Effect of spectral broadening and electron-hole scattering on carrier relaxation in GaAs quantum dots

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Luminescence efficiency in quantum dots has been a matter of some controversy recently. Theoretically, poor efficiency has been predicted owing to the phonon bottleneck in carrier relaxation, while slightly enhanced luminescence has been reported in several experiments. The approach of this letter differs from previous theoretical work in that the scattering rates are computed self-consistently accounting for the spectral broadening of the electronic spectra due to a finite energy level lifetime. Scattering of electrons and holes confined in the dot is found to be responsible for breaking the phonon bottleneck in electron relaxation reducing the relaxation time from several ns to several hundred ps. Results of a Monte Carlo simulation also including confined and interface polar optical phonon and acoustic phonon scattering for a range of quantum dot dimensions and temperatures are presented. These results may provide an explanation of the absence of a significant reduction in quantum dot luminescence compared with that from quantum wells.

In this letter, we present a theoretical analysis of electron relaxation in GaAs/Al$_{0.3}$Ga$_{0.7}$As quantum dots based on a Monte Carlo simulation of the equilibration process with the inclusion of self-consistently calculated electron-phonon and electron-hole scattering processes. An idealized flat-band potential profile is assumed, i.e., no attempt is made to describe the band-bending properties of the GaAs/AlGaAs system resulting from a particular fabrication process. The electronic spectrum is found by solving the Schrödinger equation in the effective mass approximation, and the hole states are derived by diagonalizing the four-band Kohn–Luttinger Hamiltonian, in which the standard prescription for calculating quantum confined states $[k_z \rightarrow -i(\partial/\partial x), \text{etc.}]$ is used. In order to estimate the electron-hole scattering rate, for the sake of computational simplicity, the off-diagonal coupling terms in the Hamiltonian are set to zero. The eigenenergies and eigenfunctions are found by discretizing the volume of the dot on a uniform spatial mesh terminated far away from the dot interfaces, approximating derivatives as finite differences, and solving the resulting matrix equation iteratively.

The macroscopic electrostatic model has proved useful in approximating the spatial structure and dispersion relations for phonon models in quantum wells and wires. We assume that the confined phonon modes are given by linear combinations of products of sines and cosines, whose wave vector is fixed at multiples of $\pi/L_i$, where $i=x, y, z$, by imposing the boundary condition that the phonon mode vanish near the interface with a material with a different resonant vibration frequency. Since the nonepitaxial dimensions greater than 20 nm are examined in this letter, we assume quantum well interface phonon modes varying as hyperbolic sines and cosines with the dispersion relation derivable from the boundary condition. The interface phonon energies lie near the interface with a material with a different resonant vibration frequency. Since the nonepitaxial dimensions greater than 20 nm are examined in this letter, we assume quantum well interface phonon modes varying as hyperbolic sines and cosines with the dispersion relation derivable from the boundary condition. In this letter, we present a theoretical analysis of electron relaxation in GaAs/Al$_{0.3}$Ga$_{0.7}$As quantum dots based on a Monte Carlo simulation of the equilibration process with the inclusion of self-consistently calculated electron-phonon and electron-hole scattering processes. An idealized flat-band potential profile is assumed, i.e., no attempt is made to describe the band-bending properties of the GaAs/AlGaAs system resulting from a particular fabrication process. The electronic spectrum is found by solving the Schrödinger equation in the effective mass approximation, and the hole states are derived by diagonalizing the four-band Kohn–Luttinger Hamiltonian, in which the standard prescription for calculating quantum confined states $[k_z \rightarrow -i(\partial/\partial x), \text{etc.}]$ is used. In order to estimate the electron-hole scattering rate, for the sake of computational simplicity, the off-diagonal coupling terms in the Hamiltonian are set to zero. The eigenenergies and eigenfunctions are found by discretizing the volume of the dot on a uniform spatial mesh terminated far away from the dot interfaces, approximating derivatives as finite differences, and solving the resulting matrix equation iteratively.

