Effect of Nitrogen Doping on the Photoluminescence of Amorphous Silicon Oxycarbide Films

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Received: 26 August 2019; Accepted: 26 September 2019; Published: 27 September 2019

Abstract: The effect of nitrogen doping on the photoluminescence (PL) of amorphous SiC\textsubscript{x}O\textsubscript{y} films was investigated. An increase in the content of nitrogen in the films from 1.07% to 25.6% resulted in red, orange-yellow, white, and blue switching PL. Luminescence decay measurements showed an ultrafast decay dynamic with a lifetime of ~1 ns for all the nitrogen-doped SiC\textsubscript{x}O\textsubscript{y} films. Nitrogen doping could also widen the bandgap of SiC\textsubscript{x}O\textsubscript{y} films. The microstructure and the elemental compositions of the films were studied by obtaining their Raman spectra and their X-ray photoelectron spectroscopy, respectively. The PL characteristics combined with an analysis of the chemical bonds configurations present in the films suggested that the switching PL was attributed to the change in defect luminescent centers resulting from the chemical bond reconstruction as a function of nitrogen doping. Nitrogen doping provides an alternative route for designing and fabricating tunable and efficient SiC\textsubscript{x}O\textsubscript{y}-based luminescent films for the development of Si-based optoelectronic devices.

Keywords: photoluminescence; amorphous silicon oxycarbide; nitrogen doping; defect; plasma enhanced chemical vapor deposition

1. Introduction

Efficient silicon (Si)-based luminescent materials are indispensable components to realize a cheap and complementary metal oxide semiconductor (CMOS) optical integration. Thus far, different systems of Si-based luminescent materials, such as SiO\textsubscript{x}, SiN\textsubscript{x}, SiC\textsubscript{x}, and SiN\textsubscript{x}O\textsubscript{y}, have been developed, and efforts have been devoted to understanding and ameliorating the light emission of Si-based materials [1–9]. Silicon oxycarbide (SiC\textsubscript{x}O\textsubscript{y}) has been widely explored because of its strong light emission and high solid solubility for rare earths [10–13]. SiC\textsubscript{x}O\textsubscript{y} also features a tunable band gap. As such, it is beneficial to obtaining strong white electroluminescence at a low driving voltage in SiC\textsubscript{x}O\textsubscript{y}-based light-emitting diodes [14]. In the recent reference, Gallis et al. systematically studied the white photoluminescence (PL) dynamics from SiC\textsubscript{x}O\textsubscript{y} film, where the band tail states related to the Si–O–C and/or the Si–C bonds were suggested as the sources of the luminescence [11]. Recently, optical gain was demonstrated in a-SiC\textsubscript{x}O\textsubscript{y} under ultraviolet excitation, which was attributed to the formation of a three-level luminescence model with the intermediate level related to Si dangling bond (DB) defects radiative state [15]. Furthermore, an increase in C content in SiC\textsubscript{x}O\textsubscript{y} films can cause a strong light emission ranging from near-infrared to orange regions [15]. Although performance is enhanced in SiC\textsubscript{x}O\textsubscript{y} films, progress remains slow. The main obstacle lies in the fact that the light emission efficiency generally remains too low to allow the fabrication of efficient light-emitter devices. To date, studies on the effect of doping on the optical properties of SiC\textsubscript{x}O\textsubscript{y} films have mainly focused...
on rare earth (RE) doping, such as Er and Eu doping [16–18]. However, up to now, the effect of other elements on the optical properties of SiC$_x$O$_y$ films is still unclear.

In this letter, the effect of nitrogen doping on the PL of amorphous SiC$_x$O$_y$ film was investigated. Interestingly, an increase in nitrogen content in the films induced strong red, orange-yellow, white, and blue switching PL. Combining the PL results with the analysis of the microstructure and the chemical bonding configurations within the films, it suggests that the rearrangement of chemical bonds with varying nitrogen plays an important role in the evolution of PL characteristics in the films.

2. Materials and Methods

Nitrogen-doped SiC$_x$O$_y$ films with the thickness of 550 nm were grown at 250 °C on Si substrates and quartz by radio frequency (RF) glow-discharge decomposition of SiH$_4$, CH$_4$, O$_2$, and NH$_3$ mixtures in the very high frequency plasma enhanced chemical vapor deposition (VHF-PECVD) system. The flow rates of SiH$_4$, CH$_4$, and O$_2$ were kept at 3.5, 5, and 1.2 sccm, respectively, whereas the flow rate of NH$_3$ varied from 0.5 sccm to 5 sccm to control the N content in the films. The films were named S$_x$ (x = 1, 2, 3, 4) for the NH$_3$ flow rates at 0.5, 1, 3, and 5 sccm, respectively. The RF power and the deposition pressure for the growth were maintained at 30 W and 20 Pa, respectively. The optical band gaps of the films were obtained in accordance with the Tauc plot Equation (1):

