



## Enhanced hole mobility and density in GaSb quantum wells

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### ABSTRACT

Modulation-doped quantum wells (QWs) of GaSb clad by AlAsSb were grown by molecular beam epitaxy on InP substrates. By virtue of quantum confinement and compressive strain of the GaSb, the heavy- and light-hole valence bands in the well are split and the hole mobility is thereby significantly enhanced. Room-temperature Hall mobilities as high as 1200–1500 cm<sup>2</sup>/V s were achieved for 5–10 nm QWs and biaxial strains of 1–3%. This contrasts with earlier work on GaSb/AlGaAsSb QWs on GaAs substrates in which the mobilities were found to fall off above 1% strain. Moreover, unlike in comparable InGaSb and InSb QWs, the high mobilities were maintained out to sheet densities of  $3.5 \times 10^{12}$ /cm<sup>2</sup>. As a result, the sheet resistivities observed in the GaSb/AlAsSb wells reached record levels as low as 1500 Ω/□. Modeling indicates that this performance gain is due to the larger valence band offset of the GaSb QWs and the consequent reduction in scattering because of the better confinement and the lower doping levels needed for a given sheet charge.

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### 1. Introduction

Narrow bandgap compound semiconductors exhibit high electron mobilities and peak velocities. The high velocities are reached at relatively low electric fields, enabling analog electronic devices with extremely low power consumption. For example, low-noise amplifiers (LNAs) using high-electron-mobility transistors (HEMTs) with InAs channels and AlSb barriers operate at substantially lower power than similar circuits based upon GaAs or Si [1]. Recently, there has been interest in the potential of III–V transistors for advanced logic applications which could enhance digital circuit functionality and extend Moore's law [2]. For these applications, a key to low power operation is the ability to make complementary circuits. In III–V materials, one challenge centers on maximizing the hole mobility in p-channel field-effect transistors (FETs) [3–5].

Like in Si and Ge, the hole mobilities of III–V compounds are generally low and, for this reason, researchers have exploited quantum confinement and strain as a means of splitting the heavy-hole and light-hole valence bands and giving lower in-plane hole masses and enhanced low-field mobility [3–14]. The antimonides offer the highest hole mobilities of the III–V compounds, with room-temperature values approaching 1000 cm<sup>2</sup>/V s for bulk GaSb and InSb. We previously demonstrated compressively-strained high-mobility p-type InGaSb QWs confined by AlGaSb barrier layers [7]. In this work, we investigate compressively-strained GaSb QWs confined by AlAsSb barrier layers [6,11,15]. Compared to

the InGaSb QW, the GaSb QW has a larger valence band offset and bandgap, and may also be advantageous in being a binary and so not subject to alloy scattering in the well. The structures are grown on metamorphic AlAsSb buffer layers on InP substrates. This combination is found to yield room-temperature mobilities as high as 1500 cm<sup>2</sup>/V s and sheet resistivities as low as 1500 Ω/□, with the latter value a factor of two lower than the best previously achieved for III–V p-type quantum wells.

### 2. Experimental procedures

The heterostructures studied here are grown by molecular beam epitaxy (MBE) on semi-insulating (001) InP substrates using a Riber Compact 21T MBE system. A cross-section is shown in Fig. 1a. The first layer is 160 nm of In<sub>0.52</sub>Al<sub>0.48</sub>As lattice matched to InP, followed by a ~1.3 μm buffer layer of AlAs<sub>x</sub>Sb<sub>1-x</sub>, an optional 4 nm Be-doped AlAs<sub>x</sub>Sb<sub>1-x</sub> donor layer, a 9.5 nm AlAs<sub>x</sub>Sb<sub>1-x</sub> spacer, a 5–10 nm GaSb QW, a 10.5 nm AlAs<sub>x</sub>Sb<sub>1-x</sub> barrier, a 4 nm In<sub>0.2</sub>Al<sub>0.8</sub>Sb etch-stop, and a 2 nm InAs cap. The AlAs<sub>x</sub>Sb<sub>1-x</sub> layers were grown as a digital alloy by toggling the As and Sb shutters while the Al shutter and the As and Sb valves remained open, allowing better control of composition compared to random alloys [16,17]. The AlAs mole fraction was adjusted by changing the length of time the As shutter is open relative to the Sb shutter. As shown in the calculated band-structure, Fig. 1b, there is a large valence band discontinuity between the AlAsSb and the GaSb, providing for good confinement of holes in the GaSb. The AlAs<sub>x</sub>Sb<sub>1-x</sub> is almost fully relaxed; its composition dictates the amount of biaxial

