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InAs/InAsSb type-II strained-layer superlattices for mid-infrared LEDs

J A Keen¹, D Lane¹, M Kesaria², A R J Marshall¹ and A Krier¹

¹ Physics Department, Lancaster University, Lancaster, LA1 4YB, United Kingdom
² Electronic and Electrical Engineering Department, University of Sheffield, Sheffield, S3 7HQ, United Kingdom

E-mail: j.keen@lancaster.ac.uk

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Abstract

InAs/InAsSb type-II strained-layer superlattice (SLS) and multiple quantum well (MQW) structures have been studied for their suitability in the active region of mid-infrared LEDs operating at room temperature. A series of InAs/InAs₁₋ₓSbₓ superlattices with low antimony content (x = 3.8–13.5%) were grown by MBE on InAs substrates and characterised using x-ray diffraction and photoluminescence (PL). The 4 K PL spectra of these samples exhibit the expected peak shift to longer wavelength and a reduction in intensity as the Sb content is increased. Band structure simulations highlight the effects of changing the antimony content and the layer thicknesses, to tailor the overlap of the electron and hole wavefunctions and maximise the radiative recombination rate. Analysis of the temperature dependence of the PL emission spectra enabled the extraction of quenching energies that demonstrate some suppression of Auger recombination in both the MQW and SLS structures. The MQW samples exhibit a changeover in the dominant radiative recombination process above ~100 K associated with thermal emission of holes into the InAs barriers; this behaviour was not observed in the SLS samples. These SLS structures have the potential for use as the active region in room temperature mid-infrared LEDs.

Keywords: InAs/InAsSb, superlattice, type-II, mid-infrared, LED

(Some figures may appear in colour only in the online journal)
narrowed InAs$_{1-x}$Sb$_x$ SLS structures were grown, comprising 50 periods of 14 nm InAs and 14 nm InAsSb with antimony content $x = 3.8$–13.5%, on n-InAs(100) substrates in a VG-V80H MBE system. Two multiple quantum well (MQW) InAs/InAsSb structures (40 nm InAs barrier, 10 nm InAsSb QW) containing $x_{\text{Sb}} = 3.7$, 4.3% were also grown based on our earlier work [17]. Valved cracker cells were used to provide As and Sb fluxes and a thermal effusion K-cell was used to provide the In flux. During the growth surface reconstructions were monitored by in situ reflection high energy electron diffraction (RHEED). Substrate temperature was measured using an infrared pyrometer and back-calibrated by monitoring surface reconstructions. The growth rates were calibrated by monitoring RHEED spot intensity oscillation using a photomultiplier tube. The substrate is first outgassed in the preparation chamber and oxide desorption is carried out in the growth chamber by gradually heating up to 520 °C under As flux until the weak $\times 3$ RHEED pattern transforms to the $\times 2$ pattern. The substrate temperature is lowered to 480 °C to grow an InAs buffer layer of thickness 600 nm and then further reduced to 450 °C to carry out growth of the InAs/InAsSb SLS and MQW. To obtain abrupt interfaces between InAsSb and InAs, As–Sb exchange is done by exposing the InAs surface to Sb flux for 10 s prior to InAsSb QW and SLS layer growth and before InAs barrier growth the InAsSb surface is exposed to As flux for 20 s. Antimony composition of the InAsSb layers was varied by controlling the temperature of the Sb cell, starting at 550 °C to achieve Sb content of 3.7%, and increased up to 580 °C to achieve Sb = 13.5%. Details are of the samples are given in table 1.

All the samples were characterised using high resolution x-ray diffraction (XRD) using a Bruker D8 Advance system to obtain $\omega - 2\theta$ scans. The radiation was detected using a 77 K InSb photodetector and lock-in amplifier.

### Results

**InAs/InAsSb multiple quantum wells (MQW)**

The $\omega - 2\theta$ XRD spectra obtained from the (004) x-ray rocking curves of the two MQW samples are shown in figure 1. These were peak matched with theoretical scan data simulated by RADS Mercury software. The InAs barrier and InAsSb well thickness were obtained as 40 nm and 10 nm respectively and the Sb content was determined to be 3.7% and 4.3% in Sample 1 and Sample 2 respectively, in close agreement with the target design. The strong Pendelloesung fringes are evidence of good structural layer quality and sharp growth interfaces.

