Quantum Well Carrier Sweep Out: Relation to Electroabsorption and Exciton Saturation

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Abstract—We have studied the effects of changing the barrier design of GaAs-AlGaAs quantum wells on the electroabsorption, exciton saturation, and carrier sweep-out times. Five samples have been studied with x values ranging from 0.2 to 0.4, and barrier thicknesses from 35 to 95 Å. Within this range, we find that the electroabsorption is not very sensitive to the barrier thickness, but that the ionization field of the excitons approximately doubles for an increase of x from 0.2 to 0.4. The samples with high, thick barriers have lower internal quantum efficiencies than those with low, thin barriers. We find that the exciton saturation intensity increases with increasing applied field, and decreasing barrier thickness or height. These findings are adequately explained by the field and barrier dependence of the thermionic emission and tunneling sweep-out rates. Time-resolved electroabsorption measurements confirm our understanding of the variation in sweep-out rates between samples, and indicate that the escape mechanism at low field is probably a thermally-assisted tunneling process.

I. INTRODUCTION

The physical mechanisms by which carriers escape from quantum wells under the influence of electric fields are an important factor in determining the performance of quantum well electroabsorptive devices such as the self-electrooptic effect device (SEED) [1], [2]. These are devices which operate by the quantum confined Stark effect, in which the excitons red-shift and broaden when an electric field is applied perpendicular to the quantum well layers [3]. If the carriers are swept out of the wells too rapidly on increasing the field, the broadening can increase. From this point of view, it is desirable to have quantum wells with relatively long sweep-out times, so that the excitons are sharper and can persist up to higher field levels. On the other hand, other physical properties of the quantum wells are favored by short sweep-out times. For example, there may be applications which require very fast switching times from single devices, and this would clearly be favored by having a faster sweep-out rate. Boyd et al. have recently demonstrated that the switching time of symmetric SEED devices can be made as short as 33 ps by using appropriately designed quantum wells with very fast carrier sweep-out times [4]. Another important property which is affected by the sweep out is the exciton saturation intensity. This, in fact, turns out to be an extremely important consideration in the design of SEED systems, because these systems tend to run at intensity levels much greater than that required to switch a single device. The reason for this is the need to pass on sufficient energy to the next device in the system to switch it after allowing for losses between consecutive devices. At these high power levels, the exciton absorption saturates, and this puts an effective upper limit on the maximum intensity which can be used. The system speed is determined by the time to integrate sufficient energy to switch the state of the devices, and therefore exciton saturation effects become the limiting factor in determining the maximum bit rate [5]. In a recent publication we have shown how the saturation intensity is strongly affected by the design of the quantum well structure, and that this effect is most likely caused by the change in carrier sweep-out times which accompanied the change in design [6].

In this paper we consider the physical processes which affect the carrier sweep-out rate in quantum wells subject to a perpendicular electric field, and we discuss how the sweep-out time is related to the electroabsorption and the exciton saturation intensity. We present a series of results on five carefully designed GaAs-AlGaAs quantum well structures which have allowed us to understand better the key physical mechanisms of thermionic emission and tunneling. We have thus been able to measure the dependence of the electroabsorption and exciton saturation intensity as a function of barrier design, electric field, and temperature. We have also measured the sweep-out times directly for two of the samples. From these measurements we have been able to come to a fairly clear understanding of the main trends in the sweep-out rate, and our conclusions have general applicability to other quantum well material systems, provided due consideration is given to the important material-dependent parameters. The paper is organized as follows. In Section II we discuss the basic physics that determines the sweep-out rate. In Section III we give details of the samples studied. In Sections IV and V we describe, respectively, the electroabsorption performance and exciton saturation intensity measured for the five samples. In Section VI we describe our direct measurements for the field and temperature dependence of the sweep-out times for two of the samples, and compare it to a third reference sample. In Section VII we gather together the various results and come to our conclusions.

Manuscript received December 7, 1990; revised April 30, 1991.
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IEEE Log Number 9101810.
II. CARRIER SWEEP-OUT MECHANISMS

There are three physical mechanisms that contribute to the field dependence of the carrier lifetime $\tau$ in a quantum well in an electric field: recombination, thermionic emission, and tunneling. In Fig. 1 we give a schematic diagram that illustrates these three processes. On the assumption that the three mechanisms are independent of each other, we may write:

$$\frac{1}{\tau} = \frac{1}{\tau_r} + \frac{1}{\tau_e} + \frac{1}{\tau_T}$$  \hspace{1cm} (1)

where $\tau_r^{-1}$, $\tau_e^{-1}$, and $\tau_T^{-1}$ are the recombination, thermionic emission, and tunneling rates, respectively. As will become clearer, $\tau$ is determined by a number of key parameters. These include: intrinsic material properties such as the carrier effective masses and band discontinuities; sample dependent factors such as the quantum well and barrier thickness; and external factors, such as the temperature and field strength. The three different mechanisms respond differently to changes in the various parameters, and so we shall consider each separately.

We consider first the carrier recombination rate. This is not in fact a sweep-out mechanism at all, and therefore does not contribute to the current measured in a photocurrent measurement. In good samples, the recombination rate is basically determined by the radiative lifetime, which is closely related to the electron-hole overlap and the occupancy factors [7]:

$$\frac{1}{\tau_r} \propto \sum_{n,m} \left| \langle \phi_m | \phi_n \rangle \right|^2 \int G_{md}(E) f_i(1 - f_e) E \, dE$$  \hspace{1cm} (2)

where $\tau_r$ is the bimolecular recombination lifetime. The summation is over all electron and hole sublevels in the quantum well. $\langle \phi_m | \phi_n \rangle$ is the electron-hole overlap (the interband matrix element is assumed to be independent of energy). $G_{md}$ is the reduced density of states, and $f_i, f_e$ are the conduction and valence band occupancies. The main functional dependence of $\tau_r$ is with respect to the temperature and the field strength. With increasing temperature, the radiative lifetime increases on account of the thermal spread of the carriers in the bands [8], while with increasing field the electron-hole overlap diminishes resulting again in an increase in $\tau_r$ [9]-[11]. In samples of poorer purity, effects such as trapping become increasingly more dominant. In most of our samples, it appears that essentially all of the photocarriers are collected except at very low fields in the samples with high or thick barriers. This tells us that $\tau_r$ is usually negligible in comparison to the faster tunneling and thermionic emission rates.

