PHOTOLUMINESCENCE SPECTRAL FEATURES OF SILICON NANOWIRES

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Abstract

Computer simulation using matlab programming approach is carried out to study the photoluminescence (PL) properties of silicon (Si) nanowires (NWs) with diameter between 1.5 and 5.8 nm. An integrated hybrid model comprising of quantum confinement, surface states, and exciton binding is developed to calculate the size, wavelength and photon energy dependent PL intensity. The influence of size and passivation on the band gap energy and PL spectra of Si nanowires (NWs) are examined. It is observed that all the model parameters for quantum confinement, localized surface states, and exciton energy are responsible for the changes in the electronic and optical properties of Si NWs. The simulated data are compared with experimental findings. The admirable features of the results suggest that the present model is significant for understanding the mechanism of visible PL from Si NWs. The model can be extended to study temperature dependent PL for other nanostructures of different shapes and size.

Keywords: Photoluminescence; quantum confinement; nanowires; passivation; radiative recombination

Abstrak

Simulasi komputer menggunakan pendekatan pengaturcaraan MATLAB dijalankan untuk mengkaji photoluminescence (PL) Hartanah silikon (Si) nanowires (NWS) dengan diameter antara 1.5 dan 5.8 nm. Model hibrid bersepadu yang terdiri daripada kurungan kuantum, negeri permukaan, dan exciton mengikat dibangunkan untuk mengira saiz, panjang gelombang dan tenaga foton bergantung intensiti PL itu. Pengaruh saiz dan pasif kepada jurang jalur tenaga dan PL spektrum nanowires Si (NWS) akan ditekiti. Adalah diperhatikan bahawa semua parameter model untuk berpantang kuantum, negeri permukaan setempat, dan tenaga exciton bertanggungjawab ke atas perubahan dalam sifat-sifat elektronik dan optik Si NWS. Data simulasi dibandingkan dengan hasil uji koji. Ciri-ciri terpuji keputusan menunjukkan bahawa model ini adalah penting untuk memahami mekanisme dilihat PL dari Si NWS. Model ini boleh dilanjutkan untuk mengkaji suhu bergantung PL untuk nanostruktur lain bentuk dan saiz yang berbeza.

Kata kunci: Photoluminescence; quantum confinement; nanowires; passivation; radiative recombination

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1.0 INTRODUCTION

Developing novel electronic and optoelectronic devices by nanostructuring semiconductors is ever-demanding. The observation of room-temperature visible photoluminescence (PL) from silicon (Si) nanostructures (quantum dots or wires) stirred tremendous attention towards Si nanomaterials [1]. Recent semiconductor research is exploring the feasibility of tuning the optical response of Si nanostructures via band gap engineering. Experimentally observed visible PL can be explained to some extent by quantum confinement (QC) effect that modifies the energy band gap [2]. Lately, Si nanostructures with hydrogen and oxygen passivated surface became attractive due to enhanced light emission. However, the mechanism of visible PL remains unclear despite several studies using models, experiments, and simulations approaches [2-4].

An analytical model for silicon nanocrystals (Si NSs) photoluminescence of quantum confinement effect and localized surface state have been proposed by Ding et al. [5] and Estes and Model [6]. In this work, we developed a comprehensive phenomenological model by combining the effects of surface states, exciton energy, and QC to explain the PL mechanism of Si NWs. It is demonstrated that by controlling a set of parameters extracted by fitting the model with experiments, it is possible to interpret the PL spectral features accurately. The band gap is found to decrease with the increase of NWs diameter [7]. The results presented here allow us to examine the influence of size and passivation on the band gap energy and PL spectra of Si nanowires [Si-NWs].

2.0 MODELS AND METHODS

The initial and the famous explanation for the visible PL in Si-NWs is the quantum confinement of exciton in nanometer-sized silicon. The model of the quantum confinement of Si NWs is based on the electronic confinement in wire. An empirical law that links the size dependence of optical band gap of NWs is written as:

\[ E_g = E_g^{\text{bulk}} + \frac{\hbar^2}{4a^2} \]  

(1)

where \( \hbar \) and \( a \) are QC parameters and \( a \) is the diameter of Si-NW [8].

The model is further fitted with different experimental and simulation data to extract the value of this parameter. Because of quantum confinement, small-diameter wires exhibit a direct band gap that increases as the wire diameter narrows, irrespective of surface termination. The photoluminescence is attributed to energy shift of the bottom of the conduction band to high energy and of the top of the valence band to low energy [9]. The changes in wave functions explaining the dynamics of electrons and holes that modifies the density of states is due to different in quantum confinement directions.

