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Depth-resolved cathodoluminescence spectroscopy of silicon supersaturated with sulfur

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We investigate the luminescence of Si supersaturated with S (Si:S) using depth-resolved cathodoluminescence spectroscopy and secondary ion mass spectroscopy as the S concentration is varied over 2 orders of magnitude (10^{18}–10^{20} cm^{-3}). In single-crystalline supersaturated Si:S, we identify strong luminescence from intra-gap states related to Si self-interstitials and a S-related luminescence at 0.85 eV, both of which show a strong dependence on S concentration in the supersaturated regime. Sufficiently high S concentrations in Si (>10^{20} cm^{-3}) result in complete luminescence quenching, which we propose is a consequence of the overlapping of the defect band and conduction band. © 2013 American Institute of Physics. [http://dx.doi.org/10.1063/1.4788743]
Electron Microscope. The CL system is equipped with a Ge photodiode to detect near infrared emission, sensitive from 600 to 1700 nm (2.0 eV–0.7 eV). All the CL experiments were performed at liquid nitrogen temperature (77 K) with a dwell time per pixel of 67 ns and an acquisition time of 3 ms per wavelength step (1 nm). In the depth-dependent CL analyses, we used accelerating voltages of both 5 and 10 kV, keeping the total power density constant at $2 \times 10^{6}$ W/cm$^2$. The total power density was calculated as the product of the accelerating voltage and the beam current, divided by the action volume of depth-resolved CL at 5 kV and 10 kV is overlaid (right axis), showing the distribution of electron scattering events through which the electron beam loses 60% of its starting energy in the Monte Carlo simulations.

Investigating CL at both 5 kV and 10 kV allows for the probing of distinct regions of the Si:S supersaturated layer. Figure 1 shows the SIMS doping profile that forms following ion implantation of Si with S across a range of S doses ($3 \times 10^{14} \text{ cm}^{-2}$–$1 \times 10^{16} \text{ cm}^{-2}$). Both $1 \times 10^{16} \text{ cm}^{-2}$ and $3 \times 10^{15} \text{ cm}^{-2}$ samples exhibit an approximately 200 nm thick plateau at the surface, indicating that a relatively uniform concentration of S is formed at the surface. Beneath this layer, a tail region is observed in which the S concentration gradually drops off as it approaches the melt depth approximately 400 nm below the surface. Comparison of the distribution of electron scattering events during CL with the S profile (Figure 1) illustrates the ability to probe primarily the high-concentration plateau and the dopant profile tail separately by switching between 5 kV and 10 kV accelerating voltages, respectively.

An important consideration when conducting depth-resolved CL of Si:S is the influence of the changing S concentration on self-absorption, as the sub-band gap absorbance is strongly affected by the degree of S supersaturation in Si.$^{16}$ The upper-limit of these effects can be calculated using the reported concentration-dependent values of $\alpha d$ of Si:S$^{22}$ where $\alpha$ is the absorption coefficient and $d$ is the effective thickness. We use the Beer-Lambert law to estimate the changes in self-absorption when increasing from S doses of $3 \times 10^{14} \text{ cm}^{-2}$ to $1 \times 10^{16} \text{ cm}^{-2}$ and find that the maximum expected decrease in luminescent intensity due to the increased S concentration is 34% in this doping regime. The effects of self-absorption may complicate the quantification of the observed trends in luminescence, but we note that in the following investigations, all trends of modulated luminescence with changes in the doping dose (which could include self-absorption effects) significantly exceed 34%. Therefore, qualitative discussions of the observed trends in luminescence are possible in our case.

We first isolate luminescence due to PLM-induced point defects by investigating the CL of Si samples implanted with $1 \times 10^{16} \text{ cm}^{-2}$ Si compared with that of an untreated Si wafer. At 5 kV (Figure 2(a)) and 10 kV (Figure 2(b)), the pristine Si wafer shows only a peak centered at 1.1 eV corresponding to the band-to-band transition in Si (BB-line).$^{23,24}$ At both 5 kV and 10 kV, the $1 \times 10^{16} \text{ cm}^{-2}$ Si-dose sample exhibits decreased intensity of the BB-line, and a broad emission at 0.99 eV simultaneously appears. The decrease in intensity of the BB-line is attributed to increased nonradiative recombination due to PLM-induced point defects. The luminescence at 0.99 eV is identical to the band observed by Pankove and Wu$^{25}$ following ion implantation of Si with Si and annealing at 300°C in H2. This luminescent feature is attributed to phonon modes of the X-center (also known as the I$_3$ band)$^{26}$ that has been proposed to arise from a Si self-interstitial complex.$^{27}$ Increasing the accelerating voltage from 5 kV to 10 kV results in a higher relative intensity of the BB-line compared to the 0.99 eV band, which can be attributed to the increased volume of unaffected Si that is probed using 10 kV accelerating voltage (Figure 1).

