Multiple charging of InAs/GaAs quantum dots by electrons or holes: Addition energies and ground-state configurations

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Atomistic pseudopotential plus configuration interaction calculations of the energy needed to charge dots by either electrons or holes are described, and contrasted with the widely used, but highly simplified two-dimensional parabolic effective mass approximation (2D-EMA). Substantial discrepancies are found, especially for holes, regarding the stable electronic configuration and filling sequence which defies both Hund’s rule and the Aufbau principle.

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I. INTRODUCTION

One of the most spectacular aspects of quantum-dot physics is that dots can be controllably charged by either electrons or holes and that one can measure, for each of the many charged-states, both the electronic spectrum and the charging energies. This is afforded either by injecting from a tip of a scanning tunneling microscope, or by various gate structures. Since the energy scale of both single-particle levels and Coulomb interactions in quantum dots (QDs) ($10^{-4}$–$10^{-2}$ Ry) are a few order of magnitudes smaller than those of the real atoms (~1 Ry), dots can be loaded by as many as six to ten electrons in colloidal and self-assembled dots having confining dimension of $\leq$50 Å, and up to hundreds of electrons in larger 500 Å electrostatically confined dots. The “charging energy” $\mu(N)$ is the energy needed to add a carrier to the dot that is already loaded by $N-1$ carriers,

$$\mu(N) = E(N) - E(N-1),$$

where $E(N)$ is the correlated, many-body total energy of the ground state of the $N$-particle dot. The “addition energy” $\Delta(N-1,N)$ (analogous to the difference between ionization potential and electron affinity) indicates how much more energy is needed to add the $N$th carrier compared to the energy needed to add the $(N-1)$th carrier:

$$\Delta(N-1,N) = \mu(N) - \mu(N-1) = E(N) - 2E(N-1) + E(N-2).$$

The typical electron addition energies for electrostatic dots are about 1–8 meV, and the stable spin configuration follows the rules of atomic physics; that is, the $s,p,d,...$ shells are occupied in successive order with no holes left behind (Aufbau principle) and with maximum spin (Hund’s rule). Recently, it became possible to load and measure electrons and holes into much smaller, epitaxially grown self-assembled dots of InGaAs/GaAs, where electron addition energies are about 10–60 meV, and hole addition energies are between 10 and 30 meV. More interestingly, while electrons still follow the Aufbau principle, recent hole charging experiment show that holes have unusual charging patterns that defy the Aufbau principle and Hund’s rule.

Despite the importance of the dot charging problem and the great success achieved in experimentally recording the charging spectra, the theoretical understanding of charging and addition energies is still preliminary. Most theoretical works in the area were based on particle-in-a-potential model, neglecting interband (e.g., $\Gamma$-$\Gamma$) coupling, intervalley ($\Gamma$-$X$-$L$) coupling effect, and the true atomistic symmetry (e.g., $C_{3v}$ for lens of zincblende material) which is lower than the shape symmetry. The most often used potential in such approaches is the two-dimensional (2D) parabolic form, in which all of the above noted electronic structure effects are replaced by an effective mass approximation (EMA). In this parabolic 2D-EMA model, the single particle levels have equal spacing, which equals a harmonic oscillator frequency $\omega$. Because of the simplicity of the model, all the Coulomb integrals can be related analytically to a single orbital Coulomb energy $J_{sr}$, and therefore the addition energies are determined entirely by $\omega$ and $J_{sr}$. Although the parabolic 2D-EMA model can be attractive because of its algebraic simplicity and availability of fitting parameters actual self-assembled dots are significantly different from the description of EMA model, manifesting inter-band coupling and inter-valley coupling, strain effects, low atomistic symmetry, as well as specific band offset profiles, all neglected by the 2D-EMA. It indeed has been recently shown that hole ground state configurations predicted by the 2D-EMA model are qualitatively different from those measured by hole charging experiments.

