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Author(s)

Meng, Xiangeng; Fujita, Koji; Zong, Yanhua; Murai, Shunsuke; Tanaka, Katsuhisa

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Random lasers with coherent feedback from highly transparent polymer films embedded with silver nanoparticles

Xiangeng Meng,1 Koji Fujita,1,2,a) Yanhua Zong,1 Shunsuke Murai,1 and Katsuhisa Tanaka1
1Department of Material Chemistry, Graduate School of Engineering, Kyoto University, Katsura, Nishikyo-ku 615-8510, Kyoto, Japan
2PRESTO, Japan Science and Technology Agency (JST), 4-1-8, Honcho Kawaguchi, Saitama 332-0012, Japan

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We report on observations of random lasers with coherent feedback in highly transparent polymer films embedded with silver nanoparticles. The hybrid materials were fabricated via in situ synthesis method, through which silver nanoparticles were precipitated by thermal treatment. Sharp peaks with linewidth ~0.5 nm were observed to emerge on the broad emission background when the pump energy reached a threshold, together with unidirectional laser irradiation. Random lasers with coherent feedback induced by silver nanoparticles have been rarely reported, hence, we expect that this work will add an aspect to random lasers by using metal nanostructures to obtain coherent feedback. © 2008 American Institute of Physics. [DOI: 10.1063/1.2912527]

Random lasers have attracted tremendous attention since they theoretically concern the fundamental physics of Anderson localization for photon and experimentally provide potential laser devices readily accessible from a technical point of view.1–6 Random lasers operate based on light scattering, that is why the scattering strength of random media is crucial to evaluate whether a coherent feedback is involved or not. Generally, weak scattering strength results in a gradual narrowing of emission spectrum with linewidth in scale of several nanometers while strong scattering strength gives rise to discrete sharp peaks with linewidth less than 1 nm superimposed on the emission spectrum. In order to achieve coherent feedback, dielectric materials with high refractive index (e.g., GaP, TiO2, and ZnO) have been utilized for several decades because large refractive index contrast significantly contributes to the intense scattering according to the Mie scattering theory.7–9

Metal nanoparticles are potential candidates that may play an important role in coherent feedback in random lasers. The potential of metal nanoparticles toward applications in random lasers includes two aspects. First of all, metal nanoparticles have much larger scattering cross section (σr) than dielectric nanoparticles with the same dimensions. Second, metal nanoparticles are more advantageous in their unique surface plasmon resonance (SPR) property, which has proved to spatially confine light wave near particle surface so that high gain for lasing is expected. Unfortunately, metal nanoparticles toward random lasers suffer from light absorption that is detrimental to laser oscillation and negligible for dielectric nanoparticles-based random lasers. This may explain why coherent feedback random lasers have not been observed in media composed of metal nanoparticles.10,11

In this letter, we report on a breakthrough in metal nanoparticles-based random lasers: random lasers with coherent feedback have been observed in polymer films embedded with silver nanoparticles. Silver nanoparticles embedded inside polymer films were prepared via in situ synthesis.12 In the procedure, the mixture of polyvinyl alcohol (PVA), silver nitrate, and rhodamine 6G (R6G) aqueous solutions is spin coated on glass substrate followed by annealing, leading to the precipitation of silver nanoparticles. The evolution of silver nanoparticles in PVA matrix against annealing time was characterized by optical absorption spectra recorded with a JASCO-V570 ultraviolet-visible-near-infrared spectrophotometer. The samples were optically pumped by the second harmonics of a mode-locked Nd:yttrium aluminum garnet (YAG) laser (λ=1064 nm, 10 Hz repetition rate, 25 ps pulse duration) in order to conduct random laser experiments. The laser beam was focused on the samples through a cylindrical lens to form a stripe with a width of 17 μm and a length that could be varied from 0.1 to 6.0 mm. Figure 1 shows the configuration of PVA-Ag-R6G film and excitation-detection. The emission was collected along the stripe with a fiber device into a monochromator (SPEX 270M) coupled to a liquid nitrogen cooled charge couple device (CCD 3000, the spectral resolution is 0.07 nm).