The corresponding FTIR PL spectra obtained from the MQW samples obtained at temperatures in the range 4–300 K are shown in figure 2. The PL spectra are comprised of double peaks situated at wavelengths relatively close to one another which were separated by Gaussian deconvolution. At 4 K the energy separation between the two deconvoluted peaks is small, at around 3.7 meV and 5.4 meV for Sample 1 and (more clearly visible) in Sample 2 respectively which is consistent with exciton recombination in the quantum well and which is as expected approximately 4 times the exciton binding energy in the bulk semiconductor [18, 19] (assuming an exciton binding energy of 1.3 meV in bulk InAs, and is proposed to be discussed in detail in a further publication). The PL decreases in intensity as temperature is increased. The two most important non-radiative processes are SRH and Auger recombination. The rate at which SRH recombination occurs is relatively insensitive to temperature [20]. However, Auger recombination is known to be temperature dependent and follows the general relation

$$R_{\text{Auger}} \propto \exp \left( - \frac{E_a}{kT} \right) T^3$$  \hspace{1cm} (1)

where $E_a$ is the activation energy for the corresponding Auger process and the exponential term dominates [21]. The specific Auger processes each have their own activation energies, where the CHCC and CHSH processes have activation energies given by [22]
Figure 1. XRD scan comparison with simulated data for (a) Sample 1 (MQW, Sb = 3.7%) and (b) Sample 2 (MQW, Sb = 4.3%). Black line—XRD data, red line—simulation.

Figure 2. The temperature dependence of PL obtained from the two MQW samples: (a) Sample 1 containing 3.7% Sb in the MQW and (b) Sample 2 containing 4.3% Sb in the MQW. The lines are a guide to the eye, where the dotted line follows the peak from the $e_1$–$hh_1$ transition in the type-II structure, and the solid line follows the peak from recombination in the InAs layers.
$E_{\text{CHCC}} = \frac{m_e^* E_g}{m_e^* + m_{hh}^*} \quad (2)$

$E_{\text{CHSH}} = \frac{m_{SO}^*}{m_e^* + 2m_{hh}^* - m_{SO}^*} (E_g - \Delta_0) \quad (3)$

where $E_g$ is the bandgap energy, $\Delta_0$ is the spin orbit splitting energy, and $m_e^*$, $m_{hh}^*$, $m_{SO}^*$ are the effective masses of the electrons, holes in the heavy hole band and holes in the split off band respectively. It is possible to determine the dominant Auger process using,

$$\left(\frac{E_g - \Delta_0}{E_T}\right) \frac{m_{SO}^*}{m_e^*} > 1 \quad (4)$$

such that the CHCC process is dominant when this condition is satisfied. ($E_T$ is the transition energy corresponding to $e_1 - hh_1$ recombination).
The evolution of the $e_1 - hh_1$ main peak in the PL spectra of the MQW with increasing temperature shows the characteristic red-shift due to bandgap narrowing and follows closely the well-known Varshni law [23]. In both samples, above about 80 K, PL emission begins to appear from the InAs barriers originating from thermal emission of confined holes escaping from the MQW. This process continues up to room temperature where it dominates to the extent that there is no longer any observable PL emission from the MQW. This is in contrast to the behaviour in the SLS samples considered below.

**InAs/InAsSb strained-layer superlattices (SLS)**

The four SLS samples were also characterised using high-resolution XRD and the (004) x-ray rocking curves are shown in figure 3 together with the corresponding simulations. These samples contain 50 periods of InAs/InAsSb (compared with...
ten in the MQW samples) with much thinner InAs barriers—
see table 1 for details. The measured fringes are broader
compared to the simulation, but the Pendellosung fringes are
clearly visible, which indicates good layer quality and very
low Sb segregation into the InAs wells. In each case the simu-
lation gives the corresponding Sb content and the thickness of
the InAsSb QW and InAs barrier thickness as 14.0 nm. The
strain increases with Sb content and calculations using the
Matthews Blakeslee model [24, 25] revealed that the critical
thickness is just exceeded for Sample 6 containing the highest
Sb content, which means that sample this sample may contain
some dislocations. This is consistent with the reduced struc-
tural quality evident in the XRD scan where the peaks are
less well defined compared to those in the other samples with
lower Sb content.

The 4 K photoluminescence spectra of the InAs/InAs$_{1-x}$Sb$_x$
SLS structures in figure 4 show behaviour in good agreement
with previously reported PL results on similar structures
[26]. It is evident that as Sb content of the InAs$_{1-x}$Sb$_x$ layers
increases the PL emission shifts to longer wavelengths, the
intensity decreases, and the peak broadens, which is consistent
with spatially indirect transitions in type-II QW [27]. As
shown in the normalised spectra of figure 5(b) the peak inten-
sity decreases by ~13 times as Sb content is increased from
3.8% to 13.5% and the full width half maximum increases
from ~97 nm to ~270 nm.

