Temperature dependence of time-resolved photoluminescence in closely packed alignment of Si nanodisks with SiC barriers

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Abstract
We study the temperature dependence of time-resolved photoluminescence (PL) in closely packed alignment of Si nanodisks (NDs) with SiC barriers, fabricated by neutral beam etching using bio-nano-templates. The PL time profile indicates three decaying components with different decay times. The PL intensities in the two slower decaying components depend strongly on temperature. These temperature dependences of the PL intensity can be quantitatively explained by a three-level model with thermal activation energies of 410 and 490 meV, depending on the PL components. The activation energies explain PL quenching due to thermal escape of electrons from individual NDs. This thermal escape affects the PL decay times above 250 K. Dark states of photo-excited carriers originating from the separate localization of electron and hole into different NDs are elucidated with the localization energies of 70 and 90 meV. In contrast, the dynamics of the fastest PL decaying component is dominated by electron tunneling among NDs, where the PL intensity and decay time are constant for temperature.

Keywords: Time-resolved photoluminescence, Si nanodisks, Neutral beam etching, Bio-nano-templates

Background
Attractive interdisciplinary research areas between electronic and photonic materials have been developed by modern semiconductor nanotechnology. Si nanostructures are particularly important because solar cells using Si have widely been investigated [1,2], and optical interconnections among integrated Si circuits have also been proposed by developing Si-based photodiodes and optical modulators [3,4]. Therefore, many types of Si nanostructures, such as nanocrystals (NCs), nanodots, and porous nanostructures, were reported by employing various fabrication processes [5-14]. Moreover, fabrication processes of the Si nanostructures using ‘top-down’ lithography techniques were strongly motivated for the purpose of applying the Si nanostructures to electronic and photonic devices.

We have recently proposed a fabrication process of Si nanodisk (ND) arrays, where the Si NDs are formed by damage-free neutral beam (NB) etching for Si thin films covered with etching masks of Fe nanoparticles which are regularly aligned by bio-protein engineering [15-20]. This fabrication process using the bio-templates enables us to prepare closely packed high-density Si NDs with the intentionally designed precise size and spacing in a nanometric scale with flexible film stacking. We have also observed intense photoluminescence (PL) emissions in a visible light region with fast decay times ranging from 10 ps to 2 ns [20]. The fast decaying PL characteristics reflect the dynamics of photo-excited carriers in this high-density Si ND array system, in which wavefunctions of photo-excited carriers overlap among Si NDs to some extent, and the carriers can transfer among the NDs [20]. Photo-generated or electrically injected carriers need to be effectively transferred among Si NDs for the optical applications to solar cells or light-emitting diodes. The spatial transfer of the carriers in nanostructures can also be affected by thermal effects, such as thermal hopping or escape. Therefore, in this...
paper, we investigate the detailed temperature dependence of time-resolved PL and the related carrier dynamics in these high-density Si ND arrays. Different types of PL quenching mechanism can be identified, and the activation energies for the PL thermal quenching are deduced from the temperature dependences of the PL intensity. The difference of the activation energy between different decaying PL components supports the existence of different wavefunctions of the carriers in this high-density ND system. The PL quenching phenomena elucidated in this study will give us useful information about the dynamics of photo-excited carriers, such as carrier separation and transport, when we apply these Si NDs to solar cells and high-speed photonic devices.

Methods
The high-density \( (7 \times 10^{11} \text{ cm}^{-2}) \) Si ND arrays were fabricated from polycrystalline Si thin films deposited on thermally oxidized surfaces of Si substrates under ultra-high vacuum. Bio-nano-templates consisting of ferritin supramolecules containing Fe cores were used to prepare two-dimensional closely packed alignments of the Fe cores as etching masks on the surfaces of Si thin films. The size and interspacing of the Fe cores were intentionally designed by protein engineering for the ferritin supramolecules. The Si NDs were fabricated by forming SiO\(_2\) barriers around the Si NDs masked by the Fe cores using the NB etching and subsequent oxidation processes. Details of the fabrication process are described elsewhere [15-17]. The diameter, thickness, and interspacing distance of the Si NDs mainly used in this study were designed at 10, 4, and 2 nm, respectively, by the abovementioned ferritin-protein engineering. The capping and barrier layers of SiO\(_2\) were removed with NF\(_3\) treatment. Then, a 5-nm-thick SiC layer was finally deposited on the Si ND array under a high vacuum by sputtering.

