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Impact of Size, Shape and Composition on Piezoelectric Effects and the Electronic Properties of InGaAs/GaAs Quantum Dots

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Abstract

The strain fields in and around self-organized In(Ga)As/GaAs quantum dots (QD) sensitively depend on QD geometry, average InGaAs composition and the In/Ga distribution profile. Piezoelectric fields of varying size are one result of these strain fields. We study systematically a large variety of realistic QD geometries and composition profiles, and calculate the linear and quadratic parts of the piezoelectric field. The balance of the two orders depends strongly on the QD shape and composition. For pyramidal InAs QDs with sharp interfaces a strong dominance of the second order fields is found. Upon annealing the first order terms become dominant, resulting in a reordering of the electron *p*- and *d*-states and a reorientation of the hole wavefunctions.

1 Introduction

Semiconductor quantum dots (QDs) are fascinating physical subjects exhibiting electronic properties close to hydrogen in a dielectric cage, thus merging semiconductor physics with atomic physics. Their electronic properties,[1–3] are strongly geometry dependent. In particular, the influence of the QD morphology on few-particle properties like exciton (X) and biexciton (XX) binding energies, and the exciton fine-structure splitting are currently the subject of active research.[4–6] Varying QD size, shape, composition are attractive and practical means to vary the electronic and optical properties. Many applications are based on this discovery.[7–9]

The symmetry of the confinement potential is not determined by the QD geometry alone, but also by the anisotropic strain, piezoelectricity, and the atomistic symmetry anisotropy (ASA).[10] These effects arise from the lack of inversion symmetry of the underlying zinc-blende lattice. Grundmann $et\ al.$ [2] and Stier $et\ al.$ [3] discovered the linear (first order) piezoelectric effect to cause symmetry lowering from C_{4v} to C_{2v} for pyramidal QDs. Later, Bester $et\ al.$ [10] reported a similar behavior for lens-shaped QDs, where the rotational symmetry, $C_{\infty v}$, is lowered to C_{2v} as well, with similar consequences as for the pyramidal QDs. Recently Bester $et\ al.$ [11] investigated quadratic (second order) piezoelectric effects and again investigated flat lens-shaped QDs.[12] The authors reported that the linear and quadratic effects oppose and cancel each other leading to an almost field-free QD-interior, thus practically reestablishing the $C_{\infty v}$ confinement symmetry. We will show in this paper that mutual cancellation of first and second order piezoelectric effects can not be generalized to lower symmetry and more realistic QD structures.

The electron p-state splitting presents a measure for the actual confinement anisotropy between the [110] and the $[1\overline{1}0]$ directions. Bras et~al.~[13,~14] performed intraband absorption measurements and suggested an electron p-state splitting in the order of 8 meV for capped InAs QDs. As we will show later in this work, their polarization anisotropy corresponds to a lower p-state aligned along $[1\overline{1}0]$ and a higher energy state, aligned along [110]. This finding is in accord with the results of Maltezoupolos et~al.[15] They used scanning~tunneling~spectroscopy to probe uncapped QDs and found striking evidence for large p-state splitting, with identical p-state order to Bras et~al. In some cases d-states below the second p-state were observed by Maltezoupolos et~al.

The purpose of this paper is twofold. First, we will calculate the electronic and optical properties of a large number of QDs of varying size, shape (square/circular/rhomboid base, different vertical/lateral aspect ratios) and composition (homogeneous/peaked and isotropic interdiffusion), highlighting the impact of the varying first and second order piezoelectric effects on the electronic properties.

Despite tremendous advances in structural characterization the real shape and composition of capped quantum dots, which are decisive for all applications,[1] are usually not or only poorly known. Therefore, the second purpose of our work is to predict spectroscopic quantities which serve as fingerprints for specific QD structures, thus to address the inverse problem of deriving information on size, shape and composition from spectroscopic data. The difficulty of this approach is that the relation between the QD morphology and the calculated spectroscopic properties depends on the employed model and the parameters entering the model.[16]

We obtain the electronic structure using a strain-dependent eight-band $\mathbf{k} \cdot \mathbf{p}$ Hamiltonian, including first and second order piezoelectricity. This model provides, at reasonable computational cost, a fast and transparent way to connect the electronic structure of QDs with their geometry/composition and to the bulk properties of the constituent materials.

The paper is organized as follows: In section 3 the method of calculation is outlined and some general properties, like the impact of strain on the confinement potential or the symmetry properties of the single particle orbitals are discussed. In section 4 first and second order piezoelectric fields are compared as function of shape and composition and their impact on the electronic and optical properties is addressed. In the ensuing sections 5 and 6 further consequences of different shapes and composition profiles for the electron and hole energies and their excitonic spectra are investigated. The paper is concluded in section 7.

2 The Investigated structures: Variation of size, shape and compostion

Our selection of model QDs is guided by the broad variation of structures observed in experiments (see, e.g., Ref. [17, 18] and references therein).

Fig.1 gives an overview on the investigated series of model structures.

Series A: Pyramidal InAs/GaAs QDs, similar to Ref. [3], with base lengths 10.2 nm (A1), 13.6 nm (A2), 17.0 nm (A3), and 20.4 nm (A4).

Series B: Starting with the 17 - nm-base-length pyramid of series A, the vertical aspect ratio is varied between 0.5 (full pyramid) and 0.04 (very flat).

Series C: The QDs with a circular base and a vertical aspect ratio varying between 0.5 (half-sphere) and 0.17.

Series D: Starting, again, with the 17 nm base length pyramid of series A an elongation in [110] and $[1\overline{1}0]$ direction is investigated. The lateral aspect ratio (length in [110] direction divided by length in $[1\overline{1}0]$ direction) varies between 2 and 0.5 (a value of 1 corresponds to the square base).

It is important to note, that the QD volume has been kept constant throughout series B, C and D.

Series E: A homogeneous variation of the In-content for pyramidal $In_xGa_{1-x}As/GaAs$ QDs is considered. The starting point is again the $17 \, \mathrm{nm}$ base length pyramid of series A. The In content

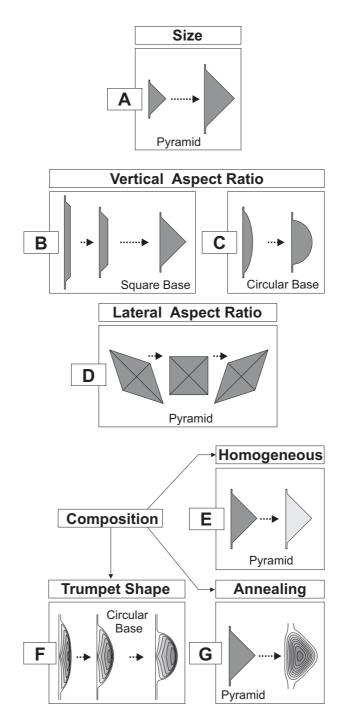


Figure 1: Structure series investigated in this paper.

decreases in steps of 10% from 100% to 70%.

Series F: The QDs of this series have a circular base together with a trumpet-shaped like InGaAs composition profile. The integral In amount of the QDs is equal to QD A3.

Series G: By applying a smoothing algorithm on structure A3 with a variable number of smoothing steps (N) the process of Fickian diffusion as a result of an annealing procedure is simulated.

Wetting layer: As indicated by figure 1 an InAs wetting layer (WL) of one monolayer thickness for series A-D is taken into account. In series E the WL average composition equals the QD composition. For the remaining series, F and G a thicker WL with a graded InAs composition profile is assumed. is taken into account too.

3 Method of calculation

3.1 Calculation of strain

Since the impact of strain on the confinement is comparable to that of the band offsets at the heterojunctions, the wavefunctions and energies are very sensitive to the underlying strain distribution. The impact of the model used for calculating the strain distribution has been analyzed in a number of publications.[3, 19] Stier $et\ al.$ [3] argue that the continuum elasticity model (CM) gives better results for QDs than the valence force field (VFF) model (Keating) in its linearized version (Kane). The major part of the differences in the strain distribution are attributed to the incorrect value of C_{44} in the VFF model and not to its atomistic character. Later, Williamson et al. [20] introduced a generalized version of the VFF model — the G-VFF model — where C_{44} is reproduced correctly. Although the potential of the Keating model in its original version is not harmonic, it has been remarked by Kane,[21] that anharmonic effects due to higher order terms are not satisfactorily treated. Therefore Lazarenkova and coworkers extended this model to include anharmonic effects correctly.[22] The same issue is addressed by Hammerschmidt $et\ al.$ [23] and Migliorato $et\ al.$ [24] who employed the Tersoff-potential method [25].