An atomistic pseudopotential description of electronic structure effects can be used instead of 2D-EMA to calculate charging energies. Here, we show that such an atomistic theory correctly reproduces the many-particle configurations as well as addition spectra for carriers in self-assembled quantum dots. We study systematically the electronic structure of self-assembled InGaAs/GaAs quantum dots, provide detailed information on the electron and hole single-particle spectrum, many-particle charging and addition spectrum, as well as ground state configurations using single-particle pseudo-potential and many-particle configuration interaction (CI) methods.

The rest of the paper is arranged as follows. In Sec. II, we introduce the basic concepts of charging and addition ener-
gies, and show how to calculate these quantities in the single-particle pseudopotential plus many-particle CI scheme. In Sec. III, we give detailed results calculated from pseudopotential-CI scheme the single-particle levels, Coulomb integrals and the ground state configurations as well as the addition energies. We contrast these results with the parabolic 2D-EMA model. We summarize in Sec. IV.

II. THEORY OF DOT CHARGING AND ADDITION ENERGIES

A. General equation for dot charging in the configuration-interaction approach

The calculation of the total-energy $E(N)$ of $N$-particle dot requires obtaining first the single-particle states from an effective Schrödinger equation, and then the many-particle state from a many-particle treatment. The first step is formulated as,

$$\left[-\frac{1}{2}\nabla^2 + V_{\text{ext}}(\mathbf{r}) + V_{\text{sc}}(\mathbf{r})\right] \psi_i(\mathbf{r}) = \epsilon_i \psi_i(\mathbf{r}),$$ (3)

where $V_{\text{ext}}(\mathbf{r})$ is the external (“bare”) potential experienced by the electrons or holes, and $V_{\text{sc}}(\mathbf{r})$ is the screening response. The single-particle orbital $\{\psi_i\}$ and energies $\{\epsilon_i\}$ are used in the second step to construct the many-particle wave functions $\{|\Psi\rangle\}$ and energies $\{E\}$ from,

$$E(N) = \langle \Psi_N | H | \Psi_N \rangle$$ (4)

where, the many-body Hamiltonian is,

$$H = \sum_{i\sigma} \epsilon_i \hat{c}_{i\sigma} \hat{c}_{i\sigma}^\dagger + \frac{1}{2} \sum_{ijkl} \sum_{\sigma_1 \sigma_2 \sigma_3 \sigma_4} \Gamma_{ijkl}^{\sigma_1 \sigma_2 \sigma_3 \sigma_4} \hat{c}_{i\sigma_1} \hat{c}_{j\sigma_2} \hat{c}_{k\sigma_3} \hat{c}_{l\sigma_4},$$ (5)

and,

$$\Gamma_{ijkl}^{\sigma_1 \sigma_2 \sigma_3 \sigma_4} = \sum_{\mathbf{s}_1 \mathbf{s}_2} \int \int d\mathbf{r} d\mathbf{r}' \psi_{i\sigma_1}^*(\mathbf{r}, \mathbf{s}_1) \psi_{j\sigma_2}^*(\mathbf{r}', \mathbf{s}_2) \psi_{k\sigma_3}^*(\mathbf{r}', \mathbf{s}_2) \psi_{l\sigma_4}^*(\mathbf{r}, \mathbf{s}_1) \frac{e^{i (\mathbf{r} - \mathbf{r}')}}{(|\mathbf{r} - \mathbf{r}'|)} e^{i (\mathbf{r} - \mathbf{r}')},$$ (6)

are the screened Coulomb and exchange integrals. In the above Eqs. (5) and (6), we use $\hat{c}_{i\sigma} = c_{i\sigma}^\dagger c_{i\sigma}$ as the field operator, whereas $c_{i\sigma}$ is a fermion operator, and $\psi_{i\sigma}(\mathbf{r})$ is the single-particle eigenfunction. Here, “$\sigma$” is a pseudospin index, i.e., an index of Kramers degenerate states, while “$s$” is the intrinsic electronic spin. For electrons in InAs/GaAs QDs, the spin-orbit interactions is extremely small and can be neglected. In this case, the pseudospin $\sigma$ and intrinsic electronic spin $s$ are equivalent. However, for holes, which have a mixture of heavy- $(H)$ light-hole $(LH)$ and split-off character, an eigenstate of $\sigma$ has both $s=\uparrow$ and $s=\downarrow$ components. The $N$-particle wave functions can be solved using, e.g., configuration interaction (CI) method,23 by expanding the $N$-electron wave function in a set of Slater determinants, $|\Phi_{e_1 e_2 \ldots e_N}\rangle = c_{e_1}^\dagger c_{e_2}^\dagger \ldots c_{e_N}^\dagger |\Phi_0\rangle$, where $c_{e_i}^\dagger$ creates an electron in the state $e_i$. The $r$th many-particle wave function is then the linear combination of the determinants,

