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Simulation of a GaAs quantum dot embedded into a AlGaAs nanowire

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Abstract—We report on a numerical model of quasi one-dimensional and quasi zero-dimensional semiconductor heterostructures. This model is strictly based on experimental structures of cylindrical nanocolumns of AlGaAs grown by molecular-beam epitaxy in the (111) direction. The nanocolumns are of 20 - 50 nm in diameter and 0.5 - 1 µm in length and contain a single GaAs quantum dot, of 2 nm in thickness and 15 - 45 nm in diameter. Since the crystal phase of these nanowires spontaneously switches during the growth from zincblende to wurzite structures we implement a continuum elastic model and 8 band $k\cdot\hat{p}$ model for polymorph crystal structures. The model is used to compute electromechanical fields, wave-function energies of the confined states and optical transitions. The model compares a pure zincblende structure with a polymorph in which the zincblende disk of GaAs is surrounded by wurzite barriers and results are compared to experimental photoluminescence excitation spectra. The good agreement found between theory and features in the spectra supports the polyphorm model.

Index Terms—Nanowire, quantum dot, polymorph materials.

I. INTRODUCTION

Since the discovery of self-assembled quantum dots (QD), the field of optoelectronics has moved interest on nanoobjects of different dimension, size and shape in the search of novel useful properties. The challenge of the QD fabrication technology is to obtain highly homogeneous arrays with little dispersion in size, composition and shape in order to maximize quantum efficiency of LEDs and lasers. This requirement has been solved only partially with self-organization[1] that allows to achieve a rather uniform distribution of the dot sizes with dispersion of 10%[2]. However, the technique has a number of essential ineradicable lacks. For instance, because of the presence of a wetting layer, it is impossible to reach high quantum efficiencies. Furthermore, this growth process offer little to no control over the dot size, an essential parameter in order to tune the emission wavelength.

Thanks to the fast development and improvement of control in the growing conditions of nanostructures research has moved to new solutions in nano-engineering quantum confined states. The growth of free standing semiconductor nanowires (NW) and nanocolumns (NC) is one of the most promising solution for application in nanoelectronics [3], and nanophotonics [4] thanks to a better relaxation of the strain field that allows more freedom in combining III-V materials. The technology of manufacturing such NWs is now intensively developing [5]-[15]. These NWs can include one or several quantum dots (or quantum disks). Modern epitaxial methods allows to control the sizes and location of NW at the nanometer scale, that favorably distinguishes such QDs from planar and is very attractive for applications in optoelectronic devices.

However, GaAs/AlGaAs NWs also possess growth mechanisms that might have strong influences on their electronic properties. It is known [16] that the crystal structure of a $<111>$ nanowire may spontaneously switch during the growth from zinc-blende (Zb) to wurzite (Wz) phases. The effect leads to a strong modification of the time resolved photoluminescence spectra in the range of the optical transitions within the barrier material [13]. Similarly, it is expected that optical transitions in the QDs are sensitive to the crystal phase of the dot itself and its surrounding media. For example the energy position of the ground and excited states in the QD may be influenced by the electric field originating from the crystal phase transition. To observe the effect of this transition it is necessary to make a comparison between experimental data obtained by time resolved photoluminescence (PL) and photoluminescence excitation (PLE) measurements and theoretical simulations.

The electronic structures and the variation of the fundamental gap of the Wz phase of these crystals have been only quite recently studied. Knowledge of the full band structure of the these polymorph materials is fundamental in order to possibly extract parameters for tight-binding [18] (TB) and $k\cdot\hat{p}$ [17] model calculations. However, at present time only a relative
small amount of parameters for bandstructure calculations are available in the literature.

Quasiparticle calculations of newly observed Wz polymorphs of InAs and GaAs has been reported in Ref.19 using the GW approximation. De et al. have presented calculations of the bulk electronic band structures of non-nitride III-V semiconductors in the Wz phase using empirical pseudopotentials including spin-orbit coupling20, while Jancu et al. have studied the electronic structure of the Wz phases of GaAs and InP using TB model and calculate the optical properties of crystal phase quantum dots formed by thin Zb layers inserted in a Wz-InP NW21. Since the Zb and Wz structures differ only by the arrangement of the second neighbors of a given atom, they used the same set of parameters for the two phases, while the atomic positions are given as input data and lattice constants are precisely determined by x-ray diffraction.