The choice of the most appropriate strain model depends on the choice of the model for the electronic structure calculations. Since we use — with the eight-band $\mathbf{k} \cdot \mathbf{p}$ model — a continuum approach, an atomistic strain model cannot unfold its full potential for two reasons:

First, the mapping of the atomic positions onto a strain tensor field is associated with a loss of information. To describe the positions of four tetrahedrally coordinated In atoms around an As atom five times the three spacial dimensions = 15 parameters are required. The strain tensor field on the other hand is described by only six independent components at each local position.

Second, the $\mathbf{k} \cdot \mathbf{p}$ model provides only a limited number of parameters to account for strain, hence the model is not sensitive to the complete information an atomistic model provides. For example, the strain tensor, derived from the CM model, for a QD with a fourfold rotational C_{4v} symmetry has C_{4v} symmetry too, in contrast to atomistic models: The tetrahedral configuration of the atoms[19] leads to C_{2v} symmetry, i.e. the strain components are different along the [110] and [1\overline{10}] directions. The resulting p-splitting, obtained by using the atomistic-model-derived strain tensor field in our $\mathbf{k} \cdot \mathbf{p}$ model, is underestimated. The structural $C_{\infty v}$ or C_{4v} symmetry is noticeably broken only in the second step by the inclusion of the piezoelectric field.

3.2 Piezoelectricity and the reduction of lateral symmetry

Piezoelectricity is defined as the generation of electric polarization by application of stress to a crystal lacking a center of symmetry. [26] The zinc-blende structure is one of the simplest examples of such a lattice and the strength of the resulting polarization is described by one parameter alone, e_{14} , for the linear case, resulting in a polarization \mathbf{P}_1 , and three parameters, B_{114} , B_{124} , and B_{156} for the quadratic case, [11] resulting in a polarization \mathbf{P}_2 . Their relation to the strain tensor field is given by:

$$\mathbf{P}_{1} = 2 e_{14} \begin{pmatrix} \epsilon_{yz} \\ \epsilon_{xz} \\ \epsilon_{xy} \end{pmatrix},$$

$$\mathbf{P}_{2} = 2 B_{114} \begin{pmatrix} \epsilon_{xx} \epsilon_{yz} \\ \epsilon_{yy} \epsilon_{xz} \\ \epsilon_{zz} \epsilon_{xy} \end{pmatrix} +$$

$$+ 2 B_{124} \begin{pmatrix} \epsilon_{yz} (\epsilon_{yy} + \epsilon_{zz}) \\ \epsilon_{xz} (\epsilon_{zz} + \epsilon_{xx}) \\ \epsilon_{xy} (\epsilon_{xx} + \epsilon_{yy}) \end{pmatrix} +$$

$$+ 4 B_{156} \begin{pmatrix} \epsilon_{xz} \epsilon_{xy} \\ \epsilon_{yz} \epsilon_{xy} \\ \epsilon_{yz} \epsilon_{xz} \end{pmatrix}.$$

$$(1)$$

Piezoelectric charges, ρ_{piezo} , arise from the polarizations:

$$\begin{array}{rcl} \rho_{\rm piezo}(\mathbf{r}) & = & -\nabla \cdot \mathbf{P} & , \\ \mathbf{P} & = & \mathbf{P}_1 + \mathbf{P}_2 & . \end{array}$$

The resulting piezoelectric potential is obtained by solving Poisson's equation taking into account the material dependence of the static dielectric constant, $\epsilon_s(\mathbf{r})$

$$\rho_{\mathbf{p}}(\mathbf{r}) = \epsilon_0 \nabla \cdot \left[\epsilon_s(\mathbf{r}) \nabla V_{\mathbf{p}}(\mathbf{r}) \right]$$

$$\Leftrightarrow$$
(2)

The first term on the right hand side of Eq. 3 refers to the true three-dimensional charge density while the second is the contribution of polarization interface charge densities due to a discontinuous $\epsilon_s(\mathbf{r})$ across heterointerfaces.

The importance of the second order term, P_2 , for In(Ga)As/GaAs(111) quantum wells (QWs) and QDs has been pointed out recently by Bester and coworkers.[11, 12] They found that in In(Ga)As QWs the linear and quadratic contributions have opposite effects on the field, and for large strain the quadratic term even dominates. For InAs/GaAs QDs, however, the situation is more complex. In addition to the large strain the QDs' three-dimensional structure comes into play: The linear term generates a quadrupole-like potential, which reduces a structural C_{4v} - or $C_{\infty v}$ -symmetry of a QD to C_{2v} .[2, 10] The effect of the quadratic term has been evaluated recently by Bester *et al.*[12] for lens-shaped QDs. It was found to cancel the first order potential inside the QD, leading to a field free QD. Here we extend this investigation to a variety of more realistic QD structures (see section 2). For a pyramidal QD with a base length of 17 nm and {101} side facets the strength

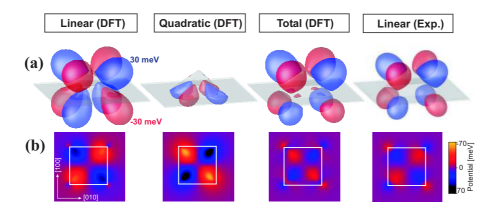


Figure 2: (Color online) (a) The piezoelectric potential isosurfaces at $\pm 50\,\mathrm{meV}$ of a pyramidal InAs QD with 17 nm base length shown for the linear, the quadratic part and for sum of both using theoretical values for the piezoelectric constants. Results obtained from experimental values are shown in the last column. (b) Contour plots of the piezoelectric potential 2 nm above the wetting layer.

and distribution of the piezoelectric potential resulting from the two orders of the piezoelectric tensor is shown in Fig. 2. Apart from the different orientation and sign of the two contributions, an important peculiarity of the second order potential is its restriction to the interior of the QD, which is in appearant contrast to the widespread first-order field. This difference is linked to the origin of the polarization P: P_1 is a function of the shear-strain components alone, whereas P_2 results mainly from the product of the diagonal and the shear-strain. However, in contrast to the shear-strain components, the diagonal elements ϵ_{ii} are large only inside the QD and its close vicinity. Therefore, P_2 -charges can only be created in this region.

3.3 Single Particle States

The energy levels and wavefunctions of bound electron and hole states are calculated using the eight-band $\mathbf{k} \cdot \mathbf{p}$ model. It was originally developed for the description of electronic states in bulk material.[36–39] For the use in heterostructures, the envelope function version of the model has been applied to QWs,[35] quantum wires, [29] and QDs.[3, 40–43] Details of the principles of our implementation are outlined in Ref. [29] and [43].

This model enables us to treat QDs of arbitrary shape and material composition, including the effects of strain, piezoelectricity, VB mixing, and CB-VB interaction. The strain enters our model via deformation potentials as outlined by Bahder.[44] Its impact on the local bandedges as a function of the QD geometry will be discussed in the next section.

The $\mathbf{k} \cdot \mathbf{p}$ model, when applied to small quantum structures, has in principle a few well-known drawbacks which have been examined in detail in Ref. [45, 46]. They are basically related to the fixed number of Bloch functions used for the wavefunction expansion, the restriction to the close vicinity of the Brillouin zone center, limited ability to account for the symmetry of the underlying lattice, and the possible appearance of spurious solutions. These problems do not arise in microscopic theories like the empirical pseudopotential method [45] (EPM) or the empirical tight-binding (ETB) method. [47, 48] Their potential, however, can only be exploited if the corresponding input parameters — the form factors in the EPM or the tight-binding parameter and their

