_4-(15$ wt%) rGO (nanospindle of copper ferrite upon rGO layers)



Fig. 6 TEM images of a rGO, b CuFe₂O₄, c CuFe₂O₄-(15 wt%) rGO, d CuFe₂O₄-(25 wt%) rGO, e CuFe₂O₄-(40 wt%) rGO nanocomposite



Fig. 7 Room temperature M–H curves of the pure $CuFe_2O_4$, $CuFe_2O_4$ –(15 wt%) rGO, $CuFe_2O_4$ –(25 wt%) rGO and $CuFe_2O_4$ –(40 wt%) rGO nanocomposite

for pure copper ferrite $(3.399 \times 10^{-3} \text{ emu}, 15.271 \text{ G})$ than for CuFe₂O₄-(40 wt%) rGO $(1.624 \times 10^{-3} \text{ emu}, 35.150 \text{ G})$. It is be noted that saturation magnetization of CuFe₂O₄ decreases upon loading in rGO. Similar superparamagnetism with higher coercivity due to increased magnetocrystallinity was reported for Fe-Al₂O₃ system [35, 45, 46]. Copper ferrites when scaled down to nanometer regime

SN Applied Sciences A SPRINGER NATURE journal alter the surface/interface properties, electronic states and magnetic properties. Nanocomposites containing both magnetic and NLO properties can offer interactions between magnetic susceptibility and nonlinear absorption in the material, which can be utilized to realize magnetocontrolled NLO materials. In such a case, by tuning the magnetic properties of the nanocomposites, the magnetic anisotropy of the magnetic ion at the interfaces can be reduced which leads to strong nonlinear optical response. Thus, with the application of external magnetic field it is possible to control the optical properties of the magnetic nanoparticles used in optical devices. Generally, the magneto-optical effect that transpires in magnetic fluids is believed to be the consequences of anisotropy in the dielectric constant (induced by the orientation of suspended particles). This induced anisotropy can also result in optical anisotropy either in the refractive index or absorbance or in both. A super paramagnetic behavior was observed mainly due to the contribution of copper ferrite nanoparticles, in which a fraction of Fe³⁺ is distributed out of the preferred octahedral A sites to tetrahedral B sites leading to the coupling between the cations. A larger extent of conjugation increases the chance for electron/energy transfer which leads to enhanced nonlinear optical response. Thus, functionalization of copper ferrite upon rGO has introduced the emergence of localized magnetic moment (super paramagnetic nature).

UV-visible absorbance data of the reduced graphene oxide, $CuFe_2O_4$ and $CuFe_2O_4$ -rGO nanocomposite are shown in Fig. 8. Upon reduction to rGO, the absorption maximum of GO was red shifted to 278 nm (238 nm) and 388 nm (290 nm). The absorptions of $CuFe_2O_4$ were observed at 445 nm (ligand field transitions) and 537 nm



(surface plasmon resonance (SPR), peak of Cu nanoparticle). The absorption peak of (CuFe₂O₄-rGO nanocomposite) at 260 nm, 394 nm, 436 nm and 541 nm is the indicative of CuFe₂O₄ deposition on rGO nanosheets. Compared to CdFe₂O₄-rGO nanocomposite [35], CuFe₂O₄-rGO shows higher absorbance at the chosen wavelength of excitation (800 nm) which makes them possible to observe nearresonant nonlinearity.

3.3 Nonlinear refraction

It is known that the ground-state absorption and ESA together can induce intensity-dependent refractive index [16]. With high repetition rate intense IR laser pulses, it can be generally expected that NLR arises mainly due to thermal lensing effects. Here, since the closed aperture transmittance is affected by the nonlinear refraction and absorption, the determination of n₂ is less straight forward from the closed aperture scans. It is necessary to separate the effect of nonlinear refraction from that of nonlinear absorption. A simple and effective method [47] to obtain purely effective n₂ is to divide the closed aperture transmittance by the corresponding open aperture scans. Figure 9 presents the ratio of closed (CA) aperture by open aperture Z-scan profiles recorded with a peak intensity of 245 MW/cm². Both pristine CuFe₂O₄ and CuFe₂O₄ functionalized upon (15, 25 and 40 wt%) rGO contents exhibit negative nonlinear refractive index-based self-defocusing nature. The value of n₂ was evaluated through theoretical fitting of closed aperture normalized transmittance