Very recently quasiparticle self-consistent GW calculations have been used to analyze the band structure of GaAs in Wz phase.22. Here, also the Kohn-Luttinger and Rashba-Shekab-Rikus effective Hamiltonian band structure parameters of Wz GaAs have been determined from these first-principles band structure calculations.

In this work, we have simulated an AlGaAs NW with an embedded GaAs QD, considering two different cases, a pure Zb case and a polymorph Wz-Zb case, respectively. Dimensions, structural and geometric parameters have been strictly derived from experimental data. We have implemented an 8-band k · p model29 for bandstructure calculations, and the parameters for the Pikus-Bir Hamiltonian of the Wz materials have been derived from the correspondent Zb materials using the cubic approximation as described by Chuang et al.32, while the stiffness parameters of Wz crystals have estimated from Martin’s transformations27, 28 using the Zb parameters.

The paper is organized as follows. In the first section we present the experimental results that have inspired our numerical investigation. In section III we describe the models and the way we have derived all the parameters needed in the simulations, while all the numerical results are discussed in section IV. Finally, we give our conclusions.

II. EXPERIMENT

In these studies we have considered an AlGaAs NW with an embedded GaAs QD. AlGaAs NWs were grown on GaAs (111)B semi-insulating substrate using an EP1203 MBE system equipped with solid sources supplying Ga and Al atoms, and an As effusion cell to produce tetramers. The Al content along the NW and within the wire cross-sections was monitored by Raman spectroscopy in order to exclude possible inhomogeneities. We have found Al concentration in the range c = 0.24 – 0.26. The growth was completed with deposition of GaAs to avoid possible oxidation of the AlGaAs shell layer. We did not perform any growth interruption at the heterointerfaces. The diameter of NWs in our samples is in the range 20-50 nm. Scanning electron microscopy (SEM) and transmission electron microscopy (TEM) characterizations were performed before the optical measurements.

The samples were excited by either second harmonic of Nd:YAG laser or doubled Ti:Sa laser with 100 fs pulse duration and 70 MHz frequency repetition. The average power of the lasers was about 10 µW, the spot diameter was 1.5 microns. Jobin-Yvon 30 cm monochromator and CCD detector were used for registration of the spectra. We used microscope to focus the laser beam and to collect the signal in the micro PL measurement of the single NW. The minimal diameter of the laser spot was 1.5 microns that allow us to extract the signal from a single NW. Additionally the set-up allows measure spectra with time resolution.

The TEM image of the single AlGaAs NW with a slice of GaAs is shown in the insert of Figure 1. The dark area corresponds to the single GaAs QD. It is clearly seen that the interfaces along the axis of growth and along the radius are rather sharp. Considering cylindrical symmetry, the GaAs QD looks like a disk surrounded by the wide-gap semiconductor AlGaAs. The diameter of the NW in our samples was 25-50 nm, the length 1000 nm, and the thickness of the QD was of several mono-atomic layers, i.e., 2-5 nm.

Figure 1 shows PL spectra taken from a single QD embedded into a NW. At low excitation intensities of ≈ 1 W/cm² a single narrow line is observed at the energy of 1.6525 eV. The width of this line is 0.2 meV and mainly limited by the spectral resolution of our spectrometer. It is natural to connect this line with the annihilation of an exciton in the single QD. This figure shows also the spectrum of photoluminescence excitation (PLE) taken from the same dot. The distance between the absorption (PLE) lines corresponds to the “transverse” carrier quantization in the dot of 15 nm in diameter. The energy of the “longitudinal” quantization is ≈130 meV, which corresponds to the QD thickness of 2.5 nm. In the PLE spectrum one can see very broad absorption lines at high energy; we connect these lines with carrier absorption in the NW itself.