The temperature dependent PL spectra from each of the
InAs/InAs$_{1-x}$Sb$_x$ SLS are shown together in figure 5. The
PL peak corresponding to the $e_1 - hh_1$ ground state trans-
sition is identifiable in all cases from 4 K up to 300 K. All
samples exhibit thermal broadening of ~1.1–1.7 $k_B T$. Unlike
the MQW samples, the holes remain strongly confined and the
PL spectra of the SLS samples do not display a peak corresponding
to the InAs barrier transition as temperature is increased.
However, an additional peak is observed in all of the SLS samples. This peak is of higher energy than the
$e_1 - hh_1$ peak and is of a different energy for each sample and
becomes visible above ~60 K in each case. The energy sepa-
ration between these peaks ranges from 15 meV to 26 meV,
consistent with $e_1 - hh_2$ transitions to the next confined hole
state in the quantum well (—dotted lines in figure 5). The peak
energy is dependent on the composition and is consistent with
the finite square well approximation for energy levels in the
type-II quantum wells. The PL linewidth of the SLS is very
similar to that of the MQW samples.

Discussion

The band structures of the InAs/InAsSb type-II structures
were calculated using Nextnano [26] software, assuming
rectangular quantum wells and periodic boundary conditions.
The strain in the structure is included within the program
according to the work of Krijn [27]. The program provides a
self-consistent solution of the Schrödinger, Poisson and cur-
cent equations. In order to find the quantization energies the
carriers are treated within the effective mass approximation
and the dependence of band offsets relies on a materials data-
base populated mostly by Vurgaftman [28]. However, more
recent experimental works have shown the parameters for the
InAsSb alloy to be inaccurate, and therefore we made appro-
priate modifications. Negligible bowing of the spin orbit splitting
energy was used in accordance with the work of Cripps
[30]. The bowing parameters used for the non-linear inter-
polation of the conduction and valence band energies were $C_{CB} = +0.65$ eV and $C_{VB} = -0.98$ eV based on a non-zero
value of $C_{VB}$ with the bowing split between CB and VB at
40:60 ratio, as suggested by Liu [17]. The overall bowing
parameter of 1.63 eV for InAsSb provides a best fit to our
experimental PL results, and is larger than the highest value of
0.938 eV previously reported by Webster [29]. The agreement
of the experimental 4 K PL data from figure 5 with the simu-
lation is shown in figure 6.

Figure 7 shows for example a comparison between the
MQW and SLS structures of similar Sb composition. The
effect of reducing the QW separation is to raise the energy of
the $e_1$ level from 1.941 eV to 1.953 eV and similarly the energy of
the $hh_1$ level is increased from 1.579 eV to 1.583 eV.
In each case the calculated transition energies are in good
agreement with the PL transitions observed at 4 K.

Considering the electron and hole wavefunctions (figure 8)
the heavy hole wavefunction is strongly localised within the
InAsSb QW regions in both the MQW and SLS structures, how-
ever in the SLS structure of thinner layers the electron wave-
function spreads out through the structure with significant
probability of residing in the QW regions. Since the overlap
of the electron and hole wavefunctions is directly related to
the radiative recombination rate it is expected that with shorter
periods the increased wavefunction overlap results in a corre-
sponding increase in PL emission intensity. Increased anti-
mongy content increases the type II behaviour resulting in the
opposite effect of reducing wavefunction overlap and is there-
fore detrimental to PL emission intensity.

The wavefunction overlap is proportional to the matrix ele-
ment $M$, which can be used as a figure of merit to compare

![Figure 6. Dependence of the 4 K PL peak energies on Sb content for the InAs/InAsSb SLS structures. The best agreement with the experimental data (black circles) was obtained with an InAsSb bowing parameter of 1.63 eV split at the ratio of 40:60 between the conduction and valence bands (red line). The dotted line represents the previously highest InAsSb bowing parameter of 0.938 eV reported by Webster [29], also split at a 40:60 ratio.](image-url)
different structures. As light propagates through the QW structure, photons are emitted by electrons of energy $E_i$ in an initial state $|i\rangle$ in the conduction band recombining with holes to a final state $|f\rangle$ of energy $E_f$ in the valence band.