The thermionic emission lifetime is determined principally by the height of the barrier over which the carriers must be emitted. Schneider and von Klitzing have derived the following expression for the thermionic emission lifetime in a quantum well [12]:

$$\frac{1}{\tau_e} = \left( \frac{k_B T}{2 \pi m_e L_w} \right)^{1/2} \exp \left[ \frac{H_i(F)}{k_B T} \right]$$  \hspace{1cm} (3)

where $Q_e/Q_h$ is the ratio of conduction to valence band discontinuities. $(Q_e + Q_h = 1)$. $\Delta E_{th}$ is the difference in band gaps between the well and barrier material, and $E_{th}^{(n)}$ is the nth subband energy relative to the center of the well. Thus as the field is increased, the emission rate increases because the barrier height decreases. Note that the emission rate of electrons and holes is very different, and in the GaAs-AlGaAs system where $Q_h = 0.65$ [13], the holes can be emitted significantly faster than the electrons due to the smaller barrier height. Note also that the emission rate can be made faster by reducing $\Delta E_{th}$ This corresponds to using low values of $x$ in the GaAs-AlGa$_{1-x}$As system.

The importance of thermionic emission for vertical transport in quantum wells has been widely studied in the context of CW transport and photocurrent measurements [14]-[16]. There is, however, very little information available about the dynamics of the thermionic emission process. It is not immediately clear whether the approach of Schneider and von Klitzing embodied in (3) is, in fact, appropriate for the SEED devices to which this paper is oriented. The reason is that (3) does not consider the time taken for the carriers to scatter from the subband in which they are excited optically to the quasi-continuum at the top of the well. In the case of SEED devices, the
observed by a number of groups [24]–[26], [32]–[34], [36]–[38]. From the point of view of devices, the enhanced tunneling rate is already potentially useful for sweeping the carriers out faster.

In writing (1) we made the approximation that the three carrier loss mechanisms contributing to the lifetime are independent of each other. This is not strictly true for thermionic emission and tunneling, because thermally assisted tunneling processes are possible. Larsson et al. proposed the thermal occupation of higher sublevels followed by rapid tunneling through the barrier as the escape mechanism for their GaAs–Al$_{0.25}$Ga$_{0.75}$As quantum well p–i–n photodetectors [17]. Schneider et al. proposed a similar process to explain the temperature dependence of the photocurrent transients in a GaAs–AlAs superlattice [24]. Moreover, near the top of the barrier, the tunneling transmission may be so high that the effective barrier height for thermal emission is reduced (Fowler–Nordheim tunneling). Since it is both a tunneling and a thermal process, the probability for thermally assisted tunneling depends strongly on the barrier thickness as well as the barrier height and temperature. The expression for the escape probability by this process would contain a product of the exponentials on the right-hand side of (3) and (5) with the appropriate activation energy as $H'(F)$ in (3) and with a reduced effective barrier height $H'_0(F)$ in (5).

On the basis of the discussion above, we can make some overall general observations about the sweep-out times. (We omit discussion of the recombination rate because, as discussed above, it is generally negligible for our samples.)

1) The sweep-out time decreases with increasing field because both the thermal emission and tunneling lifetimes decrease due to the reduction in the effective barrier height.

2) The sweep-out times increase with $L_0$ due to the increase in the tunneling time.

3) The sweep-out times increase with increasing $\Delta E_g$ due to the increase in both the thermal emission and tunneling times. For the GaAs–Al$_x$Ga$_{1-x}$As system, this implies that the sweep-out times increase with $x$.

4) The sweep-out times decrease with the temperature due to the decrease in the thermal emission time.

In trying to ascertain which of the sweep-out processes is dominant for a particular sample, it is useful to note that the hallmark of thermionic emission is its exponential dependence on $T^{-1}$, while tunneling is exponentially sensitive to the barrier thickness. Of course, a thermally-assisted tunneling process would be very sensitive both to temperature and barrier thickness.

In concluding this section it is worth emphasizing two points which are particularly relevant to the SEED and similar quantum well electroabsorptive devices. First, the sweep-out times of electrons and holes are very different due to their different effective masses and band discontinuities. This has a number of important implications for devices, especially as regards space charge effects [39]–[42]. Such space charge effects become increasingly more important at high power levels, especially in the
Ga$_{0.47}$In$_{0.53}$As–InP system, which has a much larger valence band discontinuity than GaAs–Al$_{0.3}$Ga$_{0.7}$As. The hole sweep-out times are not always much slower than the electron times in the GaAs–AlGaAs system. This follows because of the smaller band discontinuity in the valence band which favors rapid thermal emission. It is also possible that valence band mixing effects can reduce the hole tunneling times compared to those of pure heavy hole states [43]. The second point is that SEED devices generally require about an optical absorption length of quantum well material, so that the devices contain a multiple quantum well structure rather than individual wells. Equations (1)–(5) are still valid for the sweep-out time from a particular well, but it is necessary to consider what happens to the carriers once they have escaped from the wells. The field strength is generally at least 10$^8$ V cm$^{-1}$, so that the carriers will be rapidly accelerated to the saturated drift velocity of $\sim 10^7$ cm s$^{-1}$. Some of the carriers will be swept straight to the equipotentials in the contact layers, but others will be scattered and recaptured by subsequent wells. It is known that such capture events take place in less than 1 ps in InGaAs–InP quantum wells [44]. The net effect of recapture is to increase the effective sweep-out time.

