Title
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Citation
Journal of Applied Physics, 113(5): 053505

Issue Date
2013-02-07

Doc URL
http://hdl.handle.net/2115/54789

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Type
article

File Information
314-JAP13-Nahid InGaSb QW.pdf

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Carrier dynamics and photoluminescence quenching mechanism of strained InGaSb/AlGaSb quantum wells

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(Received 4 November 2012; accepted 8 January 2013; published online 5 February 2013)

GaSb based quantum wells (QWs) show promising optical properties in near-infrared spectral range. In this paper, we present photoluminescence (PL) spectroscopies of InxGa1−xSb/AlxGa1−xSb QWs and discuss the possible thermal quenching and non-radiative carrier recombination mechanisms of the QW structures. The In and Al concentrations as well as the QW thicknesses were precisely determined with the X-ray diffraction measurements. Temperature dependent time-integrated and time-resolved PL spectroscopies resulted in the thermal activation energies of ~45 meV, and the overall self-consistent calculation of the band parameters based on the measured physical values confirmed that the activation energies are due to the hole escape from the QW to the barriers. The relation of the present single carrier escape mechanism with the other escape mechanisms reported with other material systems was discussed based on the estimated band offset. The relation of the present thermal hole escape to the Auger recombination was also discussed. © 2013 American Institute of Physics. [http://dx.doi.org/10.1063/1.4789374]

I. INTRODUCTION

Near-infrared emitters can be used for a wide range of applications such as gas sensing,1 monitoring of environmental pollution such as carbon monoxide and phosphine,2–4 military scene projection, thermo-photovoltaic (TPV) devices,5,6 and low-loss fibre optical communication.7 Especially, vertical cavity surface emitting lasers (VCSELs) or vertical external cavity surface emitting lasers (VECSELs) have wide range of applications. Although InP-based edge emitting lasers have been in practical use in the telecommunication band, it is well known that high-efficiency distributed Bragg reflectors (DBRs) are difficult to grow on InP substrates. However, near-infrared lasers are generally difficult to achieve on GaAs substrates.

Carriers in such applications, GaSb has received practical interest due to its narrow band-gap properties which offer potentiality in long-wavelength optoelectronic devices. For example, InGaSb/GaAs quantum dot (QD) VCSEL was operated at room temperature (RT) employing AlAs/GaAs DBRs.11 Although Sb-based materials system is highly mismatched to GaAs substrates, interfacial 90° misfit dislocations (IMF) technique was developed,12 which localizes the dislocation generation at the GaSb/GaAs interface and prevents threading dislocations in overgrown Sb-based heterostructures. This IMF technique made RT laser operation of InGaSb/AlGaSb quantum wells (QWs) possible on (001) GaAs substrate.13 The ternary InGaSb alloys are suitable to tailor wavelengths in the near infrared and longer wavelengths over 2 μm.13–16 Recently, Lai et al. demonstrated 340-W peak output at the wavelength of 2 μm from an InGaSb/AlGaSb VECSEL operated at RT.17 Continuous-wave (CW) operation at RT was also achieved but the output power was limited to 0.12 W.18 The main factor that limits the VECSEL maximum light output was heating due to non-radiative recombination. Therefore, it is crucial to extend our understanding of the non-radiative processes. To date, numbers of works have been carried out to clarify the non-radiative processes. Some suggested that non-radiative Auger recombination is the most detrimental mechanism which limits the performance of the low band-gap long-wavelength semiconductor lasers.19–22 On the other hand, Shterengas et al.23,24 and Rainò et al.25 reported that thermally induced hole escape is the main non-radiative process which deteriorates the optical properties and performances of the GaSb-based devices at elevated temperature.

Therefore, it is the purpose of this paper to clarify the role of the carrier escape in InxGa1−xSb/AlxGa1−xSb QWs and study the relation to the Auger recombination based on transient and CW luminescence measurements. The QW samples were precisely characterized by X-ray diffraction (XRD) measurements to determine the epitaxial layer thicknesses and compositions. The temperature dependent carrier recombination process was analysed by employing CW time-integrated and time-resolved photoluminescence (PL) spectroscopy. From the temperature dependence of the steady-state PL and decay time analyses, the thermal quenching energies of the QWs were extracted. Including the measured thicknesses of the QWs, the thermal activation energies, theoretically determined band deformation potential, and the transition energies as input parameters, the quantum confinement energies were calculated by solving the Schrödinger equation for finite
potential wells. From these observations and analyses, we develop and quantify the whole energy band structure to explain the non-radiative mechanism responsible for PL quenching in our studied InGaSb/AlGaSb QW structures. Finally, the relation of the carrier escape mechanism to the Auger recombination is discussed.

