Optical excitation cross section of erbium in GaN

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Epilayers of erbium-doped GaN (GaN:Er) were synthesized by metal-organic chemical vapor deposition, and the optical excitation cross section ($\sigma_{\text{exc}}$) of Er ions in this host material were determined. Photoluminescence (PL) measurements were made using laser diodes at excitation wavelengths of 375 and 405 nm, and the integrated emission intensity at 1.54 $\mu$m was measured as a function of excitation photon flux. Together with time-resolved PL measurements, values of $\sigma_{\text{exc}}$ of Er ions in GaN:Er were obtained. For excitation at 375 nm, the observed excitation cross section was found to be $4.6 \times 10^{-17}$ cm$^{-2}$, which is approximately three orders of magnitude larger than that using resonant excitation. Based on the present and previous works, the optical excitation cross section $\sigma_{\text{exc}}$ of Er ions as a function the excitation wavelength has been obtained. The large values of $\sigma_{\text{exc}}$ with near-band-edge excitation makes GaN:Er attractive for realization of chip-scale photonic devices for optical communications. © 2013 Optical Society of America

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1. Introduction

Rare earth (RE)–doped semiconductors have received considerable attention due to the intra-4f transitions of these trivalent ions that occur in the visible and infrared spectral regions (see [1] and references therein). The outer 5s and 5p electronic shells shield the 4f electrons, and narrow spectral line emissions are allowed in the presence of the crystal field induced by the host materials [2]. The 1.54 $\mu$m emission of erbium ions (Er$^{3+}$), originating through the intra-4f transitions from the first excited manifold ($^4I_{13/2}$) to the ground state ($^4I_{15/2}$), coincides with the minimal-optical-loss band of silica fibers. Different nonsemiconductor materials have been doped with RE atoms for various optoelectronic devices, including silica for erbium-doped fiber amplifiers (EDFAs) [3]. In general, resonant excitation, at 980 or 1480 nm, provides to a small optical excitation cross section of $10^{-20}$ to $10^{-21}$ cm$^{-2}$ in optical fibers, and a long waveguide length is needed for the construction of such optical devices [4,5].

Er-doped semiconductors have been widely studied as alternative materials for optical communications. Among various semiconductor host materials, the III-nitride semiconductors exhibit superior room-temperature performance because their large energy bandgaps lead to a lower thermal quenching of Er emission [6–8]. Previously, we reported the synthesis of Er-doped GaN (GaN:Er) by metal–organic chemical vapor deposition (MOCVD) and the strong 1.54 $\mu$m emission using above-bandgap excitation [9,10]. The optical cross sections are fundamental parameters that determine the optical properties of devices made of GaN:Er materials. In this work, the optical excitation cross section $\sigma_{\text{exc}}$ and the effective decay lifetime $\tau$ of the Er emission at 1.54 $\mu$m in GaN:Er epilayers have been investigated. Photoluminescence (PL) measurements of the integrated emission intensity at 1.54 $\mu$m were made as a function of excitation photon flux. Coupled with time-resolved PL measurements, values of the $\sigma_{\text{exc}}$ of Er ions under near-bandgap excitation were obtained.

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2. Experimental Details

GaN:Er epilayers with a thickness of 500 nm were grown on AlN/Al2O3 templates by MOCVD at 1020°C. Trimethylgallium (TMGa), tris-isopropylcyclopentadienylerbium (TRIP Er), and ammonia (NH₃) were used as the group-III, Er dopant and group-V precursors, respectively, and were carried into the MOCVD reactor by H₂ gas. For the infrared PL measurements, two laser diodes having excitation wavelengths (λexc) of λexc = 375 nm (LDCU 8646) and λexc = 405 nm (Thorlabs CPS-405) were used as excitation sources. A thermoelectric-cooled infrared spectrometer (Bayspec deep-cooled InGaAs NIR spectrometer) was used to disperse the PL emission spectra. To probe the rise and decay kinetics of the 1.54 μm emission line, time-resolved PL measurements were conducted, and a square electrical pulse with 20 ms width and 10 Hz frequency was used to drive the diode lasers. A low-power germanium photodetector, used to measure the time-dependent 1.54 μm emission, was connected to a low-noise current preamplifier and a digital oscilloscope to convert the optical emission intensity into an electrical signal. The overall system response time is around 50 μs.