### III. Sample Design

From the discussion in the previous section [points 1–4], it is clear that the sweep-out times depend critically on the electric field strength, the barrier thickness $L_b$ and barrier height (which is proportional to $x$). We therefore grew five p-i-n GaAs–Al$_{x}$Ga$_{1-x}$As multiple quantum well (MQW) samples by molecular beam epitaxy with the quantum wells as the intrinsic region of the diode. A controllable field could then be applied to the quantum wells by applying reverse bias to the diode. We designed the samples with the same well thickness (95 Å), but with varying barrier thickness $L_b$ and barrier Al concentration $x$. Table I gives details of the samples. Samples I, III, and V have the same barrier thickness (65 Å), but have the $x$ value varying from 0.2 to 0.4, while samples II, III, and IV have constant $x$ (0.3) but have $L_b$ varying from 35 to 95 Å. Note that we have varied the number of periods where necessary in samples II and IV, so that the total MQW thickness remains constant at 1.0 μm. This means that a given applied voltage corresponds to the same nominal field strength for all the samples. We regard sample III as a reference, since it has the same barrier design as many SEED devices.

The samples were grown on n-doped GaAs substrates. The detailed layer structure, starting from the substrate up, was as follows. 0.2 μm GaAs buffer layer ($n = 2 \times 10^{18}$ cm$^{-3}$), 1.2 μm Al$_0.3$Ga$_{0.7}$As stop etch ($n = 5 \times 10^{17}$ cm$^{-3}$), 0.24 μm Al$_{0.5}$Ga$_{0.5}$As superlattice ($n = 5 \times 10^{17}$ cm$^{-3}$), 1.0 μm intrinsic MQW structure (see Table I), 0.75 μm Al$_{0.5}$Ga$_{0.5}$As ($p = 5 \times 10^{17}$ cm$^{-3}$), 0.1 μm GaAs ($p = 2 \times 10^{18}$ cm$^{-3}$). The samples were processed into 200 μm x 200 μm mesa areas, and after depositing gold contact pads on the top p$^+$ layer, the optical window was approximately 120 μm x 200 μm. For the room temperature electroabsorption measurements, tungsten probes were used to make contact to the samples, but wire bonding was required for the low temperature work. In the transmission measurements, the samples were mounted on sapphire or BK7 disks with clear epoxy, and the GaAs substrates were etched away using a jet etcher.

In the last four columns of Table I we indicate the values of $\tau_e$ and $\tau_f$ calculated for electrons and heavy holes at room temperature for our five samples from (3) and (5). We assumed a field strength of $1.5 \times 10^8$ V cm$^{-1}$, which corresponds approximately to the built-in field of the diode at 0 V applied bias. As expected, the escape time for samples with low or thin barriers is lower than for those with high or thick barriers. It is clear that the dominant escape mechanism for the holes is thermal emission, while for electrons, the dominant mechanism depends on the details of the sample. Note how short the hole thermal emission times are. In particular, it is not generally true to presume that the hole escape rate is very much slower than the electron escape rate. This will depend on the sample design, the temperature, and the field strength.

### IV. Electroabsorption

In Fig. 2 we show representative photocurrent spectra for the five samples at room temperature. These spectra were taken using a tungsten lamp light source and 0.25 m monochromator with a resolution of ~2 meV. The voltages shown are the reverse bias applied to the diode. The
five graphs are laid out so that x varies vertically while \( L_w \) varies across the figure. We notice immediately that all five samples show very good quantum confined Stark effect performance. Let us first consider the variation in electroabsorption with barrier thickness at constant \( x \). For \( x = 0.3 \), we measure only small variations in the exciton linewidth and red-shift on varying \( L_w \) from 95 to 35 Å. As discussed above, the principal effect of reducing the barrier thickness is to reduce the tunneling time. The reason why we do not see any large effect is that at room temperature, the linewidth is dominated by LO-phonon broadening and inhomogeneous broadening due to interface roughness and alloy disorder [45]. The thermal lifetime of the excitons is known to be only 300 fs at room temperature [46], which is much less than the 5 ps electron tunneling time calculated for the 35 Å barrier sample at 0 V (see Table I). However, there must ultimately come a point at which the thinness of the barrier does cause significant exciton line broadening, because the tunneling time continues to decrease exponentially with the barrier thickness as the barrier is thinned further. In fact, we have also investigated samples grown with 25 and 15 Å Al\(_{0.3}\)Ga\(_{0.7}\)As barriers. In these samples, the 0 V exciton linewidth was significantly larger than for samples II-IV, most noticeably for the 15 Å barrier sample, where the linewidth was broader by a factor \( \sim 1.3 \) at 5 V. From the point of view of devices, there appears to be no significant loss in electroabsorption involved in reducing the barrier thickness from 65 to 35 Å.

We now consider the variation of electroabsorption with the \( x \) value of the barriers. On changing the \( x \) value, we change the confinement energy of the carriers. This does appear to have a noticeable effect on the electroabsorption. For sample I with \( x = 0.2 \), the heavy hole exciton is barely resolved at 15 V, whereas for sample V with \( x = 0.4 \), the exciton is still well resolved at 30 V. The 80 meV exciton red-shift for sample V at 30 V bias (field \( \sim 3 \times 10^5 \) V cm\(^{-1}\)) is particularly noteworthy. This red-shift corresponds to about ten exciton binding energies. The results imply that the field for exciton ionization at least doubles on increasing \( x \) from 0.2 to 0.4. The reason why the excitons are less stable to field in the samples with smaller \( x \) is most likely that the carriers can be thermally emitted very rapidly over the relatively low barriers, which would give rise to lifetime broadening. Note that the electroabsorption of sample I with \( x = 0.2 \) is still very useable from the point of view of SEED devices. These devices require a large contrast between the low and high field absorption at the exciton wavelength, so that the main criterion is the 0 V exciton linewidth rather than the stability of the excitons to high fields. In fact, it is actually preferable that the excitons begin to broaden at more modest voltages, since then the voltage required for a particular contrast ratio decreases.
In order to get a meaningful comparison of the electroabsorption performance, we have scaled up the 0 V spectra for samples III–V by the factors shown in the figure. This was necessary because we took photocurrent rather than absorption spectra. The photocurrent \( i \) generated for incident optical power \( P \) is given by

\[
i = P \frac{\eta(1 - R)}{h\omega} \frac{\varepsilon}{[1 - \exp(-\alpha L)]}.
\]  