II. STRUCTURAL STUDY OF InGaSb/AlGaSb QWs BY X-RAY DIFFRACTION

InGaSb QWs embedded in AlGaSb barrier matrix studied in this work were grown by solid-source molecular-beam epitaxy (MBE) on GaSb(001) substrates. The InxGa1−xSb QWs of around 3-nm thickness are surrounded by barrier layers of around 50-nm-thick AlxGa1−xSb and capped by 10-nm-thick GaSb layers on top. To examine the precise thicknesses and compositions of the QWs and barriers, XRD measurements were carried out using the Bruker D8 Discover high-resolution XRD system. The simulations of the 2θ-ω scans reproduced the measurements very nicely as shown in Fig. 1(b), and the thicknesses of 2.9 nm and 3.4 nm with the respective indium (In) concentration of 32% and 36% for the two InxGa1−xSb QWs and the aluminium concentrations of 49% and 50% for the AlxGa1−xSb barrier layers were derived (the result on one of the samples is shown in Fig. 1). The derived parameters for one of the QWs are shown in Fig. 1(a).

III. EXPERIMENTAL SET UP FOR PL MEASUREMENTS

Time-integrated PL measurements were performed using non-resonant CW excitation at the low average power density of 5 W/cm² using a frequency doubled Nd:YAG laser emitting at 2.33 eV (532 nm). A He-Ne laser at 1.95 eV (633 nm) was also used for the excitation. For the temperature dependent PL spectroscopy, a 30-cm monochromator with a rotating 200 grooves/mm grating and a liquid Nitrogen cooled InGaAs photodiode detector with the extended wavelength range up to 2.2 μm were used in the lock-in operation and the excitation laser was modulated by a mechanical chopper.

Time-resolved PL spectroscopy was performed to investigate the temperature dependent carrier dynamics in the QWs. The ∼5-ps optical pulses generated by a mode-locked Ti:sapphire laser tuned at 1.59 eV (780 nm) with a repetition rate of 76 MHz were used for the excitation. A near-infrared streak camera (Hamamatsu C11293-01) combined with a 150 groves/mm monochromator was used for the detection of the time-resolved PL spectra with 20-ps time resolution. Because of the detection sensitivity limited to the wavelength below 1700 nm, the present InGaSb/AlGaSb QW samples were designed to emit in the 1550-nm band.

IV. PL MEASUREMENTS AND DISCUSSIONS

To study the thermal quenching mechanism of luminescence at elevated temperature, precise measurements and clear identification of its origin are inevitable. The carrier dynamics in the QWs is discussed quantitatively with the measurements of transient PL, together with the temperature dependent steady-state PL measurements in this section. Based on these results and their analyses, the fundamental QW band structures are determined and the luminescence quenching mechanism is discussed.

A. Time-integrated PL measurements

CW time-integrated PL measurements were performed on the two InGaSb QWs at 10 K under the low excitation power of 5 W/cm² using the YAG laser (2.33 eV) and the measured spectrum of the In0.32Ga0.68Sb QW is shown in Fig. 2(a). The PL peak energies to be discussed hereafter in this paper are obtained by employing Gaussian line-shape functions in multi-peak fitting. The dominant emission peak at 1508 nm (822 meV) is attributed to the optical transition in the In0.32Ga0.68Sb strained QW together with the later discussion. The emission peak centered at 1593 nm (778.3 meV) is associated with the donor-acceptor pair (DAP) transition usually observed in GaSb substrates26,27 and its temperature dependence will be discussed later. The small emission peak at around 1557 nm (796.1 meV) is attributed to the GaSb exciton bound to neutral acceptor (BE4-A0) transition commonly reported in previous publications.26

To confirm the PL peak assignments associated with impurities in the GaSb substrate, we replaced the YAG laser with a He-Ne laser with the lower photon energy of 1.95 eV. The InGaSb absorption coefficient at the photon energy of ~2 eV is estimated to be ∝10⁵ cm⁻¹, and then the laser penetration depth is 1/α ~ 100 nm. Our present QW samples have the total AlGaSb barrier thickness of ~100 nm that is the same order with the laser penetration depth. Therefore, the injection of photo-generated carriers into the GaSb substrate and the observation of the PL from the GaSb substrate

