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Enhancement of nanograting formation assisted by silver ions in a sodium gallophosphate glass

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We investigated the influence of silver ions during the direct femtosecond laser induced formation of nanogratings (NGs) at the surface of a tailored sodium gallophosphate glass. We observed that the silver ions had a remarkable influence because the silver containing glass showed (1) lower fluence thresholds for the formation of the NGs; (2) much smoother NG shapes; and (3) a bifunctional behavior because fluorescence from laser induced silver clusters occurs prior to the formation of the NGs. We demonstrate for the first time, to our knowledge, the formation of NGs assisted by noble metal ions, such as ions playing a kind of catalytic like role that enhances and improves the NG formation and its incubation process. Our innovative approach provides promising potential for further improvements in processes for NG formation.

OCIS codes: (100.2540) Fluorescent and luminescent materials; (140.3900) Laser materials processing; (140.3450) Laser induced chemistry; (190.4180) Multiphoton processes; (310.6628) Subwavelength structures, nanostructures.

Femtosecond (fs) laser manufacturing has led to continuous innovation in the last decade, especially in the inscription of structures with inner dimensions smaller than those of the considered beam size, such as the formation of nanogratings (NGs), which shows great potential for optical waveguides, solar cells, and biological devices [1,3]. The spatial distributions of NGs in bulk silica glass have been reported in detail in pioneer and recent works [4-7], both for bulk and surface NGs. Different mechanisms and descriptions have been proposed, including the incubation model [8]. While NGs have mostly been studied in silica by precisely tuning the laser irradiation parameters, very little attention has been paid to the material itself until now. No effort has been clearly made to adapt the considered glass matrices, either by exploring exotic compositions or by engineering a photosensitized material with the introduction of noble metal ions [9]. While silica cannot stand large amounts of silver ions [10], phosphate glasses can sustain high concentrations, allowing the direct laser writing (DLW) of strong and stable linear and nonlinear optical properties, such as silver cluster fluorescence, third-harmonic generation, or electric field-induced second-harmonic generation [11,16]. In this framework, the photochemistry of silver ions triggered under fs laser irradiation has never been explored for NG formation.

In this Letter, we study the role of silver ions incorporated in a tailored sodium gallophosphate matrix during the NG formation at its surface by means of fs laser irradiation. Accordingly, we compare the behaviors of such a sodium gallophosphate glass matrix with and without silver ions. In the silver-containing glass, we observed significant improvements due to the silver ions, such as lower fluence thresholds for the NG formations and much smoother NG shapes (also much smoother than the usual nanostructures, as in fused silica [17]). We also observed that fluorescent silver clusters are created prior to NG formation at lower fluencies, leading to spatially correlated fluorescent patterns and NGs. Moreover, material removal was eased in the case of the silver-containing glass, with cleaner final NGs. Our innovative approach, based on noble metal doping, creates the potential for the optimal engineering and generation of NGs.

Glass samples with the initial composition 20.0Ga2O3-40.0Na2O-40.0P2O5 (mol. %) and 97[20.0Ga2O3-40.0Na2O-40.0P2O5]-3Ag2O (mol. %), respectively labeled GPN and GPN-Ag, were synthesized using the conventional melt-quenching technique. The proper amount of Ga2O3, NaPO3, and Ag2O powders were thoroughly mixed, placed in a 100% Pt platinum crucible, heated up to 1100°C in ambient air, and kept at maximum temperature for 30 min before quenching. The glass samples obtained were annealed for 6 h at 40°C below the glass transition temperature of each sample. Further description of the preparation and characterizations of the sodium gallophosphate glasses can be found in [18]. The GPN and GPN-Ag samples were cut into pieces with 1 mm thickness. Finally, one of each were bound together to a silica glass substrate and polished to optical quality. This allows the two material samples to share the same interface, which guaranteed identical focusing conditions during laser structuring.