(6)

Here, \( \alpha \) is the absorption coefficient at the incident photon energy of \( h\omega \), \( R \) is the reflectivity of the front surface, and \( L \) is the total thickness of quantum well material. \( \eta \) is the internal quantum efficiency of the photodiode; that is, the number of electrons collected per absorbed photon. The principal reason why \( \eta \) would be different from unity is that the sweep-out time is comparable to the recombination time, in which case, a significant number of photoelectrons are lost from the current through recombination. The photocurrent spectra have the same general shape as the absorption spectra, with peaks at the same wavelength, but on making comparisons at different voltages, it is necessary to consider whether \( \eta \) is unchanged with voltage. At 0 V we observed that the whole spectrum for samples III–V was scaled down compared to the spectrum at 5 V. This was not true for samples I and II, where, for example, we observed a monotonic decrease in the peak photocurrent at the heavy hole exciton as a function of voltage. By contrast, the heavy hole exciton photocurrent peak for samples III–V increased at first as the voltage was applied, reaching a maximum between 2.5 and 5.0 V, and thereafter decreased with voltage. The reason for this behavior is that \( \eta \) is less than unity for samples III–V at low voltages because of their long sweep-out time. As the voltage is increased, \( \eta \) increases to unity because the sweep-out time decreases compared to the recombination time. The scale-up factors in Fig. 2 were included to compensate for the nonunity \( \eta \). In samples I and II, we observed that the photocurrent in the continuum states immediately above the heavy hole exciton was practically independent of voltage up to 5 V. We chose the scale-up values by comparing the magnitude of the photocurrent for samples III–V in the same spectral region at 0 and 5 V, i.e., we assume that only \( \eta \) in (6) changes between 0 and 5 V at this photon energy. Thus the reciprocal of the scale-up factor gives an approximate measure of \( \eta \) for the particular sample at 0 V. Note that this suggests that \( \eta \) decreases with barrier thickness and with \( \chi \), which is entirely consistent with points 2) and 3) in Section II.

We were able to verify directly that \( \eta \) is indeed unity to within experimental error for sample I. This was done by measuring the photocurrent \( I-V \) characteristic of the diode at low temperatures using a Hewlett Packard HP4145B semiconductor parameter analyzer, when the diode was illuminated with a known amount of laser power. Fig. 3 shows a schematic of the apparatus used. (The acoustooptic modulator was superfluous for this measurement, but was needed later for the saturation measurements.) For this measurement, the sample was mounted in a continuous flow Helium cryostat. Fig. 4 shows representative \( I-V \) characteristics at 30 K for two different wavelengths at an incident power level of 3.8 \( \mu \)W. Consider first the \( I-V \) for 807 nm excitation (dotted line). The laser wavelength is longer than the exciton line at 0 V, and as the voltage is increased, first the heavy hole exciton and then the light hole exciton is swept through the laser wavelength. Consider now the \( I-V \) for 796 nm excitation (solid line). This wavelength is closer to the exciton at 0 V, and so the voltage required to bring the heavy hole exciton into resonance with the laser is smaller (2 V). As for 807 nm excitation, we see peaks corresponding to both the heavy and light hole excitons. There is a third peak, which we assign to resonant tunneling effects. Low temperature photocurrent spectra of this sample reveal an anomalous line broadening around 5 V corresponding to the resonant tunneling of electrons from the first sublevel in one well to the second sublevel in the adjacent well [47]. Returning to the heavy hole peak at \( \sim 2 \) V, we notice that this heavy hole peak appears flattened. The reason for the flattened peak is that at these low temperatures and voltages, the exciton lines are very narrow, with a larger peak absorption height compared to room temperature. The absorbance \( [1 - \exp(-\alpha L)] \) is insensitive to \( \alpha \) when \( \alpha \) becomes very large, and so we see that the photocurrent becomes independent of voltage near the heavy hole peak. We can thus take \( [1 - \exp}
equal to unity at the flattened peak, and thereby obtain a value of $\eta$ from (6), given that $P$ is known, and $(1 - R) = 0.7$ for GaAs. In this way we arrive at a value of unity for $\eta$ at 30 K and 2 V to within the estimated 20% absolute accuracy of the power meter. We were able to repeat this procedure up to 200 K, and found no significant departure of $\eta$ from unity in this temperature range. Above 200 K, the exciton peaks in the $I$-$V$ characteristics were rounded at all voltages, so that we could not deduce $\eta$ because $[1 - \exp(-\alpha L)]$ was not known. Throughout this paper we have assumed that $\eta$ remains essentially unity at and slightly above room temperature, as has been observed previously [1], [3], and confirmed in recent quantum efficiency measurements in samples with similar designs to ours [48]. This assumption is consistent with the fact that the peak exciton photocurrent measured in the spectra for samples I and II was a monotonically decreasing function of voltage, as discussed above. On the basis of this assumption, we could measure the room temperature $I$-$V$ characteristic of any of the five samples for varying laser wavelength and convert it to an $\alpha$-$V$ characteristic. We thus could obtain the values of the exciton absorption coefficient at room temperature as a function of voltage. The values of $\alpha$ deduced by assuming unity $\eta$ are in good agreement with direct measurements in similar conditions [3]. Given the discussion above, this conversion is less reliable for samples III-V at low voltages, and we have to make an estimate of the actual value of $\eta$ based on the electroabsorption scale-up factor.

