Near-field optical spectroscopy of localized and delocalized excitons in a single GaAs quantum wire

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Excitons in a GaAs quantum wire are studied in high-resolution photoluminescence experiments performed at a temperature of about 10 K with a spatial resolution of 150 nm, and a spectral resolution of 100 μeV. We report an observation of quasi-one-dimensional excitons which are delocalized over a length of up to several microns along the quantum wire. Such excitons give rise to a 10-meV broad luminescence band, representing a superposition of transitions between different delocalized states. In addition, we find a set of sharp luminescence peaks from excitons localized on a sub-150-nm length scale. Theoretical calculations of exciton states in a disordered quasi-one-dimensional potential reproduce the experimental results.

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The interplay between localized and delocalized electronic excitations underlies the fundamental optical and transport properties of low-dimensional systems such as conjugated polymers, electrons at surfaces, and semiconductor nanostructures. In quasi-two-dimensional (2D) semiconductor quantum wells (QW’s) and quasi-one-dimensional quantum wires (QWR’s), the optical spectra in the range of the fundamental band gap are determined by excitons showing a binding energy on the order of 10 meV which increases upon reducing the dimensionality of the semiconductor. In the simplest picture, neglecting any structural disorder in a QW or QWR, such excitons are fully delocalized in space. In contrast to this expectation, extensive experiments with narrow QW’s gave evidence of exciton localization in individual minima of a disorder potential originating from structural imperfections of the material. Recent progress in optical microprobing by microphotoluminescence (μ-PL) and near-field optical spectroscopy has allowed one to resolve the PL spectra from single localized excitons in QW’s and QWR’s, displaying a set of sharp emission lines when recorded with a submicron spatial resolution. The spectral envelopes and spatial properties of such line spectra, which reflect a quasi-zero-dimensional behavior of localized excitons, depend largely on the scale length of interface roughness in the particular system, and have been described by detailed theoretical models.

So far, evidence of the coexistence of and the interplay between localized and delocalized excitonic states has remained scarce. Recent studies of thin QW’s of substantial interface roughness by μ-PL suggest the occurrence of both types of excitons. Spatially resolved PL studies on different QWR structures did not show evidence of delocalized excitonic states, but suggested an optical behavior of QWR’s arising from closely spaced localized, quasi-zero-dimensional excitons. This makes these QWR’s similar to a chain of quantum dots. It thus seems particularly important to clarify the role that delocalized, truly 1D excitonic states play for the optical and transport properties of such nanostructures.

In this paper we report a near-field PL study of excitons confined in a single GaAs QWR. By combining high spatial and spectral resolution, we resolve the characteristic emission features of excitons in a complex quasi-1D disorder potential. We demonstrate the coexistence of single localized excitons and of excitons that are delocalized over a length of up to several microns along the QWR. Our experimental results are well reproduced by theoreical calculations of excitonic states in a disordered QWR.

We investigate a GaAs/(AlGa)As coupled QWR-dot nanostructure grown by molecular-beam epitaxy on a patterned GaAs(311)A substrate. On the substrate, a 15-nm-high mesa with sidewalls alternatingly misaligned by ±30° with respect to the [01-1] direction (zigzag pattern) is prepared by wet chemical etching. The QWR structure consists of a nominally 3-nm-thick GaAs QW layer clad between 50-nm-thick Al0.5Ga0.5As barriers. QWR’s are formed due to the preferential migration of Ga adatoms from the mesa top and bottom toward the sidewalls of the mesa, resulting in a local increase of QW thickness up to 4.5 nm and a lateral confinement. Figure 1(a) shows a schematic top view of the structure in which the QWR follows the zigzag pattern. Every 4.5 μm two QWR segments merge with an angle of 120°. Every other corner the mesa top surrounds the merging of the quantum wires with an angle of 240°. At these positions, an extra local accumulation of GaAs gives rise to the formation of dots. The confinement potential of this type of QWR was studied in earlier photoluminescence (PL) and photoluminescence excitations measurements. The lateral width of the confined region and the quasi-1D confinement energy of excitons are 70 nm and 55 meV, respectively.

A near-field scanning optical microscope for cryogenic temperatures was used in an illumination/collection geometry with combined spatial and spectral resolutions of 150 nm and 100 μeV, respectively. Spatially resolved images were recorded by scanning the probe tip over the sample. For high-resolution measurements, the sample was excited with either a HeNe laser (1.96 eV) or a tunable narrow-band Ti:sapphire laser (bandwidth <200 μeV). PL spectra were re-
made by chemically etching single mode optical fibers. 