$$|\Psi_N^{(r)}\rangle = \sum_{e_1 e_2 \ldots e_N} A_r(e_1, e_2, \ldots, e_N) |\Phi_{e_1 e_2 \ldots e_N}\rangle.$$ (7)

Once $|\Psi_N\rangle$ is known, we can then calculate the corresponding total energies for the ground states as well as excited states using Eq. (4). Once we solve the CI problem, we get the order of total CI energy for various holes or electron configurations, so we can see if Hund’s rule or the Aufbau principle or spin-blockade occurs. For example, Hund’s rule states that degenerate single-particle levels are occupied with maximum number of unpaired electrons, while the Aufbau principle states, nondegenerate single-particle levels are occupied in order of increasing single-particle energy.

We construct all possible Slater determinants corresponding to $N$ electrons or $N$ holes [i.e., we ignore the excitonic (electron+hole) excitations], using only the bound states of the dots, (i.e., we neglect all continuum states). The underlying electrons that are not considered explicitly by this approach are represented by the dielectric screening function $e(r-r')$ in Eq. (6).

B. The Hartree-Fock equations for charging and addition energies

The addition energies at CI level can be written as the Hartree-Fock (HF) addition energies plus the correlations, i.e.,

$$\Delta_{\text{CI}}(N-1,1) = \Delta_{\text{HF}}(N-1,1) + \Delta_{\text{corr}}(N-1,1),$$ (8)

where $\Delta_{\text{corr}}$ is the correlation energy correction to the addition energy calculated in HF. Since the HF equations are used by many experimentalists to deduce Coulomb energies,3,8,11 we review it here. In the Hartree-Fock approximation, where the effect of correlations is neglected but the direct Coulomb and exchange interactions are retained, simple expressions can be derived for the addition energies. The total energy of $N$ electrons is simply,

$$E_{\text{HF}} = \sum_{i\sigma} \epsilon_{i\sigma} + \sum_{i\sigma, j'\sigma'} (J_{i\sigma, j'\sigma'} - K_{i\sigma, j'\sigma'}),$$ (9)

where, $i$ is the single-particle level index of all occupied states, and $\epsilon_{i\sigma}$ is the corresponding single-particle energy. The $J_{i\sigma, j'\sigma'} = \Gamma_{j'\sigma',i\sigma}^{\sigma} \Gamma_{i\sigma,j'\sigma'}$ and $K_{i\sigma, j'\sigma'} = \Gamma_{j'\sigma',i\sigma}^{\sigma} \Gamma_{i\sigma,j'\sigma'}$ in Eq. (9) are Coulomb and exchange integrals, respectively. Since the spin index “$\sigma$” is not an actual electronic spin, but rather an index for two Kramers degenerate states, in principle the exchange integrals $K_{i\sigma, j'\sigma'}$ are not simply diagonal in $\sigma$, $\sigma'$, as has been widely used in the dot charging literature.3,11,15,16 However, adopting the literature approximation, $K_{i\sigma, j'\sigma'} = K_{i\sigma,j'\sigma'}$, considering the $s$ and two $p$ orbitals, $\epsilon_{p_1} < \epsilon_{p_2}$ and assuming the particle filling order follows Hund’s rule, as shown in Fig. 1, the total energies for $N=1,2,3,4$ electrons in the Hartree-Fock approximation are

$$E_{\text{HF}}(1) = \epsilon_s,$$

$$E_{\text{HF}}(2) = 2\epsilon_s + J_{ss},$$

$$E_{\text{HF}}(3) = 2\epsilon_s + \epsilon_{p_1} + J_{sp_1} + 2J_{sp_1} - K_{sp_1},$$