In this model, we consider Si NWs of well-defined diameter distribution as an ensemble of nanoscale spherical particles. In view of this consideration, the intensity of PL at particular photon energy becomes proportional to the population of occupied surface states and the oscillator strength. The number of surface states in a crystallite is proportional to the number of atoms on the surface and hence, surface area \( A \) of the crystallite. Considering \( N_s \) as the total number of surface states \( N_s \propto A \). Again, if we assume that each atom in a crystallite contributes at least one photo excited carrier to the crystallite, the number of photo excited carriers \( N_p \) in a crystallite is proportional to its volume \( V \), hence, \( N_p \propto V \). Furthermore, since the rate of transition from an excited carrier to the localized surface states is proportional to the product of the number of excited photo carriers and the number of available empty surface states in steady state condition, the population \( N_s \) of photo carriers in surface states participating in PL processes becomes proportional to the product of \( N_t \) and \( N_p \). That is,

\[ N_c \propto N_tN_p \propto a^2d^3, \]

(2)

where \( a \) is the diameter of the crystallite wire.

The rate of radiative transition depends on the oscillator strength \( f \). The oscillator strength in nanocrystalline materials varies as inverse power law and can be approximated as \( f \sim \frac{1}{d^a} \), where power exponent \( a \) depends on the material properties as well as the range of crystallites sizes being used. Taking the oscillator strength into account, the radiative transition probability in nanocrystallites of diameter becomes,

\[ P(d) \propto N_c f \propto a^{5-a}. \]

(3)

The PL intensity from an ensemble of crystallite of Si NWs having size distribution will be obtained by summing the contributions from all the crystallites having size diameter \( d \) [10]. The PL intensity with respect to crystallites of size \( d \) is given by,

\[ I(d) \propto P(d)\rho(d). \]

(4)

Then, the emitted photon energy from nanocrystallites wire will be lower than the energy band gap of the crystallite by an amount of the localization energy \( E_s \) of the surface states and the exciton binding energy \( E_{ex} \) which are functions of crystallites size. The emitted photon energy from crystallites of the quantum dot is given as

\[ E_p = E_g^{\text{bulk}} + \Delta E - E_s - E_{ex}. \]

(5)

where the amount of band gap up shift due to QC in the nanocrystallites wire is given by \( \Delta E \) and \( E_g^{\text{bulk}} \) is the energy band gap corresponding to the bulk crystalline material.

By transforming Eq. (4) from \( d \) to \( \Delta E \) using standard Fourier Transform one obtains,
\[ I(\Delta E) = \int I(d) \delta (\Delta E - \frac{b}{2\sigma}) d\sigma \propto \int d^{5-d} \rho(d) \delta \left( \Delta E - \frac{b}{2\sigma} \right) dd. \quad (6) \]

By considering a normal distribution of crystallite diameters in nanocrystallites silicon wire, then

\[ \rho(d) = \frac{1}{\sqrt{2\pi} \sigma} \exp \left( -\frac{(d-L_0)^2}{2\sigma^2} \right) \quad (7) \]

where \( L_0 \) and \( \sigma \) are the mean crystallite size of NWs and standard deviation, respectively. Substituting Eq. (7) into Eq. (6), we obtain the analytical expression of PL intensity as

\[ I(\Delta E) \sim \frac{1}{\sqrt{2\pi} \sigma} \left( \frac{b}{\Delta E} \right)^{(6-d+\theta)} \exp \left[ -\frac{(\frac{b}{\Delta E} - L_0)^2}{2\sigma^2} \right] \quad (8) \]

Eq. (8) indicates that the PL profile varies strongly with the QC parameters \( \theta \) and \( \delta \). Therefore, good care should be taken in using the correct QC model for band gap up shift estimation.

The oscillator strength and the exciton binding energy \( E_b \) both are complicated functions of the size of nano crystallites and their surrounding media. Following careful calculations of Proot et al. for crystalline silicon, we adopted a ranging value of 0.01 - 0.22 eV for \( E_b \), which is a good average value for the range of crystalline sizes from \( \sim 5.8 \) nm to \( 1.5 \) nm [11]. The localization energy \( E_{ss} \) is taken to be the order of phonon energies which is about 0.05eV for optical phonons and \( E_g \) as 1.12 eV for crystalline silicon at room temperature.