Fig. 8 UV-visible absorption spectra of \bm{a} rGO, CuFe_2O_4–(15 wt%) rGO, CuFe_2O_4–(25 wt%) rGO, CuFe_2O_4–(40 wt%) rGO and \bm{b} pure CuFe_2O_4

Fig. 9 CA Z-scan pattern of rGO, CuFe₂O₄–(15, 25 and 40 wt%) rGO and CuFe₂O₄ at an intensity of 245 MW cm⁻² of Ti:Sapphire (800 nm, 150 fs and 80 MHz) laser. The scattered circles represent the experimental data, and the solid line is the theoretical fit as proposed by Sheik-Bahae

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 (T_{CA}) [48]. As expected, pure rGO with its planar structure and capacity to dissipate heat quickly possesses stronger nonlinear refraction than pure CuFe₂O₄. Here rGO $(-2.8 \times 10^{-16} \text{ m}^2/\text{W})$ possesses two times higher n₂ than $CuFe_2O_4$ (-1.4×10⁻¹⁶ m²/W). In the $CuFe_2O_4$ -rGO nanocomposite, the increase in the content of graphene boosts the nonlinear refractive index. The nonlinear refractive index of CuFe₂O₄-(15, 25 and 40 wt%) rGO nanocomposite was estimated to be -1.7, -3.3 and -3.8×10^{-16} m²/W. The π -electron distribution and the free carriers of the sp² domain in graphene sheets are the prime reason for the occurrence of NLR in rGO. Here excitation with ultrafast lasers on the sample leads to transfer of carriers in the NP's state. Enhancement in carrier density is observed in the composite system due to the low lifetime of the excited carriers from the NP's to the valence band of rGO leading to the improvement in NLR. Strong thermal conductivity due to the planar structure of rGO and high repetition rate fs laser excitation results in dominant thermal contribution and high n₂. It is pertinent to note here that the excitation of the laser involves high repetition rate (80 MHz) which is much higher than the thermal diffusion time and there is a possibility that the cumulative thermal effects also contribute to the Z-scan signal. Excess energy from thermal dissipation redistributes within the solute molecules which turn them into intramolecular heat and may also further raise the solution temperature by dissipating eventually into the photon. Thus, the change of refractive index due to diffusion effect has considerable applications in optical communication systems that are operating at high repetition rate. As a result of the contribution of thermal nonlinearity, the NLO coefficients of the investigated materials were found to be higher than most of the earlier reports. Measurements with fs, kHz pulses will separate the thermal contribution, and these will be taken up in near future.

3.4 Intensity-dependent nonlinear absorption

Nonlinear optical absorption is broadly classified as two classes: Actual transitions give resonant nonlinearity which is slow and virtual transitions generate non-resonant nonlinearity that is fast in nature. Linear absorption is a predominant mechanism in the resonant region, while it is multi-photon processes in the non-resonant region. To understand the nonlinear absorption process of the prepared sample, the acquired data were fitted using the standard relation for open aperture normalized transmittance [49]. It is to be noted that the match found between numerical and experimental data just ascertains the reliability of the experiment performed and it does not confirm or indicate the occurrence of 2PA, to be genuine or sequential two-step ESA. Change

in the nonlinear absorption coefficient corresponding with input intensity can provide information on the mechanism involved. Genuine 2PA occurs when β-value remains constant for varied input intensity as weak 2PA cannot drain the population of electrons in the ground state. However, in ESA process due to substantial depletion of ground-state population nonlinear absorption coefficient either will increase or decrease as function of peak intensity. The OA data recorded for rGO and $CuFe_2O_4$ (Fig. 10a-f) show RSA pattern with minimum transmittance at the focus. As depicted in the linear absorption spectrum, materials possess small absorption in the domain of excitation wavelength, and hence, the observed nonlinearity is expected to be of resonant type. The changes in input intensity on the interaction with the material play an important role in 2PA process [50]. The measured data were found to be fitting well with the theoretical 2PA equation. The possible mechanisms for 2PA in such materials process are depicted in Fig. 11. Here, for pure rGO and CuFe₂O₄ 2PA coefficient was found to be increasing with input peak intensity. This clearly suggests that the observed nonlinearity arises due to ESA process, i.e., 1PA + ESA = 2PA. The source of the witnessed nonlinearity in rGO has occurred due to the process mentioned below: As witnessed in the ground-state (UV-Visible) absorption pattern of rGO and CuFe₂O₄, the materials possess considerable absorption in excitation region (800 nm, 1. 55 eV), and thus, the observed nonlinearity is of resonant type. Under NIR excitation (800 nm), electrons in the ground state absorb two photons (800 nm, 1.55 eV) through sequential process and transit to the excited state of rGO (388 nm, 3.2 eV) and pure copper ferrite (445 nm, 2.79 eV).