A near-field PL spectrum was recorded at a fixed position on the QWR. The spectral widths of the PL peaks range from 200 to 350 μeV.

Two-dimensional near-field images, recorded at the spectral position of the sharp lines [Fig. 2(b)] show that these luminescence contributions stem from regions resolution limited in size. Thus they are attributed to the emission of localized excitons out of local potential minima that arise from monolayer-height fluctuations of the QWR profile. When the detection energy is tuned to the low-energy side of the broad continuum (out of the sharp resonances), the PL spatial distribution reveals the existence of regions with a larger average extension of 400–600 nm in diameter [Fig. 2(c)]. For detection energies in the high-energy part of the emission band, a completely different situation appears. Here the two-dimensional images [Fig. 2(d), detection at 1.674 eV, and excitation at 1.96 eV] indicate a uniform PL distribution delocalized along the QWR axis on a length scale of more than 2 μm. An even larger extension is measured in the region where the QWR’s are not interrupted by dots [Fig. 2(c), excitation at 1.70 eV]. In this case the spatial distribution of the PL signal extends over the whole QWR across the corner.

Delocalized PL components are observed for a broad range of detection energies. The corresponding spectral dependence is shown in Fig. 2(a) by the dotted curve. The shape of this spectrum is close to a Gaussian centered at 1.6740 eV with a full width at half maximum of 10 meV. Similar spectra were measured in different regions of the sample, and are independent of the excitation photon energy in the range between 1.70 and 2 eV.

Figure 3 shows the power dependence of the near-field PL spectra recorded at a fixed position on the QWR. The excitation power was varied over more than three orders of magnitude with values between $P=0.002$ and $5.6 \mu W$, corresponding to carrier densities between $1.4 \times 10^2$ and $4 \times 10^4$.

As the combined spatial and spectral resolution is decreased below 200 nm/100 μeV, the broad PL bands break up into a set of sharp and intense emission peaks superimposed on a broad continuum [Fig. 2(a)]. The near-field PL spectrum was recorded at a fixed position on the QWR. The spectral widths of the PL peaks range from 200 to 350 μeV.

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sites with less than 150-nm diameter, spectral and spatial features: reveal three emission components with distinctly different significantly to the PL spectra even at the lowest excitation length $L_c$ and a Gaussian correlation function in space of correl- $s$ and 5.6 $m$ $E_X$ $\sigma$ of excitons which are delocalized along the QWR over a length of up to several microns and cover a 10-meV range of emission energies. This delocalized component is present even at very low excitation densities, where only a minor fraction of the available exciton states in the QWR is populated and—thus—nonlinear effects of state and/or band filling can safely be neglected.

The occurrence of the three PL components clearly demonstrates that disorder induced local variations of the 1D confinement potential of the QWR and the interplay between localized and delocalized states play a central role for the electronic properties of this type of QWR structure. We performed a theoretical study of the excitonic spectra in order to develop a quantitative understanding of the disordered QWR structure. In this model, we consider a QWR oriented along the $x$ axis with a lateral confinement potential $V(y)$ of a width of 70 nm and a confinement energy of 55 meV, superimposed by a randomly generated potential [Fig. 4(a)]. The random potential is characterized by a Gaussian energy distribution of width $\sigma$ and a Gaussian correlation function in space of correlation length $L_c$. In the $(x,y)$ plane, excitons are subject to local energy fluctuations both along the QWR axis ($x$) and perpendicular to it ($y$). Following the procedure outlined in Ref. 9, the in-plane exciton center-of-mass (COM) eigenenergies and wave functions $\psi(R)$ [$R=(X,Y)$ is the COM coordinate] are calculated by numerically solving the two-dimensional Schrödinger equation for the COM motion within the local potential. As discussed in Ref. 19, this approximation of a factorized COM motion is appropriate for disorder strengths of less than 10 meV, i.e., the typical exciton binding energy $E_X$ in thin quantum-well and wire nano-

structures. Independent experimental evidence for the validity of this assumption is obtained from the power-dependent PL spectra [Fig. 3]. These spectra show no signatures of charged exciton transitions, that are expected in the limit $\sigma \gg E_X$, where electrons and holes localize separately.