The orientation dependent parameters are found to be \( \theta = 1.40 \) and \( \delta = 3.37 \) eV. This is obtained by fitting the model with experimental values of size dependent band gap of Bruno et al. [12] using quantum electronic gap expression (Eq.1) as shown in Figure 1. Our model expression (Eq. 8) is used to generate PL spectra using these quantum confinement parameters with the correction of \( \pm 0.5 \) for comparison and further analyses. PL spectra are simulated using a matlab program. The band gap energy is found to increase as the NWs diameter are decreased [13].

![Figure 1](image1.png)  
**Figure 1** Band gap variation of Si-NWS with diameter [6].

The analytical expressions (Eq.1) for the size dependent band gap from the model are fitted to the pseudo-potential approximation (PPA) and tight binding method (TBM). Fig. 2 and Fig. 3 exhibit the fitted electronic gap [14]. The \( \theta \) and \( \delta \) values of the quantum confinement parameters for PPA are found to be 10.5 and 1.39 and for TBM it is 13 and 1.39, respectively. These fitted energy values are used in our model Eq. (8) to generate PL spectra of nanocrystalline-silicon structure.

![Figure 2](image2.png)  
**Figure 2** Model fitted to Pseudo-Potential Approximation (PPA).

![Figure 3](image3.png)  
**Figure 3** Model fitted to Tight Binding Method (TBM).

### 3.0 RESULTS AND DISCUSSION

PL intensity versus photon energy with varying standard deviation is shown in Fig. 4. It indicated that the PL peak shift towards higher energy (blue energy) as the silicon nanowire standard (sigma) deviation parameters decreases. This validated the quantum confinement model. Furthermore, a red shift is observed with the increase of standard deviation \( \sigma \). The PL spectra broaden as well as shifts towards low photon energy accompanied by a decrease in relative PL intensity. This indicates that the amount of size dispersion affects both the PL peak energy and it width.
Fig. 4 reveals the variation in the PL intensity against photon energy with varying Si-NWs diameter. It depicts the red-shift of PL peak position with the increasing mean crystallite size for fixed size dispersion. The PL peak is blue shifted as the Si NWs diameter is decrease. This confirms the validation of the quantum confinement model [4]. The increase in PL intensity with the increase in Si NWs diameter is attributed to the larger number of luminescent center and recombination effects.

It is clear from Fig. 4 and Fig. 5 that the presence of exciton energy in the forbidden region contributed for the observed blue shift. Hence, the integrated surface states effect, exciton states, and quantum confinement effect model can explain accurately the experimentally observed PL spectra of Si NWs and the blue shift in PL peak.

Definitely, this model is able to predict the already observed experimental PL data on Si NWs produced by a variety of techniques. Figure 6 and 7 compares the simulation results (6b and 7b) with the experimental (6a and 7a) data from Ledoux et al., (2000). Clearly, a blue shift in the PL peak is evidenced with the decrease in mean diameter of Si NWs. Furthermore, the PL intensity is greatly influenced by the size distribution of NWs.

Furthermore, PL spectra were generated using parameters found from size dependent band gap (PPA and TBM) in Fig. 8 and Fig. 9. The results revealed that
surface passivation enhances the band gap energy of the Si NWs structure which is in conformity with surface state effects.

**Figure 8a** NWs size dependent PL Spectra from Pseudo potential approximation

**Figure 8b** NWs size dependent PL Spectra from Tight binding model

**Figure 9a** Wavelength dependent PL spectra from PPA

**Figure 9b** Wavelength dependent PL spectra from TB

### 4.0 CONCLUSION

The mechanism of photoluminescence and the enhancement of band gap in Si NWs are investigated using a hybrid phenomenological model. The effects of quantum confinement, localized surface states and exciton binding energy are integrated in the model. The results exhibit that by controlling a set of parameters extracted by fitting this developed model with experimental finding, it is possible to interpret the observed PL spectral features. It is shown how mean diameter of NWs affects the PL intensity and band gap energy. The importance of localized surface states in predicting the PL data from Si-NWs using the developed model is demonstrated. Furthermore, the band gap is found to increase on passivating the NWs surface with oxygen more than hydrogen. We affirm that both QC and surface passivation together with exciton effects determine the optoelectronic properties of Si NWs. The mechanism of PL emission and enhancement of band gap is understood. In general, this developed model is able to explain the experimental observation of visible PL from Si-NWs as well as others Si nanostructures (quantum dots and wells).

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### References


