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Langbein, Wolfgang Werner; Gislason, Hannes; Hvam, Jørn Märcher; Zeman, J

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Anisotropic carrier and exciton confinement in T-shaped quantum wires revealed by magneto-photoluminescence

W. Langbein, H. Gislason, J.M. Hvam, J. Zeman* Mikroelektronik Centret, The Technical University of Denmark, Building 345 east, DK-2800 Lyngby, Denmark; E-mail: langbein@mic.dtu.dk

The realization of one-dimensional (1D) semiconductor nanostructures with large confinement energies is of importance for device applications. Different techniques such as growth on tilted substrates (Serpentine superlattices) or prepatterned substrates (V-groove quantum wires) and the cleaved-edge overgrowth of T-shaped structures have been demonstrated. For the T-shaped structures, the confinement energy has recently increased to above 2kT at room temperature by optimizing structure parameters. A decreased exciton diameter in the [110] direction in the wire compared with the [001] well was revealed by magneto-photoluminescence. We determine here the extension of the T-shaped quantum wire (T-QWR) state in both confining directions [110] and [001] to verify its 1D character, as shown for crescent-shaped wires.

The investigated optimized T-QWR sample with 37 meV confinement energy consists of a 18-nm [001] Al0.92Ga0.08As multiple quantum well (QW) structure intersecting with a 4.3-nm [110] GaAs QW, within Al0.92Ga0.08As barriers. It was excited by an Ar+ laser at 514 nm using a multimode fiber. The photoluminescence (PL) at temperatures between 5 K and 30 K was collected by the fiber, dispersed in a monochromator, and detected by a gallium arsenide (GaAs) photomultiplier. A magnetic field from 0 to 26 T was applied by a superconducting and a resistive magnet.

The PL spectra for the field direction [001] are shown in Fig. 1a. It is seen that the T-QWR PL shows a smaller shift to high energies with increasing field than the [001] QW PL. The shift of the T-QWR PL is shown in Fig. 1b for the three principal directions of the field. It shows three distinctly different magnitudes, according to the different projected sizes of the exciton states.

To quantitatively analyze the data, we apply two different models. Using an anisotropic 3D exciton model, we can deduce the three exciton Bohr radii (\(a_{[001]}\), \(a_{[110]}\), and \(a_{[011]}\)) and the exciton binding energy \(E_B\) using the diamagnetic shifts (see Table 1). The result shows an anisotropic exciton wave function and an exciton binding energy of 11 meV, in agreement with recent calculations.

Using a free particle model, we can determine the extension \(w_{[001]}\), \(w_{[110]}\) of the electron wave function along the two confinement directions [001] and [110] (see Table 1).

QWA4 Table 1. Deduced Parameters of Excitonic Transition of T-QWR Sample

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a_{[001]})</td>
<td>2.1 nm</td>
<td>anisotropic</td>
</tr>
<tr>
<td>(a_{[110]})</td>
<td>5.6 nm</td>
<td>anisotropic</td>
</tr>
<tr>
<td>(a_{[011]})</td>
<td>10.2 nm</td>
<td>exciton</td>
</tr>
<tr>
<td>(E_B)</td>
<td>11 meV</td>
<td>confined</td>
</tr>
<tr>
<td>(w_{FWHM} [001])</td>
<td>6.5 nm</td>
<td>carrier</td>
</tr>
<tr>
<td>(w_{FWHM} [110])</td>
<td>10.3 nm</td>
<td>carrier</td>
</tr>
</tbody>
</table>

The QWR and the [001] QW transitions are visible. (b) Energy shift of the QWR PL as a function of the field strength for field directions as labeled.
A. Richter, M. Siiptitz, Ch. Lienau, schemes is a particularly promising technique of near-field scanning optical microscopy (NSOM), offering subwavelength spatial resolution. Near-field scanning optical microscopy (NSOM), offering subwavelength spatial resolution in the 100-nm range in combination with time-resolved detection schemes is a particularly promising technique for such experiments. Here we report on the first time-resolved near-field spectroscopic investigation of the carrier dynamics in single gallium arsenide (GaAs)/(AlGa)As sidewall quantum wires.