The samples of the Si ND array were placed on a cold finger cooled by a closed He compressor in a vacuum cryostat with quartz windows. The time-resolved PL spectra were observed at various temperatures by combining the excitation of second harmonic femtosecond pulses with the wavelength of 400 nm, pulse width of 150 fs, and repetition rate of 76 MHz of a mode-locked Ti-sapphire laser, with the detection of a synchroscan streak camera (Hamamatsu Photonics, Hamamatsu, Japan). A spot diameter of the laser light focused on the sample surface was 100 \( \mu \text{m} \). The excitation power density was 8.4 mJ cm\(^{-2}\). The number of electron–hole pair generated per one ND was calculated to be less than 1, taking the sheet density of ND into account. Therefore, the multiple exciton generation or Auger process were not induced. The time width of the instrumental response curve was less than 15 ps, and the time resolution of 5 ps was obtained after deconvolution with the instrumental response.

Results and discussion
Time-integrated PL spectra of the Si ND array at various temperatures are shown in Figure 1a. PL emission bands with the wavelengths of 655 nm \((1.89 \text{ eV}, E_1 \text{ band})\) and 564 nm \((2.22 \text{ nm}, E_2 \text{ band})\) are visible for the whole temperature range. The observed PL cannot be attributed to the indirect bandgap emission affected by a quantum confinement effect, which was often reported in small Si NCs with diameters of 2 to 5 nm. These confined emission energies increased up to 1.7 eV, as reported in Si NCs with the diameter of 2.5 nm [6]. The optical bandgap energy of our Si ND system with the thickness of 4 nm and diameter of 10 nm has been calculated to be ca. 1.5 eV from the one-band Schrodinger equations with classic envelope function theory [19]. However, in our case, the PL peak energy is markedly higher than these energies. Moreover, as described later, decay times of the observed PL are ranging from 10 ps to 2.0 ns, which are much shorter than those in the microsecond-scale characteristic for the indirect bandgap recombination of carriers or defect-related emissions. There are several reports for surface-related emissions in the visible light region, which have been confirmed by PL measurements of samples with different surface treatments [10]. The spectral widths of the PL bands are less than 200 meV. The spectral

![Figure 1](http://www.nanoscalereslett.com/content/8/1/223)
lineweights of single Si nanocrystals were reported to be 100 meV or more [5,21], which were also dependent on the fabrication method and surface conditions. In our case, the size of the Si ND was precisely controlled by the diameter of the Fe core formed in a cavity of the ferritin molecule. The size uniformity of 8% was confirmed from the statistical analysis of SEM images [17]. Therefore, an effect of inhomogeneous broadening due to the size distribution on the PL spectral shape is estimated not to be significant. This estimation is supported by a fact that no remarkable spectral diffusion, which is a time-dependent redshift of the PL spectral energy, was observed for both PL bands in the time-resolved PL spectra. Time-dependent redshifts due to thermal hopping of carriers or energy transfer were frequently observed in systems of high-density quantum dots with significant size distributions.

Temperature dependences of the spectral shape and energy were not seen. Both PL bands exhibit similar temperature dependences of the intensity. The PL intensity of the $E_2$ band is much weaker than that with the SiO$_2$ barrier, which was previously reported [22]. Therefore, we consider that this $E_2$ band originates from oxygen-related surface or interface states of the Si NDs, and we would like to discuss mainly about the $E_1$ emission. In the low-temperature regime below 150 K, the PL intensity is almost constant. The intensity increases toward 200 K and peaks at a maximum around 250 K. The PL intensity then decreases monotonically with further increasing temperature. Quantitative discussion about these temperature dependences of the PL intensity will be made later.