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Similarly, we can calculate the addition energies Eq. (1),

\[ E_{HF}(4) = 2\epsilon_s + \epsilon_{p_1} + \epsilon_{p_2} + J_{ss} + 2J_{sp_1} + 2J_{sp_2} + J_{p_1p_2} - K_{sp_1} - K_{sp_2} - K_{p_1p_2}, \]

\[ E_{HF}(5) = 2\epsilon_s + 2\epsilon_{p_1} + \epsilon_{p_2} + J_{ss} + 4J_{sp_1} + 2J_{sp_2} + 2J_{p_1p_1} + 2J_{p_1p_2} - 2K_{sp_1} - K_{sp_2} - K_{p_1p_2}, \]

\[ E_{HF}(6) = 2\epsilon_s + 2\epsilon_{p_1} + 2\epsilon_{p_2} + J_{ss} + 4J_{sp_1} + 4J_{sp_2} + J_{p_1p_1} + J_{p_1p_2} + 4J_{p_1p_2} - 2K_{sp_1} - 2K_{sp_2} - 2K_{p_1p_2}. \]

We can then readily calculate the charging energies Eq. (1) in this approximation,

\[ \mu_{HF}(1) = \epsilon_s, \]

\[ \mu_{HF}(2) = \epsilon_s + J_{ss}, \]

\[ \mu_{HF}(3) = \epsilon_{p_1} + 2J_{sp_1} - K_{sp_1}, \]

\[ \mu_{HF}(4) = \epsilon_{p_1} + 2J_{sp_1} + J_{p_1p_1} - K_{sp_1} - K_{p_1p_2}, \]

\[ \mu_{HF}(5) = \epsilon_{p_1} + 2J_{sp_1} + J_{p_1p_1} + J_{p_1p_2} - K_{sp_1}, \]

\[ \mu_{HF}(6) = \epsilon_{p_1} + 2J_{sp_1} + 2J_{p_1p_2} + J_{p_1p_2} - K_{sp_1} - K_{p_1p_1}. \]

Similarly, we can calculate the addition energies Eq. (2) as follows,

\[ \Delta_{HF}(1, 2) = J_{ss}, \]

\[ \Delta_{HF}(2, 3) = (\epsilon_{p_1} - \epsilon_s) + 2J_{sp_1} + J_{ss} + K_{sp_1}, \]

\[ \Delta_{HF}(3, 4) = (\epsilon_{p_1} - \epsilon_{p_2}) + 2J_{sp_1} - 2J_{sp_2} + J_{p_1p_1} + K_{sp_1} + K_{sp_2} + K_{p_1p_2}, \]

\[ \Delta_{HF}(4, 5) = (\epsilon_{p_1} - \epsilon_{p_2}) + 2J_{sp_1} - 2J_{sp_2} + J_{p_1p_1} - K_{sp_1} + K_{sp_2} + K_{p_1p_2}. \]

From above equations, we see that to predict \( \Delta(N-1, N) \) for the first two cases \( (N = 3) \), we need to know 4 parameters \( (J_{ss}, J_{sp_1}, K_{sp_1}, \epsilon_{p_1} - \epsilon_s) \), but for \( N = 6 \), we need to know 9 parameters. Because of the large number of parameters needed, the analysis of charging effects in the literature resort to additional approximations aimed at reducing the number of parameters using simplified effective mass models. Equations (12) are going to be used below to contrast the charging spectra deduced from simplified literature models versus our more complete treatment.

\[ \Delta_{HF}(5, 6) = (\epsilon_{p_1} - \epsilon_{p_2}) + 2J_{sp_1} - 2J_{sp_2} + J_{p_1p_1} + J_{p_1p_2} + K_{sp_1} - K_{sp_2} - K_{p_1p_2}. \]