In Figure 2 a TEM image of one of our AlGaAs NW clearly shows the evidence of phase transitions from Zb to Wz crystal...
The total mechanical and electrical free energy change $dU$ of a piezoelectric medium can be written as [24]:

$$dU = dU^{\text{mech}} + dU^{\text{elec}} = TdS + \sigma_{ik}d\varepsilon_{ik} + E_idD_i,$$

where $T$, $S$, $\sigma_{ik}$, $\varepsilon_{ik}$, $E_i$, and $D_i$ are the temperature, entropy, stress tensor, strain tensor, electric field, and electric displacement, respectively.

With isentropic conditions ($dS = 0$) and using crystal symmetry considerations the elastic energy for a crystal with Zincblende (Zb) symmetry read [25]:

$$U^{\text{mech}}_{\text{Zb}} = \frac{1}{2} \left[ C_{11,\text{Zb}}(\varepsilon_{xx}^2 + \varepsilon_{yy}^2 + \varepsilon_{zz}^2) + 2C_{12,\text{Zb}}(\varepsilon_{xx}\varepsilon_{yy} + \varepsilon_{xx}\varepsilon_{zz} + \varepsilon_{yy}\varepsilon_{zz}) + 4C_{44,\text{Zb}}(\varepsilon_{xy}^2 + \varepsilon_{xz}^2 + \varepsilon_{yz}^2) \right],$$

where $C_{ij,\text{Zb}}$ are the linearly independent stiffness constant parameters for a Zb structure.

Similarly, for a crystal with Wurtzite (Wz) symmetry we have for the elastic energy [25]:

$$U^{\text{mech}}_{\text{Wz}} = \frac{1}{2} \left[ C_{11,\text{Wz}}(\varepsilon_{xx}^2 + \varepsilon_{yy}^2 + \varepsilon_{zz}^2) + 2C_{12,\text{Wz}}(\varepsilon_{xx}\varepsilon_{yy} + 2C_{13,\text{Wz}}(\varepsilon_{xx} + \varepsilon_{yy}) + 4C_{44,\text{Wz}}(\varepsilon_{xy}^2 + \varepsilon_{xz}^2 + \varepsilon_{yz}^2) + 2(C_{11,\text{Wz}} - C_{12,\text{Wz}})\varepsilon_{xy}^2 \right],$$

(3)

where $C_{ij,\text{Wz}}$ are the linearly independent stiffness constant parameters for a Wz structure.

While the stiffness parameters for Zb GaS, AlAs and AlGaAs alloy can be found in literature [26], we do not have available the correspondent Wz parameters, so we have estimated them from Martin’s transformations [27], [28] using the Zb parameters:

$$C_{11,\text{Wz}} = \frac{1}{6}(3C_{11,\text{Zb}} + 3C_{12,\text{Zb}} + 6C_{44,\text{Zb}}) - 3\Delta^2/(C_{11,\text{Zb}} - C_{12,\text{Zb}} + 4C_{44,\text{Zb}})$$

$$C_{12,\text{Wz}} = \frac{1}{6}(C_{11,\text{Zb}} + 5C_{12,\text{Zb}} - 2C_{44,\text{Zb}}) + 3\Delta^2/(C_{11,\text{Zb}} - C_{12,\text{Zb}} + 4C_{44,\text{Zb}})$$

$$C_{13,\text{Wz}} = \frac{1}{6}(2C_{11,\text{Zb}} + 4C_{12,\text{Zb}} - 4C_{44,\text{Zb}})$$

$$C_{33,\text{Wz}} = \frac{1}{6}(2C_{11,\text{Zb}} + 4C_{12,\text{Zb}} + 8C_{44,\text{Zb}})$$

$$C_{44,\text{Wz}} = \frac{1}{6}(2C_{11,\text{Zb}} - 2C_{12,\text{Zb}} + 2C_{44,\text{Zb}}) - 6\Delta^2/(C_{11,\text{Zb}} - C_{12,\text{Zb}} + 4C_{44,\text{Zb}}),$$

(4)

where

$$\Delta = \frac{\sqrt{2}}{6}(C_{11,\text{Zb}} - C_{12,\text{Zb}} - 2C_{44,\text{Zb}}).$$

(5)

The piezoelectric field is included directly in our fully-coupled continuum model [23], and the not available Wz piezoelectric parameters have been approximated using parameters of well known Wz materials, as GaN, AlN and AlGaN alloy.