	Quantity	Unit	GaAs	InAs	Interpolation	Reference
Lattice constant	a	A	5.6503	6.0553	linear	[27]
Fundamental gap	E_g	meV	1518.0	413.0	$1518 - 1580c + 475c^2$	[27]
Averaged VB edge[28]	$E_{v}^{'}$	meV	-6920	-6747	$-6920 + 231 c - 58 c^2$	[27, 29]
Spin-orbit coupling energy	\triangle_0	meV	340	380	$340 - 93c + 133c^2$	[30]
Optical matrix parameter	E_p	meV	28000	22204	$(1.238 - 0.2095 c) \frac{1 - m_e}{m_e} \frac{3E_0(E_0 + \Delta_0)}{3E_0 + 2\Delta_0}$	[31, 32]
CB effective mass	m_e	m_0	0.067	0.022	$0.0667 - 0.0419 c - 0.00254 c^2$	[33]
Luttinger parameter	γ_1		7.1	19.7	1/[(1-c)/7.1 + c/19.7]	[30, 34]
Luttinger parameter	γ_2		2.02	8.4	1/[(1-c)/2.02 + c/8.4]	[30, 34]
Luttinger parameter	γ_3		2.91	9.3	1/[(1-c)/2.91 + c/9.3]	[30, 34]
Kane parameter	B	$ m meVnm^2$	0	0	linear	[29, 35]
CB-VB coupling by strain	b'	meV	0	0	linear	[29, 35]
CB hydrostatic def. pot.	a_c	meV	-8013	-5080	linear	[33]
Gap hydrostatic def. pot.	a_g	meV	-8233	-6080	linear	[33]
VB shear def. pot. [100]	b_v	meV	-1824	-1800	linear	[30]
VB shear def. pot. [111]	d_v	meV	-5062	-3600	linear	[30]
Elastic compliance	C_{11}	GPa	118.8	83.3	linear	[30, 34]
Elastic compliance	C_{12}	GPa	53.8	45.3	linear	[30, 34]
Elastic compliance	C_{44}	GPa	59.4	39.6	linear	[30, 34]
Static dielectric constant	ϵ_s		13.18	14.6	linear	[33]
Piezoelectric constants						
Linear (exp.)	e_{14}	$\mathrm{C/m^2}$	-0.16	-0.045	linear	[33]
Linear (calc.)	e_{14}	C/m^2	-0.230	-0.115	linear	[11]
Quadratic (calc.)	B_{114}	C/m^2	-0.439	-0.531	linear	[11]
Quadratic (calc.)	B_{124}	C/m^2	-3.765	-4.076	linear	[11]
Quadratic (calc.)	B_{156}	C/m^2	-0.492	-0.120	linear	[11]

Table 1: Material parameters for 6.5 K used in this work. The theoretical values for the linear and quadratic piezoelectric coefficients (C/m²) are taken from Bester *et al.*[11] Symmetry considerations for the zinc-blende crystal structure imply that there are only 24 non-zero elements of the $B_{\mu jk}$ tensor, which can be reduced to three independent elements, B_{114} , B_{124} , and B_{156} .

strain dependence in the ETB — are known with sufficient accuracy. Reliable generation of these parameters, however, is highly nontrivial. One of the most appealing features of the $\mathbf{k} \cdot \mathbf{p}$ model, in contrast, is the direct availability of all parameters entering the calculations. Additionally, the required computational expense of the method is comparatively small. The material parameters used in this work are taken from Ref. [3] and are listed in Tab. 1 together with the appropriate interpolation rules. As mentioned earlier in Ref. [3] the treatment of the thin wetting layer is difficult because its thickness usually is represented by a too small number of voxels. Two-dimensional 2D states confined in the WL but not in the QD appear at higher energies and indicate the end of the zero-dimensional 0D spectrum. The effort expended at modeling the WL only aims to obtain the transition from 0D to 2D states at plau- sible energies. The bound states obtained from our method are accurate apart from the discretization error discussed in Ref. [3], but the numerical 2D-like states are not realistic and therefore not included in this paper.

3.3.1 Strain versus Confinement Profile

In this section we address the shape dependence of the strain field and the resulting confinement potential. For this purpose we consider a full and a truncated pyramid (aspect ratio = 0.21) from series B. The local band edges (Fig. 3 (b) and (d)) are obtained by pointwise diagonalization of the Hamiltonian H at k=0. For the sake of clarity we resort to a simpler approximation of the local band edges employing the following formulas for the CB, the heavy hole (HH) and the light hole (LH) bands (thus ignore shear strain induced HH-LH coupling and split-off band contributions [32]):

$$V_{\text{CB}}(\mathbf{r}) = E_{\text{CB}} + a_{\text{c}} \, \epsilon_{\text{H}}(\mathbf{r}) ,$$

$$V_{\text{HH}}(\mathbf{r}) = E_{\text{VB}} + a_{\text{v}} \, \epsilon_{\text{H}}(\mathbf{r}) + \frac{b}{2} \, \epsilon_{\text{B}}(\mathbf{r}) ,$$

$$V_{\text{LH}}(\mathbf{r}) = E_{\text{VB}} + a_{\text{v}} \, \epsilon_{\text{H}}(\mathbf{r}) - \frac{b}{2} \, \epsilon_{\text{B}}(\mathbf{r}) ,$$

$$(4)$$

where the hydrostatic strain $\epsilon_{\rm H}$ and the biaxial strain $\epsilon_{\rm B}$ are defined as

$$\begin{split} \epsilon_H(\mathbf{r}) &= \epsilon_{xx}(\mathbf{r}) + \epsilon_{yy}(\mathbf{r}) + \epsilon_{zz}(\mathbf{r}) \ , \\ \epsilon_B(\mathbf{r}) &= \epsilon_{xx}(\mathbf{r}) + \epsilon_{yy}(\mathbf{r}) - 2\,\epsilon_{zz}(\mathbf{r}) \ . \end{split}$$

 $V_{\rm HH}$ and $V_{\rm LH}$ are the heavy-hole and light-hole bands, $a_{\rm c}$, $a_{\rm v}$, b are the deformation potentials, and $E_{\rm CB/VB}$ is the unstrained band edge energies. In this simplified picture the hydrostatic strain shifts the CB- and VB-edge and the biaxial strain introduces a splitting between the heavy and light-hole band edge. That means — provided the hydrostatic strain remains constant or a_v is very small — that an increasing biaxial strain reduces the energy gap. Figure 3 provides more insight in how the corresponding strain quantities affect the confinement potential. The larger hydrostatic strain inside the full pyramid increases the CB potential more than for the truncated pyramid. The smaller biaxial strain and its sign change at the QD-center, on the other hand, lead to a smaller splitting and a crossing of HH and LH band edges for the full pyramid. Both effects lead to a deeper confinement of both, electrons and holes, with decreasing aspect ratio.

Lets now have a look at the deformation of a lattice unit cell at different positions of the QD for different geometries:

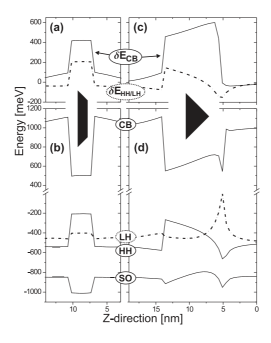


Figure 3: Impact of strain on the local band edges for the cases of a full and a truncated pyramid. (a) and (c): The dotted lines indicate the energy shift of the VB imposed by biaxial strain: $\delta E_{\rm HH/LH} = b/2\,\epsilon_{\rm b}({\bf r})$. The solid line marks the contribution of the hydrostatic strain $\delta E_{\rm c} = a_{\rm c}\epsilon_{\rm h}({\bf r})$. The resulting local band edge positions are shown in panels (b) and (d). While the HH-band inside the truncated QD is always above the LH-band, a crossing between HH and LH band edge occurs in the full pyramid (d).

(a) A unit cell in the center of a *cubic* InAs/GaAs-QD is a cube. Since the strain components ϵ_{xx} , ϵ_{yy} , and ϵ_{zz} have the same value, no biaxial strain is present. The hydrostatic strain is compressive, i.e., negative. (b) If the cubic QD gets flattened, the lateral pressure exceeds the vertical one and the unit cell extends in z-direction. Hence, the ϵ_{zz} component is larger than ϵ_{xx} and ϵ_{yy} , the biaxial strain is negative and the HH edge resides above the LH edge. (c) The pyramidal case is more complex: A unit cell at the center of the QD-base is elongated in z-direction, leading to negative biaxial strain. Near the tip, however, the lateral forces become smaller than the vertical (directed towards the cell) forces. Consequently, the unit cell becomes flat with positive biaxial strain and the light-hole band edge moves on top. Thus, the QD aspect ratio determines the lateral and the vertical pressure proportion acting on a unit cell.

3.3.2 Symmetry properties of the single particle states

Let us first recapitulate some basic properties which apply to almost all QDs considered here. The confined electron states can be classified according to the number of nodal planes of the envelope function: 0 planes correspond to s-like, 1 plane to p-like, 2 planes to d-like states etc. An alternative description links their transformation behavior under the QDs symmetry operations to the irreducible representations of the symmetry group. This is exemplified in Fig. 4 for the s, p, and d-shell for three typical QDs: a flat lens $(C_{\infty v})$ and a full pyramid in absence (C_{4v}) and presence of piezoelectricity (C_{2v}) . Since only the last case — with C_{2v} confinement symmetry — is of practical interest, Stier et al. [3] introduced a labeling scheme $|abc\rangle$ where a, b, and c are the number of nodes in $[1\overline{10}]$, [110], and [001] directions respectively. The correspondence of this scheme to the irreducible representations of C_{2v} is shown in Fig. 4.