C. Atomic treatment of the single-particle problem

The most general treatment of the single-particle problem of Eq. (3) describes both \( V_{ext} \) and \( V_{scr} \) atomistically, much in the same way as molecules are treated quantum-mechanically. In this description \( V_{ext}(\mathbf{r}) \) is a superposition of the ionic potential of individual atoms of type \( \alpha \) at lattice site \( n \),

\[ V_{ext}(\mathbf{r}) = \sum_{\alpha} \sum_{n} v_{\text{ion}}^{(\alpha)}(\mathbf{r} - \mathbf{r}_n), \]

and \( v_{\text{ion}}^{(\alpha)} \) is \(-Z_{\alpha}/r\) in an all-electron (core+valence) treatment (where \( Z_{\alpha} \) is the atomic number) or \( v_{ps}^{(\alpha)}(\mathbf{r}) \) in the pseudopotential (valence-only) scheme [where \( v_{ps}^{(\alpha)}(\mathbf{r}) \) is the ionic pseudopotential]. The potential \( V_{ext} \) naturally contains the correct point-group symmetry of the object, through the atomic position vectors \( \{\mathbf{r}_n, \mathbf{R}_\alpha\} \), and includes atomic relaxation if appropriate (again, through the atomic positions), as well as chemical inhomogeneity (alloying) or surface-passivation effect. The screening response \( V_{scr}(\mathbf{r}) \) of Eq. (3) is in general a functional of the density-matrix \( \rho(\mathbf{r}, \mathbf{r'}) \) and can be described, e.g., via Hartree-Fock or the density-functional theory, both requiring a self-consistent (iterative) solution to Eq. (3). These approaches are currently limited to small dots, relative to the \( 10^3-10^5 \) atom dots which charging experiments exist. Furthermore, local density approximation (LDA) suffer from the famous “LDA error,” whereby the band gap and effective masses are badly underestimated. Higher-order method such as GW approximation or time-dependent density functional theory (DFT) have yet to demonstrate applicability to large dots for which high-quality experiments exist.

An approximation to the screening \( V_{scr}[\rho(\mathbf{r}, \mathbf{r'})] \), which allows calculation on large dots, and fixes the LDA error, is provided by the “screened pseudopotential approach,” where it is assumed that \( V_{scr}(\mathbf{r}) \) can be described as a superposition of screening potentials \( v_{scr}^{(\alpha)}(\mathbf{r}) \) of the individual atoms, and lumping together \( v_{\text{ion}}^{(\alpha)}(\mathbf{r}) + v_{scr}^{(\alpha)}(\mathbf{r}) \) to yield a screened atomic pseudopotential \( v_{ion}^{(\alpha)}(\mathbf{r}) \), such that,
\[ V_{\text{ext}}(\mathbf{r}) + V_{\text{scr}}(\mathbf{r}) = V_{\text{so}}(\mathbf{r}) + \sum_{\alpha} \sum_{n} v_{\text{epm}}^{(\alpha)}(\mathbf{r} - \mathbf{R}_{\alpha}). \]  

(14)

Here, \( v_{\text{epm}}^{(\alpha)}(\mathbf{r}) \) is determined semiempirically. Unlike the classic empirical pseudopotential method, which fitted only to eigenvalues, here we require that when Eq. (3) is applied to the underlying bulk periodic solids containing atom \( \{\alpha\} \), \( \{v_{\text{epm}}^{(\alpha)}\} \) reproduce the measured band energies, effective-mass tensors, deformation potentials, and the single-particle wave functions have a large overlap with the corresponding LDA wave functions. In Eq. (14), a nonlocal potential \( V_{\text{so}}(\mathbf{r}) \) is also added to the total potential to represent the spin-orbit interaction. In our approach, the potential of an As atom depends on the number of Ga and In atoms around it as

\[ v_{\text{As}}(\text{Ga}In_{4-n}) = \frac{n}{4} v_{\text{As}}(\text{Ga}) + \frac{4-n}{4} v_{\text{As}}(\text{In}), \]  