The transient PL for the $E_1$ band emission as a function of temperature in the Si ND array is shown in Figure 1b. The temporal evolution of each PL profile cannot be expressed by a single exponential function. The best fit was obtained typically using a triple exponential function as shown in Figure 1c, which is common for all array samples of the high-density Si NDs. From this fitting, we have identified three PL decaying components with different time constants $\tau_1 = 770$ ps, $\tau_2 = 110$ ps, and $\tau_3 = 15$ ps, respectively, for this case at 250 K as an example. Several papers have demonstrated ultrafast PL in a sub-picosecond region for Si NCs by means of up-conversion PL. The ultrafast emission ranging 2.0 to 2.4 eV was observed, which was attributed to the pseudodirect gap emission from the core states of Si NCs [11,12]. In contrast, the PL components observed in our samples show time constants ranging from 10 ps to 1 ns, where values are much higher than those of the above pseudodirect gap emissions. Therefore, the most probable origin of the $E_1$ emission is emissive surface states weakly located at the interfaces of Si NDs. Dhara and Giri have reported the PL emission with the wavelength of about 600 nm with decay times of several nanoseconds [13]. They assigned this PL to the quasi-direct bandgap emission in heavily strained Si NCs because of their unique preparation of the NCs by milling. Sa’ar reviewed recent developments in the PL studies of various Si nanostructures and suggested that neither quantum confinement model nor surface chemistry model can solely explain the entire spectrum of emission properties [14].

The three PL components with different decay times imply three different types of emissive sites in the present ND array. We assigned these three decaying components from the disk density and excitation power dependences of the PL decay time and intensity [20]. The emission with the slowest decay time $\tau_1$ on the order of 1 ns was interpreted by electron–hole pairs or excitons localized at individual NDs, because this PL component was dominant in the case of low-density dispersive NDs with the disk interspacings larger than 40 nm. The emission with the decay time $\tau_2$ was understood by recombination of an electron–hole pair or exciton not strongly localized in each ND, where each wavefunction of the carrier spreads over neighboring NDs to some extent due to periodic regular alignment of the ND separated by ultrathin potential barriers. The fastest PL component with $\tau_3$ was attributed to the recombination which was strongly affected by the electron tunneling among the NDs. In other words, this fastest PL was quenched by the electron transfer. The latter two faster PL components appeared only at high excitation densities in the high-density ND arrays. The interpretation of the electron transfer was also supported by the conductance measurement for sister samples [23].

The time-integrated PL intensities of the three decaying components were deduced by fitting the PL decay curves with the triple exponential function. The PL intensities are plotted as a function of temperature in Figure 2. As can be seen, time-integrated intensities of the two slower decaying components ($I_1$ and $I_2$, corresponding to the PL components with the decay times $\tau_1$ and $\tau_2$) depend strongly on temperature, while the fastest decaying component ($I_3$ with $\tau_3$) is almost constant for temperature.

We analyzed these temperature dependences of PL intensities of the $I_1$ and $I_2$ components by a thermal quenching model taking an existence of ‘middle state’ into account [24]. In our calculation, we assumed that the time-integrated intensity of the observed PL was equivalent to that measured by the steady-state excitation because the PL decay times in the present Si ND system are below 2 ns. In this model, we considered three levels schematically shown in Figure 2b. The emissive excitonic level denoted by $E_x$ is assumed to exist between the barrier level for thermal escape of photo-excited carriers from individual NDs and the lower-energy level $E_0$. This $E_0$ level is possibly due to localization at trap states formed by spatial displacements of wavefunctions of an electron and hole in
the ND system. The electronic states in the Si NDs can largely be affected by the interfacial bonding states of Si atoms. Therefore, radiative interfacial states \(E_s\) and deeper trap levels \(E_0\) can be formed. The PL intensity from this middle state is basically proportional to the number of electron–hole pair or exciton at this level and thus dependent on a thermal escape rate beyond the barrier as well as on a thermal excitation rate from the lowest trap level. In this case, the PL intensity can be described as follows:

\[
I(T) = I(0) \left(1 + \frac{D \exp(-E_{\text{low}}/k_BT)}{1 + C \exp(-E_{\text{act}}/k_BT)}\right)
\]

(1)

where \(E_{\text{act}}\) and \(E_{\text{low}}\) are activation energies for the thermal escape and thermal excitation, respectively. \(C\) and \(D\) are proportionality factors. The calculations using Equation 1 are fitted to experimental values and shown by solid lines in Figure 2a.