For the confined hole states such a classification is more complex due to the mixing of HH an LH parts, which can have different symmetry properties.[49]

3.3.3 Heavy hole - Light hole coupling

For the understanding of the polarization properties of the excitonic absorption spectra the HH-LH coupling and the resulting state mixing is of largest importance. There are three main mechanisms that finally determine the HH/LH ratio of the hole wavefunctions: First, the relative position of the local HH and LH bandedge, second their masses relative to each other, and third the coupling strength among the VBs, which is mainly a function of the shear strain (see Bahder [44]). Heavyhole and light-hole states are decoupled if $\epsilon_{xx} = \epsilon_{yy}$ and the shear strain components ϵ_{xy} , ϵ_{xz} , ϵ_{yz} are zero.

These mechanisms are tightly linked to the QD shape: In the first case the inhomogeneity of the strain (especially the biaxial strain, see Eq. 4) leads to a splitting of the HH-LH band edges [Fig. 3(b)] and to a crossover of the two bands if the biaxial strain changes its sign as in the case a full pyramid [Fig. 3(d)]. In the second case the shear strain itself is large if the QD has a large aspect ratio. For flat QDs in contrast the shear strain becomes small, as does the HH-LH coupling. As an example, the LH fraction of the hole ground state — an indicator for the strength of the coupling — varies in square-based QDs (series *B*) from 9.2 % (for the full pyramid) to 2.8 % (for the most truncated pyramid). For circular based QDs (series *C*) we observe a variation between 14.4 % (half sphere) and 2.8 % (flat lens). The dependence on the aspect ratio is similar to that of the (first order) piezoelectric field, since both, the HH-LH coupling and the (first order) piezoelectric field, originate from the presence of shear strain.

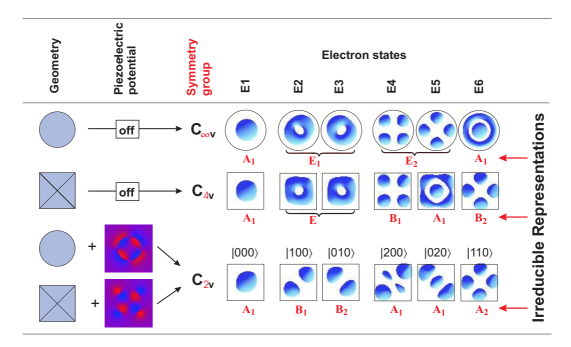


Figure 4: (Color online) Electronic states classified according to the irreducible representations of the symmetry groups $C_{\infty v}$, C_{4v} and C_{2v} . The symmetry lowering from $C_{\infty v}$ down to C_{4v} and C_{2v} can arise from a change of geometry, from a piezoelectric field, or from the ASA effect.

The magnitude of the light hole projection also depends heavily on the QD size, as can be seen in Fig. 5. In average, the LH percentage increases for higher excited hole states and – except for the hole ground state – it increases for decreasing QD size, when the hole energies approach the GaAs VB edge.

3.4 Optical Properties

3.4.1 Interband spectra

The interband absorption spectra are calculated by Fermi's golden rule applied to excitonic states calculated by the configuration interaction method.[49]

There exist no strict selection rules for the decay of excitons. As a rule of thumb one can say that those transitions have a large oscillator strength where electron and hole state share the same symmetry properties *and* have a sizable spatial overlap. However, since the hole states consist of HH and LH parts, each with its own symmetry, they have finite recombination probabilities with a number of different electron states.

3.4.2 Intraband spectra

In contrast to the excitonic decay, the CB intraband transitions follow strict transition rules: for two electron states, $|a\rangle = |ijk\rangle$ and $|b\rangle = |i'j'k'\rangle$, it is a necessary condition that at least one of the expressions i-i', j-j' or k-k' is an odd number. For example the transition between the electron states $|100\rangle \rightarrow |000\rangle$ has a sizeable oscillator strength in contrast to $|200\rangle \rightarrow |000\rangle$, which is

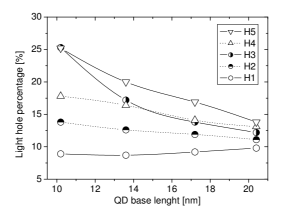


Figure 5: The light hole fractions of the first five hole states shown for the *series A* (pyramids of different size) as function of the QD base length.

a forbidden transition. In general the oscillator strength of intraband transitions is smaller than for excitonic transitions and the anisotropies between absorption in [110] and $[1\overline{1}0]$ direction are much larger. The latter is an ideal fingerprint of the electronic spectrum of QDs and the associated symmetry properties of the wavefunctions.

3.5 Numerical Aspects

For the calculation of strain and piezoelectricity the extension of the simulation box is seven times larger than the QD in vertical and four times larger in lateral direction. The QD is placed in the center. The grid resolution is half the lattice constant of GaAs. For the eight-band $\mathbf{k} \cdot \mathbf{p}$ calculations, a smaller region of this box is cut out, thereby doubling voxel size to 0.5656 nm. The box is large enough to not affect the investigated energy levels and their symmetry. For example, a pyramid of 17 nm base-length is calculated on a 60 x 60 x 60 grid, using Dirichlet boundary conditions.

4 The Impact of the piezoelectric field

4.1 First and second order piezoelectricity as function of size, shape and composition of the QD

In this section we explore the balance between linear and quadratic piezoelectric terms as a function of shape and composition of the QDs. In Fig. 6 the piezoelectric potentials of five different QDs are shown as a lateral 2D slice taken at the z-component of the QD barycenter. For all shown QDs, the first and second order term are quadrupole-like, but oriented in opposite directions. The appearance of the total potential at a larger distance from the QD is dominated by the first order term, a possibly important issue for stacked QDs. Which of the terms prevail inside the QD strongly depends on the actual geometry and the composition profile as will be detailed now.

(a) Flat, lens-shaped QDs. The interior of flat lens-shaped QDs is almost field-free [see Fig.6(a)], due to the cancellation of the first and second order piezoelectric contributions. These findings

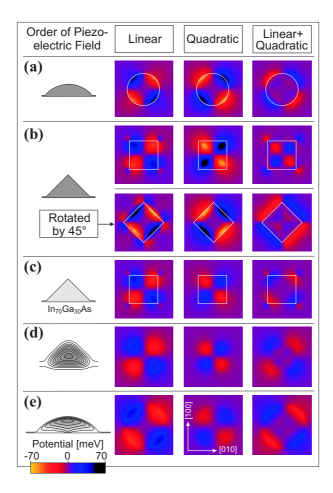


Figure 6: (Color online) Lateral scans through the piezoelectric potential: linear part (left), quadratic part (middle), and the sum of both (right) shown for different variations of the QD morphology. (a) A lens-shaped QD (from series C vertical aspect ratio, $ar_V = 0.21$), (b) a full InAs pyramid (from series A) with a base length of 17.2 nm: In the first case the base edges are oriented along [100] in the second the pyramid is rotated by 45° , hence the edges are oriented along [110]. (c) The In fraction of the pyramid from (b) is decreased down to 70 % (from series E). (d) An isotropic diffusion procedure is applied to QD (b) (from series E). (e) Here we show results for a nonisotropic internal InAs composition profile taken from series E.

confirm the results of Bester et al.[12]

- (b) Pyramidal QDs. In contrast to the lens-shaped QDs, for pyramidal InAs QDs with $\{101\}$ facets the quadratic term strongly dominates inside the QD [see Fig. 6(b1) and Fig. 2]. A rotation by 45° [see Fig. 6(b2)], however, leads to a balance of both contributions and results in a field-free QD as in case (a).
- (c) Homogeneous alloying. Now we consider an $In_{70}Ga_{30}As$ pyramidal QD with $\{101\}-facets$. The linear piezoelectric potential remains unaffected: It is almost as large as for the pure InAs QD in (b1). This seems surprising at first sight, since the shear strain components become smaller for rising Ga content. This decrease is compensated by the larger first order piezoelectric constant e_{14} , which is linearly interpolated between the values of InAs and GaAs (see table 1). The second order field, however, drops drastically for two reasons: First, the diagonal and the nondiagonal strain components, entering the calculation as products in Eq. 1, both diminish. Second, the parameter B_{124} , which leads the largest contribution of P_2 , is smaller. As a result the interior piezoelectric field of the QD is dominated by the first order term.
- (d) Annealed pyramid. The first order potential of an annealed structure shows only minor changes in the field distribution compared to the unannealed InAs pyramid (b1). The second order potential is drastically reduced similar to case (c). This strong decrease occurs already for the first annealing step. Thus, the second order term is very sensitive to the degree of interface abruptness. In the resulting overall potential only a very small portion of the QD interior is still dominated by the second-order field.
- (e) Trumpet-shaped composition profile. Here we consider a different In distribution compared to case (d) but the field distribution is very similar to the case of the annealed pyramid.