(15)

where \( n \) is the number of Ga atoms around the As atom. In this atomistic approach, one assume that \( v_{\text{epm}}^{(\alpha)}(\mathbf{r}) \) is transferable to different environments. Note that a fixed \( v_{\alpha} \) is a good approximation if the dot has no free surfaces (as is the case in self-assembled dots, where only a strained interface between chemically-similar materials is present). For surface atoms in free-standing dots, a separate \( v_{\text{epm}}^{(\alpha)}(\mathbf{r}) \) is fitted to LDA surface calculations. For InAs/GaAs dots, we use the pseudopotentials of Ref. 19. These pseudopotentials have been tested not only for the InAs and GaAs binaries, but also for alloys and superlattices of the corresponding ternaries.19

Once \( \{v_{\text{epm}}^{(\alpha)}(\mathbf{r})\} \) is known, one can solve Eq. (3) for the bulk solid, quantum wells superlattices, quantum-wires or quantum dots by adopting a supercell approach where the respective objects are placed. In our case, Eq. (3) is solved using the “linear combination of Bloch bands” (LCBB) method, where the wave functions \( \psi_{\mathbf{k}}(\mathbf{r}) \) are expanded as,

\[ \psi_{\mathbf{k}}(\mathbf{r}) = \sum_{n,k} \sum_{\lambda} C_{n,k}^{(\lambda,k)} \Phi_{n,k,\lambda}(\mathbf{r}). \]  

(16)

In the above equation, \( \{\Phi_{n,k,\lambda}(\mathbf{r})\} \) are the bulk Bloch orbitals of band index \( n \) and wave vector \( \mathbf{k} \) of material \( \lambda \) (=InAs, GaAs), strained uniformly to strain \( \bar{\varepsilon} \). The inclusion of strain-dependent basis functions improves their variational flexibility. We use \( \bar{\varepsilon}=0 \) for the (unstrained) GaAs matrix material, and an average \( \bar{\varepsilon} \) value from valence force field (VFF) method for the strained dot material (InAs). For the InAs/GaAs system, we use \( n=8 \) (including spin) for electron states on a \( 6 \times 6 \times 16 \) k-mesh. Note that the potential \( V_{\text{ext}}(\mathbf{r}) + V_{\text{so}}(\mathbf{r}) \) contain full strain effects through the use of relaxed atomic positions, in addition to the explicit strain19 and alloy composition29 dependence.

In the atomistic approach to the single-particle problem, one includes (i) multi-band coupling [different \( n \) in Eq. (4)]; (ii) inter-valley coupling [different bulk k-points in Eq. (4)]; (iii) spin-orbit coupling \( V_{\text{so}} \) in Eq.(14)]; (iv) the proper strain profile [by relaxing \( \{\mathbf{R}_{\alpha}\} \) in Eq. (14) to minimize strain]; (v) realistic chemical profile [distributing the species \( \{\alpha\} \) as in a random alloy, or interdiffused interfaces]. The ensuing single-particle orbitals \( \{\psi_{\mathbf{k}}\} \) transform like the representations of the point-group created by the ionic positions. These underlying atomistic structures could break the symmetry represented by the macroscopic shape of the quantum dots. For example, a lens-shaped dot has a macroscopic cylindrical symmetry with [110] and [110] being equivalent, but if the dot is made of a zincblende material, the real symmetry is \( C_{2v} \), where, [110] and [\( \bar{1} \)110] are not equivalent. Yet the continuum models do not “see” the atomistic symmetry (the “farsightedness effect”). Therefore, as discussed in Ref. 31, in reality the atomistic wave function need not to be simple “pure” s-like or p-like. As a result, the Coulomb energy \( J_{ij} \) and exchange energy \( K_{ij} \) obtained with atomistic single-particle orbital \( \{\psi_{\mathbf{k}}\} \) do not have simple relationships as predicted by the parabolic 2D-EMA model. The deviations of the atomistic calculated \( J_s \) from the simplified 2D-EMA ones are going to lead to new physical behavior (e.g., new ground state symmetry of the many-particle state), as illustrated below.