The \(E_{\text{act}}\) values, which express PL quenching slopes in the high-temperature region, were determined to be \(E_{\text{act1}} = 490\) meV and \(E_{\text{act2}} = 410\) meV for the time-resolved \(I_1\) and \(I_2\) components, respectively. The \(E_{\text{act}}\) values represent the energy difference between the emissive state \(E_s\) and the transfer channel among NDs for thermally activated electron hopping. These values of \(E_{\text{act1}}\) and \(E_{\text{act2}}\) agree well with the barrier height of 0.44 eV for the electron, taking into account the SiC bandgap energy of 3.0 eV obtained from reflectance measurements of a thick SiC film fabricated by the same deposition method and the PL energy of 1.9 eV for the Si NDs. The valence band offset is assumed to be 0.6. The valence band offset for the Si/SiC interface has not been known as far as we know, but that of the Si/SiO\(_2\) has been recently determined to be 0.6 [25]. Therefore, we interpret that the obtained \(E_{\text{act}}\) values show a potential height for the electron at the Si ND/SiC interface. The difference in the above two activation energies \((E_{\text{act1}} - E_{\text{act2}} = 80\) meV) indicates the energy difference between the two emissive states showing the different decay times of \(\tau_1\) and \(\tau_2\), if we assume that the potential height responsible for the thermal escape is identical for both PL components. On the other hand, the \(E_{\text{low}}\) values, which describe negative quenching slopes of the gradual increases in the PL intensity at low temperatures, are obtained to be \(E_{\text{low1}} = 70\) meV and \(E_{\text{low2}} = 90\) meV for the time-resolved \(I_1\) and \(I_2\) components, respectively. At low temperatures, the electron and hole responsible for the \(I_1\) emission are separately trapped at different shallow potential minima within each ND, and the recombination rate significantly decreases. These spatially isolated electron and hole are thermally depopulated within the ND and can recombine at higher temperatures, which results in increases in the PL intensities. This lowest energy level is efficiently not emissive, but the carriers are not extinct via defect-related non-radiative centers because the thermally excited carriers from these states emit strong PL at higher temperatures.

The electron and hole responsible for the \(I_2\) emission can be localized at different NDs in the low-temperature regime and then the electron and hole pair are spatially separated. Therefore, the recombination probability decreases significantly (non-emissive). Increasing the temperature, the electron and hole are thermally excited to the free-like state (emitting \(I_2\)), and the recombination of the electron and hole can take place again. We find that the \(E_{\text{low2}}\) value indicating the activation (localization) energy of the \(I_2\) emission at low temperatures agrees well with the energy difference between \(E_{\text{act1}}\) and \(E_{\text{act2}}\); \(E_{\text{act1}} (490\) meV) - \(E_{\text{act2}} (410\) meV) = 80 meV \((E_{\text{low2}} = 90\) meV). From this correspondence, we attribute the gradual increase in the \(I_2\) emission at the low-temperature region to the thermal excitation of the
carriers from the dark state, where the electron and hole are localized in different individual NDs, to the free-like state among neighboring NDs.

Figure 3 shows PL decay times of the three PL components as a function of temperature. The decay times of $\tau_1$ and $\tau_2$ decrease monotonically with increasing temperature in the high temperature region above 240 K. This temperature-induced lifetime shortening coincides well with the abovementioned thermal quenching due to the electron escape from individual NDs through the transfer channel. Therefore, we conclude that the PL decay characteristics at the high-temperature region are significantly affected by the thermal escape of electrons. In contrast, the PL decay time of $\tau_3$ is almost constant for temperature. This fact infers that electron tunneling through thin barriers play a significant role for the decay characteristics of this fastest PL component rather than the thermal hopping. The picture of ultrafast tunneling of the electron has been discussed in our recent paper and is supported by an experimental fact that the fastest PL component with $\tau_3$ appears only when high-density excitations are made for the dense ND system [20]. The electron tunneling process will be important when we consider applications of superlattices composed of the present high-density Si NDs to solar cells with high efficiencies because a photo-excited electron–hole pair can be immediately separated by this tunneling process before the radiative recombination takes place. Further efforts to enhance the tunneling process will be performed by designing proper barrier materials and the spatial alignment of NDs.