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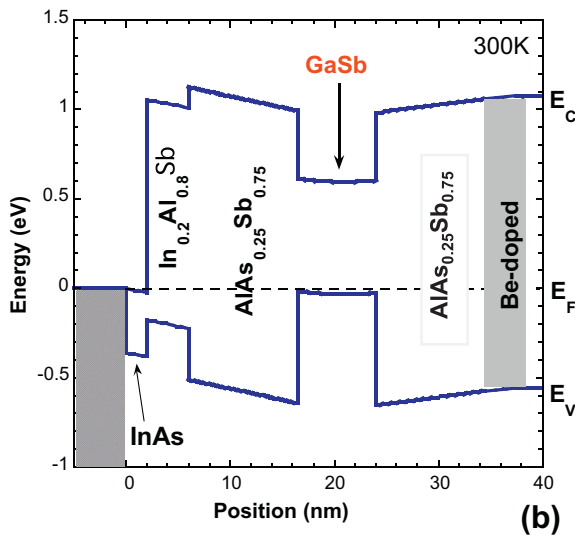
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InAs	2 nm
In <sub>0.2</sub> Al <sub>0.8</sub> Sb	4 nm
AlAs <sub>x</sub> Sb <sub>1-x</sub>	10.5 nm
GaSb	5–10 nm
AlAs <sub>x</sub> Sb <sub>1-x</sub>	9.5 nm
AlAs <sub>x</sub> Sb <sub>1-x</sub> (Be)	4 nm
AlAs <sub>x</sub> Sb <sub>1-x</sub>	1.3 μm
In <sub>0.52</sub> Al <sub>0.48</sub> As	0.16 μm
SI InP (001)	substrate

**(a)**



**Fig. 1.** (a) Cross-section of GaSb/AlAsSb QW heterostructures. The AlAs<sub>x</sub>Sb<sub>1-x</sub> layers are composed of AlSb/AlAs short-period superlattices. (b) Calculated band-structure for a GaSb/AlAs<sub>0.25</sub>Sb<sub>0.75</sub> QW with a hole sheet density of  $1.0 \times 10^{12}/\text{cm}^2$ .

strain in the GaSb QW. For a buffer layer of AlAs<sub>x</sub>Sb<sub>1-x</sub>, the in-plane biaxial strain associated with the lattice mismatch between the well and buffer layer is given by:

$$\varepsilon = \frac{6.0954}{6.1355 - 0.4736x} - 1 \quad (1)$$

The MBE growth temperature is near 450 °C for the InAlAs. The temperature is then raised to 520 °C for the AlAs<sub>x</sub>Sb<sub>1-x</sub> buffer layer, and then reduced back to 450 °C for the GaSb QW and subsequent layers. The GaSb QW and InAs cap were grown at a rate of 0.2 monolayers (ML)/s, and other layers were grown near 1.0 ML/s as calibrated from reflection high-energy electron diffraction oscillations.

Based upon previous results on this MBE system, we expect the layer thicknesses to be uniform to within 1% across the 76-mm-diameter substrate if the wafer is rotated. Table 1 includes relevant parameters for the nine MBE growths in this study. For six growths (#1–5 and 8), the wafers were rotated during all the layers except the AlAs<sub>x</sub>Sb<sub>1-x</sub> buffer layer. This resulted in AlAs mole fractions (and hence GaSb strain) which varied by more than a factor of two across the wafer. Several  $5 \times 5$  mm squares were then cleaved from different locations on each wafer to provide a range of samples with varying strain in a highly efficient manner. For three

**Table 1**

Growth parameters for the samples in this study. Resistivity values are from contactless wafer mapping, and are not well-calibrated above 15,000 Ω/□.

Growth#	QW thickness (nm)	Dopant	Substrate	Rotate-buffer	Rotate-doping	Resistivity Ω/□
1	10	None	InP	N	Y	7100–18,000
2	7.5	None	InP	N	Y	7700–17,000
3	5	None	InP	N	Y	6500–27,000
4	7.5	Be	InP	N	Y	2100–4200
5	7.5	Be	GaAs	N	Y	2400–6700
6	7.5	Be	InP	Y	N	2200–4500
7	7.5	Be + Be	InP	Y	N	1800–3900
8	7.5	Si	InP	N	Y	4100–>50,000
9	7.5	Si	InP	Y	N	10,000–>50,000

growths (#6, 7, and 9), the wafers were rotated for all layers except the 4 nm layer(s) of AlAs<sub>x</sub>Sb<sub>1-x</sub> doped with Be or Si. This resulted in a gradient in the dopant concentration and hence hole sheet density across the wafers.

Hall/van der Pauw transport measurements were performed on a total of 79 samples at 300 K, using magnetic fields of 0.37, 0.55 and 1.0 T. Measurements were performed at two or more current levels at each B field, and average values are given in this paper with standard deviations usually less than 5%. Room-temperature, 55-point resistivity maps were generated for each wafer from eddy-current measurements using a contactless Leighton 1500 system [18]. Atomic force microscopy (AFM) measurements were performed on selected samples to yield root-mean-square (rms) roughness over  $5 \times 5$  μm regions. X-ray diffraction (XRD) measurements were made on a double-crystal system using Cu-Kα radiation and compared to simulations using dynamical diffraction theory.

