Observation of excitonic transitions in InSb quantum wells


Department of Physics and Astronomy and Laboratory for Electronic Properties of Materials, The University of Oklahoma, Norman, Oklahoma 73019

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We report the observation of interband exciton transitions in InSb/Al$_x$In$_{1-x}$Sb multi-quantum-well samples. The exciton peaks are identified with the use of a simple quantum well model. The strain present in the InSb wells alters the spectrum significantly from that for unstrained III–V materials and makes it possible to use the exciton spectrum in determining the band offset. © 1998 American Institute of Physics. [S0003-6951(98)01934-2]

Since the advent of the first band-gap engineered materials, optical transitions corresponding to interband excitations have been highly useful in understanding confinement-induced band structure in semiconductors.\textsuperscript{1,2} In this letter we report the first observation of the interband exciton spectrum in InSb quantum wells. Because of its narrow energy gap and small effective mass, InSb has the highest intrinsic electron mobility of all binary III–V compounds; thus InSb quantum wells can potentially be fabricated into specialized photonic and high speed electronic devices. Of special interest in our strained-layer samples are the qualitative changes in the exciton spectrum compared to unstrained III–V quantum wells. The altered exciton spectrum is similar to that observed in the strained In$_{x}$Ga$_{1-x}$As/GaAs system.\textsuperscript{4}

Our samples are multi-quantum-well (MQW) InSb/Al$_x$In$_{1-x}$Sb heterostructures grown on GaAs substrates.\textsuperscript{5} For the measurements described here, we have chosen a sequence of 25 nominally undoped InSb MQWs with well thickness $d = 250$ Å. The alloy barriers are 500 Å thick with Al concentration $x = 0.09$. We use a Fourier transform infrared (FTIR) spectrometer to monitor the sample transmission as a function of photon frequency $\nu$. To reduce unwanted Fabry–Perot interference, we have coated the top surface of the sample with an antireflection NiCr film.\textsuperscript{6} Figure 1 shows the transmission, $T_{\text{MQW}}$, of a MQW sample referenced to the transmission, $T_{\text{sub}}$, of a sample with a similar substrate and epitaxial layer but lacking the InSb MQW sequence. Above the InSb band-gap energy (236 meV in unstrained bulk InSb),\textsuperscript{7} the spectrum shows absorption structures characteristic of excitonic continuum absorption between the stepped density of states for two-dimensional confined valence and conduction band states and the discrete exciton peaks at the step onsets.

The onset structures are enhanced by plotting the difference between a 4.2 K spectrum and that at a higher temperature (22 K), as in Fig. 2. This differential spectrum is made possible by the temperature dependence of the InSb band gap. The 4.2 K gap of 236 meV is reduced to 228 meV at 80 K, thus for the temperature difference of Fig. 2, a linear variation of energy gap with temperature predicts that the energy gap differential is about 2 meV. This differential enhancement technique gives essentially the negative derivative of the exciton structure in Fig. 1. We have checked this by numerically calculating the derivative of the 4.2 K transmission curve of Fig. 1. The 3D peak for a thick (5.7 μm) InSb epitaxial layer on GaAs is also shown in the figure. This peak is observed at the accepted 236 meV value of the band-gap energy since the exciton binding energy is expected to be small (~0.4 meV) due to the small effective mass in the conduction band.

In our MQW structures, the bulk InSb band gap is altered because of the strain caused by the pinning of the in-plane lattice constant in the InSb layers to the lattice constant of the barriers. Assuming a linear variation with alloy concentration, the strain is given by

$$\epsilon_i = \frac{a' - a}{a} x,$$

where $a$ and $a'$ are the InSb and AlSb lattice constants, 6.47 and 6.136 Å, respectively.\textsuperscript{7} The strain modifies the unstrained band-gap $E_g$ to the values $E'_g$ given to first order in $\epsilon_i$ by\textsuperscript{8}

![FIG. 1. Transmission spectrum of InSb/Al$_{0.09}$In$_{0.91}$Sb MQW sample at 4.2 K. The ratio of the transmission of the MQW sample to that of a reference sample lacking the MQWs is plotted vs. photon energy. Calculated transition energies are marked.](image-url)
where a and b are the formation potentials, \( C_{ij} \) and the Landolt-Börnstein values for all these quantities.