Quantum wires (QWR) with a lateral width of 60 nm and a thickness of up to 13 nm embedded by a 6-nm GaAs quantum well (QW) were grown by molecular beam epitaxy of GaAs/(AlGa)As multilayer structures on patterned GaAs(311)A substrates along the [01-1] direction (Fig. 1a). The confinement potential of the QWR and the embedding QW as directly extracted from NSOM photoluminescence excitation (PLE) spectra at 77 K, Fig. 1b, reveals shallow asymmetric energetic barriers with a height of about 18 meV on the mesa top and 14 meV on the mesa bottom in the vicinity of the QWR.

In our experiments, electron-hole pairs are created by transmitting a 6-fs Ti:Sapphire laser pulse at 1.614 eV through an NSOM fiber probe with an aperture diameter of about 200 nm. The laser spectrum overlaps with the excitonic absorption band of the GaAs QW. QWR luminescence at 1.54 eV is detected in the far-field as a function of tip position using time-correlated single-photon counting as the tip is scanned perpendicularly to the QWR. The time-resolved data directly reveal the delayed onset of QWR luminescence for excitation positions between 1 and 3 μm. The experiments directly demonstrate the pronounced influence of spatial bandgap variations on a subwavelength scale on the real-space carrier transfer within the two-dimensional (2D) continuum and thus on the carrier trapping dynamics into the 1D QWR states.

QWA5 Fig. 2. (a) Time resolved QWR luminescence for spatially resolved excitation of the excitonic absorption band of the GaAs QW through the near-field probe tip as a function of tip position along the lateral axis at a sample temperature of 10 K. White regions are regions of high photoluminescence at the detection wavelength. (b) Cross sections through Fig. 2a at fixed excitation positions. Trapping of carriers generated within the flat area QW is strongly suppressed.

The confinement potential of the QWR and the embedding QW as directly extracted from NSOM photoluminescence excitation (PLE) spectra at 77 K, Fig. 1b, reveals shallow asymmetric energetic barriers with a height of about 18 meV on the mesa top and 14 meV on the mesa bottom in the vicinity of the QWR.

At a sample temperature of 10 K, QWR luminescence is observed for localized excitation in a narrow region around the QWR, in between the two energetic barriers separated by about 800 nm. In this region, the capture into the QWR and the energetic relaxation within the QWR is fast, so that with the temporal resolution of our experiment of 250 ps, the temporal dependence of the QWR luminescence is dominated by the QWR recombination lifetime of 0.9 ns, independent of excitation position. For excitation outside the barriers, the diffusive transport of excitons across the barriers is strongly suppressed and trapping of carriers generated within the flat area QW is prohibited.

At a temperature of 100 K a drastically different behavior is found. QWR luminescence is observed for QW excitation at distances of several micrometers away from the QWR. This occurrence of QWR luminescence involves ambipolar drift-diffusion across the barriers. The time-resolved data directly reveal the delayed onset of QWR luminescence for excitation positions between 0.5 and 3 μm.

The experiments directly demonstrate the pronounced influence of spatial bandgap variations on a subwavelength scale on the real-space carrier transfer within the two-dimensional (2D) continuum and thus on the carrier trapping dynamics into the 1D QWR states.

QWA5 Fig. 3. (a) Time resolved QWR luminescence at a sample temperature of 100 K. Experimental conditions as in Fig. 2. (b) Cross sections through Fig. 3a at fixed excitation positions. The time-resolved data directly reveal the delayed onset of QWR luminescence for excitation positions between 1 and 3 μm.

We report here the realization of large third-order nonlinear susceptibility, χ(3) from Audiedielectric (SiO2 and TiO2) composite films near the Au percolation threshold. The composites films were prepared by co-sputtering.

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Large third-order optical nonlinearity in Audiedielectric composite films in femtosecond time scale

H.B. Liao, R.F. Xiao, H. Wang, K.S. Wong, G.K.L. Wong, Department of Physics, Hong Kong University of Science and Technology, Clear Water Bay, Kowloon, Hong Kong

We report here the realization of large third-order nonlinear susceptibility, χ(3) from Audiedielectric (SiO2 and TiO2) composite films near the Au percolation threshold. The composite films were prepared by co-sputtering.