D. Continuum treatment of the single-particle problem: EMA

It is sometimes customary to avoid an atomistic description of the single-particle problem in favor of a single-band particle-in-a-box model. In this approach, one sets \( V_{\text{scr}}=0 \) and replaces \( V_{\text{ext}} \) of Eq. (3) by a pure external potential, describing the macroscopic shape of the object, e.g., a box represents a quantum well, or a sphere with finite or infinite barriers represents a quantum dot. In some cases, this \( V_{\text{ext}} \) is calculated “realistically” from a combination of band offset, the gate potential and the ionized impurities, but it is treated nevertheless as a macroscopic external field. Under this approximation, simple results can be obtained for cylindrical QDs, where the angular momentum \( L \) and \( \ell \), are good quantum numbers. The electron (hole) single-particle levels have well defined shell structures, with nondegenerate s shell, twofold degenerate p shell and d shell, etc. However, if the confining potential is parabolic, there is another state that is accidentally degenerate with the two \( d \) states. We denote these three states \( d_{1x}, d_{2x}, d_{3y} \) for the parabolic 2D-EMA model. Furthermore, for parabolic confining potentials, the single-particle levels have equal spacing between two adjacent shells, e.g., \( \epsilon_{p}-\epsilon_{s}=\epsilon_{d}-\epsilon_{p} \). Another reason for the attractiveness of the single-band particle-in-a-box approach to the single-particle problem [Eq. (3)], is that the many-particle problem [Eq. (5)] becomes simple. For example, the quantum Monte Carlo (QMC) approach is currently applied only within the single-band particle-in-a-box approach for such large objects as self-assembled quantum dots. In a parabolic 2D-EMA model all the Coulomb integrals needed for charging and addition energy can be all related to \( J_{ss} \):

\[ J_{ab}=\epsilon_{ab} J_{ss}, \]

\[ K_{ab}=\epsilon_{ab}^2 J_{ss}. \]  

(17)

For example, \( J_{sp}=3/4 J_{ss}, K_{sp}=1/4 J_{ss} \). Therefore, the charging/addition energies and ground state configurations
are totally determined by $J_{ss}$ and the single-particle energy spacing $\omega$.

However, real self-assembled quantum dots grown via the Stranski-Krastanov techniques, are not well-described by the single-band particle-in-a-box approaches, despite the great popularity of such approaches in the experimental literatures. The model contains significant quantitative errors and also qualitative errors, whereby cylindrically symmetric dots are deemed to have, by symmetry, no fine-structure splitting, no polarization anisotropy, and no splitting of (twofold degenerate) $p$ levels and $d$ levels, all being a manifestation of the “farsightedness effect.”

III. RESULTS

Using the pseudopotential approach for single-particle and configuration interaction approach for the many-particle step, we studied the electron or hole addition energy spectrum up to 6 carriers in lens-shaped InAs dots embedded in a GaAs matrix. We study dots of three different base size, $b$ = 20, 25, and 27.5 nm, and for each base size, two heights, $h = 2.5$ and 3.5 nm. To study the alloy effects, we also calculated the addition spectrum for alloy dots In$_{1-x}$Ga$_x$As/GaAs of $h/b = 3.5/25$ nm dots, with Ga composition $x = 0$, 0.15, 0.3, and 0.5. In this section, we give detailed results of the single particle energy levels and Coulomb integrals, and the addition energy. We also compare the results with what can be expected from the parabolic 2D-EMA model.

A. Single-particle level spacing: Atomistic versus 2D-EMA description

1. Electron levels

We depict in Fig. 2 the calculated energy-level diagram of a pure lens-shaped InAs/GaAs quantum dot, with height $h = 2.5$ nm and base $b = 20$ nm. Figure 2 shows that the electron confinement energy is 230 meV, somewhat larger than the hole confinement energy (190 meV). The $p$ levels are split as are the $d$ levels, even though the dot has macroscopic cylindrical symmetry (see below).

The pseudopotential calculated electron single-particle energy spacings are summarized in Table I for QDs of different heights, bases, and alloy compositions. Table I gives the fundamental exciton energy $E_x$ calculated from CI approach for each dot. These exciton energies are between 980 and 1080 meV for pure InAs/GaAs dots, and can be as large as 1297 meV for In$_{1-x}$Ga$_x$As/GaAs alloy dots. This range agrees very well with experimental results for these classes of dots, ranging from 990 to 1300 meV.