### 3. Results and discussion

In Fig. 2, we show the resistivity map for growth #4, a Be-doped sample with a 7.5 nm GaSb QW, grown on an InP substrate. The resistivity varied from 2060 to 4240 Ω/□. A 5-mm wide strip was cleaved as indicated, yielding twelve  $5 \times 5$  mm samples for characterization. The transport results will be discussed later.

AFM data was collected on several samples with a variety of strains and hole densities. The rms roughness was between 0.5 and 1.0 nm, with no clear correlation to mobility or other parameters (substrate, QW thickness, doping). These values are as good or better than what was measured for InAs-channel HEMT structures on Al(Ga)Sb buffer layers and GaAs or InP substrates and are fully compatible with monolithic-microwave integrated circuit (MMIC) processing [19,20].

In Fig. 3, we show the XRD scan for sample 4L, the piece labeled L in Fig. 2. The buffer layer consisted of 888 periods of (4.0 s AlSb/1.0 s AlAs). Peaks are visible for the InP substrate and the short-period superlattice ( $n = -2, -1, 0,$  and  $+1$ ). A simulation was generated by adjusting the superlattice thicknesses to match the experimental peak positions, and is shown below the experimental data in Fig. 3. The layer thicknesses were 0.33 nm AlAs and 1.30 nm AlSb, yielding a period of 1.63 nm. The lattice relaxation for the superlattice was assumed to be 100%. Based upon the nominal

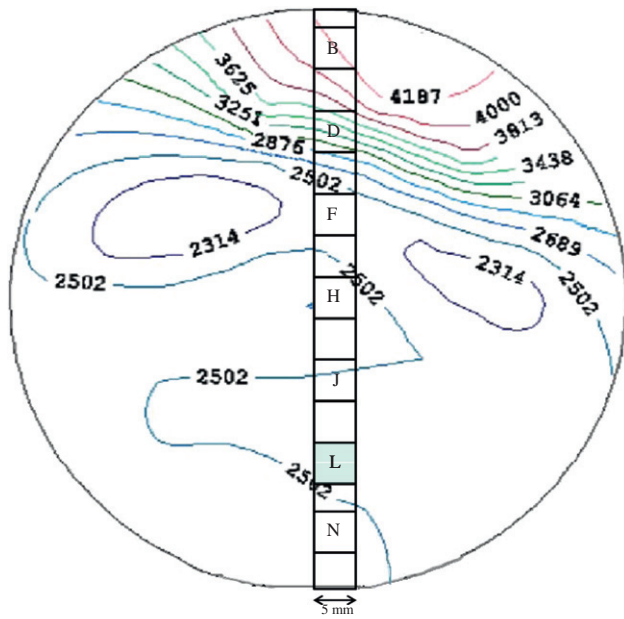


Fig. 2. Resistivity map in units of  $\Omega/\text{sq}$  for growth #4. The gradient across the 76-mm wafer is a result of varying strain and mobility in the GaSb QW.

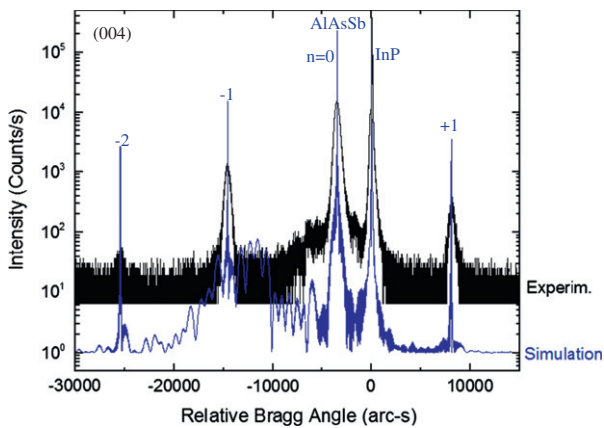


Fig. 3. Double-crystal X-ray diffraction data for sample 4L (upper curve) and simulation (lower curve). AlSb and AlAs thicknesses in the buffer layer short-period superlattice were varied to obtain a good fit to the experimental data.

growth rates, we expect a period of 1.50 nm in the center of wafer #4; L is in a region of higher Al flux than the center. The buffer layer thickness is  $1.45 \mu\text{m}$  ( $888 \times 1.63 \text{ nm}$ ). The epilayer peaks are all broadened compared to the simulation. For example, the full-width at half-maximum for the  $n = 0$  superlattice peak is 520 arc-s. This is a result of a high density of misfit dislocations required to relax the lattice mismatch. Using the superlattice layer thicknesses and Vegard's law, we calculate the effective ternary composition to be  $\text{AlAs}_{0.202}\text{Sb}_{0.798}$ . Using Eq. (1), the compressive biaxial strain in the GaSb is 0.92%. Samples with higher AlAs mole fractions generally exhibited weaker satellite peaks. This indicates that the superlattices are not as well defined but does not imply lower crystalline quality.