The OA pattern in Fig. 10b, c of the 15 wt% decorated system shows a combined SA and RSA pattern (W-pattern) for low input intensity ($I_0 = 245 \text{ MW cm}^{-2}$). The W-pattern with the 2PA process is formed due to the additional hump peak (Z=0) flanked between the two valleys representing SA peak. And the nonlinear absorption coefficient is affected by the existence of the two competing effects (RSA and SA). As the general open aperture equation for 2PA process cannot be modeled for W-type nonlinear absorption because of expected inhomogeneous broadening of transitions in these conditions, the 2PA coefficient is obtained by fitting the experimental data with following equations [51]

$$\beta(l) = \beta_{\mathsf{RSA}}(l) + \beta_{\mathsf{SA}}(l) = \frac{\beta_{\mathsf{RSA}}}{1 + \frac{l}{l_{\mathsf{S}}^{\mathsf{RSA}}}} + \frac{\beta_{\mathsf{SA}}}{1 + \frac{l}{l_{\mathsf{S}}^{\mathsf{RSA}}}}$$

2PA coefficients obtained for the best fits to the experimental data are presented in Table 1. The experiments

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Fig. 10 Intensity-dependent OA Z-scan curve of pure **a** rGO, **b** CuFe₂O₄–(15 wt%) rGO (245 MW/cm²), **c** CuFe₂O₄– (15 wt%) rGO (245–735 MW/ cm²), **d** CuFe₂O₄–(25 wt%) rGO, **e** CuFe₂O₄–(40 wt%) rGO and **f** CuFe₂O₄. The experimental OA data (represented as scattered circles) were fitted (solid line) with the theoretical equations as proposed by Sheik Bahae





Fig. 11 Possible electronic transitions for observed 1PA+ESA=2PA in pure and decorated systems. Here 2PA involves $n-\pi^*$ transition state of rGO and CuFe₂O₄-rGO nanocomposite and SPR peak of Cu nanoparticles for pure CuFe₂O₄

were performed on an automated Z-scan experimental setup. The errors in estimation of focal spot size, peak intensities, fitting procedures and calibration of intensity are major sources of error in the estimation of the NLO coefficients. Based on several measurements, the maximum error was estimated to be \pm 5%. As the error correction is employed uniformly, the change in NLO coefficients of different samples was directly compared.

It is evident that since the sample shows negligible linear transmittance at 800 nm, the contribution of linear absorption coefficient (α_0) is neglected. The dominance of RSA pattern was observed for the higher excitation intensity, thus providing stronger 2PA coefficient, while for copper ferrite-decorated (25 wt% and 40 wt%) reduced graphene oxide systems it displayed RSA (valley) at lower input intensity ($I_0 = 245$ MW cm⁻²). Therefore, it can be

NLO parameters	800 nm, 150 fs, 80 MHz excitation					
	Nonlinear absorption coefficient (β_{2PA}) × 10 ⁻¹⁰ m/W			Onset optical limiting threshold (µJ/cm ²)		
	$I_{\rm o}$ =245 MW/cm ²	$I_0 = 490 \text{ MW/cm}^2$	$I_{\rm o}$ =735 MW/cm ²	$I_0 = 245 \text{ MW/cm}^2$	$I_0 = 490 \text{ MW/cm}^2$	$I_0 = 735 \text{ MW/cm}^2$
rGO	2.1±0.10	2.5 ± 0.12	2.7±0.13	38.45	35.66	31.42
CuFe ₂ O ₄	0.7 ± 0.03	1.0 ± 0.05	3.3 ± 0.16	57.50	54.12	27.53
CuFe ₂ O ₄ –(15 wt%) rGO	0.17 ± 0.01	0.9 ± 0.04	1.1 ± 0.05	-	56.73	55.59
CuFe ₂ O ₄ –(25 wt%) rGO	1.0 ± 0.05	1.4 ± 0.14	6.7±0.33	57.50	45.22	22.29
CuFe ₂ O ₄ -(40 wt%) rGO	1.3 ± 0.06	1.7 ± 1.70	12.7 ± 0.63	52.38	35.36	12.45