(a) GaSb caplayer (10 nm)
Al0.49Ga0.51Sb (49 nm)
In0.32Ga0.68Sb (2.89 nm)
Al0.49Ga0.51Sb (49 nm)
GaSb

(2θ-ω scan using the Bruker D8 Discover high-resolution XRD system exemplarily shown for the case of In0.32Ga0.68Sb/Al0.49Ga0.51Sb QW.)
will be sensitively dependent on this penetration depth and the diffusion of photo-generated carriers. Under this condition, the PL was measured at the higher excitation power of 3000 W/cm² as displayed in Fig. 2(b) and the DAP and BE4-A⁰ peaks originating from the GaSb substrate became more intense. In addition to the three PL peaks found in Fig. 2(a), two more PL peaks are observed at 1544 nm (802.9 meV) and 1530 nm (810.3 meV) in Fig. 2(b), which are ascribed to another GaSb exciton bound to neutral acceptor (BE2-A⁰) transition and the free-exciton (EF) transition, respectively. The free exciton energy of GaSb together with its binding energy results in the band gap energy of 811.7 meV which agrees well with the previously reported values. Therefore, it can be concluded that the additional peaks of the DAP, BE2-A⁰, BE4-A⁰, and EF are emerged from the GaSb substrate due to the carrier diffusion.

The dominant PL peak in Fig. 2(a), which was attributed to the QW emission, was further studied on its temperature dependence as shown in Fig. 3(a). The temperature dependence of the measured luminescence peak position of the In₀.₃₂Ga₀.₆₈Sb QW is displayed by the red circles in Fig. 3(a) and was fitted using the Varshni empirical expression. The temperature dependent integrated PL intensities of In₀.₃₂Ga₀.₆₈Sb (diamonds) and In₀.₃₆Ga₀.₆₄Sb QWs (circles) fitted by Arrhenius law result in the activation energies of 45 and 47 meV, respectively.
\[ E(T) = E_0 - \frac{xT^2}{\beta + T}, \] (1)

where \( E_0 \) is the 0 K energy gap, \( T \) is the temperature in K, \( x \) and \( \beta \) are the fitting parameters, respectively. The solid line results from the fit using the Varshni parameters of \( x = 0.39 \text{ meV/K} \) and \( \beta = 180 \text{ K} \) reported on InGaSb. The DAP transition shows very weak temperature dependence up to 90 K as displayed by the diamonds connected with the dashed line in Fig. 3(a). For the higher temperature, the DAP peak intensity decreases rapidly, while the QW excitonic emission remains dominant. Since DAP undergoes thermal ionization easily due to the small binding energy, the DAP emission intensity rapidly decreases for the higher temperature. Based on these studies, it can be unambiguously inferred that the PL emission appearing at the wavelength of 1508 nm (822 meV) at low temperature is the exciton emission in the InGaSb QW.

The temperature dependences of the PL spectra measured on the In\(_{0.32}\)Ga\(_{0.68}\)Sb and In\(_{0.36}\)Ga\(_{0.64}\)Sb QWs are shown in Figs. 3(b) and 3(c), respectively. The QW emission was fitted with the Gaussian line-shape function on each measured spectrum, and its spectrally integrated PL intensity was plotted as a function of inverse temperature in Fig. 3(d). The closed diamonds are for the In\(_{0.32}\)Ga\(_{0.68}\)Sb QW and the closed circles are for the In\(_{0.36}\)Ga\(_{0.64}\)Sb QW. The solid (black) lines are fitting to the experimental data using the Arrhenius equation

\[ I(T) = \frac{I_0}{1 + B \exp\left(-\frac{E_a}{kT}\right)}, \] (2)

where \( I_0 \) is the integrated PL intensity at 0 K, \( E_a \) is the thermal activation energy, \( k \) is the Boltzmann constant, and \( T \) is the sample temperature in K. \( B \) is the dimensionless coefficient which is defined simply as the ratio of carrier capture time and carrier escape time of the related non-radiative process. For the In\(_{0.32}\)Ga\(_{0.68}\)Sb QW, the best fit of Eq. (2) yields the activation energy \( E_a = 45 \pm 9 \text{ meV} \) and the corresponding coefficient \( B = 1270 \pm 210 \). For the In\(_{0.36}\)Ga\(_{0.64}\)Sb QW, the best fit results in the activation energy of \( E_a = 47 \pm 11 \text{ meV} \) with the corresponding coefficient of \( B = 615 \pm 93 \).