The experimental setup of the NG formation, as detailed elsewhere [8], is composed of a commercial Ti:Sapphire chirped-pulse amplification laser system (Spitfire, Spectra-Physics) with a maximum output energy of 2 mJ per pulse at the central wavelength λlaser = 800 nm, with a pulse duration τ = 45 fs (FWHM) and a repetition rate frep = 1 kHz. The beam was tightly focused at the surface sample with a microscope objective (Mitutoyo, 50×, N.A. = 0.42), which was adjusted by maximizing the retro-reflection from a He Ne laser beam. The diameter of the focal spot was estimated at 2 × ω0 = 2 μm. The samples were mounted on motorized three-dimensional (3D) translation stages (Melles Griot). The
considered writing speeds, with sample motion at constant speed from \( v = 10 \) to 100 \( \mu \text{m/s} \), allowed the estimation of the number of pulses \( N \) that overlapped at each position during laser structuring, as \( N = 2 \times \omega_o \times f_{rep} / v \). The laser beam was linearly polarized perpendicular to the sample translation direction. The energy per pulse \( (E_p = 10 \) to 100 nJ) was adjusted with a half-wave plate and a Glan polarizer and was measured after the objective lens. The estimated pulse fluence is \( F_p = E_p / \pi \times \omega_o^2 \), whereas the effective beam fluence is defined as \( F = N \times F_p \). After irradiation, the samples were imaged first by standard bright-field and fluorescence microscopy. Then, the samples were gently cleaned in an ultrasonic bath with distilled water for 4 min and characterized by a scanning electron microscope (SEM in the secondary electron emission mode) (FEI, model Quanta 3D FEG).

Both the silver-free and silver-containing samples were laser-processed and subsequently characterized under the same experimental conditions to ensure that the differences between GPN and GPN-Ag can only be attributed to the presence of silver ions.

The refractive indices at \( \lambda_{\text{near}} \) of the GPN and GPN-Ag samples were estimated at 1.536 and 1.544, respectively, as inferred from the Sellmeier fitting of M-lines prism coupling measurements (Metricon 2010) at different wavelengths (532, 633, 972, 1308, and 1538 nm). The close values of the refractive indices of both the Ag-doped and nondoped GPN samples ensure that the focusing conditions will be similar for both glasses. Optical transmission spectroscopy of the two pristine glasses led to the estimation of the linear absorption coefficient in the UV-visible range. As seen in Fig. 1(a), the transmission cutoff of the GPN-Ag is significantly shifted toward longer wavelengths, suggesting a potentially much larger nonlinear three-photon absorption cross section (at longer wavelengths, suggesting a potentially much larger absorption coefficient to the Ag\(^{+} \) insertion. Normalized excitation (emission wave length at 412 nm) and emission fluorescence (excitation wave length at 272 nm) spectra of the pristine GPN Ag glass. Absence of emission in the nondoped GPN sample confirms that these fluorescent properties result from the silver ions.

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Fluorescence spectroscopy revealed the silver-doping influence because the pristine GPN-Ag sample showed a strong excitation band centered at 272 nm, related to a broad emission band centered at 412 nm, as shown in Fig. 1(b). Such fluorescence behavior is typical of silver ion-containing glasses [19], whereas the nondoped sample showed no fluorescence response. The excitation band of silver ions is to be correlated with the shift of the transmission cutoff in the GPN-Ag, strongly suggesting that the introduction of such ions is responsible for enhanced nonlinear absorption during the laser structuring process.

Laser irradiations were first performed at constant speed (20 \( \mu \text{m/s} \)) by increasing the pulse energy from 10 to 65 nJ. Features were observed under bright-field microscopy and luminescence imaging (excitation by a laser diode at 483 ± 35 nm and collection after a band-pass filter at 536 ± 40 nm). The GPN sample showed a damage threshold at 35–40 nJ per pulse with bright-field microscopy [Fig. 2(a)], but it showed no sign of a fluorescence response [Fig. 2(c)]. Comparatively, the GPN-Ag sample showed a much lower threshold, with optical damages starting from 25–30 nJ [Fig. 2(b)]. Additionally, GPN-Ag led to a fluorescent emission [Fig. 2(d)], which is typical of the laser-induced creation of silver clusters [12], with an increasing fluorescence emission while increasing the input pulse energies. Probing the associated migration of silver ions during the creation of silver clusters is still in progress. Fluorescent clusters appeared prior to the damage threshold in GPN-Ag, demonstrating the photoreactivity of the silver ions in the early stages of the glass matrix modifications toward NG formation. Finally, no evidence of metallic silver nanoparticle formation was observed under such laser irradiances, suggesting their absence in these experimental conditions.