V. EXCITON SATURATION

Exciton saturation measurements were made using the apparatus shown in Fig. 3. The excitation source was either a CW Styryl 9 dye laser or a titanium sapphire laser. We used an acoustooptic modulator to produce a train of square wave pulses of 100 ns duration at a repetition rate of 1 MHz. This low 10% duty cycle was necessary in order to reduce thermal effects in the sample. Spot radii on the sample ranged from 2-30 $\mu$m. The experiment consisted in measuring the $I$-$V$ photocurrent characteristic of the diodes as a function of incident laser power and wavelength with an HP4145B semiconductor parameter analyzer. Fig. 5 shows typical results for two different wavelengths in sample V at room temperature. (Sample V is the one that saturates most easily because of its high barriers.) We have divided the photocurrent by the laser power to obtain the responsivity of the quantum wells, considering them as a photodiode. In this figure, positive voltage corresponds to reverse bias on the diode. The laser spot was Gaussian with a measured $e^{-2}$ radius ($w$) of 4.5 $\mu$m. In Fig. 5(a), the laser wavelength is 840 nm, approximately 5 nm longer than the heavy hole exciton at 0 V. The responsivity shows peaks at the voltages where the applied field exactly red-shifts the excitons to the laser wavelength, because the absorption has a maximum at the exciton wavelengths. Thus in Fig. 5(a) we first see a peak at 6 V corresponding to the heavy hole exciton, and then at 16 V for the light hole exciton. The solid curve shows the responsivity measured at an incident power level of 0.4 $\mu$W, while the dotted curve shows the responsivity at 40 $\mu$W. The 40 $\mu$W curve follows the 0.4 $\mu$W curve fairly closely, except in the vicinity of the heavy hole peak. (The small down scaling of the 40 $\mu$W curve is probably just the experimental accuracy of our power measurements.) We measure less responsivity at the heavy hole peak at the higher power level, which we interpret as being caused by saturation of the excitons. In Fig. 5(b) we show the responsivities measured at the same power levels as for Fig. 5(a), but with the laser tuned to 846 nm. Now we must apply 12 V reverse bias to red-shift the heavy hole exciton to the laser wavelength, and the light hole is outside the range of the 20 V sweep. We observe negligible saturation at the heavy hole peak at the higher power level, which tells us that the saturation intensity has increased with the voltage. This is also why we do not observe any saturation at the light hole peak in Fig. 5(a).

In order to obtain a value for the heavy hole exciton saturation intensity at a particular voltage we proceed as follows. First we convert the responsivity at the heavy hole peak to an absorption coefficient using (6) with the assumption of $\eta = 1$. In Fig. 6 we show the values of $\alpha L$ deduced in this way as a function of average incident laser intensity ($P/\pi w^2$). The data is for the heavy hole exciton at 6 V, as in Fig. 5(a). In the intensity range where the saturation varies linearly with intensity, the averaged intensity correctly accounts for the variation in the saturation across the spot size due to the Gaussian profile of the laser. If we assume a phenomenological absorption saturation dependence of the form

$$\alpha(I) = \frac{\alpha_0}{1 + (I/I_s)}$$

Fig. 5. Room temperature responsivity curves of sample V for two different wavelengths at incident power levels of either 0.4 $\mu$W (solid line) or 40 $\mu$W (dotted line).
then at low intensity $\alpha$ is a linear function of $I$, and $I_0$ is the extrapolated intercept on the intensity axis. We have shown the extrapolation of $\alpha(I)$ in Fig. 6, which gives an intercept at 3.7 kW cm$^{-2}$. In order to obtain the value of $I_0$, from the experimental data, we have to allow for the fact that the intensity $I$ in (5) is the local intensity inside the sample, while the data is measured against incident intensity. In the limit of small changes in $\alpha$, where the effective interaction length approximation is valid [45], $I_0$ can be obtained by multiplying the experimental intercept by $(1 - R)(1 - \exp(-\alpha_0d))/\alpha_0L$. We thus obtain a value of 1.8 kW cm$^{-2}$ for the heavy hole exciton saturation intensity of sample V at 6 V and room temperature.

In Fig. 7 we show the 300 K saturation intensities of the heavy hole excitons as a function of reverse bias for the five samples. The values were obtained exactly as described in the previous paragraphs, with the same spot size throughout. The average field is determined by the applied voltage, the built-in voltage of the diode, and the intrinsic region width, and is therefore approximately the same for all the samples at a given voltage. From Fig. 7, we can draw three clear conclusions about $I_s$.

1) $I_s$ increases with applied field.
2) $I_s$ decreases with the barrier thickness $L_0$.
3) $I_s$ decreases with the barrier Al concentration $x$.

It is clear that thin barriers with low Al concentration give the best saturation performance for SEEDs, and that careful design of the quantum well structure can lead to improved saturation performance.

We now discuss the physics behind the experimental behavior 1)–3). Exciton saturation in quantum wells has been extensively studied experimentally and theoretically in a variety of material systems for the case of zero electric field [45, 46, 49–58]. In the GaAs–AlGaAs system, for example, the heavy hole excitons are known to saturate at 300 K with a microscopic saturation areal density of $N_e \approx 3 \times 10^{11}$ cm$^{-2}$ for the case of zero applied field [45, 49]. The case of finite electric field has received less attention, with only a few studies published at present [6, 31, 42]. A quantitative analysis of $I_s$ at finite applied field is complicated by the fact that $\tau_s$ and $\tau_r$ are different, as discussed in Section II. Since the steady state areal carrier populations generated at a CW intensity $I$ are given by

$$N_e = \frac{I \tau_s (I_s + L_0)}{\hbar \omega}$$

we have therefore to consider the effect of having different electron and hole densities within the sample. There are two main consequences of this. First we must consider the different saturating efficiencies of the two particle types, and second, we have to consider the effects of space charge buildup. These two effects are discussed in more detail below. However, before going into details we can make some general observations which give the main ideas to explain the results. The principal point is that (8) shows that the carrier densities vary strongly with the sweep-out rate through $\tau_s$. This means that by reducing the sweep-out times we can reduce the carrier densities generated at a given intensity. Both space charge effects and microscopic exciton saturation depend on the carrier density, and thus by reducing the sweep-out time, we should be able to increase the exciton saturation intensity. The close parallel between 1)–3) above and points 1)–3) in Section II, confirms this supposition, and strongly suggests that the variation in $I_s$ between the samples is caused by differing tunneling and thermionic emission rates. This conclusion does not depend on the detailed microscopic mechanism of the exciton saturation.