**B. Time-resolved PL measurements**

For further investigating the carrier dynamics in the QW structures, time-resolved PL measurements were performed with the 20-ps time resolution. In the following, the measured results on the In\(_{0.32}\)Ga\(_{0.68}\)Sb QW are presented and discussed (the In\(_{0.36}\)Ga\(_{0.64}\)Sb QW shows qualitatively similar results).

The excitation power dependence of the transient PL decay measured on the In\(_{0.32}\)Ga\(_{0.68}\)Sb QW at 4 K is shown in Fig. 4(a). It is found that the PL transients are well fitted by mono-exponential decay superposed with background noise for all the excitation power. This fitting resulted in the PL lifetime of \( \sim 600 \text{ ps} \) irrespective of the excitation power. The temperature dependent time-resolved PL measurements were also performed and the measured results at the temperatures of 4 K and 215 K are exemplarily shown in Fig. 4(b). The temperature dependence of the PL lifetime obtained from the
measurements on the In$_{0.32}$Ga$_{0.68}$Sb QW is summarized in Fig. 4(c). The lifetime remains almost the constant value of 600 ps in the low temperature range below 100 K, while it decreases at the higher temperature. The observed temperature dependence of the lifetime was fitted with the following equation:25

$$
\tau(T) = \frac{\tau_d}{1 + (\tau_d/\tau_e)\exp(-E_a/kT)},
$$

(3)

where $T$ is the temperature, $k$ is the Boltzmann constant, $\tau_d$ is the recombination lifetime extracted from Fig. 4(a) at low temperature, $E_a$ is the activation energy, and $\tau_e$ is the escape time of charge carriers from the QW to the barriers.22 The fitting of the experimental data results in the escape time of 93 ± 8 ps and the activation energy of about 43 meV ± 11 meV. The activation energy of 43 meV derived from the time-resolved measurements is very close to the one of 45 meV previously deduced from the time-integrated PL measurements.

C. Calculation of QW band structures based on PL measurements

The agreement of the activation energies derived from the time-resolved and time-integrated PL measurements together with the fact that the lifetime does not depend on the excitation power suggests that this thermal activation process is the carrier escape from the QW to the barriers and is responsible for PL quenching. For the complete and self-consistent understanding of the measured results, the QW confinement energies were calculated by solving the Schrödinger equation for the finite potential wells. First, the deformation potential and the strain contribution to the band structure were considered. The valence-band structure near the zone centre ($k_z = 0$) can be significantly reshaped with strain. The total Hamiltonian which describes the band structure of the strained QW can be expressed as,

$$
H = H_0 + H_s,
$$

where $H_0$ and $H_s$ are the Luttinger-Kohn (LK) Hamiltonian and the strain Hamiltonian, respectively. When a QW is under biaxial strain in the (001) QW plane, the strain components are written as

$$
\varepsilon_{xx} = \varepsilon_{yy} = \varepsilon_{II} = \frac{\Delta a}{a},
\varepsilon_{zz} = -\varepsilon_{II} = \frac{2C_{12}}{C_{11}},
$$

$$
\varepsilon_{xy} = \varepsilon_{yx} = \varepsilon_{zz} = 0,
$$

(4)

where $\Delta a/a$ is the strain due to the lattice mismatch at the hetero-interface and $C_{11}$, $C_{12}$ are the stiffness constants. The QW plane was considered to be the x-y coordinate plane. For this condition, the strain Hamiltonian $H_s$ is expressed as34

$$
H_s = -E'_H - \frac{3}{2m^*}E_U \left( \frac{\hbar^2}{2m^*} \right),
$$

(5)

where

$$
E'_H = 2a \left( \frac{C_{11} - C_{12}}{C_{11}} \right) \varepsilon_{II},
$$

$$
E_U = -b \left( \frac{C_{11} + 2C_{12}}{C_{11}} \varepsilon_{II} \right) \varepsilon_{II},
$$

(6)