Influence of Wetting layer The presence of the thin wetting layer has no sizable impact on the magnitude and the symmetry of the piezoelectric field. A (001) quantumwell in the zinc-blende system on its own can not produce any piezoelectric field, due to the absence of shear strain. Therefore, for the very flat QDs of series B, where the QD height approaches the WL thickness, both orders of the piezoelectric field become extremely small ', and their impact on the electronic states is negligible.

To conclude this section, we find that the results of Bester *et al.*[12] — the cancellation of first and second order piezoelectric effects inside the QD — is correct only for the specific QD structure considered in their work (lens-shaped QDs with a vertical aspect ratio smaller than 0.35). Their findings can not be generalized to all QD geometries and composition profiles. In particular, the second-order field is very sensitive to the exact QD shape and the composition profile.

4.2 Impact on the single-particle states

Electron and hole states respond in different ways to the piezoelectric potential. For the electron states it is just a minor perturbation lifting possible degeneracies of p- and d-states. The shapes of the wavefunctions themselves are hardly affected. The hole states, in contrast, follow very closely almost any potential variation to minimize their potential energy. This behavior is connected to the very different effective masses of both carrier types, which plays the role of a weighting factor in the trade-off between kinetic and potential energy.

For the electrons, their small effective mass translates into a steeply rising dispersion $E(\mathbf{k})$. Since the kinetic energy in \mathbf{k} -space is given by $E_{\rm kin} \approx \int_{V_b} E(\mathbf{k}) \Psi(\mathbf{k}) \, d\mathbf{k}$, a steep $E(\mathbf{k})$ makes it very

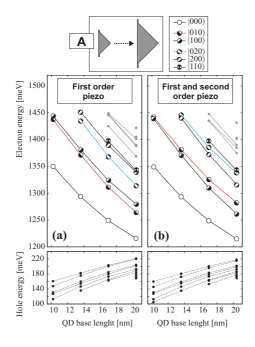


Figure 7: (Color online) Single-particle electron and hole energies for pyramidal QDs with different size (series A). In the left panel (a) only the first order piezoelectric effect is accounted for, using the experimental value of ϵ_{14} . In the right hand panel also the second order is taken into account, using the piezoelectric constants from [11].

sensitive to a k-space wavefunction spreadout. As a result, the wavefunction in real-space is very stiff and reluctant to adapt to small potential fluctuations, because otherwise it would produce $\Psi(\mathbf{k})$ components far from the Γ -point.

For the hole states, especially for those with a large HH-fraction, the situation is reversed. The $E(\mathbf{k})$ function is very flat and a larger $\Psi(\mathbf{k})$ spreadout does result only in a small gain of kinetic energy. Therefore, the hole states adapt to tiny details of the potential landscape in order to minimize their potential energy.

Some principal features of the first and second order piezoelectric potential have already been described in section 3.2. Here we proceed by studying their impact on the single-particle states. For this purpose we have calculated the single-particle energies and wavefunctions for our structures using two piezoelectric models: (1) we used the classical experimental value e_{14} and omitted any quadratic effect and (2) we used the values of e_{14} and $B_{\mu jk}$ from Ref. [11], thus accounting for linear and quadratic piezoelectric terms.

The electron states $|ijk\rangle$ with $i \neq j$ respond very sensitively to the piezoelectric field, whereas those with i=j are almost unaffected. Therefore, to probe the impact of the different orders of piezoelectricity, we monitor the p-states $|100\rangle$ and $|010\rangle$ and the d-states $|200\rangle$ and $|020\rangle$ as a function of the order of the piezoelectric tensor [case (1) or (2)].

Pyramidal and truncated pyramidal QDs (Series A/B). The strength of the first order piezoelectric field and the resulting p and d-state splitting is a function of the QD-height [3] [see also Fig. 7(a)]. As can be seen from Fig. 7(b), this applies even more if the second order is also taken into account; the p-state order, however, is reversed compared to case (1). The same applies to the $|200\rangle$ and $|020\rangle$ d-states.

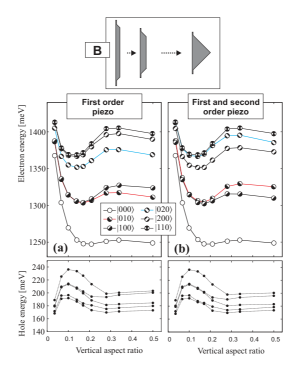


Figure 8: (Color online) Single-particle electron and hole energies for truncated pyramids (series B) as function of the vertical aspect ratio ar_V . In the left panel (a) only the first order piezoelectric effects are accounted for, using the experimental value of ϵ_{14} . In the right panel (b) also the second order is taken into account, using the piezoelectric constants from Ref. [11].

Lens-shaped InAs QDs (Series C). Compared to the series A and B the second order piezoelectric potential exceeds the first order terms inside the QD only in the case of a halfsphere, as can be seen from the reversed p-state splitting in Fig. 9 at $ar_v = 0.5$. For smaller aspect ratios, $ar_v < 0.5$, the p-states and the $|200\rangle$ and $|020\rangle$ d-states are degenerate, indicating that first and second order effects compensate each other with respect to their impact on the electronic states. If we consider the first order piezoelectric field alone [case (2)], even for the structural $C_{\infty v}$ QDs a p- and d-state splitting is found. For the hole orbitals we do not observe any degeneracy neither for case (1) or (2).

Comparison of wavefunction orientation between square and circular based InAs QDs (series B and C).

Figure 10 shows the order and orientation of the wavefunctions for a truncated pyramid from series B and a lens-shaped QD from series C, both sharing the same vertical aspect ratio of 0.21. As long as only the first order terms of the piezoelectric fields are considered, all electron wavefunctions have the same orientation, except for the state $|e_5\rangle$. If the second order is also taken into account, the p- and d-orbitals of the truncated pyramid change their orientation. For the lens-shaped QD the p-orbitals keep their orientation, although they are almost degenerate in energy, as are the d-orbitals.

Elongated QDs (Series D). In this series two of the main lateral anisotropy sources are contrasted: The elongation in [110] and $[1\overline{1}0]$ direction and the piezoelectric effect. To separate their influence on the single-particles we first calculate the electronic states in absence [Fig. 11(a)] and in presence of the piezoelectric field [Fig. 11(b,c)].

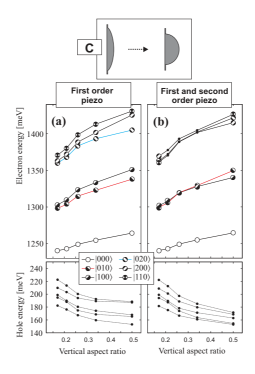


Figure 9: (Color online) Single-particle electron and hole energies for lens-shaped QDs (series C) versus the vertical aspect ratio ar_V . In the left panel (a) only the first order piezoelectric effect is accounted for, using the experimental value of ϵ_{14} . In the right hand panel (b) also the second order is taken into account, using the piezoelectric constants from Ref. [11].

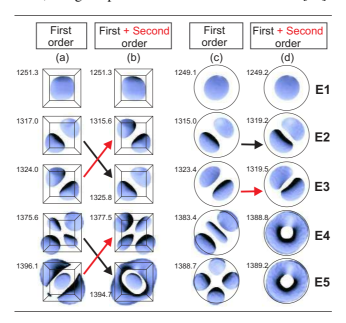


Figure 10: (Color online) Comparison of the electron wavefunction shapes and state ordering for first and second order piezoelectric effects. On the left hand side a truncated pyramid from series B and on the right hand side a flat lens-shaped QD from series C are considered. Results for the electron states in absence of a piezoelectric field are shown in Fig. 4. The use of first order experimental constants lead to the same symmetry properties and state ordering as for the calculated values. Energy values in meV are given with respect to the unstrained VB-edge.