The saturation of excitons by a gas of thermalized carriers, but with differing electron and hole populations, can be anayled by an extension of the methods of Schmitt-Rink et al. [57], [58]. At room temperature, where the carrier gases have Boltzmann distributions, the exciton saturation is essentially proportional to the number of carriers in the region of $k$ space up to $\sim 1/a_r$, where $a_r$ is the exciton radius. This implies that the saturation density for a single thermalized carrier type at high temperatures is proportional to its effective mass [see (2.3.37b) in [58]]. (The heavier carriers are distributed over a larger region of $k$ space.) The fractional exciton absorption saturation $\Delta \alpha / \alpha_0$ is then given by

$$\frac{\Delta \alpha}{\alpha_0} = \frac{I}{I_s} = \frac{N_e}{N_e} + \frac{N_h}{N_h}$$

Fig. 6. Intensity dependence of the heavy hole exciton absorption in sample V at 6 V reverse bias.

Fig. 7. Heavy hole exciton saturation intensity at room temperature as a function of applied reverse bias for the five GaAs–AlGaAs, As quantum well p-i-n samples.
When \( N_e = N_h = N \), the absorption saturation is simply 
\(-\alpha \chi / \alpha_0 = N / N_e \). Therefore we deduce that the individual 
particle saturation densities \( N_e \) are greater than \( N \), by 
a factor \( m_e / m_h \), where \( \mu = m_e m_h / (m_e + m_h) \) is the reduced mass. Taking this into account, we can combine (8) and 
(9) to obtain

\[
I_e = \frac{\hbar \omega N_e}{\alpha_0 (L_e + L_a)} \frac{m_e + m_h}{m_e \tau_e + m_h \tau_h}.
\]

In principle, (10) allows us to evaluate the exciton saturation intensity at finite applied field. In order to do this, 
we need detailed knowledge of the field dependence of both \( \tau_e \) and \( \tau_h \), and also of \( N_e / \alpha_0 \), \( N_h / \alpha_0 \) may change 
somewhat as the field is applied. \( N \) is inversely proportional to \( \tau_e \) [57], and \( \alpha_0 \) has been calculated to increase 
by 25-40% on increasing the field from 0 to 10^5 V cm^{-1} in \(
\sim 100 \) Å quantum wells [3]. [59]. [60]. This would 
cause a reduction in \( N_e \) by a factor of \( \sim 2 \) at most. This 
decrease in \( N_e \) is offset, to a large extent, by the accompanying 
decrease in \( \alpha_0 \). Therefore it is very unlikely that the field and barrier dependence of \( N_e / \alpha_0 \) can explain the 
experimental results. Therefore we expect the strong field 
and barrier dependence of \( \tau_e \) and \( \tau_h \) to dominate the field 
dependence of \( I_e \). A quantitative analysis of (10) requires 
measurements of both \( \tau_e \) and \( \tau_h \) as a function of voltage, 
and this data is not yet available. However, our basic 
conclusion that the saturation intensity is dominated by the 
lifetimes is fully consistent with (10).

As mentioned above, the other consequence of having 
different electron and hole densities in the sample is the 
buildup of space charge as the intensity is increased. For 
everything, if the electrons escape faster than the holes, then 
the holes are left behind and generate the space charge. 
The space charge can screen the applied field and generate 
field nonuniformities. The result is a reduction of the 
exciton red-shift produced by the external field, and also 
exciton broadening. Both these effects would change the 
absorption at the laser wavelength as the intensity is 
increased, and might therefore look like exciton saturation. 
Moreover, (3)-(5) shows that \( \tau_e \) and \( \tau_h \) can begin to depend 
on each other and on the laser intensity by means of the 
space charge modifications to the field. Steady-state 
space charge effects have been observed in InGaAs-InP 
[39]-[41], and Wood et al. have recently given a self-
consistent analysis of the intensity dependent space-charge 
effects in this material [42]. These space charge effects 
are particularly acute in InGaAs-InP, where the hole 
sweep-out times are very slow because of the large band 
discontinuity for that system. It is reasonable to expect that 
steady state space charge effects in the GaAs-AlGaAs 
system will be less severe than in InGaAs-InP, because 
the band discontinuities are smaller, and the hole 
thermal emission times in GaAs-AlGaAs are expected to be fast 
(see Table I). We were able to see some evidence for space 
charge effects at low voltages in sample V, which has the 
largest band discontinuity of our five samples. Fig. 8 
shows the responsivity of this sample with the laser tuned 

to 838 nm, such that the heavy hole exciton was red-
shifted to the laser wavelength with 4 V reverse bias. As 
for Fig. 5, the solid and dotted lines are for 0.4 and 
40 \( \mu \)W incident power. We notice that at the higher power 
level the heavy hole peak has saturated and shifted to 
about 5 V. This shift might well be caused by the buildup 
of space charge, since this effect would tend to reduce the 
field inside the sample, and thus we would have to apply 
more external voltage to bring the average internal field 
up to the level required to red-shift the exciton to the laser 

dependence of \( I_e \). A quantitative analysis of (10) requires 
measurements of both \( \tau_e \) and \( \tau_h \) as a function of voltage, 
and this data is not yet available. However, our basic 
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more external voltage to bring the average internal field 
up to the level required to red-shift the exciton to the laser 