(7)

where $a$ and $b$ are the hydrostatic and shear deformation potentials, respectively. The matrix element derived from Eq. (5) results in the heavy-hole (HH) band that is uncoupled to the other bands, while the coupling of light-hole (LH) and spin-orbit split-off (SO) bands occurs by the matrix element $\sqrt{2}E_U$.35 The shear deformation potential term $E_U$ is proportional to the in-plane strain $\varepsilon_{II}$ as is shown in Eq. (7) and is larger for the larger strain. At $k_z = 0$, the $8 \times 8$ Hamiltonian including the strain terms becomes doubly degenerate $4 \times 4$ matrix. The conduction band in this matrix is decoupled and the Eigen value equation results in the strain contribution to the HH valence-band maximum (VBM) as35,36

$$
E_d = -E_H - E_U.
$$

(8)

The details of the analysis are narrated in references by Pollak and Cardona35 and Suemune.36 To evaluate Eq. (8), the material parameters, e.g., the lattice constant, energy gap, and the values of $a$, $b$, $C_{11}$, and $C_{12}$ were deduced by the linear interpolation of the corresponding values established for InSb, GaSb, and AlSb.37 Using Eqs. (4), (6), (7), and (8), the strain contribution to the HH-VBM of the In$_{0.32}$Ga$_{0.68}$Sb and In$_{0.36}$Ga$_{0.64}$Sb QWs is calculated as $E_d = -E_H - E_U = -72$ meV and $E_d = -E_H - E_U = -78$ meV, respectively.

We have precisely determined the well thickness and the alloy compositions with the XRD measurements in Sec. II, and therefore, the unstrained energy gaps of the In$_{0.33}$Ga$_{0.67}$Sb QW and the Al$_{0.49}$Ga$_{0.51}$Sb barriers are given as 538 meV and 1203 meV, respectively, from the reported materials data.37 Including the calculated strain contribution, the QW energy gap increases to 610 meV. In the present InGaSb/AlGaSb common-anion heterostructures, it is known that the band offset is localized to the conduction band. Then it is natural to assume that the thermal activation energy of 45 meV for the carrier escape from the QW to the barriers is dominated in the valence band. Under the present low excitation condition, it will favourably be the hole escape from the QW ground energy state to the AlGaSb barriers. The measured QW exciton emission energy was 822 meV. Consideration of the small exciton binding energy (1.14 meV)$^{38,39}$ leads to the transition energy of 823 meV from the conduction-band to the valence-band QW energy states. Based on the above available information, we have calculated the QW confinement energies by solving the Schrödinger equation by varying the band offset values. Using the iteration method, it is found that the small valance-band offset of 13% gives the most precise agreement with the experimentally observed QW emission energy. This small valence-band offset is comparable to the previously published values.40 Our calculated band parameters are summarized in Fig. 5 and the calculated transition energy is given by

$$
E_T = E_e^1 + E_g - E_d + E_{hh}^1
= (183 + 538 + 72 + 30) \text{meV} = 823 \text{meV},
$$

(9)

where $E_e^1 = 183$ meV is the ground-state electron confinement energy, $E_{hh}^1 = 30$ meV is the ground-state HH confinement energy, $E_g = 538$ meV is the energy gap of the In$_{0.32}$Ga$_{0.68}$Sb QW, and $E_d = -72$ meV is the strain contribution.
where the ratio of 41–44 the carrier escape mechanism has been discussed with the valence-band QW energy state to the barriers. The nature of the PL quenching mechanism is the hole escape from the energy states. Equation (10) classifies three different scenarios measured on the QW sample did not show any change with increasing the excitation power, which apparently implies that the QW does not follow the typical density dependence of the Auger recombination. However, the hole leakage that limits the maximum hole concentration is closely related to this phenomenon. In Fig. 3(a), the integrated PL intensity initiates to quench above 50 K, while the PL decay time remains the same up to 100 K. This difference suggests that the hole concentration is kept nearly the same in spite of the hole escape at around 100 K, which keeps the Auger recombination rate nearly constant. At the higher temperature, it is probable that the energy broadening increases the Auger recombination rate as well as the thermally activated hole escape. However, from the temperature dependent time-resolve PL measurements, it is noticed that the PL decay time decreased from 600 ps at 4 K to 510 ps at 220 K, which is rather small variation in comparison to the typical reported 3–4 times drastic decrease of the Auger recombination time with temperature. This will be attributed to the limited hole concentration by the hole escape in the present situation.