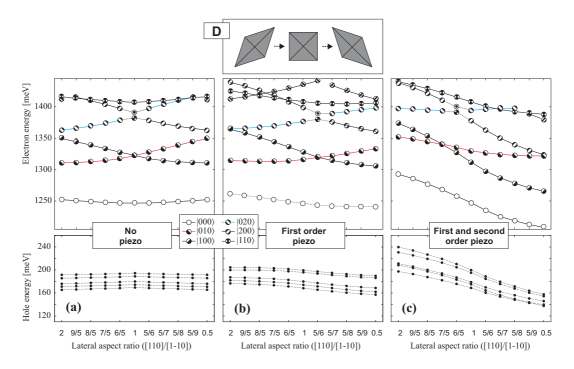


Figure 11: (Color online) Single-particle electron and hole energies for elongated QDs (series D) versus the lateral aspect ratio $\operatorname{ar_L}$. No piezoelectricity is included for the results in panel (a). In panel (b) only the first order piezoelectric effect is accounted for, using the experimental value of ϵ_{14} . In the right hand panel (c) also the second order is taken into account, using the piezoelectric constants from Ref. [11].

In absence of the piezoelectric field the electronic states are insensitive to the orientation of the QD. The spectrum shown in Fig. 11(a) is symmetric and the electron p-states are degenerate at a lateral aspect ratio $\operatorname{ar_L} = 1.0$. The electronic d-states and all hole states, in contrast, show no degeneracies at all. Moreover, the d-states show an anticrossing behavior upon changing the lateral aspect ratio from values smaller than one to values larger than one.

Taking the piezoelectric field into account complicates the picture a lot [Fig. 11(b,c)]. Again, the second order piezoelectric field dominates over the first order field. This can be seen from the p-level crossing point, which is at 1.2 for case (1) (first order only) and -1.4 for case (2) (first and second order). The second remarkable observation is the large energy shift of all electron and hole states for case (2) [Fig. 11(c)], which is much more pronounced than for case (1). However, since the shift applies to both particle types in the same way, it is not expected to be visible in the transition energies, but rather in the activation energies.

Variation of InGaAs composition and its distribution (Series E/F/G). In these series different manifestations of composition changes are investigated. First, the average composition determines the accumulated hydrostatic strain inside the structure. It is, in general, larger for larger In integral content. Since the second order piezoelectric tensor couples diagonal strain components with shear strain components, the quadratic part of the piezoelectric potential is expected to be very sensitive to composition changes.

In series E the average In content in a pyramid is varied homogeneously. The p-state splitting resulting from the first order parts alone are not affected by the composition variation [Fig. 12(a)], since the decreasing shear strain is compensated by an increasing piezoelectric constant e_{14} ,

resulting from the linear interpolation between the values of GaAs ($e_{14} = -0.16$) and InAs ($e_{14} = -0.045$). The second order contributions, in contrast, decrease rapidly for increasing Ga content, as can be seen in Fig. 12(b), where the *p*-states cross at a Ga fraction of 20 %. This is due to the decreasing magnitude of the diagonal components of the strain tensor, which enter only the second order terms, but not the first order ones.

In series *F*, a trumpet-shaped composition profile is investigated for different vertical aspect ratios. Since the In atoms are spread over a wider range in a larger QD with only a small region of high In concentration, the local strain (especially the diagonal components) is smaller. This translates into a smaller first order and into a *much* smaller second order piezoelectric potential. The order of the electron states remains unchanged upon changes of the aspect ratio and introduction of second-order piezoelectricity (Fig. 13).

The most interesting series in this context, however, is series G. Here we applied a couple of annealing steps to the QD interfaces simulated by Fickian diffusion. We found that the strength of the second order piezoelectric potential is extremely sensitive to this procedure, as can be seen from the crossing of the p and d-states ($|200\rangle$ and $|020\rangle$) after two annealing steps [see Fig. 14(lower panel)]. The first order potential in contrast is not affected at all. This behavior can be related to the strong abrupt decrease of the exciton fine-structure splitting, which has been observed experimentally for a first very modest annealing step.[6] The fine-structure splitting in QDs is related to the same sort of anisotropy which causes the p-state splitting.[5]

4.3 Impact of the piezoelectric field on the inter- and intraband spectra

The optical spectra are determined by energy, shape and orientation of the electronic states. Especially the electron p-shell is sensitive to the various sources of lateral anisotropy: QD elongation and the ASA effect both stretch the first electron and hole p-states into the same direction. A linear piezoelectric field, in contrast, results in an alignment in opposite directions (see Fig.15).

Interband absorption spectra. In Fig. 16(a1) and (b1) we compare the excitonic absorption spectra of a pyramidal QD (17.2 nm base length) for the two considered cases of the piezoelectric effect. The labels indicate the symmetry of the electron state (Eijk) and the hole state Hl, which take part in the absorption process. At first sight the spectra look very similar apart from small differences in the polarization. But some of the peaks have changed their origin, e.g., the E010-H1 and the E100-H1 absorption peak have reversed order in both plots.

Intraband spectra. The situation is different for CB intraband transitions [Fig.16(a2) and (b2)]. Here the spectroscopic signature is completely different in terms of the peak energies and in particular with respect to the polarization. The transitions E010-E000 and E100-E000, being the acid test for the sequence of the p-states order have reversed order. In contrast to the interband peaks the *p*-state transitions are clearly distinguishable by their polarizations: E010-E000 is polarized along [110] and E010-E000 along [110].

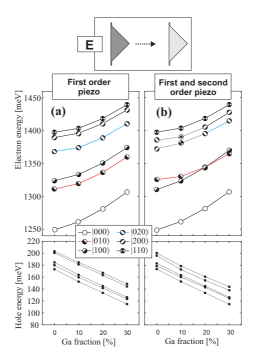


Figure 12: (Color online) Single-particle electron and hole energies for series E versus the Ga fraction inside the InGaAs QD. In the left panel (a) the first order piezoelectric effect is accounted for using the experimental value of ϵ_{14} only, whereas for the right hand panel also the second order is taken into account, using the piezoelectric constants from Ref. [11].

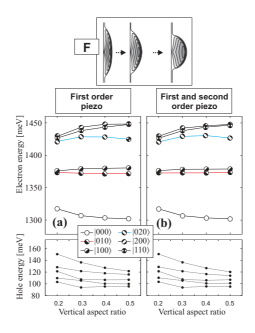


Figure 13: (Color online) Single-particle electron and hole energies for series F versus the vertical aspect ratio ar_V . In the left panel (a) the first order piezoelectric effect is accounted for using the experimental value of ϵ_{14} only, whereas for the right hand panel also the second order is taken into account, using the piezoelectric constants from Ref. [11].

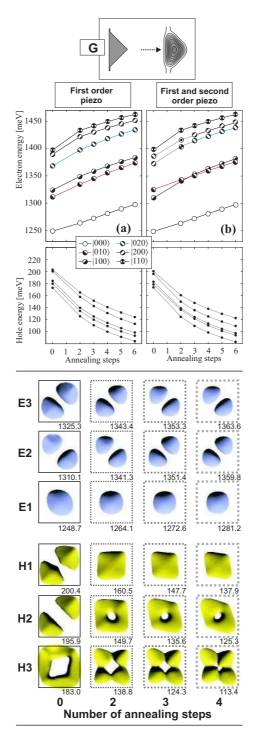


Figure 14: (Color online) (upper panel) Single-particle electron and hole energies for series G versus the number of annealing steps. In the left panel (a) the first order piezoelectric effect is accounted for using the experimental value of ϵ_{14} only, whereas for the right hand panel also the second order is taken into account, using the piezoelectric constants Ref. [11]. (lower panel) Probability density (isosurface at 65 %) shown for the first three bound electron and hole orbitals as a function of annealing steps. An electron p-states reordering occurs between annealing step two. Only a small degree of annealing is necessary to significantly change the hole wavefunction shape. Energy values are given with respect to the unstrained VB-edge.