3. Results and Discussions

Figure 1 shows typical PL spectra near 1.54 μm of GaN:Er epilayers at an excitation power of ~1.6 mW. It is apparent that the 1.54 μm emission is stronger for λexc = 375 nm than for λexc = 405 nm by a factor of about four, which is consistent with our previous results using light-emitting diodes with various λexc as excitation sources [11]. Figure 2(a) shows the 1.54 μm emission spectra, and Fig. 2(b) shows the integrated 1.54 μm emission intensity Iint measured from a GaN:Er epilayer as a function of excitation photon flux ϕexc with λexc = 375 nm. As expected, Iint increases with an increase of ϕexc. The optical excitation process of Er³⁺ atoms can be expressed by

\[ \frac{dN_{Er}^+}{dt} = \sigma_{exc}\phi_{exc}(N_{Er} - N_{Er}^+) - \frac{N_{Er}^+}{\tau}, \]

where NEr⁺ is the excited Er³⁺ concentration and NEr is the total concentration of the optically active Er³⁺ centers [12–15]. As the steady state is reached, Iint can be expressed as

\[ I_{int} \propto \frac{N_{Er}^+}{\tau_{rad} + \frac{1}{\tau_{nonrad}}} = \frac{\sigma_{exc}\phi_{exc}}{\sigma_{exc}\phi_{exc} + 1} \frac{N_{Er}^+}{\tau_{nonrad}}, \]

where τrad refers to the radiative decay lifetime and can be described as 1/τ = 1/τ_rad + 1/τ_nonrad (where τ_nonrad refers to the nonradiative decay lifetime). Equation (2) describes the measured data very well. The solid curve in Fig. 2(b) is the least squares fit of the measured Iint as a function of ϕexc using Eq. (2). The best fit was found with σexcτ = 1.5 × 10⁻¹⁹ cm⁻² s for λexc = 375 nm.

The rise and decay kinetics of the 1.54 μm emission line were probed using time-resolved PL measurements to provide the values of σexc and τ separately for λexc = 375 nm. Figure 3(a) shows time-resolved PL measurement data for λexc = 375 nm for different ϕexc. When the excitation is on (t = 0 – 20 ms), the 1.54 μm emission intensity increases with time.

![Fig. 1. (Color online) Infrared PL spectra near 1.54 μm measured from a GaN:Er epilayer at an excitation power ~1.6 mW for λexc = 375 and 405 nm. The schematic structure diagram of the GaN:Er epilayer grown on an AlN/sapphire template is shown in the inset.](image1)

![Fig. 2. (Color online) (a) Infrared PL emission spectra near 1.54 μm and (b) integrated 1.54 μm emission intensity Iint measured from a GaN:Er epilayer as a function of the excitation photon flux ϕexc for λexc = 375 nm. The solid curve is the least squares fit of data using Eq. (2).](image2)

![Fig. 3. (Color online) (a) Rise and decay kinetics of the 1.54 μm emission line probed using time-resolved PL measurements and (b) reciprocal value of the characteristic rising time τpump as a function of the excitation photon flux ϕexc for λexc = 375 nm measured from a GaN:Er epilayer sample. The solid curves are the least squares fit of data using Eqs. (3)–(5).](image3)
and approaches a saturation level for $t > 10$ ms. A further increase in $\sigma_{\text{exc}}$ only leads to a slightly higher saturation level. The measured $I_{\text{int}}$ as a function of time in the turn-on (excitation) time period can be described as

$$I(t) = I_0 \left(1 - \exp \left(-\frac{t}{\tau_{\text{pump}}} \right) \right).$$