Table 1 Estimated nonlinear absorption coefficient and onset optical limiting threshold values

deduced that as the size of CuFe₂O₄-rGO increases, the effect of RSA increased suppressing SA behavior. This might be due to greater cross section of excited states than ground state for larger particles. For all higher intensities ($I_0 = 490 \text{ MW cm}^{-2}$ and 735 MW cm $^{-2}$), decorated rGO systems exhibited reverse saturable absorption. Here the prepared reduced graphene oxide contains both sp² (Pristine graphene honeycomb lattice) and sp³ (defects state and oxygen-containing functional groups) hybridized domains. sp³ hybridization in the carbon units of rGO leads to 2PA at different ($I_0 = 245 \text{ MW cm}^{-2}$, $I_0 = 490 \text{ MW cm}^{-2}$ and $I_{o} = 735 \text{ MW cm}^{-2}$) intensities provided by ultrafast pulses. It is to be mentioned that decorated systems depict linear absorption maxima at 394 nm due to $n-\pi^*$ transition in the rGO. As the composites possess energy states near to excitation wavelength, the resonant nonlinearity gives rise to sequential 2PA (two photons were sequentially absorbed) rather than genuine 2PA. The incorporation of $CuFe_2O_4$ on the rGO system results in the participation of σ -states (sp³) in 2PA process. The increase in irradiation of input fluence to 735 MW/cm² on the decorated system leads to very deep decrease in the transmittance as presented in Fig. 10e which is a sign of strong reverse saturable behavior. The present results of nonlinear absorption coefficient were compared with the other well-known system in Table 2 reported at same experimental setup. With increasing β for different samples considered, the difference between the valley depth and the peak height in nonlinear refraction varies. And the amplitude of the valley is almost as large as peak, referring the involvement of Kerr-type nonlinearity [52]. Thus, considering the above observation along with fact that the repetition rate of the laser that has been used in the experimental procedure is 80 MHz, the influence of thermal effects along with the electronic effects cannot be ignored.

3.5 Optical limiting behavior

The changes in transmission of the prepared hybrids were recorded against input fluence using open aperture

SN Applied Sciences A Springer Nature journal Table 2Comparison of the two-photon absorption coefficientsobtained for the carbon and metal/semiconductor nanoparticles[Ti:Sapphire (~800 nm)]

Samples	Nonlinear absorp- tion coefficient, β_{2PA} (m/W)		
CdFe ₂ O ₄ -rGO [26]	10 ⁻¹⁵		
ZnFe ₂ O ₄ –rGO [<mark>48</mark>]	10 ⁻¹⁵		
GO and rGO [55]	10 ⁻¹³		
SiO ₂ –Se [56]	10 ⁻¹⁵		
CuS nanoparticles [57]	10 ⁻¹⁵		
Ni–Ag [58]	10 ⁻¹⁵		
CuO/f-HEG [59]	10 ⁻¹⁴		
Fe ₂ O ₃ [60]	10 ⁻¹⁵		

Z-scan experiment. This experiment is capable of recording the variation in light fluence [F(z)] for corresponding beam radius $\omega(z)$ [53, 54]. The optical limiting threshold is defined as the incident fluence at which the transmittance falls to 50% of the linear transmittance. Here all the optical limiting curves were extracted from the open aperture Z-scan data and the fluence range $(3-60 \mu J/cm^2)$ is limited to the designed Z-scan geometry. As direct measurement of output with change in input intensity is not measured, the curve has the limitation of extending (or extrapolating the data) until the transmittance falls off to 50%. Hence, as an alternate approach, onset optical limiting threshold is reported which is the incident fluence at which optical limiting activity starts. The OL curve derived from the open aperture pattern demonstrates the changes in transmittance value with respect to input fluence as represented in Fig. 12. The ability to deliver ultrafast pulses with controlled energy through flexible fibers can open up a host of possibilities toward developing robust ultrafast laser microsurgery systems. It is to be noted that as short pulse lasers allow high precision microsurgery with minimal invasiveness, high repetition rate femtosecond laser is under prime consideration. Hence, femtosecond laser microsurgery has emerged strongly in tissue ablation,