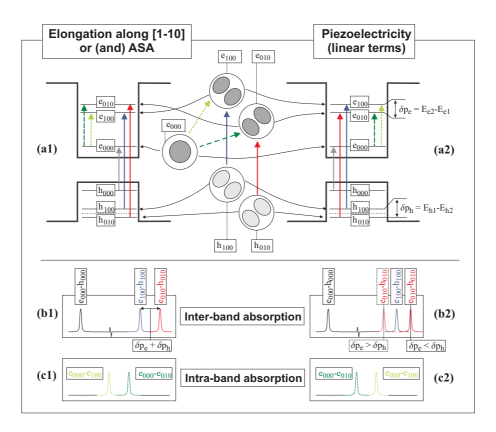


Figure 15: (Color online) A possible QD elongation or the effect of the atomistic symmetry anisotropy (ASA) has a qualitatively different impact on the level ordering than a piezoelectric field (here schematically shown for first order terms only). In the latter case the first electron and hole p-states are oriented in orthogonal direction. Panel (b) schematically shows the resulting interband absorption spectra. In (b1) the p-channel splitting is the sum of the respective electron and hole p-state splitting ($\delta p_e + \delta p_h$), whereas in case (b2) the splitting is determined by the difference ($\delta p_e - \delta p_h$). Since the polarization (not shown here) of these peaks is rather weak, a distinction between the two p-channel peaks might be difficult in experiment. An additional hindsight can provide intraband transition spectra (c). These peaks are nearly 100% polarized and allow a clear assignment of the transition type.

5 The vertical and lateral aspect ratio

5.1 Vertical aspect ratio (series B and C)

Since the relation between the vertical aspect ratio and piezoelectric field has already been discussed in Sec. 4.2, we continue with the discussion of the influence of the vertical aspect ratio on the single-particle states, using series B (square base) and C (circular base) and highlighting the following results:

- (i) As long as the aspect ratio is between 0.15 and 0.5, the localization of electron and hole states is either constant (electrons in series A) or increases (energy of electron states decrease and of hole states increase) with decreasing aspect ratio. This unusual finding is in contrast to what a simple particle-in-a-box model suggests and is related to the redistribution of strain inside the QD. In general two competing processes determine the localization energies. The first one is the rising quantization in z-direction, resulting in a decreasing localization. This effect, however, becomes dominant only very flat QDs [ar_v < 0.15 (0.1) for electrons (holes), see Fig. 8]. The second one is a redistribution of strain from being dominant hydrostatic to more biaxial [Fig.3 (a) and (c)]. This process reduces the local band gap [Fig.3(b) and (d)], thus, increasing the localization energy, as discussed in section 3.3.1. As a result, the excitonic absorption spectrum of a flat QD, as shown in Fig.16(c1), is red-shifted by ca. 30 meV compared to the pyramid in panel (b1).
- (ii) The increase of the localization energy with decreasing aspect ratio is much more pronounced for circular-based QDs than for square-based ones. This points at a larger degree of strain redistribution in lens-shaped QDs.
- (iii) For very flat QDs, the the z-quantization becomes dominant; the critical aspect ratio, however, is different for electron and hole in series B:

Electron shift: For $ar_V < 0.15$ the decreasing height causes an energy shift of 120 meV for the electron ground-state and 80 meV for the first excited electron state. At first, for $0.1 \le ar_V < 0.15$, the wavefunctions are compressed, thus accumulating kinetic energy. Later, for $ar_V < 0.1$, when the z-confinement becomes too strong, they evade by spreading into the surrounding matrix, increasing the potential energy of the state. Both effects result in an increased electron energy.

Hole shift: Due to their larger mass, hole states in general prefer to adapt their wavefunction shape close to the details of the VB-confinement profile. Thus the major part of the hole wavefunctions remain inside the QD until the aspect ratio becomes smaller than 0.1. Below that value, the energy decreases by 50 meV for the ground state and 43 meV for the first excited state. These shifts are attributed to the gain of kinetic energy, rather than to a barrier penetration. Even for the flattest QD, 65 % of the hole ground state wavefunction remains inside the dot, in contrast to 47% for the electron ground state.

5.1.1 Excitonic absorption spectra

A comparison of the excitonic absorption spectra of the full pyramidal and the truncated QD reveals significant differences, reflecting the different strain distributions and piezoelectric fields.

The magnitude of the shear strain components decreases with decreasing vertical aspect ratio, resulting in a smaller piezoelectric field and reduced HH-LH coupling. The former reduces the electron p-splitting leading to a degeneracy of the E100-H1/E010-H1 transitions, the latter re-

duces the LH-percentage of the hole states. Thus, the number of absorption channels is decreased from 10 in the case of the full pyramid to 7 in the case of the flat truncated pyramid. The E000-H2 peak of the pyramid in Fig. 16(b1), for instance, is only visible because the sizeable LH-part of H2 has the same s-like symmetry as the E000 state. The HH-part of H2, in contrast, carries $|110\rangle$ -symmetry and can therefore not interact optically with the E000 state, as its overlap integral vanishes for symmetry reasons. For the flat truncated QD in Fig. 16(c1) the LH-percentage of H2 is small. Consequently, the E000-H2 peak vanishes.

The overall appearance of the absorption spectra are quite different for different aspect rations and should allow a discrimination in PLE experiments.

5.2 Lateral aspect ratio (series D)

A QD elongation is often discussed as a possible source of the exciton fine-structure splitting, since it introduces a symmetry reduction from C_{4v} to C_{2v} already on the level of the QD structure. However, as long as no piezoelectricity (and/or the ASA in the case of atomistic models) is included, there is no distinction possible between elongations along [110] and [110] on the basis of the single particle energies or the peak energies of the excitonic spectra. In this case only the peak polarization delivers the information on the QD orientation.

In Fig. 17 interband and the CB intraband spectra are shown for two QDs with the same shape but aligned along different directions. Their lateral aspect ratios are $(ar_L=5/7)$ and $(ar_L=7/5)$ respectively. Linear and quadratic terms of the piezoelectric field are included in the calculation. The excitonic absorption spectra differ in their peak positions, their intensity, and, in particular, in their polarization degree. The polarization is even more pronounced in the intraband transition spectra: For a QD elongated along $[1\overline{1}0]$ ($ar_L=5/7$) only those lines are visible in the considered energy range, which are polarized in $[1\overline{1}0]$ direction. The piezoelectric effect and the elongation sum up with respect to their impact on the *p*-state splitting. For the other case, a QD elongated along [110], the piezoelectric effects and the elongation are compensating each other to a large degree and, as a result, the transitions E010-E000 and E100-E000 are very close in energy. Hence, the two orientations are clearly distinguishable by energy and polarization of their intraband spectra.

6 Varying composition profiles

6.1 Inverted cone like composition profile

The experiments of Fry *et al.* [50], employing Stark effect spectroscopy, have shown that, for certain QDs, the hole groundstate is localized towards the top of the dot, slightly above the electron. Barker *et al.* [51] have conjectured that this observation can be attributed to an InAs gradient towards the upper part of the QD. Later on, Sabathil *et al.* [52] studied the impact of different composition profiles on the behavior of the electron-hole dipole in presence of a lateral electric field.

These investigations were accompanied by structural investigations using X-TEM[53] and X-STM[17, 54], guiding us to the choice of the composition profile of series *F*, which will be discussed in this paragraph.

In order to identify the consequences of the inhomogeneous composition profile like in series F, we compare the flattest QD of this series ($\operatorname{ar_v}=0.2$) (further referred to as $\operatorname{QD}_{F-inhom}^{0.2}$) to the pure InAs, lens-shaped QD from series C with the same vertical aspect ratio (further referred to as $\operatorname{QD}_{C-hom}^{0.2}$). Both QDs contain the same integral amount of In. Compared to the archetype pyramidal QD, the electron-hole alignment of $\operatorname{QD}_{F-inhom}^{0.2}$ is indeed reversed and their barycenters are separated by 0.2 nm. We would like to point out that this separation is very sensitive to the choice of the VB-deformation potential parameter a_v . Literature values of a_v scatter remarkably between $a_v = 1 \, eV$ and $a_v = -1 \, eV$ (see Vurgaftman $et \, al.$ [55]).

Comparing single particle energies (Figs. 9 and 13) and absorption spectra (Fig. 19) of $QD_{C-hom}^{0.2}$ and $QD_{F-inhom}^{0.2}$ yields the following results:

- (i) The electron and hole ground state energies are shifted by 70 and 80 meV to higher energies for $QD_{F-inhom}^{0.2}$, resulting in a 150 meV larger groundstate exciton absorption energy.
- (ii) The peak order in the excitonic absorption spectra remains unchanged (Fig. 19 left panel).
- (iii) The electron s-p sublevel spacing remains the same, other than the p-d sublevel spacing. As a result, the E010-E000 intraband transition appears at almost the same energy, but the d-p transitions of $\mathrm{QD}_{F-inhom}^{0.2}$ are shifted to lower energies (Fig. 19 right panel).