Expressions from 1 to 5 have been used to derive the governing equations of the electromechanical fields of the crystal [25].

### B. $\vec{k} \cdot \vec{p}$ method

We have used for band structure calculations 8-band $\vec{k} \cdot \vec{p}$ model [29], which describes electron, heavy-hole, light-hole, and spin-orbit split-off bands around the $\Gamma$ point of the Brillouin zone and treats all other bands as remote bands. An $8 \times 8$ effective-mass Hamiltonian has been implemented, based on Foreman’s application of Burt’s exact envelope function theory to planar heterostructures [30].

The wave function of a state $n$ with energy $E_n$ is given by a linear combination of the eight Bloch parts weighted by the respective envelope functions,

$$\psi_n(r) = \sum_{i=1}^{8} \phi_i(r) u_{iT}(r),$$

(6)

where $\phi_i$ are the envelope function and $u_i$ are the Bloch states at $\vec{k}=0$ [31].
The Luttinger-Kohn parameters the Zb \( \vec{k} \cdot \vec{p} \) Hamiltonian (\( L_1, L_2, M_1, M_2, M_3, N_1 \), and \( N_2 \)) have been taken from Ref.[26], while the \( A_i \) parameters in the Pikus-Bir model for Wz have been derived from the correspondent Zb materials using the cubic approximation as described in Ref.[32]:

\[
A_1 = -\gamma_1 - 4\gamma_2, \quad A_2 = -\gamma_1 + 2\gamma_2, \quad A_3 = 6\gamma_2, \\
A_4 = -3\gamma_2, \quad A_5 = -3\gamma_3, \quad A_6 = -3\sqrt{2}\gamma_3,
\]

GaAs Pikus-Bir parameters are in a good agreement with the corresponding parameters given by GW calculations in Ref.[22]. For the band offset of the GaAs Wz material we refer to Ref.[16], and we have assumed an equivalent offset for the AlAs Wz material.

Results for electromechanical fields with the continuum model are included in the framework of the \( \vec{k} \cdot \vec{p} \) as it is usually done via deformation potentials [33].

The complete set of parameters for the Wz materials we have used for our simulations is given in Table I, with the same usual notation as in Ref.[26].

<table>
<thead>
<tr>
<th>Parameters</th>
<th>GaAs Wz</th>
<th>AlAs Wz</th>
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<td>( a ) (nm)</td>
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<td>0.4002</td>
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<td>( c ) (nm)</td>
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<td>( E_0 ) (eV)</td>
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<td>( \Delta_{\sigma} ) (eV)</td>
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<td>( \Delta_{\pi} ) (eV)</td>
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<td>0.019</td>
</tr>
<tr>
<td>( m_| ) (eV)</td>
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<tr>
<td>( m_\perp ) (eV)</td>
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<td>0.320</td>
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<td>( A_2 )</td>
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<td>( A_3 )</td>
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<td>( A_4 )</td>
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<td>( A_5 )</td>
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<td>-4.26</td>
</tr>
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<td>( A_6 )</td>
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<tr>
<td>( VBO ) (eV)</td>
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<tr>
<td>( D_3 ) (eV)</td>
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<tr>
<td>( D_4 ) (eV)</td>
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<td>3.9</td>
</tr>
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<td>( D_5 ) (eV)</td>
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<tr>
<td>( D_6 ) (eV)</td>
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<tr>
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<td>( C_{12} ) (GPa)</td>
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<td>( C_{13} ) (GPa)</td>
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<td>( C_{33} ) (GPa)</td>
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<td>( C_{44} ) (GPa)</td>
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<td>( e_{33} ) (C/m²)</td>
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<tr>
<td>( e_{15} ) (C/m²)</td>
<td>-0.11</td>
<td>-0.24</td>
</tr>
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Table I

| Parameters for the Wz Structures Used in the Simulations. |

Optical properties are calculated in the dipole approximation, starting from the dipole matrix elements \( \mu_{nm} \), between states \( n \) and \( m \), respectively. These are calculated from the momentum matrix elements which is represented in \( k \)-space as [34]

\[
\vec{p}_{nm} = \frac{m_0}{\hbar} \frac{\partial H_{nm}}{\partial \vec{k}}.
\]