studying neural injury, and cleaving microtubules which require energy pulses on the order of microjoules. Thus, the observed optical limiting of $CuFe_2O_4$ -decorated (40 wt%) rGO with onset limiting threshold in the order of few µJ/cm² is of potential use in biomedical applications. Here, all the samples exhibit optical limiting (due to 2PA and SDF) pattern expect $CuFe_2O_4$ -(15 wt%) rGO at 245 MW/cm² due to the existence of dominant SA process (W-pattern). As the nonlinear absorption coefficient is intensity dependent, here 2PA coefficient increases with the increase in the number of photons used for excitation. Pure rGO possesses higher 2PA coefficient compared with pure copper ferrite. At 245 MW/cm², copper ferrite possesses higher 2PA than rGO. This combination of two extreme 2PA absorbed resulted in the variation in 2PA coefficient with change in rGO content. Among the composite, higher 2PA coefficient was observed for (40 wt%) rGO followed by (25 wt% and 15 wt%) rGO. It is interesting to note that at lower input intensity excitation rGO possesses stronger 2PA coefficient compared with the nanocomposites. Here the lowest nonlinear absorption coefficient was witnessed for (15 wt%) rGO due to the combined presence of stronger SA and weaker RSA. As witnessed in TEM, higher content of CuFe₂O₄ upon rGO has suppressed the contribution of rGO and thus possesses lower nonlinear absorption coefficient compared to pure rGO and CuFe₂O₄. However, (25 wt% and 40 wt%) rGO has a stronger 2PA coefficient than its pure counterparts. The maximum 2PA

coefficient was witnessed for $CuFe_2O_4$ -(40 wt%) rGO at 735 MW/cm² excitation due to the combined contribution of rGO and $CuFe_2O_4$ which is four times higher than pure rGO and $CuFe_2O_4$. The presence of layer like $CuFe_2O_4$ nanoplatelets upon the layer rGO offers higher advantage in enhancing the nonlinear absorption coefficient. As discussed in the literature, similar tuning of NLO properties with respect to the grain size of the material was observed in CuCl nanocrystals, ZnO nanocolloids, ZnFe₂O₄ and CdO [61–63]. The decoration of copper ferrite with different concentrations upon rGO sheets with varied morphology results in acquiring tunability of NLO properties. Thus, nanoplatelet $CuFe_2O_4$ -(40 wt%) rGO can be potential energyabsorbing ESA-based optical limiters against Ti:Sapphire (150 fs, 800 nm and 80 MHz) laser pulses.

4 Conclusions

CuFe₂O₄ was made to anchor upon (15, 25 and 40 wt%) rGO contents by hydrothermal method where electrostatic force plays an important role in distributing and holding various sizes of particles. The decoration of CuFe₂O₄ nanostructures upon rGO nanosheets was confirmed by analyzing their XPS, XRD, FTIR, VSM, Raman, SEM, TEM and UV-Vis absorption studies. Higher content of rGO with CuFe₂O₄ nanoplatelets exhibited superior self-defocusing (NLR) and 2PA (NLA) process than other lower rGO content hybrid structure. Intensity-dependent nonlinear absorption studies showcase the following features of CuFe₂O₄-rGO composites (1) CuFe₂O₄–(15 wt%) rGO at lower intensity of excitation exhibits W-pattern with dominant saturable absorption behavior arising from the influence of Pauli blocking rGO, (2) CuFe₂O₄-rGO composite exhibits nearresonant reverse saturable absorption and is ascribed to be sequential 2PA (1PA + ESA) process and (3) CuFe₂O₄ nanoplatelets decorated upon 40 wt% rGO exhibited superior self-defocusing (NLR) and 2PA (NLA) process than other lower rGO content hybrid structure due to the interaction between the layered morphologies of host and decorating molecules. The prepared CuFe₂O₄ nanoplatelet-decorated rGO hybrid layer structure provides an insight on the possible way to enhance NLO properties by anchoring layered complex oxides upon carbonaceous-layered materials.

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Compliance with ethical standards

Conflict of interest The author(s) declare that they have no competing interests.

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