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Fröhlich form of the interaction Hamiltonian from the Fermi Golden Rule:

\[
W_{if}^{\text{POP}} = \frac{2\pi}{\hbar} \sum_{l,m,n} \frac{e^2 \hbar \omega_{\text{LO}}}{2L_x L_y L_z} \left( \frac{1}{\varepsilon_r} - \frac{1}{\varepsilon_0} \right) \\
\times \left[ \left( \frac{\pi l^2}{L_x} + \frac{\pi m^2}{L_y} + \frac{\pi n^2}{L_z} \right)^2 \right]^{-1} I_{lf}^2 \\
\times \left( \frac{n(\omega_{\text{LO}}) + \frac{1}{2} \pm \frac{1}{2}}{M_{if}^{\text{POP}}(l,m,n)} \right)^2 \\
\times \delta(E_f - E_i \pm \hbar \omega_{\text{LO}}),
\]  

where \( I_{lf} \) is the overlap integral for the cell periodic parts of the electron wave function, presumed unity in this work, and the rest of the symbols have their usual meaning. The upper sign corresponds to phonon emission, the lower sign to absorption. Taking an arbitrary vertex of the box as origin, the overlap integral for the envelope parts of the wave function can be written down simply as

\[
M_{if}^{\text{POP}}(l,m,n) = \int d^3r \varphi_i^*(r) \sin \left( \frac{mlx}{L_x} \right) \sin \left( \frac{mx}{L_y} \right) \sin \left( \frac{nx}{L_z} \right) \varphi_f(r). 
\]  

The delta function in Eq. (1) is replaced by a Gaussian line shape function:

\[
\delta(E_f - E_i \pm \hbar \omega_{\text{LO}}) \rightarrow \frac{1}{\sqrt{1.44 \sigma^2}} \exp \left( -\frac{(E_f - E_i \pm \hbar \omega_{\text{LO}})^2}{1.44 \sigma^2} \right). 
\]  

An iterative self-consistent calculation of the scattering rates and the linewidth \( \sigma = \hbar / \tau \), where \( \tau \) is taken to be the reciprocal of the scattering rate, is necessary to determine \( W_{if}^{\text{POP}} \) from Eqs. (1)–(3). The scattering rates have also been calculated for interface phonon modes and have been found to be lower by two orders of magnitude than the rates for confined bulklike phonons. Consequently, even for phonon energies in the range between 33 and 36 meV, confined phonon scattering dominates, with the energy difference made up by the broadening of the electronic spectrum. By assumption, the lattice is maintained at a constant temperature.

In quantum wires, acoustic phonon (AP) scattering is one of the mechanisms responsible for electron relaxation for quantum wires of cross sections less than 100 Å×100 Å. In quantum dots, the AP scattering rate can be calculated using the Fermi Golden Rule and the deformation potential theory:

\[
W_{if}^{\text{AP}} = \frac{\pi}{\rho \omega q} \sum_q D^2 q^2 \frac{1}{\omega q} \left( n(\omega_q) + \frac{1}{2} \pm \frac{1}{2} \right) \\
\times \left| \int d^3r \varphi_i^*(r) \exp(\pm iq \cdot r) \varphi_f(r) \right|^2 \delta(E_f - E_i \pm \hbar \omega_q),
\]  

where \( \rho \) is the mass density of GaAs, \( D_A \) is the acoustic deformation potential, and \( q \) is the phonon wave vector with a continuous spectrum. We include acoustic phonon scattering in our Monte Carlo simulation, although in small quantum dots, the energy levels are separated by energies available only to short-wavelength AP. The matrix element for short phonon wavelength averages out to nearly zero, and the AP rates are by several orders of magnitude smaller than POP modes. Due to the weaker coupling of acoustic phonons to electrons, the broadening linewidth is also much smaller than that for POP.