Finally, oscillator strengths for the optical transitions are given by [36]:

\[
\bar{F}_{nm} = \frac{2\pi e^2 \hbar^2}{\epsilon_0 m_e^2 V} |\hat{e} \cdot \vec{p}_{nm}|^2,
\]

where \( \hat{e} \) is the direction unit vector of the electric field of the linearly polarized incident light, \( V \) is the quantum dot volume, \( e \) is the electron charge, and \( \epsilon_0 \) is the vacuum permittivity.

C. Geometry and structure

In our simulations we have considered an Al,Ga\(_{1-c}\),As nanowire (\( c = 0.25 \)) with a diameter of 40 nm and height of 50 nm, with an embedded GaAs QD with a diameter of 20 nm and an height of 5 nm. Two different cases have been studied and compared: a pure Zb case, where the whole structure is assumed to be grown in pure Zb crystal phase, and a mixed structure as shown in Figure 3, where the GaAs QD is sandwiched by two adjacent 8 nm high Wz layers and surrounded by a Zb AlGaAs shell. The mixed structure is completed by Zb layers at the top and at the bottom.

We will refer in the following to this structure as the mixed (Mx) case.

IV. Results

A. Electromechanical fields

The electromechanical fields (strain and piezoelectric fields) are evaluated as described in section III-A. In Figure 4 we plot the magnitude of the absolute value of the strain field for the pure Zb and the Mx structures, respectively.

It is interesting to notice that in the Mx case, besides the usual strain field inside and in the proximity of the QD, given by the lattice mismatch between the two materials, we can observe a weaker strain field at the interfaces between the two crystal phases of AlGaAs. This field is also observed in the TEM image shown in Figure 2. Despite its weakness, which
was indeed expected from the small lattice mismatch between the GaAs and the Al$_{0.25}$Ga$_{0.75}$As ($c = 0.25$) and the even smaller crystal mismatch between Zb- and Wz- AlGaAs, the strain field is non zero practically everywhere in the nanowire. This also affects the electric properties of the structure.

The electric field originating from the piezoelectric effect in the Mx case is relatively strong and presents an unusual shape, different from the typical results obtained for Zb QDs. This can be observed in Figure 5, showing the $z$ component of the electric field for both Zb and Mx structures.

Instead of the usual quadrupole shape obtained for Zb QDs, the Mx case gives a Wz-like shape of the electric field close and inside the dot, with its typical dipole characteristics, and opposite signs inside and just outside of the top and the bottom of the dot. This occurs even though the central volume of the column is in a pure Zb crystal phase and owes to the stronger magnitude of the piezoelectric effect induced by the Wz layers, which leads to stronger a electric field. A stronger magnitude of the piezoelectric effect in Wz heterostructures compared to Zb heterostructures has been already reported in the literature [25].

B. Results for confined states

The quantum confinement in the QD is mainly given by the band offset between the two materials and between the two different crystal structures of the AlGaAs in the Mx case, since the impact of the observed strain fields on the confinement via the deformation potentials[33] is not particularly relevant. In Figure 6 we plot the conduction ($E_c$) and valence ($E_v$) band along the $z$ direction for the considered structures, i.e., Zb and mixed.

The usual case of a Zb QD embedded in a Zb matrix has been considered as a useful comparison in order to better understand the results for the energy levels. For the Mx conduction band, the well of the QD is surrounded by the energetically higher barrier of the Wurtzite layers, higher as
compared to the lower potentials of the Zb AlGaAs. This results in an increased quantum confinement for the electron states. On the other hand, the hole bands of the Mx structure exhibit a sort of double well with the potential profile of the QD and the two adjacent Wurtzite layers acting as a second wider well, that can contain several confined states. The form of the potential has a notable impact on the hole wave functions.