Finally, we discuss about the temperature dependences of the PL decay time based on the abovementioned non-radiative decaying processes possibly caused by the thermal quenching beyond the barriers and energy relaxation to the localization or trap states. The PL decay times of the $I_1$ and $I_2$ components can be separated into a radiative lifetime $\tau_r$ and non-radiative lifetime $\tau_{nr}$ if we assume that the internal quantum efficiency of each PL component is 1 at the temperature showing the maximum PL intensity. The $\tau_r$ and $\tau_{nr}$ were calculated using the
following equations:
\[
\tau_r(T) = \tau_{PL}(T) \frac{I_{\text{max}}}{I(T)}
\]
\[
\tau_{nr}(T) = \tau_{PL}(T) \frac{I_{\text{max}}}{I_{\text{max}} - I(T)}
\]

where \(\tau_{PL}\) is the PL decay time measured, and \(I\) and \(I_{\text{max}}\) are the PL intensity at a certain temperature \(T\) and the maximum PL intensity, respectively. If the quantum efficiency at the temperature showing the maximum PL intensity is smaller than 1, absolute values of both the \(\tau_r\) and \(\tau_{nr}\) varies. However, the trends of the temperature dependences of the \(\tau_r\) and \(\tau_{nr}\) should be similar because the PL intensity shows non-monotonic temperature dependence. The \(\tau_r\) and \(\tau_{nr}\) lifetimes deduced for the \(I_1\) and \(I_2\) components are plotted as a function of temperature in Figure 4a, b, respectively, together with the measured \(\tau_{PL}\).

At the lower temperature region below 200 K, the \(\tau_{nr}\) value decreases with decreasing temperature, and the \(\tau_{PL}\) becomes dominated by the \(\tau_{nr}\). This trend can be understood by the existence of non-emissive localized or trap states as discussed above. The \(\tau_{nr}\) value increases toward the maxima with increasing temperature because of the thermal excitation of the carriers from the localized or trap levels to the emissive ones. In contrast, in the high-temperature regions toward room temperature, the \(\tau_{nr}\) decreases with increasing temperature because of the thermal escape from the emissive level beyond the barriers. These PL dynamics for the two slower decaying PL components of \(I_1\) and \(I_2\), expressed by the temperature dependences of the \(\tau_r\) and \(\tau_{nr}\) agree well with the thermal quenching and excitation processes elucidated by the temperature dependences of intensities of these PL components.

Conclusions

We have studied temperature dependences of time-resolved PL in the two-dimensional high-density Si ND arrays fabricated by NB etching using bio-nano-templates, where the PL time profiles with various temperatures are fitted by triple exponential decay curves. We find that the time-integrated PL intensities in the two slower decaying components depend strongly on temperature, which is attributed to PL quenching due to thermal escape of electrons from emissive states of individual NDs in addition to thermal excitations of carriers from localized or trap states in the individual NDs to the emissive ones. The temperature dependences of the PL intensity were analyzed by the three-level model. The following thermal activation energies corresponding to the thermal escape of the electron are obtained to 410 and 490 meV, depending on the PL components. In addition, we find dark states of photo-excited carriers, which can be attributed to the separate localization of the electron and hole into different NDs with the localization energies of 70 and 90 meV, depending on the PL components. The PL decay times of these two decaying components ranging from 70 to 800 ps are also affected by this thermal escape at high temperatures from 240 to 300 K. The fastest decaying component shows a constant decay time of about 10 ps for various temperatures, in which the decay characteristic is dominated by the electron tunneling among NDs.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

TK and AM conceived the spectroscopic study, participated in its design and coordination, and drafted the manuscript. TK and YM carried out the time-resolved PL measurement and analyzed the data. MI, CH, and SS conceived the fabrication process and participated in its design and coordination. MI and CH fabricated the Si-ND array sample. All authors read and approved the final manuscript.

Acknowledgments

This work is supported in part by the Japan Society for the Promotion of Science, Grant-in-Aids for Scientific Research (S) No. 22221007.

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Received: 4 April 2013 Accepted: 1 May 2013
Published: 10 May 2013

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