Next, we elucidate the effects of S-supersaturation by comparing Si implanted with Si to Si implanted with S at an identical dose ($1 \times 10^{16} \text{ cm}^{-2}$). The CL spectra of the $1 \times 10^{16} \text{ cm}^{-2}$ S-supersaturated sample taken at 5 kV (Figure 2(a)), which probing the most highly doped region (Figure 1), exhibit significantly reduced overall intensity without clearly resolvable luminescent bands. Increasing the accelerating voltage to 10 kV, thus probing the tail of the doped profile, results in the appearance of the BB-line at 1.1 eV and a CL band centered around 0.85 eV. The luminescence at 0.85 eV does not appear in either the 5 kV or 10 kV spectra from the Si-implanted sample and is therefore related to the presence of S. Here, we note that the S-related luminescence at 0.85 eV observed in our supersaturated samples is different from the metastable luminescence centers at 0.82 eV (1.5 μm) and 0.94 eV (1.32 μm) that have been observed in Si containing equilibrium concentrations of S (concentrations 1-3 orders of magnitude lower than investigated here)$^{24,26}$.

To better understand the effect of S-supersaturation on the implantation-related band at 0.99 eV and the S-related band around 0.85 eV, the implanted S dose was decreased from $1 \times 10^{16} \text{ cm}^{-2}$ to as low as $3 \times 10^{14} \text{ cm}^{-2}$ and the
resulting luminescence was measured at 5 kV (Figure 3(a)) and 10 kV (Figure 3(b)). When the S dose is decreased to $3 \times 10^{15}$ cm$^{-2}$, the band at 0.99 eV appears, which has been attributed to Si self-interstitials. Recall this band was observed in the $1 \times 10^{16}$ cm$^{-2}$ Si-implanted sample but absent in the S-implanted sample of the same dose. Furthermore, we observe a strong increase in the intensity of the 0.99 eV band with decreasing S dose. Lower doses of S than used in this study ($1 \times 10^{14}$ cm$^{-2}$), followed by PLM, have been used to fabricate light emitting diodes based on zero-phonon luminescence from the W-line (1.018 eV), which is also attributed to Si self-interstitials caused by the implantation process, but the dependence of this luminescence on S concentration was not investigated. It has been previously reported that the intensity of the 0.99 eV band increases upon annealing of ion-implanted Si in H$_2$ ambient and this behavior was attributed to the ability of H to passivate competing nonradiative recombination pathways. S impurities have been shown to provide both radiative and non-radiative recombination pathways, and thus we attribute the decreasing 0.99 eV luminescence with increasing S concentration to the associated non-radiative recombination pathways.

The general trends in the luminescence spectra of Si:S samples at 10 kV (BB-line, 0.99 eV band, and 0.85 eV band) are similar to those collected at 5 kV. The band centered at 0.85 eV appears in the $3 \times 10^{15}$ cm$^{-2}$ S-dose sample at 5 kV, increases in intensity as the dose is decreased further to $1 \times 10^{15}$ cm$^{-2}$, and remains constant as the S dose is further decreased to $3 \times 10^{14}$ cm$^{-2}$, which indicates that this luminescence band is relatively insensitive to S concentration in this regime. This is in stark contrast to the concentration dependence of the 0.99 eV band, which monotonically decreases with increasing S concentration.

Using ion implantation and PLM to synthesize Si:S, we have identified a S-related luminescence at 0.85 eV. There are two known S states with energy levels that align well with the 0.85 eV band: a neutral S substitutional atom or a charged S dimer, with energies 320 meV and 371 meV below the conduction band, respectively. However, S-related luminescence at these energies has not been previously reported. One possibility is that increased impurity-impurity interactions at supersaturated concentrations lead to changes in the luminescent behavior of S impurities, but we would expect such a phenomenon to show strong concentration dependence. Alternatively, it is possible that the band at 0.85 eV is related to a metastable S complex arising from the PLM process. The melting and ultrafast resolidification during PLM represents a unique processing regime that, as evidenced by the supersaturated concentrations of S achieved, produces metastable impurity distributions. Furthermore, metastable luminescent S-complexes have been previously observed, though at different energy levels: S complexes observed at dilute S-concentrations at 0.82 eV and 0.94 eV form only after heating with a blow torch and quenching in ethylene glycol. Further investigations will be necessary to understand the source of the S-related luminescence at 0.85 eV.