In Figure 7 we plot all the confined states for the electrons and the holes for both Zb and mixed situations. All states are doubly degenerate for spin inversion due to time reversal symmetry.

![Figure 7](image.png)

Figure 7. The calculated states for electrons (top) and holes (bottom). The x scale enumerates the number of the correspondent state (different scale for electrons and holes), while the energy in the y axis is referred to the Fermi level as in Figure 6.

As far as electron states are concerned, the main difference we observe between the Zb case and the Mx case is a higher degree of quantum confinement in the latter, which slightly increases the energy levels and gives raise to two further confined states, i.e., states 11 and 12 are delocalized in the Zb model, while confined within the QD in the Mx case.

The shape and the symmetry of the electron wave functions do not significantly differ between the two models, implying that they are mainly dependent on the ZB crystal structure of the QD, and are barely affected by the electric field and its symmetries. However, the actual lack of knowledge of the real values of piezoelectric coefficients for the Wz phase of these materials makes uncertain the impact of the electric field on the symmetry of the wave functions.

The higher electron states are completely delocalized in the Zb layers adjacent to the Wz layers. These states are not optically active, since they have almost zero overlap with the hole wave functions confined in the QD.

The situation for the hole states is notably different and quite interesting. Comparing the curves plotted in the bottom part of Figure 7, it is clearly possible to see that, while the first confined states of the two models are very close in energy (also showing a similar trend), higher energy states (in absolute value) display different trends. The states of the Mx model are systematically lower in energy. The onset of this change corresponds to states that start to delocalize in the Wz layers, adjacent to the QD. The increased available volume explains their lower energy compared to the case of a pure Zb structure. This is confirmed by the contour plots of the probability density which we show in Figure 8. Here we present a slice of the xz plane, with z = 30 nm and z = 40 nm, showing basically the center of the nanowire with the dot and the two adjacent Wz layers.

Figure 8 (left) shows the probability density of the holes ground state, which, as expected, is completely confined in the QD. Figure 8 (right) shows the probability density of the hole state number 32, according the bottom y labeling of Figure 7. This state is mainly confined in the Wz layers. A similar situation, with states confined in the volume formed by the QD and the two adjacent Wz layers, is observable for a considerable number of the higher energy hole states. All these states are optically active, since they have a non-zero overlap with the electron states confined in the dot.

C. Oscillator strengths results

We finally present results for the oscillator strengths, according dipole approximation. In Figure 9 we show the optical transitions.

Despite some resemblances in the distributions of the peaks, we can notice an effect which confirms what we have discussed in the previous section. We observe a blue shift of the lowest transitions of the Wz instance, namely for the ground state - ground state transition and the transitions between the
more confined energy levels. This is clearly due to the higher energy of the electrons in the Mx case. But, on the other hand, for the peaks at higher energy we notice a red shift of the lines correspondent to the Mx models, due to the difference in energy between the less confined hole states, up to 20 meV as observable in Figure 7.

Finally, in Figure 10, the lowest range of the spectrum is compared with the positions of three peaks given by experimental results from a specific nanowire sample with approximately the same dimensions of our model.

Here, the peak correspondent to the lowest transition, namely the ground state - ground state transition, has been measured by a PL experiment, while the other two positions we show are given by a PLE data set from the same sample. Although our Mx model cannot be completely validated due to the fluctuation of measurement data from different samples, the agreement for this particular sample is very good.

V. CONCLUSION

We have presented numerical simulations for polymorph Wz-Zb cylindrical AlGaAs NW with an embedded GaAs QD. Firstly, the connection with experimental structures has been shown, and consequently suitable continuum elastic and 8 band $k \cdot \overrightarrow{p}$ models have been derived. All the necessary numerical parameters have been determined by appropriate approximations, where not available in the previous literature.

The results for the strain fields are in a good agreement with the experimental data, showing, besides the usual field given by the lattice mismatch between different material, a not-zero field at the interfaces between the two crystal phase of AlGaAs.

The analysis of the confined energy levels, wave functions and optical transitions clearly show the differences between an uniform crystal phase and a polymorph heterostructure. The comparison of numerical results and experimental spectra for optical transitions confirm the validity of our polymorph numerical model.

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