$$\alpha(h\nu)^{1/2} = A(\frac{h\nu}{E_{opt}})^{1/2}$$

where $\alpha$ is the absorption coefficient, $A$ is a coefficient quantifying the slope of the absorption edge, and $h\nu$ is the photon energy [19]. The calculated $E_{opt}$ increases linearly from 2.83 eV to 3.66 eV as the NH$_3$ flow rate increases from 0.5 sccm to 5 sccm. This finding demonstrates that N doping can widen the bandgap of SiC$_x$O$_y$ films, which may result from the substitution of stronger Si–N bonds for weak Si–Si bonds or Si–C bonds. The comparison of PL with $E_{opt}$ results indicated that the value of PL peak energy of all the films was obviously smaller than the corresponding $E_{opt}$, suggesting that the origin of PL was not from the band-to-band recombination.

The microstructure of the SiC$_x$O$_y$:N films was examined using Raman spectra (Figure 2a) to further understand the origin of PL characteristics. All the SiC$_x$O$_y$:N films exhibited similar line shape characteristics typical of amorphous silicon-based materials. A broad Raman band, which was ascribed to the transverse optical (TO) vibration mode of amorphous silicon, peaked at ~470 cm$^{-1}$ for all the SiC$_x$O$_y$:N films. These results showed that all the SiC$_x$O$_y$:N films had a uniform amorphous structure without the presence of Si nanocrystals [20]. Furthermore, there was no obvious change in the surface morphology of the films prepared at different NH$_3$ flow rates, as was revealed by atomic force microscopy (Figure 2b).
The FTIR spectra of the SiCO:N films were obtained to study the local bonding changes in the films grown at different NH$_3$ flow rates (Figure 4). In the S1 film, the vibration modes related to Si–N, C–Si–O, Si–H, and C–H bonds could be clearly observed. A small band shoulder at 1265 cm$^{-1}$ was assigned to the Si–CH$_3$ stretching vibration [25].

The films were measured by time-resolved PL to obtain further insights into the PL mechanism of the SiCO:N films (Figure 3). The decay curves could be fitted with a double exponential function:

$$I(t) = A_1 \exp\left(-\frac{t}{\tau_1}\right) + A_2 \exp\left(-\frac{t}{\tau_2}\right)$$

(2)

where $A_i$ and $\tau_i$ (i = 1, 2) are the normalized amplitudes of the components and the time constants, respectively [21]. The obtained average lifetime of the SiCO:N films was about 1 ns. The luminescent dynamic behavior was similar to that observed in defect-related luminescent Si-based materials, such as SiN$_x$O$_y$ and SiC$_x$O$_y$ [19,21]. Furthermore, it was also found that the luminescence decay lifetimes in our case were shorter than those in the band-tail recombination model where a broader band-tail brought a longer lifetime, as the photogenerated carriers could be thermalized into deeper localized states [22]. Therefore, the results suggested that the light emission of the SiCO:N films originated from the defect luminescent centers in the films.

The FTIR spectra of the SiCO:N films were obtained to study the local bonding changes in the films grown at different NH$_3$ flow rates (Figure 4). In the S1 film, the vibration modes related to Si–C, Si–N, C–Si–O, Si–H, and C–H bonds could be clearly observed. The bands centered at 860 and 1039 cm$^{-1}$ could be ascribed to Si–N and C–Si–O stretching modes, respectively [23,24]. Additionally, a band at 1265 cm$^{-1}$ was assigned to the Si–CH$_3$ stretching vibration [25]. A small band shoulder at
800 cm$^{-1}$ was observed and was assigned to the Si–C stretching vibration [24]. A distinct absorption peak at 2170 cm$^{-1}$ and a weak band located at 2965 cm$^{-1}$ were attributed to the Si–H and the C–H stretching vibrations, respectively [26]. A weak band around 3375 cm$^{-1}$ was associated with the N–H stretching mode [23]. The most important feature for the FTIR spectra was the strong dependence of major bands on NH$_3$ flow rates. As the NH$_3$ flow rates increased, the intensity of C–Si–O bonds gradually decreased, and the peak gradually became red shifted. As the NH$_3$ flow rate increased to 3 sccm, this band broadened and red shifted to ~1010 cm$^{-1}$ with a shoulder at ~940 cm$^{-1}$, which was assigned to the N–Si–O vibration [27]. As the NH$_3$ flow rate further increased to 5 sccm, the band of the N–Si–O vibration became dominant, indicating that the silicon oxycarbide-dominant phase of the film transformed into silicon oxynitride. Apparently, the increase in the NH$_3$ flow rate resulted in chemical bond reconstruction in the films. Based on the FTIR spectra, the evolution of PL characteristics could be suggested from the chemical bond reconstruction in SiCO:N films.