6.2 Annealed QDs

In series G we simulate the effect of annealing on the electronic properties using a pyramidal QD, with a base length of 17.2 nm as model structure. We find the following peculiarities as result of the annealing procedure:

- (i) The electron and hole groundstate energies shift by 50 and 70 meV to higher energies, resulting in a blue-shift of the exciton groundstate absorption energy of ca. 120 meV [see Figs. 14 and 18 (b1)].
- (ii) The *s-p* level spacing and the hole sublevel spacing increases, resulting in a larger separation of the s-channel from the p-channel transitions. This finding is surprising at first sight, since high excitation PL results for annealed QD ensembles reveal a reduced *s-p* channel spacing.[56] These experiments, however, were performed in an energy range closer to the GaAs bandgap. In this energy range the excited states start to delocalize, resulting in a decreasing electron *s-p* splitting.
- (iii) Due to the increasing *s-p* level spacing, the intraband transition shift to higher energies [Fig. 18 (b2)].

6.3 InGaAs QDs with uniform composition

The choice of the average $In_xGa_{1-x}As$ composition of QDs is often employed to taylor the emission wavelength. The impact of the Ga content (1-x) on the electronic properties is investigated using series E. From Figs. 12 and 18 we can derive the following properties:

- (i) An increase of the Ga content by 30% shifts the electron and hole groundstate by 55 meV each to higher energies, resulting in a blue-shift of the exciton groundstate absorption energy of ca. 110 meV [Fig. 18 (a1)].
- (ii) In contrast to our findings for the annealing series G, the separation of the s- and p-shell electrons decreases. Hence, by comparing Figs. 18(a1) and (b1), we find the p-channel transitions

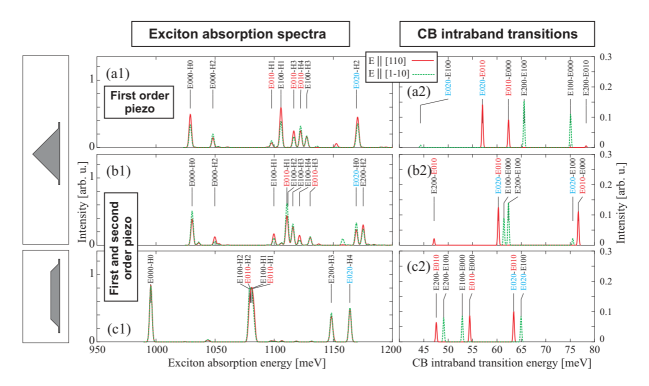


Figure 16: (Color online) Excitonic absorption spectra (left panel) and CB intraband transition spectra (right panel) shown for a full InAs pyramid and a flat truncated pyramid both taken from series *B*. In (a) and (b) the results are contrasted for the two approaches for calculating the piezoelectric field.

much closer to the s-channel transitions than in series G. The resulting intraband transitions are shifted to lower energies [Fig. 18 (a2)].

7 Conclusions

We investigated systematically the correlation between structural QD properties like size, shape, and composition and their electronic and optical properties.

- (i) We compared the electronic properties of round and square based InAs/GaAs QDs with constant volume but varying aspect ratios. We found a large redistribution of strain from dominantly hydrostatic, if the aspect ratio is 0.5 (full pyramid and half sphere, respectively), to pronounced biaxial for flat dots. The reduced average energy gap in case of flat QDs results in an increasing electron and hole localization, a finding that is more prominent for QDs with a circular base. When the QD height becomes smaller, the HH-LH coupling decreases, since the shear strain components tend to disappear. As a result, the LH-percentage of the hole ground-states drops from around 10 % down to 2.8 %.
- (ii) The transition energies of pure InAs QDs (series C) are smaller than those with Ga incorporation (see series F), even if the integral amount of In is the same in both cases. QDs with nonuniform composition profiles (like series F) have larger transition energies.
- (iii) We simulated the effect of annealing on a pyramidal QD. The electron and hole groundstate energies are shifted by 50 and 70 meV to higher energies, resulting in a blue-shift of the exciton

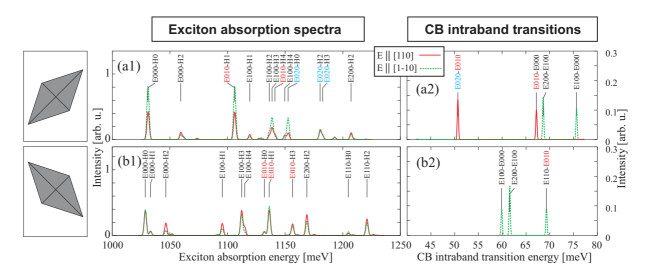


Figure 17: (Color online) Excitonic absorption spectra (left panel) and CB intraband transition spectra (right panel) shown for parallelepipedal elongated QDs from series D having different lateral orientations with $(ar_L=5/7)$ and $(ar_L=7/5)$ respectively. First and second order piezoelectric effects are included.

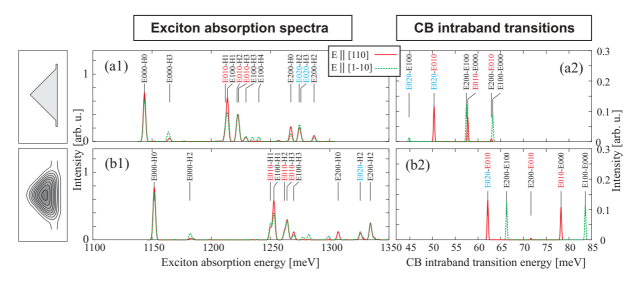


Figure 18: (Color online) Excitonic absorption spectra (left panel) and CB intraband transition spectra (right panel) shown for the unannealed and the strongest annealed QD of series G. First and second order piezoelectric effects are included.

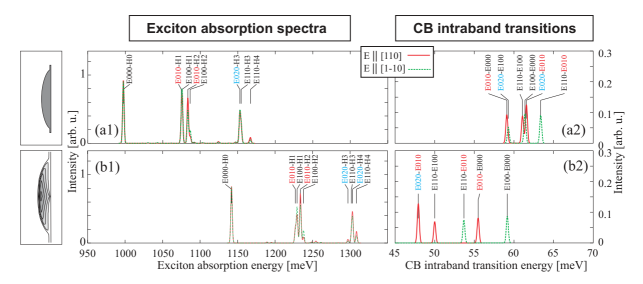


Figure 19: (Color online) Excitonic absorption spectra (left panel) and CB intraband transition spectra (right panel) shown for two QDs having a circular base. The first one is taken from series C and the other from series F, both having the same total amount of InAs inside the QD structure and a similar vertical aspect ratio of $\operatorname{ar}_V = 0.21$ (a) and $\operatorname{ar}_V = 0.2$ (b) respectively. First and second order piezoelectric effects are taken into account.

groundstate transition energy of ca. 120 meV. The *s-p* level spacing and the hole sublevel spacing increase, resulting in a larger separation of the *s*-channel from the p-channel transitions. The corresponding electron and hole wavefunctions are stronger localized.

- (iv) We compared the classical approach of calculating the piezoelectric field in QDs [2] to the one recently developed by Bester *et al.*,[11, 12] which takes into account the second order piezoelectric field, for a wide range of different QD structures. We found the potential arising from the quadratic terms to be very sensitive to the base shape, the vertical aspect ratio, and the composition. Its orientation inside the QD is reverse compared to the first order potential and can surpass it, resulting in a reversal of the electron *p* and *d*-state ordering and a reorientation of the hole wavefunctions, as observed for InAs pyramidal QDs. In this case the quadratic terms exceeds the linear contributions inside the QD. Upon gradual annealing of this QD, thus introducing an more isotropic composition gradient, the second order field decreases dramatically, leaving the first order contribution as the dominant part. As a result, the electron *p* and *d*-state order and the hole wavefunction orientations change. The same is observed for a pyramid with increasing, homogeneously inserted, Ga content.
- (v) We calculated the excitonic absorption spectra and the CB-intraband transitions. The former are very sensitive to almost all applied structural changes and to the piezoelectric field, leading to strong modification of the p- and d-channel transitions. Upon changing the model for the piezoelectric field calculations, a peak reordering within these absorption channels can occur and the polarization anisotropy between $e \parallel [110]$ and $e \parallel [1\overline{10}]$ can change. CB-intraband-transition spectra are strongly polarized either along [110] or $[1\overline{10}]$ and very sensitive to changes of the piezoelectric field or any other anisotropy.

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