MBE growths #1–3 did not include a doping layer. The growths were nominally identical except for the thickness of the channel layer. There was no rotation during the growth of the AlAsSb buffer layer. All the samples from these three growths were p-type, with hole sheet concentrations of  $1.0 \pm 0.3 \times 10^{12} \text{ cm}^{-2}$ ; these results

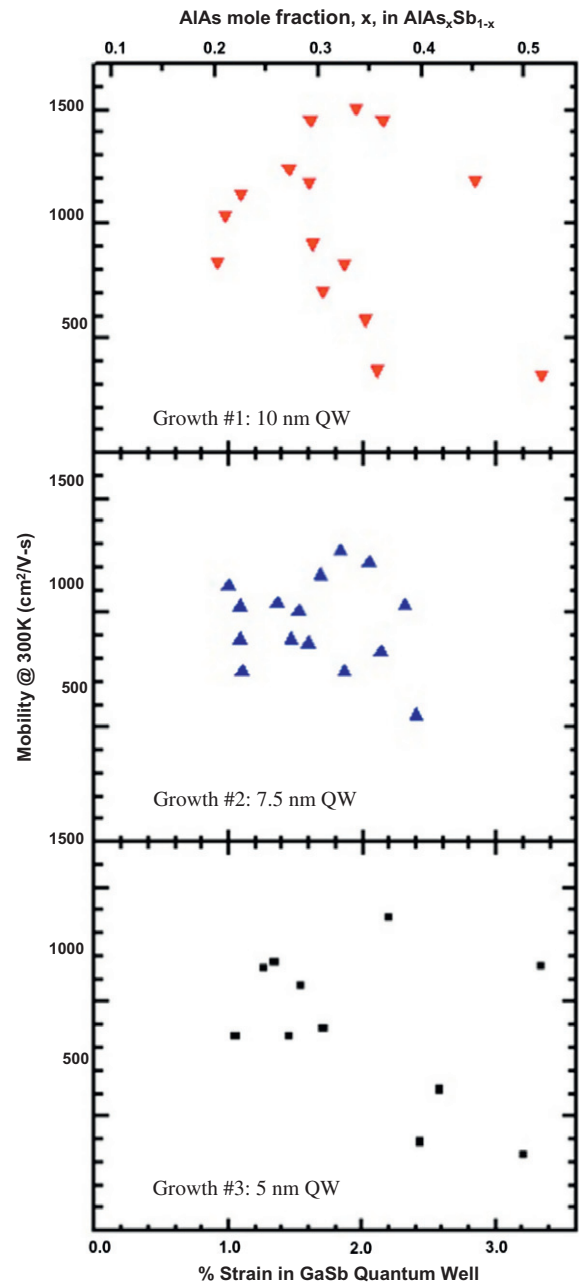
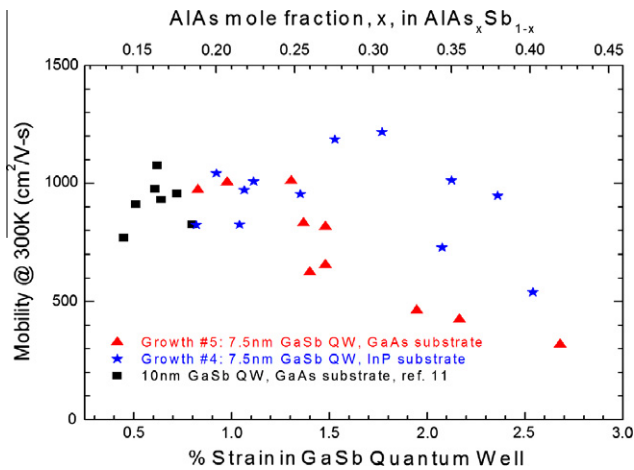


Fig. 4. Room-temperature mobility as a function of GaSb QW strain (or buffer layer composition) for growths #1–3, differing only in QW thickness. The samples were not intentionally doped.

are consistent with previous results for undoped GaSb and InGaSb quantum wells (QWs) [6,7,9]. In Fig. 4, we plot the measured room-temperature mobility as a function of strain for (a) growth #1, 10 nm QW, (b) growth #2, 7.5 nm QW, and (c) growth #3, 5 nm QW. For each growth, we observe considerable strain-induced mobility enhancement compared to unstrained GaSb QWs ( $\mu = 200\text{--}400 \text{ cm}^2/\text{V s}$ , see Refs. [6,21,22]). For growth #1, we observe an increase in mobility as strain increases from 1.0 to 1.5%. For higher strains, there are substantial variations in the mobility. One sample with a strain of 2.0% has a mobility of  $1500 \text{ cm}^2/\text{V s}$ . This is a record value for GaSb and matches the highest values for InGaSb [7]—the largest of any III–V material at room temperature. The three samples with mobilities of  $1400\text{--}1500 \text{ cm}^2/\text{V s}$  and strains of 1.6–2.1% were adjacent to one another on the wafer. The four samples in the same strain range and mobilities of