Figure 2(a) describes the evolution of optical absorption spectra for PVA-Ag-R6G films as a function of annealing time at 110 °C. An absorption band at around λ=420 nm

FIG. 1. (Color online) Schematic illustration of configuration of PVA-Ag-R6G film and excitation-detection. The PVA acts as a matrix while silver nanoparticles as scattering centers and R6G as a light emitting material. The film thickness is about 4 μm. The sample is pumped by the second harmonics of a mode-locked Nd:YAG laser (λ=1064 nm, 10 Hz repetition rate, 25 ps pulse duration). The inset is a photograph of the sample (top view).

*a)Electronic mail: fujita@dipole7.kuic.kyoto-u.ac.jp.
emerged when the annealing time reached 30 min and then gradually rose with the increase in annealing time and finally saturated after 420 min annealing. This band can be ascribed to SPR signal of spherical silver nanocrystals. During the thermal treatment, the PVA reduces Ag\(^+\) ion while the alcohol is oxidized to ketone.\(^{13}\) The absorption band at around \(\lambda=530\) nm is due to the electronic transition from the ground state \(S_0\) to the first singlet excited state \(S_1\) of R6G. Figure 2(b) presents the difference in absorption spectra between annealed samples and as-prepared one along with the simulation (solid lines) by the Mie scattering theory and Drude model. The sample was annealed at 110 °C.

The extinction cross section of one particle consists the scattering and absorption cross section \(\sigma_e\) and \(\sigma_a\), namely, \(\sigma_e=\sigma_s+\sigma_a\). The scattering cross section of one Ag particle is given by the Mie scattering theory and Drude model. The scattering and absorption cross sections \(\sigma_s\) and \(\sigma_a\) are calculated by using the formulas:

\[
\sigma_s = \frac{8\pi a^2}{k^2} \left(1 - \frac{1}{k^2}ight) \left(1 - \frac{4a^2}{k^2}ight) \left(1 - \frac{3a^2}{k^2}ight)
\]

\[
\sigma_a = \frac{8\pi a^2}{k^2} \left(1 - \frac{1}{k^2}ight) \frac{1}{\epsilon-1}
\]

where \(a\) is the radius of Ag nanoparticles, \(k\) is the wave number of light, and \(\epsilon\) is the complex dielectric constant of Ag nanoparticles.

The difference spectra \(\Delta\sigma = \sigma_{\text{annealed}} - \sigma_{\text{as-prepared}}\) between the annealed samples and as-prepared one along with the simulated curves by the Mie scattering theory and Drude model is shown in Fig. 2(b). The curves at longer wavelength are not involved in the simulation process since they are significantly affected by thermal treatment, the PVA reduces Ag\(^+\) ion while the alcohol is oxidized to ketone.\(^{13}\) The absorption band at around \(\lambda=530\) nm is due to the electronic transition from the ground state \(S_0\) to the first singlet excited state \(S_1\) of R6G. Figure 2(b) presents the difference in absorption spectra between annealed samples and as-prepared one, as well as the simulation curves by using the Mie scattering theory and Drude model.\(^{14}\) The curves at longer wavelength are not involved in the simulation process since they are significantly affected by the absorption band at around \(\lambda=530\) nm of R6G. The evolution of SPR signal is very obvious with an increase in the annealing time. The simulation result indicates that the average size of Ag nanoparticles is \(~1.0\) nm in radius while the volume fraction of silver nanoparticles relative to PVA is \(~3.8 \times 10^{-4}\). The particle size is consistent with the previous report.\(^{12}\) It should be noted that no change of appearance has been found in the sample during the 532 nm laser irradiation in laser experiment, indicating that the pump laser has no influence on silver nanoparticles in PVA matrix.