D. Discussion

The present self-consistent results confirm that the dominant PL quenching mechanism is the hole escape from the valence-band QW energy state to the barriers. The nature of the carrier escape mechanism has been discussed with the ratio of

\[ \nu = \frac{E_a}{\Delta E} \]  

where \( E_a \) is the measured thermal activation energy and \( \Delta E \) is the optical gap between the emitting states and the higher energy states. Equation (10) classifies three different scenarios of charge carrier escape. For \( \nu = 1 \), an escape process of fully correlated electron-hole pairs, i.e., bound excitons, is described, while \( \nu \) less than 1/2 is described for single carrier escape. In the case of correlated electron-hole (e-h) escape, \( \nu = \frac{1}{2} \) is satisfied. In the case of InAs/InGaAlAs emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape. In our present InGaSb/AlGaSb, QW sample also emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape. In our present InGaSb/AlGaSb, QW sample also emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape. In our present InGaSb/AlGaSb, QW sample also emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape. In our present InGaSb/AlGaSb, QW sample also emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape. In our present InGaSb/AlGaSb, QW sample also emitting in the 1.55-μm band, the authors concluded \( \nu = \frac{1}{2} \), that is, the mechanism behind the thermal escape was suggested to be a correlated e-h pair escape.

Finally, we want to discuss the relation to the Auger recombination process which is commonly considered for narrow-band-gap semiconductors. This non-radiative Auger recombination is a density dependent mechanism which usually increases with decreasing band-gap energy and is typically found to be increased monotonically with temperature and carrier density. Main Auger recombination processes are divided into two processes of the conduction (C)-C-hole(H) (CCCH) transition and C-H-H-SO(S) (CHHS) transition. The CCCH process involves one electron transition within the conduction band and another electron transition to the valence band. The CHHS process involves one hole transition to the conduction band and another hole transition to the SO band. In the case of the Sb-based semiconductors, the SO energy is very large, such as 820 meV for GaSb, and is compatible to their energy gaps, and the CHHS process dominates because the conservation of energy and momentum for the Auger transition is easier. This CHHS process is critically dependent on the hole concentration.

From the time-resolved PL measurements discussed in Sec. IV B, the PL lifetimes measured on the QW sample did not show any change with increasing the excitation power, which apparently implies that the QW does not follow the typical density dependence of the Auger recombination. However, the hole leakage that limits the maximum hole concentration is closely related to this phenomenon. In Fig. 3(a), the integrated PL intensity initiates to quench above 50 K, while the PL decay time remains the same up to 100 K. This difference suggests that the hole concentration is kept nearly the same in spite of the hole escape at around 100 K, which keeps the Auger recombination rate nearly constant. At the higher temperature, it is probable that the energy broadening increases the Auger recombination rate as well as the thermally activated hole escape. However, from the temperature dependent time-resolve PL measurements, it is noticed that the PL decay time decreased from 600 ps at 4 K to 510 ps at 220 K, which is rather small variation in comparison to the typical reported 3–4 times drastic decrease of the Auger recombination time with temperature. This will be attributed to the limited hole concentration by the hole escape in the present situation.

V. CONCLUSIONS

Photoluminescence quenching mechanism of In\(_x\)Ga\(_{1-x}\) Sb/Al\(_{0.493}\)Ga\(_{0.507}\)Sb QWs was investigated. The In and Al concentrations as well as the QW thicknesses were precisely determined with the XRD measurements. Temperature dependent time-integrated and time-resolved PL spectroscopies resulted in the thermal activation energies of \( \sim 45 \) meV, and the overall self-consistent calculation of the band parameters based on the measured physical values confirmed that the activation energies are due to the hole escape from the QW to the barriers. The relation of the present single carrier escape mechanism with the other escape mechanisms reported with other material systems emitting in the 1.55-μm band was discussed based on the estimated band offset. The relation between the present thermal hole escape and the Auger recombination was also discussed.
ACKNOWLEDGMENTS

This work was partially supported by the Grand-in-Aid for Scientific Research (S), No. 24226007 from the Ministry of Education, Culture, Sports, Science and Technology-Japan, and SCOPE (Strategic Information and Communications R&D Promotion Programme) from the Ministry of Internal Affairs and Communications, Japan. N.A.J. acknowledges financial support via a MEXT scholarship.


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