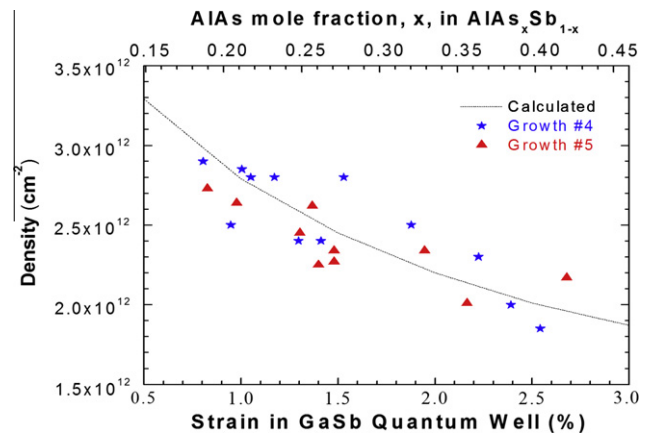


**Fig. 5.** Room-temperature mobility as a function of GaSb QW strain (or buffer layer composition) for samples with Be modulation doping. Note the difference between InP and GaAs substrates.

400–800  $\text{cm}^2/\text{V s}$  were from a different area of the wafer. Similar trends were observed for growths #2 and 3. This suggests that one or more parameters other than strain are important. Possibilities include buffer layer growth rate and thickness, V:III flux ratios, and growth temperature. The XRD measurements allowed us to determine the buffer layer growth rate and thickness for each sample. There was not a clear correlation with mobility. We expect significant variations in flux ratios and substrate temperature across the wafer during the buffer layer growth when it is not rotating, but cannot measure these values across the wafer. There is an apparent correlation between the sheet density and mobility for samples with strain near 2% on all three growths, with the high-mobility samples having densities about 20% lower than the low-mobility samples. Although higher density could cause lower mobility, it seems much more likely that the two effects arise from a common underlying mechanism (e.g. defects). Given the fact that high-mobility samples were found for each of the three growths for strains near 2%, we expect that if this strain regime was targeted and growths were performed with substrate rotation throughout, one could identify growth conditions that ensure consistently high mobilities.

Growth #4 was identical to #2 (7.5 nm QW) except that Be doping was added to achieve higher sheet concentrations. The results are plotted as stars in Fig. 5. The average sheet density was  $2.5 \times 10^{12} \text{ cm}^{-2}$ , with the particular values clearly correlating with strain and ranging from  $2.9 \times 10^{12} \text{ cm}^{-2}$  for 0.8% strain to  $1.9 \times 10^{12} \text{ cm}^{-2}$  for 2.5% strain. The transport data is consistent with the resistivity data in Fig. 2. It is important to point out that the results for growths #1–4 contrast with earlier work from our group as well as that of Tokranov et al. [11] in not showing a consistent degradation in GaSb mobility for higher strains. Our earlier work [6] (not shown here for clarity) found a drop above  $\sim 1\%$  strain, while Tokranov observed the mobility to peak at about 0.6% strain as shown in Fig. 5. A possible explanation for this discrepancy is that both sets of earlier studies used GaAs substrates, whereas the new experiments used InP substrates. To confirm this, we grew a wafer (#5) identical to #4 except that the substrate was GaAs and the InAlAs buffer layer was replaced by epitaxial GaAs. The results, plotted as triangles in Fig. 5, show decreasing mobility for strains above 1.3% in rough agreement with the earlier experiments.

To aid in understanding the results of this paper, we have performed a set of  $8 \times 8 \text{ k.p}$  calculations using the nextnano program [23]. We used the heterostructure shown in Fig. 1 with a 7.5 nm



**Fig. 6.** Channel density as a function of QW strain for samples from wafers #4 and 5. Solid line represents calculated values.

QW to match growths #4 and 5. The background density was set at  $7 \times 10^{11} \text{ cm}^{-2}$  and the Be dopant concentration was adjusted to yield  $2.2 \times 10^{12} \text{ cm}^{-2}$  at a 2% compressive strain for the GaSb. In Fig. 6, we plot the channel density as a function of QW strain. We also include the data points from wafers #4 and #5. The good agreement between the calculated and experimental data indicates that the drop in density is due to a reduced transfer of charge from the doped layer caused by the rise in the (hole) energy of the valence band states in the QW induced by the increasing compressive strain. This agreement also suggests that we understand our system both from experimental and theoretical viewpoints. With respect to the former, it is interesting to note that it was the use of multiple samples from a single MBE growth that made this comparison possible. Had these been separate growths, the unintentional day-to-day variations in MBE conditions would have made it difficult to observe such small density changes.