The matrix element for this transition is defined as:

$$M = f |x|i = \int \psi^*_f(r)\psi_i(r)d^3r$$  \hspace{1cm} (5)

which can be separated into two terms:

$$M = M_{cv}M_{nv'}$$  \hspace{1cm} (6)

where $M_{cv}$ is the valence-conduction band dipole moment:

$$M_{cv} = u_e |x|i,\hspace{1cm} (7)$$

and $M_{nv'}$ is the electron–hole overlap:

$$M_{nv'} = e n' |n|h = \int_{-\infty}^{\infty} \psi^*_e(z)\psi_h(z)dz.$$  \hspace{1cm} (8)

Since electric dipole transitions between the conduction and valence bands are strongly allowed then it can be assumed that $M_{cv}$ is non-zero, hence the matrix element $M$ for optical transitions is proportional to the overlap of the electron and hole states [32].

Considering the ground state transition, electrons in the $n' = 1$ state in the conduction band recombine with holes into the $n = 1$ state in the valence band. Furthermore, the wavefunction overlap can be considered for a single period of the periodic SL structure spanning from $z = -\frac{P}{2}$ to $z = +\frac{P}{2}$ containing a single QW region. Therefore, the expression for the electron–hole wavefunction overlap can be simplified:

$$M_P = e |h| = \int_{-\frac{P}{2}}^{+\frac{P}{2}} \psi^*_e(z)\psi_h(z)dz.$$  \hspace{1cm} (9)

The wavefunction overlap was calculated for each structure and decreases with increasing Sb in both the MQW and SLS structures, giving a reduction of $\sim 2.1\%$ in the SLS and a decrease of $\sim 1.7\%$ in the MQW for a change of $1\%$ of the antimony content in the InAsSb well in each case. The radiative recombination rate is proportional to the matrix element squared $M_P^2$ which was calculated for each sample and the values are given in table 2. The reduction of $13 \times$ in the experimental PL spectra of the SLS samples as the antimony content increases in the QW layers (figure 1) is more than the calculated approx. $5.4 \times$ decrease in $M_P^2$ shown in the table. Consequently, we attribute the remaining reduction to non-radiative recombination mechanisms which are dominated by Auger processes.

The calculated ground state transition energy $(e_1 - \hbar h_1)$ and the split off energy $(E_T - \Delta_{SO})$ used to determine the dominant Auger process in the MQW and SLS structures as well as the corresponding activation energies are given in tables 2 and 3. In order to consider the non-radiative processes the principal Auger activation energies were determined using the transition energies and spin orbit split-off energies calculated using Nextnano. The values obtained are given in table 3 alongside the experimentally determined activation energies obtained from Arrhenius plots for each sample. Increasing the spin orbit split off energy such that it becomes larger than the band gap ($\Delta_{SO} > E_T$) suppresses the CHSH process [32]. This excess is larger in the SLS samples than in the MQW samples, so CHSH Auger recombination is more suppressed in the SLS samples than in the MQW samples of similar antimony composition and CHCC dominates instead. For the SLS samples the condition given by equation (4) is satisfied for all samples except Sample 3, which has the lowest antimony content. Hence, CHCC Auger recombination is the dominant recombination process for the SLS samples and the calculated activation energy for CHCC is found to decrease with antimony content. The corresponding experimental values have uncertainties that arise due to CO absorption which complicates Gaussian deconvolutions at high temperatures. However, the general trend is that activation energy decreases with increasing antimony content. Note that the sample which appears to have larger activation energy with increased antimony content (Sample 6) has a thickness that exceeds the critical thickness for this structure. It is therefore reasonable to say that the overall trend of the results is consistent with the calculated CHCC Auger activation energies which dominate the transitions.
for the SLS samples. Meanwhile, for the MQW samples, the calculated CHSH Auger activation energy in the higher antimony sample (Sample 2) is larger, consistent with CHSH being suppressed due to $E_T \sim \Delta_0$ detuning, which is strongly dependent on antimony content. The experimentally determined activation energy is larger than both the calculated activation energies for the CHCC and CHSH processes, which indicates that Auger recombination is reduced in these type-II MQW structures. The experimental activation energies for the SLS samples are also significantly higher than those calculated for CHCC based on Nextnano. However, the calculated activation energies do not include Coulombic effects or band bending and more detailed calculations which account for the difference in $e-h$ overlap and the corresponding radiative recombination are required to properly reconcile the calculated and experimental values. Nevertheless, we observed clear differences in the temperature dependent PL spectra which show that the holes remain better confined at higher temperatures in the SLS than in the MQW. We attribute this to Coulombic attraction arising from the increased $e-h$ overlap in the SLS because of the thinner InAs barriers between the InAsSb wells.