(3)

where $\tau_{\text{pump}}$ and $I_0$ refer, respectively, to the characteristic rising time and to the saturation emission intensity ($t \to \infty$) under different $\sigma_{\text{exc}}$. The solid curves in Fig. 3(a) are the least squares fit of the measured time-dependent intensity data using Eq. (3), from which $\tau_{\text{pump}}$ under various $\sigma_{\text{exc}}$ can thus be obtained. The reciprocal $\tau_{\text{pump}}$ value is plotted as a function of $\sigma_{\text{exc}}$ for $\lambda_{\text{exc}} = 375$ nm in Fig. 3(b). It was observed that $\tau_{\text{pump}}^{-1}$ values increase with an increase of $\sigma_{\text{exc}}$, which implies faster saturation processes for the 1.54 $\mu$m emission under higher $\sigma_{\text{exc}}$. By solving Eq. (1), $\tau_{\text{pump}}^{-1}$ can be expressed as

$$\tau_{\text{pump}}^{-1} = \sigma_{\text{exc}} \sigma_{\text{exc}} \frac{1}{\tau}.$$  

(4)

The solid line in Fig. 3(b) is a least squares fit of data using Eq. (4). The best fit values of $\tau$ and $\sigma_{\text{exc}}$ for $\lambda_{\text{exc}} = 375$ nm are $\tau = 3.2$ ms and $\sigma_{\text{exc}} = 4.6 \times 10^{-17}$ cm$^{-2}$, respectively. The value of $\sigma_{\text{exc}} \tau$ obtained from the PL rising kinetics is thus $\sigma_{\text{exc}} \tau = 1.47 \times 10^{-19}$ cm$^{-2}$ s, which is in excellent agreement with that obtained from the relation between the measured $I_{\text{int}}$ and $\varphi_{\text{exc}}$ shown in Fig. 2(b) using Eq. (2) ($\sigma_{\text{exc}} \tau = 1.5 \times 10^{-19}$ cm$^{-2}$ s). This provides high confidence of our analysis. On the other hand, the PL decay kinetics (during the turn-off period) of 1.54 $\mu$m emission can be described by an exponential decrease with time,

$$I(t) = I_0 \exp(-t/\tau).$$ 

(5)

as illustrated in Fig. 3(a).

For comparison, $\tau$ and $\sigma_{\text{exc}}$ of Er in the same GaN:Er epilayers for $\lambda_{\text{exc}} = 405$ nm were also estimated through PL and time-resolved PL measurements as shown in Fig. 4(a). Figure 4(b) plots the measured $I_{\text{int}}$ of the 1.54 $\mu$m emission as a function of $\varphi_{\text{exc}}$ for $\lambda_{\text{exc}} = 405$ nm. As expected, an increase of $\varphi_{\text{exc}}$ leads to a stronger 1.54 $\mu$m emission. The line with open squares in Fig. 4(b) is the least squares fit of data using Eq. (2). The best fit value of $\sigma_{\text{exc}} \tau$ with $\lambda_{\text{exc}} = 405$ nm is $\sigma_{\text{exc}} \tau = 7.1 \times 10^{-20}$ cm$^{-2}$ s. To obtain the values of $\sigma_{\text{exc}}$ and $\tau$ separately with $\lambda_{\text{exc}} = 405$ nm pumping, we plot the measured $\tau_{\text{pump}}^{-1}$ from time-resolved PL as a function of $\varphi_{\text{exc}}$ for $\lambda_{\text{exc}} = 405$ nm in Fig. 4(b). Similarly, an increase of $\varphi_{\text{exc}}$ also leads to a shorter rising time to reach the saturation level, as described by Eq. (4). The line with solid circles in Fig. 4(b) is the least squares fit of data with Eq. (4). The fitted values of $\tau$ and $\sigma_{\text{exc}}$ are $\tau = 3.8$ ms and $\sigma_{\text{exc}} = 1.7 \times 10^{-17}$ cm$^{-2}$. Consequently, the value of $\sigma_{\text{exc}} \tau$ is $\sigma_{\text{exc}} \tau = 7.1 \times 10^{-20}$ cm$^{-2}$ s for $\lambda_{\text{exc}} = 405$ nm. This smaller value of $\sigma_{\text{exc}}$ for $\lambda_{\text{exc}} = 405$ nm compared with that for $\lambda_{\text{exc}} = 375$ nm explains the results of lower optical pumping efficiency of the 1.54 $\mu$m emission for $\lambda_{\text{exc}} = 405$ nm shown in Fig. 1.