The poorer quality GaSb seen at high strains when on a GaAs as opposed to an InP substrate could certainly result from the former substrate having a higher defect density. However, the fact that high mobilities were previously obtained for strains up to 2% for InGaSb [7] and InSb QWs [10] despite using GaAs substrates suggests that the trouble is not with the substrate per se. Instead we speculate that the substrate dependence for GaSb QWs arises from defects in the AlAsSb buffer layer. For layers above the thermodynamic critical layer thickness, the degree of lattice relaxation may depend on the particular defects in the layer on which the strained layer is grown [24–26]. If there is a higher density (or a different type) of dislocations for AlAsSb on GaAs compared to AlAsSb on InP, then the GaSb films could relax more in the former case for equivalent strains. In the simplest case, if the misfit dislocation types were the same for fully relaxed AlAsSb buffer layers grown on InP and GaAs, then the layers grown on GaAs would have a factor of two higher misfit dislocation density, reflecting the factor of two larger mismatch between the buffer layer and substrate. This could lead to a peak in mobility at lower strains for growth on GaAs. In the case of InSb and InGaSb QWs, their use of different buffers layers (InAlSb and AlGaSb, respectively) means their dependence on the substrate can be entirely different.

In Fig. 7, we plot hole mobility as a function of sheet concentration for InSb [10] and InGaSb [7] QWs. The hole mobilities for both channel materials decrease as carrier density increases. To explore this relationship for GaSb/AlAsSb/InP heterostructures, we grew two wafers, #6 and #7, in which the samples were rotated during all layers except the AlAsSb(Be) doping layer(s). As discussed earlier, this should yield a gradient in Be concentration across the wafer and hence allow a range of hole densities to be investigated

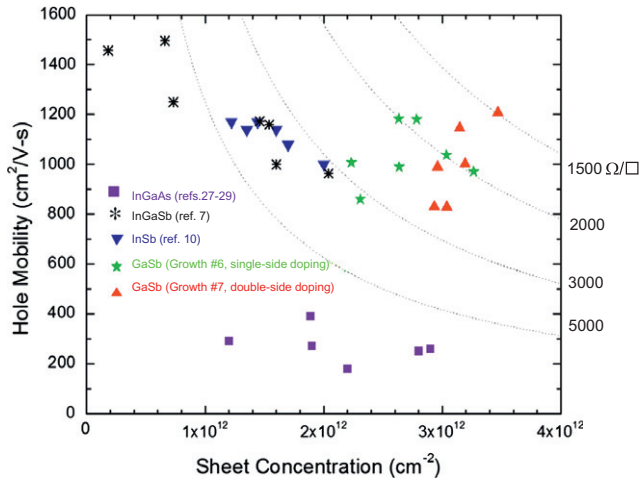


Fig. 7. Room-temperature hole mobility as a function of sheet concentration for growths #6 and 7 as well as data from the literature for InGaSb, InSb, and InGaAs.

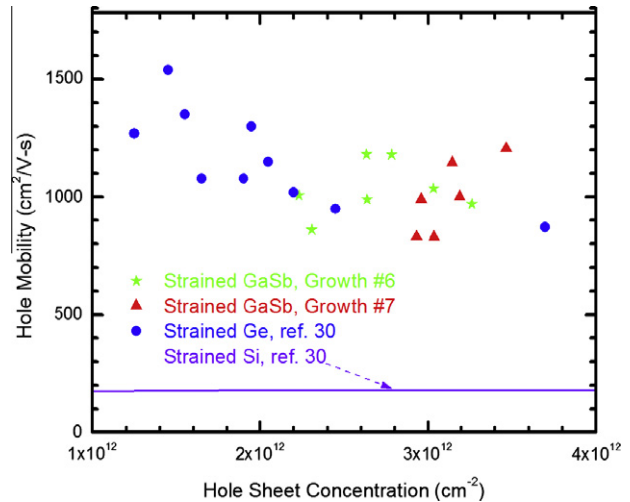


Fig. 8. Room-temperature hole mobility as a function of sheet concentration for growths #6 and 7 as well as data from the literature for strained Ge and strained Si.

with a single growth. A buffer layer composition of  $\text{AlAs}_{0.29}\text{Sb}_{0.71}$ , corresponding to a GaSb strain of 1.7% was chosen. The GaSb thickness was 7.5 nm. The Be cell temperature (879 °C) was the same as in wafers #4 and #5. The data for wafer #6, with single-sided Be doping as shown in Fig. 1, are plotted as stars in Fig. 7. The sheet densities vary by a factor of 1.5 across the wafer. Wafer #7 had double-side doping, with 4 nm layers of  $\text{AlAsSb}(\text{Be})$  on both sides of the QW and 5.4 nm spacer layers. The Be temperature was again 879 °C. The results, shown as up-triangles in Fig. 7, show less variation in density across the wafer than for #6. Simulations discussed below show that this is likely due to a saturation in the ability of the modulation doping to transfer charge. Most interesting is the fact that, unlike InSb and InGaSb QWs, the mobility of GaSb QWs does not degrade as the density increases over the range studied. As a result, we observe sheet resistivities to drop as low as  $1500 \Omega/\square$ , a factor of two improvement over the best values for InSb and InGaSb, and much lower than other III–Vs. Our previous results for GaSb QWs on GaAs substrates are not shown on Fig. 7 for clarity, but the lowest resistivity achieved was  $3100 \Omega/\square$  [6]. We also include data from the literature for p-InGaAs QWs. The lower mobilities result in resistivities greater than  $7000 \Omega/\square$  for InGaAs [27–29].