Finally, the concentration dependence of the luminescence from supersaturated Si:S provides evidence of the S-induced changes in the band structure as concentrations increase. This is in stark contrast to the concentration dependence of the 0.99 eV band, which monotonically decreases with increasing S concentration.

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approach 1 at.%. In the 5 kV spectra (Figure 3(a)), each luminescent band (BB-line, 0.99 eV, 0.85 eV) shows a unique concentration dependence as the dose is increased from $3 \times 10^{14}$ cm$^{-2}$ to $3 \times 10^{15}$ cm$^{-2}$, but are suppressed when the dose is further increased to $1 \times 10^{16}$ cm$^{-2}$. The lack of luminescence from the $1 \times 10^{16}$ cm$^{-2}$ S-implanted sample at 5 kV suggests that a S concentration around $3.5 \times 10^{19}$ cm$^{-3}$ (0.7 at.%) quenches luminescence from Si:S (Figure 1). Furthermore, the onset of quenching is visible in the $3 \times 10^{15}$ cm$^{-2}$ spectra at 5 kV where the intensity of the 0.85 eV and the BB-line is significantly less than the level they maintained from $3 \times 10^{14}$ cm$^{-2}$ to $3 \times 10^{15}$ cm$^{-2}$. Within the resolution of this characterization, we can say that the onset of quenching occurs around or just above the peak concentrations of S in the $3 \times 10^{15}$ cm$^{-2}$ sample ($1 \times 10^{20}$ S cm$^{-3}$ or 0.2 at.%).

Previous investigations into the optoelectronic properties of supersaturated Si:S indicate the emergence of an impurity band from impurity levels in the gap with increasing chalcogen concentration, leading to drastic changes to the band structure at a concentration threshold similar to where we observe luminescence quenching ($\geq 1 \times 10^{20}$ cm$^{-3}$). Combined computational and experimental investigations of the effect of increasing chalcogen concentration on the band structure suggests an insulator-metal transition due to the overlapping of the defect and conduction bands at S concentrations above $1 \times 10^{20}$ cm$^{-3}$, and this model accurately predicts the observed increased sub-band gap absorptance. The concentration thresholds at which the chalcogen-induced defect band is predicted to overlap with the conduction band ($1 \times 10^{20}$ cm$^{-3}$) and at which a S-induced insulator-metal transition has been observed experimentally agree well with the onset of quenching that we observe using depth-resolved CL spectroscopy ($1 \times 10^{20}$ cm$^{-3}$). Thus, our investigation of the concentration-dependence of luminescence agrees with the band-overlap model, as the impurity and conduction band overlapping could feasibly result in luminescent quenching near the band edge. Finally, although the sub-band gap absorptance and impurity-mediated conduction increases with increasing S concentration, we have shown that S-induced changes in the band structure at concentrations above $1 \times 10^{20}$ cm$^{-3}$ are not conducive to enhancing sub-band gap luminescence. These investigations highlight, however, the orders of magnitude of S-supersaturation ($10^{18}$–$10^{19}$ cm$^{-3}$) that could potentially allow for simultaneous sub-band gap absorptance, increased gain and extended response in photodiodes, and luminescence.

In summary, we have combined depth-resolved CL and SIMS to elucidate the luminescent behavior of Si:S and the effect of varying the concentration within the supersaturated regime. We find that band-band luminescence is reduced by the presence of S and further decreases with increasing S concentration. Luminescence at 0.99 eV, attributed to Si self-interstitial clusters, shows a pronounced increase in intensity with decreasing S concentration which we attribute to a reduction in competing non-radiative recombination pathways. We report S-related luminescence around 0.85 eV that corresponds well with the energy levels of a substitutional S atom or a charged S dimer in Si, though further studies are needed to elucidate its exact nature and its appearance only at supersaturated concentrations of S. Above $1 \times 10^{20}$ S cm$^{-3}$, we observe quenching of all luminescence, the onset of which correlates well with predicted overlapping of the defect band and conduction band. This work demonstrates the efficacy of depth-resolved CL for probing the depth dependence of luminescence in alloys fabricated using ion implantation and pulsed laser melting and reveals the concentration-dependent luminescent properties of S-supersaturated Si.

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