With the above- or near-bandgap excitation, the dominant excitation mechanism of the 1.54 $\mu$m emission in GaN:Er is due to the exchange of energy between the electrons captured by the complexes involving Er and nitrogen vacancies (Er$_{\text{Ga}}$-V$_{\text{N}}$) and the 4$f$ core states of Er [10,15]. Thus, the presence of a higher photoexcited electron density translates to a higher optical cross section. Because the relative optical absorption coefficient and hence the photoexcited carrier density decrease exponentially with excitation photon energy near the band edge [11,15], the excitation cross section $\sigma_{\text{exc}}$ at $\lambda_{\text{exc}} = 375$ nm is expected to be higher than that at $\lambda_{\text{exc}} = 405$ nm, which is what we observed. Based on Eq. (4), a larger excitation cross section naturally leads to a shorter effective decay lifetime at $\lambda_{\text{exc}} = 375$ nm than at $\lambda_{\text{exc}} = 405$ nm.

Based on the present and previous studies [15,16], we have obtained the optical excitation cross section $\sigma_{\text{exc}}$ of Er ions in GaN host as a function of the excitation wavelength $\lambda_{\text{exc}}$, and the results are summarized in Fig. 5. Under a resonant excitation ($\lambda_{\text{exc}} = 980$ or 1480 nm), the 1.54 $\mu$m emission is limited by the small absorption cross section of Er$^{3+}$ atoms [16]. On the other hand, with the above or near-bandgap excitation, Er$^{3+}$ atoms are excited through carriers generated by optical absorption in the GaN host. In this case, $\sigma_{\text{exc}}$ is determined by the absorption coefficient of GaN, which is very large and decreases exponentially with a decrease of the excitation photon energy [15]. This result of exponential decrease of $\sigma_{\text{exc}}$ with an increase of $\lambda_{\text{exc}}$ explains very well our earlier results, which showed an approximately exponential decrease of the 1.54 $\mu$m emission intensity with a decrease of the excitation photon energy in the band-edge region [11].
4. Summary and Conclusions

In summary, the optical excitation cross section $\sigma_{exc}$ of Er in GaN epilayers grown by MOCVD has been studied. The determined value of $\sigma_{exc} = 4.6 \times 10^{-17}$ cm$^{-2}$ s for $\lambda_{exc} = 375$ nm is approximately three orders of magnitude larger than that using a resonant excitation ($\lambda_{exc} = 980$ nm or 1480 nm). While EDFAs are an established technology, they are difficult to integrate with many other functional photonic devices because a long waveguide length is needed for resonant excitation due to the small excitation cross section and limited Er doping concentration. In contrast, Er-doped III-nitride semiconductors with above- or near-band-edge excitation via laser diode pump or current injection could be an attractive alternative approach to achieve chip-scale infrared emitters and optical amplifiers for optical communications [17]. If realized, GaN:Er-based devices would lead to integration with other functional optical devices, such as wavelength routers, optical switches, light sources, and detectors, to build monolithic photonic integrated circuits. The materials growth effort is supported by JTO/ARO (W911NF-12-1-0330), and the optical characterization work is supported by NSF (ECCS-1200168). HXJ and JYL thank the AT&T Foundation for the support of the Ed Whitacre and Linda Whitacre endowed chairs. JMZ acknowledges support from NSF under the IR/D program. Any conclusions in this paper do not necessarily reflect the views of NSF/JTO/ARO.

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