It is also instructive to compare our p-channel GaSb results to strained Si, currently used for CMOS, and strained Ge, proposed for future CMOS nodes. Fig. 8 includes strained Si data, recent strained Ge data from Intel [30] plus our GaSb data. The GaSb QWs, like Ge QWs, offer substantially higher mobilities and lower sheet resistivities than strained Si. Based upon results from gated hall structures on InGaSb QWs, we expect the GaSb advantage over Si to be maintained at higher carrier densities [31].

Dopants other than Be may be feasible for antimonide p-FET structures. Liao and Cheng investigated carbon doping in InGaSb/AlGaSb QWs [9]. They achieved controllable sheet densities ranging from  $0.7$  to  $4.1 \times 10^{12}/\text{cm}^2$ . Silicon is an amphoteric dopant in III–V compounds. It is generally n-type for arsenides and InSb, but p-type for GaSb and AlSb [32]. Growth #8 was the same as #4 except that Be was replaced by Si. The resistivity map was very non-uniform, with values ranging from  $4100$  to  $67,000 \Omega/\square$ . Transport measurements in the low-resistivity region yielded a mobility of  $430 \text{ cm}^2/\text{V s}$  and a sheet density of  $3.4 \times 10^{12}/\text{cm}^2$ . The results in a more resistive region were:  $230 \text{ cm}^2/\text{V s}$  and  $0.8 \times 10^{12}/\text{cm}^2$ . For growth #9, we changed the dopant layer from AlAsSb to AlSb to avoid electrons from Si dopant atoms in the AlAs. We also rotated the sample during the buffer layer, yielding uniform QW strain

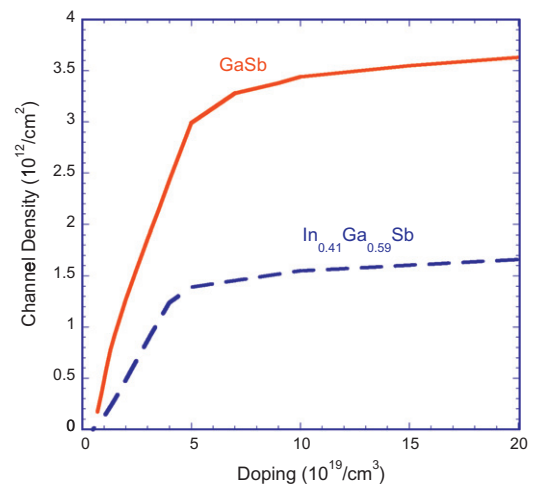
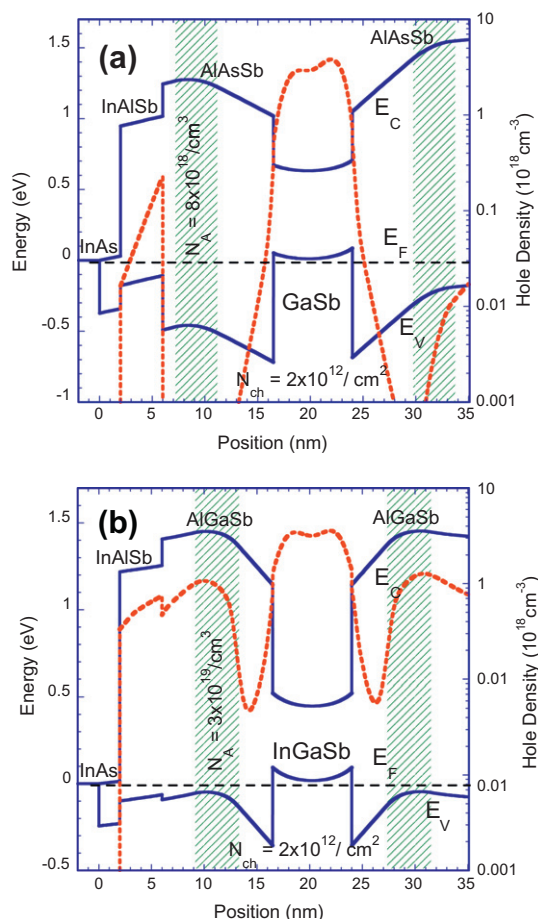


Fig. 9. Calculated hole density in channel as a function of total doping for GaSb and  $\text{In}_{0.41}\text{Ga}_{0.59}\text{Sb}$ .

across the sample, and did not rotate during the AlSb(Si) layer, yielding a gradient in Si concentration. The resistivity map was again very nonuniform, but all values were above  $10,000 \Omega/\square$ . For both growths, extremely large anisotropies were observed in the transport measurements, in contrast to the Be-doped samples. It is clear that Si was creating p-type modulation doping in portions of the wafers, but we do not understand the low mobilities or large anisotropies and non-uniformities. In contrast, Be doping in this study and in our previous work on InGaSb- and GaSb-channel pFETs was always well behaved. We reached similar conclusions when comparing Be and Si as dopants for thick layers of AlSb [33].