The band-edge masses are taken to be 0.0139, 0.015, and 0.47 for the electrons, light holes, and heavy holes, respectively, in free electron units. Band-edge effective mass values in the alloy are assumed to vary with concentration \( x \) according to the Kane model. The energy gap in the alloy is taken to vary with concentration \( x \) as \( E_g(x) = E_g(0) + (2.0 \text{ eV})x \). The uncertainty in the slope, 2.0 eV, is about 5%; thus the gap value for \( x = 0.09 \) has an uncertainty of about 2%.

With the selection rule, \( 
\Delta n = 0 \) (where \( n \) is the subband index), the calculation predicts three transitions: three heavy-hole transitions to the three conduction-band subbands and one transition from the single light-hole subband to the conduction band. The predicted transition energies shown in Fig. 3 are also marked in Figs. 1 and 2. The close agreement with the experimental peaks suggests that we have correctly identified the transitions. Because the highest lying subband is so weakly bound, we do not expect to be able to distinguish transitions to this subband from transitions to the continuum; indeed, the predicted position of the HH3-CB3 transition marks the onset of a broad absorption band. We ascribe the absorption structure near 430 meV to interband transitions across the gap of the Al\(_{0.09}\)In\(_{0.91}\)Sb barrier. If this value corresponds to the alloy gap, the \( E_g(x) \) relation, given above from Refs. 7 and 11, predicts and Al concentration of 9.7%. We intend a careful future study of the alloy absorption structure as a function of concentration.

Especially noteworthy is the position of the light-hole transition (LH1-CB1). This transition is seen in GaAs quantum wells very near the heavy-hole ground-state subband transition (HH1-CB1). Its shift to near the HH2-CB2 position is a consequence of the strain-induced difference between the band gaps for the light and heavy holes. As in the similar In\(_n\)Ga\(_{1-x}\)As/GaAs system, Eq. (2) (corrected by the small, quadratic term in \( \epsilon_i \)) predicts that, for offset ratios larger than 0.77, there is no potential well for light holes in the InSb. Thus for offset ratios larger than 0.77, the system of light holes and electrons is predicted to become type II. The offset ratio at which the transition occurs is nearly independent of concentration \( x \) since both the strain \( \epsilon_i \) and the alloy band-gap \( E_g(x) \) vary linearly with \( x \) and thus only

![FIG. 2. Differential transmission spectrum of an InSb/Al\(_{0.09}\)In\(_{0.91}\)Sb MQW sample. The transmission difference, \( T_{\text{MQW}}(4.2 \text{ K}) - T_{\text{MQW}}(22 \text{ K}) \) relative to the average transmission \( T \) is plotted vs photon energy. Calculated transition energies and band gaps are indicated.](Image 62x477 to 286x738)

![FIG. 3. Conduction and valence band edges for an InSb/Al\(_{0.09}\)In\(_{0.91}\)Sb quantum well with a band offset ratio of \( Q_c = 0.70 \).](Image 349x528 to 526x738)
through the small correction term to Eq. (2), which is quadratic in $\varepsilon_i$, does a weak concentration dependence occur.

The uncertainties in the offset ratio and in the alloy band gap give an associated uncertainty in the confinement potentials for electrons and holes. In view of this, using a more sophisticated model\(^2\) for the confined states is not warranted at this time, but we hope to be able to determine the alloy band gap more accurately as a function of concentration using transmission measurements in the MQW samples. The sensitivity of the LH1-CB1 transition to the offset\(^1\) as the thickness $d$ is varied should allow us to determine the offset-ratio for the InSb/Al\(_{x}\)In\(_{1-x}\)Sb system in conjunction with a four-band calculation.\(^2\) This will require line shape identification of the exact transition energies, a subject we have avoided here. Early analysis with the four-band calculation suggests that the band offset ratio is lower than the value, 0.70, used here and could be as low as 0.55. Another study we hope to report in the near future is that of the exciton spectrum for doped samples.

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