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image.png}
\caption{Responsivity of sample V for 838 nm excitation at an incident power level of either 0.4 \( \mu \)W (solid line) or 40 \( \mu \)W (dotted line). The shift in the heavy hole peak to higher voltage with increasing power may be an indication of the importance of space charge effects in this sample at low voltages.}
\end{figure}
allowing for the linear $T$ dependence of $N_t$ at high temperatures [57]. The applied voltage was 8 V. The measurement could not be made at lower voltage because the value of $[1 - \exp(-\alpha_0 L)]$ was too close to unity to give meaningful results from photocurrent measurements (see Fig. 4). We interpret the results as follows. $(\alpha L T^{-1})$ is essentially independent of $T$ at low temperatures because the lifetimes are dominated by tunneling. As the temperature is raised, the thermal emission process becomes activated. The error bars on the measurements are unfortunately too large to obtain reliable values of the activation energy. The important result is the strong sensitivity to temperature, which confirms our intuition that the high saturation intensity measured at room temperature is caused by rapid thermal emission.

We close this section with a brief discussion of the estimated accuracy of our values of $I_i$. The Gaussian beam averaging and effective interaction length correction are accurate up to first order in $\delta \alpha/\alpha_0$. This is also true of the theoretical models used to write (9). We could not reliably detect values of $\delta \alpha/\alpha_0$ less than 2–3%, and our values of $I_i$ are based mainly on the intensities required to give $\delta \alpha/\alpha_0$ of ~5–10%. The first order approximation should be reasonable at these intensity levels. Also, at these relatively low power levels, our neglect of higher order many-body effects such as band-gap renormalization should be reasonable. Such effects become increasingly more important at higher intensities, and may be the reason why, for example, the data in Fig. 6 do not fit very accurately to the two-level saturable atom approximation of (7) at high intensity. We do not think that thermal effects are significant in our data. These would have shown up in the data such as Figs. 5 and 8 as a shift to lower voltage at high powers, which was never observed with the 10% high frequency duty cycle. We assumed that the internal quantum efficiency was unity and invariant with intensity. There is no reason to expect $\eta$ to vary with intensity in the limit of small $\delta \alpha/\alpha_0$. This follows because $\eta$ is determined by the ratio of the sweep-out time to the recombination time, and this should be unaffected by intensity to first order. The assumption of $\eta = 1$ affects the values of $\alpha L$ determined from the photocurrent spectra, but the values of $I_i$ deduced from extrapolating $\alpha(I)$ are not very sensitive to $\eta$ for the values of $\eta$ encountered in the samples (minimum value ~60%), provided $\eta$ is unchanged with intensity. Our absolute accuracy is affected by the accuracy of our value of the beam radius, and also the calibration of the power meter. A reasonable estimate for the absolute accuracy is a factor of 2. The principal result of our work is the variation of $I_i$ between the samples and with the applied voltage. This is a comparative study, with the measurements performed indendently for all the samples and voltages, and so the basic result is valid, irrespective of the validity of any of our approximations.

VI. CARRIER SWEEP-OUT MEASUREMENTS

We made direct measurements of the sweep-out times of samples I and II by performing pump-probe experiments on transmission samples made from the same wafer as the previous measurements. A schematic diagram of the apparatus is given in Fig. 10. The samples were mounted in a continuous flow Helium cryostat for the temperature dependent measurements. The laser pulse-width was 0.5 ps, tunable from 790-870 nm, and the estimated maximum carrier density was $5 \times 10^{15}$ cm$^{-3}$. The focused spot radius was typically 15–20 μm. The combination of lock-in techniques and aperturing of the transmitted beams ensures that we only detect the change in probe transmission induced by the pump. We used a camera imaging system to check that the pump and probe spots were overlapping on the sample. In these conditions, the rise time of the detected nonlinear signal is directly related to the sweep-out time [61]–[63].

In Fig. 11 we show our results for the sweep-out times measured for samples I and II, together with our earlier $p-i-n$ quantum well sample [32]. This earlier sample had 0.9 μm of GaAs–Al$_{0.3}$Ga$_{0.7}$As quantum wells with a well thickness of 65 Å and $L_a = 57$ Å, and therefore has barriers similar to those of sample III. The values for the escape time given are the 10–90% rise time of the differential electroabsorption signal, as has been used previously [32], [63]. On comparing the results for the three samples, we can come to general conclusions about the field and barrier dependence of the escape times and compare them to our understanding based on (1)–(5).

1) All three samples show a general trend to a reduction in escape time as the voltage is increased. The minima at 5, 7, and 10 V for the three samples are well explained by resonant tunneling of electrons between the first and second sublevels of adjacent quantum wells. The arrows in Fig. 11(a) and (b) show the resonant voltage deduced from independent spectroscopic studies on these same samples [47], [64], and also from low temperature photocurrent $I$–$V$ characteristics (see Fig. 4). The coincidence of the resonant voltages measured by independent techniques and the minima in the escape time confirms our assignment of the escape time minima to electron resonant tunneling, and also indicates that space charge effects are negligible at the carrier densities gen-
erated. At fields well above the resonance, the escape time limits out to a value of around 10 ps for samples I and II. This time corresponds to the average transit time of the carriers through the 1 μm intrinsic region at a saturation drift velocity of ~10^7 cm s^{-1}, and represents an effective lower limit on the escape times that can be measured by this technique. The reduction of escape time with applied field can be caused by either an increase in the thermionic emission rate or the tunneling rate [see (3)–(5)]. These measurements are consistent with the measured increase in saturation intensity with voltage.

2) On comparing Fig. 11(a) and (c), we see that a reduction in x at approximately constant L_b leads to a reduction in the escape time. Again, this is consistent with either enhanced tunneling or thermionic emission, although the latter would seem more likely. These results confirm that the most likely cause of the increase in I, on reducing x is the reduction in the sweep-out time.