To investigate the improved high-density performance of GaSb QWs, we have performed additional  $8 \times 8 \text{ k.p}$  calculations in which we compare GaSb/AlAsSb QW devices with  $\text{In}_{0.4}\text{Ga}_{0.6}\text{Sb}/\text{AlGaSb}$  QW devices. In Fig. 9 we illustrate the effectiveness of doping in the two devices by plotting the channel charge vs. doping concentration for double-sided doping (as in wafer #7) with 5.4 nm spacers and 4 nm doped layers. As the figure shows (and as mentioned



**Fig. 10.** Calculated band structure and hole density for (a) double-sided doping of a GaSb QW (growth #7) and (b) double-sided doping of  $\text{In}_{0.41}\text{Ga}_{0.59}\text{Sb}$ . A hole density of  $2 \times 10^{12}/\text{cm}^2$  is assumed for both heterostructures.

earlier), the larger valence band offset and hence greater well depth of the GaSb QW results in it having a more efficient transfer of charge into the well from the modulation dopant. This explains the saturation of well charge in GaSb at about  $3.5 \times 10^{12}/\text{cm}^2$  as seen in the experimental data of Fig. 7, although higher values could presumably be reached by decreasing the spacer thickness. In comparison, the saturation level for InGaSb is much lower (unless the dopant is moved closer to the well). The doping efficiency of GaSb is a primary reason why it can reach much lower levels of sheet resistance than InGaSb (or InSb).

A second comparison between GaSb and InGaSb devices is shown in Fig. 10 where we plot their band diagrams and hole density profiles at 300 K. For the devices shown we employ double-sided doping so as to achieve an integrated hole density of  $2 \times 10^{12}/\text{cm}^2$  in both devices. For the densities considered, the primary subbands occupied are the two highest heavy hole (heavy out-of-plane mass) subbands [6]. Under all these conditions, the lighter in-plane subbands dominate at a sheet density of  $2 \times 10^{12}/\text{cm}^2$  with the heavier in-plane subbands being separated in energy by 60 meV at  $-0.5\%$  strain to 260 meV at 3% strain and the occupancy (in the former case) being 2.7%. Again the smaller well depth of the InGaSb QW results in it having a reduced charge transfer efficiency into the well; this is seen in Fig. 10 where to reach the same integrated hole density, the InGaSb device requires that the doping level be both higher ( $3 \times 10^{19}/\text{cm}^3$  vs.  $8 \times 10^{18}/\text{cm}^3$  for the GaSb device) and closer to the channel (4 nm vs. 5.4 nm).

Also evident in Fig. 10 is the fact that the lower valence band offset and smaller heavy-hole mass of the InGaSb device results in a stronger penetration of the hole wave functions into the barrier (6.4% of the well charge vs. 1.8% for GaSb). Both of these factors – the lower doping and decreased wave function tailing of the GaSb device – will tend to reduce barrier-associated scattering by ionized impurities and/or disorder in the GaSb device as compared to InGaSb. Hence, for a given geometry and well charge, a GaSb device will tend to have a higher mobility and smaller sheet resistance than a comparable InGaSb (or InSb) device as reported in this paper.

#### 4. Conclusions

Quantum wells of GaSb with  $\text{AlAs}_x\text{Sb}_{1-x}$  barriers were grown on  $\text{AlAs}_x\text{Sb}_{1-x}$  buffer layers on InP substrates as opposed to the GaAs substrates used heretofore. The buffer layer composition was adjusted to yield compressive biaxial strains of 1–3% in the GaSb. The strain-induced modification of the valence band structure, combined with confinement effects, resulted in room-temperature mobilities as high as  $1500 \text{ cm}^2/\text{V s}$ . Similar hole mobilities have been observed in highly strained InSb and InGaSb QWs. For the GaSb wells, however, we found the high mobilities are sustained up to densities of  $3.5 \times 10^{12} \text{ cm}^{-2}$ . We attribute this enhancement to the GaSb/AlAsSb having the deepest well, and thereby reduced scattering due to stronger confinement and lower levels of doping needed for the same channel charge. The crucial upshot from a device technology perspective is that the sheet resistivity of the GaSb can be as low as  $1500 \Omega/\square$ , a factor of two lower than any previous III–V p-type QW. GaSb p-channel FETs can thus have lower access resistance and higher drive currents. As such, these high-performance devices together with matching n-channel FETs [34–37] might well serve as the basis for a future ultra-low-power CMOS technology, especially if high-quality gate dielectrics [38] are also incorporated. Of course which material system(s) will win out in the Moore's law endgame is a complicated question that depends on many other factors such as channel hole mobility and velocity, contact resistance, subthreshold slope,  $I_{\text{on}}/I_{\text{off}}$  ratio, and compatibility with n-FET materials [39]. In addition to CMOS, other device technologies may benefit from the advantage of p-GaSb quantum wells with low sheet resistance.

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