The Pauli exclusion principle has to be considered in the occupation of discrete quantum dot levels. However, because of the small size of dots under study, it is permissible to assume that only one electron is injected in any one quantum dot. With this assumption, the electron-electron interaction and screening can be ignored. Previous treatments of electron relaxation mechanisms in quantum dots have not included electron-hole (EH) scattering. Since the hole confinement energy is much smaller than that for the electron, the hole distribution can be assumed to be in thermal equilibrium with the lattice. Therefore, the electron may lose energy by interacting with a thermalized hole in the valence band. The matrix element can be computed in the Born approximation:

\[
M_{if}^{\text{EH}} = \int d^3r \int d^3r_h \sum_{\nu=1}^{13/2,-13/2} \varphi_j^*(r_x) \varphi_i^*(r_h) \frac{e^2}{4\pi \varepsilon_0} \\
\times \frac{1}{|r_x - r_h|} \varphi_j(r_x) \varphi_i(r_h),
\]  

where the sum runs over the \( |3/2, \pm 3/2| \) and \( |3/2, \pm 1/2| \) angular momentum states. The scattering rate is calculated by applying the Fermi Golden Rule with the broadening found in an iterative fashion as described above.

Given the relevant scattering rates, electron relaxation can be modeled by a Monte Carlo simulation. Electrons are injected at the top of the potential barrier in a thermal distribution, and their evolution is followed in energy and time. An averaging over a large number of electrons (and equivalently, dots) is necessary in order to produce statistically reliable results. In Fig. 1 the evolution of the mean energy of the electron distribution in the 50 Å×250 Å×250 Å GaAs/Al_{0.33}Ga_{0.67}As quantum dot at 300 K is shown. In absence of EH scattering, the main energy loss mechanism is POP emission with a lifetime-broadened energy spectrum. Nevertheless, POP scattering is unable to provide coupling between states whose separation is different from 36 meV by
At lower temperatures, phonon and EH scattering are radiative recombination time in most samples. However, if more than several meV. As a result, the relaxation time reaches several ns, thus becoming comparable with the non-radiative recombination time in most samples. However, if EH scattering is taken into account, the bottleneck is broken, and the electron relaxation time decreases to a little over 100 ps. At lower temperatures, phonon and EH scattering are more efficient since fewer phonons are available for absorption, and the hole distribution is sharper. Therefore, the effect of EH scattering is similar yet more pronounced, with a relaxation time of less than 30 ps at 70 K.

A brief comment on the accuracy of the assumption of the equilibrium hole distribution is in order. Since the hole confinement energies are much smaller than the electron confinement energies, and the hole density of states is much higher than the electron density of states, the hole spectrum is broadened as to be nearly continuous. In these circumstances, hole relaxation is likely to resemble relaxation of electrons in quantum wires, for which the relaxation time does not exceed 100 ps. Since we have estimated the electron relaxation times in quantum dots to be of the order of several hundred ps at room temperature. The equilibration time is reduced by lowering temperature and expanding the size of the dot. However, it remains much larger than the relaxation time in quantum wires (several tens of ps) at 300 K. Therefore, the modulation properties of quantum dot lasers, for which very low threshold currents have been predicted, can be impaired.

In brief summary, we have presented the results of a Monte Carlo simulation of electron relaxation in 20–45 nm quantum dots with the inclusion of the broadening of the energy spectrum and coupling to confined and interface phonon modes and of the consequences of electron-hole scattering. The latter is the mechanism principally responsible for breaking the bottleneck which occurs as a result of a sharp LO phonon spectrum and inefficiency of acoustic phonon scattering. The electron relaxation time in a 50 Å×250 Å×250 Å dot has been estimated to be several hundred ps at room temperature. The equilibration time is reduced by lowering temperature and expanding the size of the dot. However, it remains much larger than the relaxation time in quantum wires (several tens of ps) at 300 K. Therefore, the modulation properties of quantum dot lasers, for which very low threshold currents have been predicted, can be impaired.

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