For a disorder strength of a several meV, that is typical for the considered thin nanostructures, three types of $1s$ exciton states are found in the QWR: (i) excitons localized in a single potential minimum [Fig. 4(b)] with a nearly Gaussian shaped wavefunction $\psi(R)$. The extension of $\psi(R)$ is defined by $L_c$ along the QWR axis ($x$) and by the lateral width of the potential minimum along $y$. These localized excitons couple strongly to light with a matrix element $M \approx D | \psi(R) |$ ($D$ is the optical dipole matrix element of the excitonic $1s$ transition) and give rise to sharp emission peaks.
in the near-field spectra. The axial density of these localized excitons is about 1/4Lc, and they are distributed over an energy range of roughly 2σ. Comparison with the average number and energy distribution of emission lines in Figs. 2(a) and 3 allows one to approximate the free parameters of the model as Lc = 20 nm and σ = 5 meV. (ii) At low transition energies, we find several eigenstates with wave functions that extend over few local potential minima, thus showing extensions between 50 and 200 nm [Fig. 4(c)]. Their coupling to light is reduced with respect to that of localized excitons. Comparison to experiment indicates that this class of wave functions gives rise to the emission from localized regions extending beyond the spatial resolution of our experiment in the low-energy part of the PL spectrum [Fig. 2(c)]. (iii) We find a large density of about 10 states/ (μm meV) that are delocalized along the QWR with wave-function extensions between several 100 nm and several μm. A representative wave function with an extension of about 600 nm is shown in Fig. 4(d). The wave functions of these eigenstates show highly complex shapes which represent disorder-induced interferences between a broad distribution of plane waves with momentum Kc centered around an average (Kc). The fraction of this distribution with momentum Kc < Kmax = EX nlhc (Ex: is the exciton transition energy, n: refractive index) couples to light and contributes to absorption and emission spectra with a matrix element M that is considerably reduced compared to that of localized exciton states.21 In the optical spectra, these delocalized states give rise to a broad distribution of densely spaced weak resonances. If the linewidth of the individual resonances is broader than their energy separation, individual transitions are not resolved, and broad structureless bands occur in the absorption and—for sufficient thermal population of emitting states—also in the PL spectrum. We attribute the spectrally broad and spatially delocalized luminescence in our near-field measurements [Figs. 2(d) and 2(e)] to those delocalized excitons.

In Fig. 4(e), the overall optical spectrum calculated from the model is summarized. The square of the transition moment |M|2 is plotted as a function of energy for both localized [components (i) and (ii)] and delocalized exciton states. In agreement with our experiments, we find a broad quasi-continuous contribution from transitions between delocalized states, superimposed by individual transitions between localized levels. Both contributions occur in a similar range of transition energies. In the luminescence spectra, the relative contribution from individual eigenstates α is proportional to |Mα|2Na, with Na being the average occupation of α. In a quasiequilibrium distribution of localized and delocalized excitons, Na is determined by the energy of the exciton state and the temperature Tc of the exciton gas. The overall width of the measured delocalized PL component [circles in Fig. 2(a)] is well reproduced for a value of Tc ~ 30–40 K, i.e., somewhat higher than the lattice temperature of nominally 10 K. The elevated exciton temperature is due to the non-resonant excitation with excess energies of several 10 meV (also see Ref. 22).

A comment should be made on the scattering processes underlying the luminescence linewidths. At low temperatures of the exciton gas, acoustic-phonon scattering represents the main inelastic relaxation mechanism. A calculation of scattering matrix elements with the exciton wave functions derived in our model gives phonon emission times of 5–10 ps from delocalized exciton states for a lattice temperature of 10 K. This corresponds to characteristic line broadenings of 150–300 μeV. Such linewidths are larger than the energy separation between delocalized states, resulting in a broad structureless emission band as observed in our measurements. Phonon absorption times of localized excitons vary strongly between the different minima of the disorder potential. They are on the order of 10 to 30 ps at 10 K, corresponding to linewidths of 50–150 μeV, similar to what is found in the PL spectra.23

In conclusion, the coexistence of localized and delocalized excitons in a quasi-1D GaAs quantum wire was observed in luminescence experiments with high spectral and spatial resolution. Excitons delocalized over a length of up to several microns give rise to a broad luminescence band. In contrast, excitons localized in minima of the quantum wire disorder potential lead to a set of narrow individual emission lines. Theoretical calculations of exciton states in a disorder potential account for this behavior, and show that the observed broad luminescence band consists of many individual transitions between delocalized states, each showing a spectral broadening larger that the mutual energy separation. Coupling between localized and delocalized states plays an important role for exciton relaxation and for excitonic transport along the quantum wire, where contributions from both ballistic and diffusive motion are expected.

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23. The calculated scattering times are sufficiently high to establish a quasiequilibrium exciton distribution close to the lattice temperature within a luminescence decay time of about 1 ns (see Ref. 17).