Figure 3 depicts the evolution of emission spectra as a function of pump energy. The emission spectrum shows a broad spontaneous emission band with a linewidth of \(~32\) nm when the pump energy is low. However, when the pump energy reaches the threshold (herein \(~1.12\) \(\mu\)J), a sharp peak that centers at \(\lambda=567\) nm with a linewidth of \(~0.5\) nm suddenly emerges on the broad amplified spontaneous emission background. In contrast, the sample containing no silver nanoparticles merely showed a spontaneous emission band, indicating that silver nanoparticles have played a crucial role in the emergence of sharp peaks. With a further increase in pump energy, more sharp peaks are observed in the broad emission background. For a clear demonstration of sharp peaks, the emission spectra just below and above the pump threshold are inserted in Fig. 3. We suppose that these peaks are really due to laser oscillation by analyzing the evolution of emission intensity as a function of pump energy that clearly confirms a threshold behavior, as shown in Fig. 4. Another evidence for laser oscillation is the observation of unidirectional laser emission, as shown in the inset of Fig. 4, since the spontaneous emission is lack of unidirectional irradiation. The results in Fig. 4 resemble reported random lasers with coherent feedback.\(^{53}\) For instance, the number of laser modes is small when the pump energy is relatively low since only those cavities with gain high enough can lead to lasing. The increase in pump energy increases the number of cavities that contribute to the lasing.

The extinction cross section \(\sigma_e\) of one particle consists the scattering and absorption cross section \(\sigma_e\), namely, \(\sigma_e=\sigma_s+\sigma_a\). The scattering cross section of one Ag particle is given by the Mie scattering theory and Drude model.
nanoparticle with 1.0 nm in radius is $\sigma_s \sim 1.5 \times 10^{-24}$ m$^2$ while $\sigma_a \sim 5.3 \times 10^{-21}$ m$^2$ at $\lambda = 567$ nm. Therefore, the scattering intensity of such a silver nanoparticle is much weaker than the absorption intensity. However, we have found certain enhancement of fluorescence instead of fluorescence quenching for all the samples containing silver nanoparticles. The results indicate that the enhancement is possibly caused by SPR, as previously revealed. The scattering mean free path $l_s$ can be estimated via the Mie theory $l_s = 1/\rho \sigma_s$, where $\rho$ is the number density of silver nanoparticles in PVA host. The $\rho$ is calculated to be $\sim 9.1 \times 10^{22}$/m$^3$, thus, $l_s \sim 7.3 \times 10^{-3}$ mm. Therefore, $k l_s \sim 1.2 \times 10^8 \gg 1$, implying that the present system operates in extremely weak scattering regime. Although coherent feedback random lasers have been observed in dielectric based random media, the scattering strength as weak as that in this work has never been conducted. We surmise that the observed lasing peaks are associated with the highly localized optical modes due to SPR.

The present random media include three important characteristics that make it distinct from existing random media. First, metal nanoparticles instead of dielectric nanoparticles are used as scattering centers. Second, the particle size is much smaller than the emission wavelength, which rules out the possibility of single particle as a resonator. Third, the scattering strength is extremely weak, which makes polymer films highly transparent for efficient optical pumping. Furthermore, the number density and particle size of silver nanoparticles in PVA matrix can be facilely manipulated through annealing temperature and time, as well as the initial concentration of Ag$^+$ ions.

In summary, we have observed random lasers with coherent feedback in highly transparent polymer films embedded with silver nanoparticles. The scattering mean free path estimated by the Mie theory is about eight orders of magnitude larger than the emission wavelength, indicating that normal light scattering barely contributes to the emergence of coherent feedback. We infer that the observed random lasers with coherent feedback are closely associated with highly localized light modes caused by SPR interaction between silver nanoparticles and light. We expect that this work adds an aspect to the fields of random lasers with metal nanostructure-based random media.

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