**Conclusion**

High quality InAs/InAs$_{1-x}$Sb$_x$ ($x = 3.7-13.5$) type-II MQW and SLS structures on InAs substrates have been fabricated by MBE and investigated using XRD and PL spectroscopy as the basis for the active region of MIR LEDs operating at room temperature. The 4 K PL spectra of these samples exhibit the expected peak shift to longer wavelength and a

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**Table 2.** Calculated and experimental values for the MQW and SLS structures at 4 K.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Sb (%)</th>
<th>Exp. 4 K PL peak energy (eV)</th>
<th>Calculated $e_1 - hh_1$ transition at 4 K (eV)</th>
<th>$M_1^2$</th>
<th>$E_T - \Delta_{SO}$ (meV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (MQW)</td>
<td>3.7</td>
<td>0.376</td>
<td>0.377</td>
<td>59</td>
<td>39</td>
</tr>
<tr>
<td>2 (MQW)</td>
<td>4.3</td>
<td>0.367</td>
<td>0.369</td>
<td>45</td>
<td>54</td>
</tr>
<tr>
<td>3 (SLS)</td>
<td>3.8</td>
<td>0.367</td>
<td>0.369</td>
<td>1225</td>
<td>46</td>
</tr>
<tr>
<td>4 (SLS)</td>
<td>6.2</td>
<td>0.318</td>
<td>0.335</td>
<td>778</td>
<td>117</td>
</tr>
<tr>
<td>5 (SLS)</td>
<td>9.5</td>
<td>0.279</td>
<td>0.287</td>
<td>428</td>
<td>180</td>
</tr>
<tr>
<td>6 (SLS)</td>
<td>13.5</td>
<td>0.242</td>
<td>0.231</td>
<td>228</td>
<td>246</td>
</tr>
</tbody>
</table>

*This sample exceeds the critical layer thickness limit calculated by Mathews Blakeslee model [24, 25].

**Table 3.** A comparison of the experimentally determined activation energies and the calculated values of the main Auger recombination processes. Activation energies for Auger recombination mechanisms calculated using Nextnano compared with experimentally determined values from PL.

| Sample | Sb (%) | $|E_T - \Delta|$ (meV) | $\frac{|E_T - \Delta|}{E_T - \Delta}$ | $E_{CHCC}^a$ (meV) | $E_{CHSH}^a$ (meV) | $E_a^{expt.}$ (meV) |
|--------|--------|-------------------------|----------------------------------|------------------|------------------|------------------|
| 1 (MQW) | 3.7   | 39                      | 0.7                              | 20               | 9               | 33 ± 3           |
| 2 (MQW) | 4.3   | 54                      | 1.0                              | 19               | 13              | 46 ± 4           |
| 3 (SLS) | 3.8   | 46                      | 0.9                              | 19               | 11              | 32 ± 3           |
| 4 (SLS) | 6.2   | 117                     | 2.5                              | 17               | 27              | 28 ± 4           |
| 5 (SLS) | 9.5   | 180                     | 4.5                              | 15               | 42              | 22 ± 4           |
| 6 (SLS)  | 13.5  | 246                     | 7.1                              | 13               | 58              | 24 ± 5           |

*This sample exceeds the critical layer thickness limit.
reduction in intensity as the Sb content is increased. Band structure simulations highlight the effects of changing the structure, specifically the antimony content and the layer thicknesses, to tailor the overlap of the electron and hole wavefunctions to maximise the radiative recombination rate. Analysis of the PL data along with Nextnano modelling of the structures enabled a comparison of the experimentally derived activation energies with calculated activation energies for the characteristic non-radiative Auger processes and $e^{-}h$ overlaps. The dominant Auger process was determined to be CHCC in the SLS structures and CHSH in the MQW structures. In the SLS structures the activation energies follow a downward trend with increasing antimony content, whereas the MQW exhibited the opposite behaviour. In both cases the experimentally activation energies are larger than the calculated values, indicating some degree of Auger suppression. PL studies revealed the desired InAs to InAsSb ground state transition exists up to room temperature in the SLS structures, but not in the MQW which exhibit increasingly InAs bulk-like behaviour above ~100 K. This is attributed to an increased $e^{-}h$ overlap and a larger Coulomb attraction which keeps the holes better confined in the SLS, thus preserving the transition up to high temperatures. We consider that the SLS structures are therefore an excellent prospect for room temperature LEDs.

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ORCID iDs

J A Keen https://orcid.org/0000-0002-7065-7871
D Lane https://orcid.org/0000-0001-6647-2869
M Kesaria https://orcid.org/0000-0003-1664-0806

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