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**Figure 3.** Room temperature time resolved photoluminescence for the SiCO:N films with various NH$_3$ flow rates.

**Figure 4.** Fourier transform infrared (FTIR) spectra of the SiCO:N films grown at different NH$_3$ flow rates.
The composition of the SiCO:N films was examined through X-ray photoelectron spectroscopy (XPS) (Figure 5). The atomic percentages of Si, C, O, and N in the SiCO:N film fabricated at an NH3 flow rate of 0.5 sccm were 51.89%, 19.82%, 27.22%, and 1.07%, respectively. This finding indicated that the Si-rich silicon oxycarbide phase was dominant in the S1 film. The change in the XPS spectra was the gradual decrease in Si and C concentrations with the increase in N concentration and NH3 flow rates (Figure 5). As the NH3 flow rate increased to 5 sccm, the N concentration rapidly increased to 25.6%, whereas the Si and the C concentrations decreased to 40.8% and 8.22%, respectively. This finding was consistent with the observed results in the FTIR spectra shown in Figure 4, that is, the N–Si–O vibration band became dominant, while the C–Si–O vibration band significantly weakened as the NH3 flow rate increased to 5 sccm. This result indicated that the dominant phase in the films changed from silicon oxycarbide to silicon oxynitride when the NH3 flow rate increased to 5 sccm.

![Figure 5. The atom concentration of Si, C, O, and N of the SiCO:N films against the NH3 flow rates.](image)

The PL decay analysis (Figure 3) revealed that the luminescent dynamic behavior in the nitrogen doped SiCxOy films featured a defect-related luminescent characteristic, as observed in SiNxOy and SiCxOy films. Previous studies clarified that C-related nonbridging oxygen hole centers (NBOHC) are the principal radiative recombination centers in silicon oxycarbide, and they are responsible for light emission ranging from the green region to the red region [28]. In our case, the PL intensity in the film decreased as the NH3 flow rate increased. This change was similar to that of the intensity of the C–Si–O bonds (Figure 4). Therefore, the observed tunable light emissions from green to red may have originated from recombination through C-related NBOHC defects in SiCxOy films. The PL spectra of the S3 film could be deconvoluted into a strong green band and a weak blue band. As the NH3 flow rate increased to 5 sccm, the intensity of the green PL band decreased dramatically, whereas the blue PL band of the film S4 became dominant. This behavior could be attributed to the change in the dominant phase of the films from silicon oxycarbide to silicon oxynitride as a result of the increase in NH3 flow rate to 5 sccm (Figure 4). In the case of amorphous SiNxOy films, the blue PL could be ascribed to the radiative recombination between N–Si–O defect states and the valence band tail states [27]. Thus, the blue PL from S3 and S4 was likely from N–Si–O defect luminescent centers.

4. Conclusions

In summary, we report the effect of nitrogen doping on the PL of amorphous SiCxOy films. Nitrogen doping can induce strong red, orange-yellow, white, and blue switching PL with a recombination lifetime in nanoseconds. This process can also widen the band gap of SiCxOy films. The PL results and the FTIR analyses reveal that the switching characteristics in PL originate from the variation in defect
luminescent centers resulting from the chemical bond re-construction as a function of nitrogen doping. Apparently, nitrogen doping provides an alternative route for designing and fabricating tunable and efficient SiC\(_x\)O\(_y\)-based luminescent films for the development of Si-based optoelectronic devices.

**Author Contributions:** Data curation, J.S. and Z.L. (Zhenxu Lin); Investigation, Y.Z., Z.L. (Zewen Lin), and H.L.; Methodology, C.S., and Y.G.; All authors analyzed the experimental data; Writing–review & editing, J.S., Z.L. (Zhenxu Lin), R.H. and W.Z.; All authors participated in discussions and knew implications of the work.

**Funding:** This work was supported by National Natural Science Foundation of China (Nos. 61274140), Natural Science Foundation of Guangdong Province (2015A030313871), Young Talents in Higher Education of Guangdong (2017KQNCX129), the Distinguished Young Teacher Training Program in Higher Education of Guangdong (YQ2015112), Science and Technology Planning Project of Guangdong Province (2017B090921002) and Science and Technology Planning Project of Chaozhou (2018SS24).

**Conflicts of Interest:** The authors declare no conflict of interest.

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