3) On comparing Fig. 11(b) and (c) we see how a reduction in L_b at approximately constant x leads to a reduction in the carrier escape time. This is characteristic of a tunneling event, and cannot be explained in terms of pure thermionic emission. The result explains the increase in I, observed on reducing L_b, and also the faster SEED switching time measured for samples with 35 Å barriers compared to 60 Å [4].

4) The escape times of all three samples are very sensitive to temperature at low voltages, but practically independent of T above resonant voltage. (The fact that we give no data points in Fig. 11(c) at 100 K below 9 V indicates a very long lifetime.) This result suggests that the escape process is thermally assisted at low fields, and proceeds by tunneling at high fields. A simple explanation of why this might be so has been given by Larsson et al. [17]. They argued that at low fields a pure tunneling escape mechanism is unlikely due to the large number of
barriers that must be crossed. On the other hand, at high fields the carriers need only tunnel through a single barrier, and therefore this process has a high probability. Since the low-field escape time is also affected by the barrier width [cf. point 3 above], the most likely explanation seems to involve a thermally-assisted tunneling escape mechanism.

In order to investigate the low-field thermally-assisted escape mechanism, we examined the temperature dependence of the escape time of sample 1 at 3.0 V in more detail. Fig. 12 shows the variation of the measured sweep-out time against temperature. We find two different types of behavior. Up to 100 K, the sweep-out time is independent of the temperature, while above 100 K it decreases rapidly with T. Around room temperature, the measured escape time begins to limit out at the sample transit time of ~10 ps. These results are consistent with the temperature-dependent saturation measurements shown in Fig. 9. The results can be readily explained on the basis of (1), (3), and (5). These equations show that at low temperatures, thermal emission is frozen out, and the escape time is determined by tunneling, which depends only very weakly on T. We thus deduce a value of 330 ps for the tunneling time. At sufficiently high temperatures, thermal emission must become the dominant escape mechanism, and we can use (1) to deduce T<sub>e</sub>. To do this we subtracted the 13 ps sample transit time from the measured sweep-out time, and we assumed that T<sub>e</sub> is negligible. The simple allowance for the transit effects becomes less reliable for the data points at higher temperatures.

The deduced values of T<sub>e</sub> are plotted on a logarithmic scale against 1/T<sup>-1</sup> in the inset of Fig. 12 between 140 K and room temperature. This Arrhenius plot gives the activation energy as 49 ± 15 meV at 300 K. The ±15 meV uncertainty is based on the sensitivity of the fit to variations in the activation energy, and also on the estimated accuracy of the thermal emission times. The curve fitted to the sweep-out times in Fig. 12 was obtained from (1) with tunneling and transit times of 330 and 13 ps, respectively. The thermal emission time was taken from the straight line fit in the inset. We repeated this analysis at 3.5 V, and found a tunneling time of 100 ps and thermal activation energy of 47 meV.

In order to obtain a clearer understanding of the physical significance of the activation energy, we show in Fig. 13 a sketch of the electron and heavy hole energy levels of the sample at an applied field of 45 kV cm<sup>-1</sup>, corresponding to about 3.0 V applied bias after allowing for the built-in voltage. The levels were found by fitting the low temperature optical absorption spectra using the tunneling resonance technique to find the subband energies at finite field. The x value of the AlGaAs was determined to be 0.19 from the absorption edge of the top Al<sub>x</sub>Ga<sub>1-x</sub>As p-type contact, and also from X-ray measurements. We assumed a band offset ratio Q<sub>1</sub>:Q<sub>2</sub> of 67:33 [65]. The shaded bands indicate the energies which correspond to the experimentally determined limits of the activation energy. We see that the activation energy seems to correspond to two possible escape mechanisms: either emission of heavy holes over the top of their confining barrier, or the onset of coupling of electrons to continuum states through just one barrier. Our other results show the sensitivity of the escape process to the barrier thickness at low fields, and so the latter mechanism seems more likely.

The proposed escape mechanism via thermal occupation of higher subbands [17], [24] does not appear to be consistent with our data. Similarly, the simple emission of electrons over the top of their barrier is inconsistent with our results. There is, however, another escape mechanism which would account for the deduced activation energies, namely, phonon-assisted tunneling [31], [66]-[68]. Our measured activation energy is consistent with the energy of the AlAs-like LO-phonon mode in the AlGaAs barrier.
Further work will be needed to determine the escape mechanism more precisely. A question remains as to whether the sweep-out times measured in this way are for electrons or holes, or a convolution of both. The experiment measures the fastest escaping carriers, which are usually presumed to be the electrons. From the discussion in Section II, we do not consider it justified to make such a presumption for the GaAs—AlGaAs system because of the possibility of very rapid hole thermal emission. However, the fact that we do see a minimum at the field for resonant tunneling of electrons does seem to suggest that the sweep-out times refer to electrons only.

VII. CONCLUSIONS

We have demonstrated the importance of the carrier sweep-out physics both for the excitation saturation intensity and for the electroabsorption. We have shown that the excitation saturation intensity can be increased by using low or thin barriers, without significant loss to the electroabsorption performance. This result is particularly important for SEED systems. We have also shown that the excitation saturation intensity increases with applied field. Our time-resolved measurements indicate that the sweep-out mechanism is thermally assisted at low fields. The precise mechanism is yet to be resolved, but is probably either thermal excitation of the electrons to the point where they can couple to continuum states through just a single barrier, or alternatively, optical phonon-assisted tunneling. At high fields, the escape mechanism is probably pure tunneling. The behavior of the holes is still not fully understood, and more work will be needed in this area before arriving at a quantitative analysis of the excitation saturation intensity. An improvement in the understanding of the sweep-out physics should lead to further advances such as the recently demonstrated 33 ps optical SEED switch [4].

ACKNOWLEDGMENT

We are indebted to J. E. Henry and M. M. Becker for the processing of the samples. We would like to thank G. D. Boyd, K. Leo, T. Sizer, and D. S. Chemla for many helpful discussions.

REFERENCES


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