Once irradiated and softly cleaned with distilled water, the revealed structures were then observed with SEM, showing a NG formation threshold between 25 and 30 nJ per pulse in GPN-Ag (Fig. 3), which is significantly lower than the 35–40 nJ required in the nondoped GPN sample (Fig. 3). In addition to the lowering of the NG formation threshold, the SEM images also revealed that the NGs in GPN-Ag were impressively smoother than those in GPN. Their full width—labeled W—is also wider for given irradiation conditions, including a larger number of grooves. The width of individual grooves typically varied from 10 to 65 nJ for decreasing pulse energies between 25 and 10 nJ. Features were observed under bright-field microscopy and luminescence imaging (excitation by a laser diode at 483 ± 35 nm and collection after a band-pass filter at 536 ± 40 nm). The GPN sample showed a damage threshold at 35–40 nJ per pulse with bright-field microscopy [Fig. 2(a)], but it showed no sign of a fluorescence response [Fig. 2(c)]. Comparatively, the GPN-Ag sample showed a much lower threshold, with optical damages starting from 25–30 nJ [Fig. 2(b)]. Additionally, GPN-Ag led to a fluorescent emission [Fig. 2(d)], which is typical of the laser-induced creation of silver clusters [12], with an increasing fluorescence emission while increasing the input pulse energies. Probing the associated migration of silver ions during the creation of silver clusters is still in progress. Fluorescent clusters appeared prior to the damage threshold in GPN-Ag, demonstrating the photoreactivity of the silver ions in the early stages of the glass matrix modifications toward NG formation. Finally, no evidence of metallic silver nanoparticle formation was observed under such laser irradiances, suggesting their absence in these experimental conditions.

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spans from 44 to 290 nm for GPN-Ag compared to 146 to 204 nm for GPN (Fig. 3). These observations, especially the lower limit of 44 nm in GPN-Ag (as qualitatively visible at the edges of the NGs for 40, 45, and 50 nJ per pulse), illustrate the different dynamics of both the ionization rates and incubation processes observed in the two samples—such aspects directly influencing the morphology of the NGs, as evidenced in Fig. 4. The optical damage threshold increases with the scan speed of the sample, with higher thresholds for the nondoped sample [Fig. 4(a)]. Moreover, NGs in GPN-Ag showed a larger width and number of grooves compared to those in GPN, which were assessed from the SEM images, leading to distinct typical square root dependences versus pulse energies [Fig. 4(b)] in agreement with the incubation dynamics proposed elsewhere [8,20]. Further optical characterization of the diffraction efficiencies of the NGs will require additional work with larger structured areas.

The preceding results can be interpreted as follows. First, the pulse energy threshold for optical damage grows more or less linearly with scan speed, which is the signature of a beam fluence-dependent process. On the contrary, the fluorescence threshold is independent of scan speed and solely depends on pulse energy, which is related to the beam intensity. The insertion of a doping species (the silver ions herein) provides a higher ability to release free electrons from nonlinear absorption under fs irradiation as compared to the glass matrix itself. In this sense, the silver ion population behaves as an extra reservoir of accessible electrons, more efficiently released than those from the glass matrix. Once released into the glass matrix, this larger amount of electrons can more efficiently trigger the incubation process via the enhanced creation of defects, which explains the lowering of the NG formation threshold in GPN-Ag. Moreover, ionizing silver ions is much less destabilizing for the glass network than ejecting electrons from the glass matrix itself, which corroborates the much smoother NG structures in the doped glass. Through this approach, the initial ionization step is better controlled and, somehow, the initial homogeneous distribution of silver ions makes this first step less affected by random variations in the nondoped material. Then, in the second step, the creation of fluorescent silver clusters indicates that silver also undergoes reduction processes. Because Ag⁺ ions initially behave as potential electron donors, whereas silver clusters behave as electron acceptors, silver species globally act as a kind of catalytic-like actor during the laser structuring and the NG formation, even though silver globally undergoes irreversible reactions, with the production of new and stable silver clusters. Therefore, silver ions play an active role in the series of phenomena that occur during NG formation, most likely by providing additional free electrons and defects but also by making the series of phenomena free from the stochastic distributions of defects in the nondoped material. These considerations lead to the enhancement of NG formation with larger grooves but also to significantly improved quality and smoothness. Such an innovative approach could be extended to some other elements with the potential ability to act as homogeneously distributed defects inserted in a glassy matrix, as far as such doping species show a sufficiently high solubility in the selected glass system. To better understand the silver-mediated processes, further studies should also investigate ionic motions (for instance, the potential migration of silver ions).

In this Letter, we have reported on NGs obtained in a new phosphate glass composition. We have presented the first demonstration (to our knowledge) of the silver ion-mediated formation of NGs, which led to a significant lowering of the NG formation threshold, wider structures with a larger number of grooves correlated to silver cluster fluorescence properties, and a remarkable improvement of NG smoothness. By providing a reservoir of available free electrons, silver ions appear as electron donors that also allow overcoming the stochastic distribution of defects in the glassy matrix, enhancing the NG formation. Such a very general approach (restricted to the specific use of silver ions herein) should further
provide enhanced abilities for the manufacturing of high-quality NGs for photonic devices.

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