(a) $s$-$p$ and $p$-$d$ energy spacing: From Table I, we see that for electrons in the lens-shaped dot, the $s$-$p$ energy level spacing $\delta_{sp} = \epsilon_s - \epsilon_p$ and $p$-$d$ energy level spacing $\delta_{pd} = \epsilon_d - \epsilon_p$ are nearly equal, as assumed by the 2D harmonic model. The energy spacing $\delta_{sp}$ and $\delta_{pd}$ range from 50 to 80 meV (Fig. 2), depending on the dot geometries. The electron energy spacings decrease with increasing QD base sizes. The electron energy spacing of alloy dots are much smaller than those in pure InAs/GaAs QDs, because of reduced confine-

ment. For Ga rich dots ($x = 0.3$–0.5), the single-particle energy level spacings range from 30 to 45 meV. These values agree with the infrared absorption measurements of intra-band transitions of alloy InGaAs QDs, which give $\delta_{sp} \approx 41$–45 meV. When the Ga composition reaches $x = 0.5$, the $s$-$p$ energy level spacing $\delta_{sp}$ becomes significantly different from the $p$-$d$ energy level spacing $\delta_{pd}$, thus deviating from harmonic potential approximation.

(b) Shell definition: Figure 2 shows that the energy levels of electrons in a lens-shaped dot have well defined $s$, $pd$ shell structure. However, while effective mass and $k$-$p$ models predict degenerate $p$ and $d$ levels, for cylindrically symmetric (e.g., lens-shaped) QDs, atomistic calculations show that even in perfect lens-shaped dots, the $p$-$p$ and $d$-$d$ levels are split by 2–4 meV (Fig. 2) due to the actual $C_{2v}$ symmetry. We denote the two $p$ levels as $p_1$ and $p_2$, and similarly, the three $d$ levels as $d_1$, $d_2$, and $d_3$, in increasing order of energy. The results listed in Table I show that $\delta_{pp} = \epsilon_{p_2} - \epsilon_{p_1}$ and $\delta_{dd} = \epsilon_{d_3} - \epsilon_{d_1}$ are very sensitive to the aspect ratio of the dots while not being very sensitive to the alloy compositions. Figure 3 depicts $\delta_{pp} = \epsilon_{p_2} - \epsilon_{p_1}$ versus dot heights [Fig. 3(a)] and bases sizes [Fig. 3(b)]. In general, we see that $\delta_{pp}$ increases with increasing dot height, and it decreases with increasing dot base size.

2. Hole levels

In contrast to electrons, hole single-particle levels (Table I) display a much more complicated behavior that is totally beyond the EMA description.

(a) $s$-$p$ spacing: As one can see from Table I, the hole
TABLE I. Summary of the pseudopotential-calculated single-particle level spacing (in meV) of In\textsubscript{1−x}Ga\textsubscript{x}As/GaAs quantum dots of different heights, base sizes, and Ga compositions. \(e_i\) and \(h_i\) are the \(i\)th electron and hole single-particle energy levels. \(E_X\) is the lowest exciton energy.

<table>
<thead>
<tr>
<th>Height (nm)</th>
<th>2.5</th>
<th>3.5</th>
<th>2.5</th>
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<td>25</td>
<td>25</td>
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<tr>
<td>(e_{p1}−e_i)</td>
<td>77.0</td>
<td>72.0</td>
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<td>57.0</td>
<td>51.5</td>
<td>44.2</td>
<td>33.2</td>
<td>54.9</td>
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<tr>
<td>(e_{d1}−e_{p2})</td>
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<td>73.6</td>
<td>64.0</td>
<td>58.7</td>
<td>50.7</td>
<td>43.2</td>
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<td>996</td>
<td>1095</td>
<td>1188</td>
<td>1297</td>
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</tbody>
</table>

\(s-p\) energy spacing ranges from 10 to 18 meV for dots of sizes we studied. These energy spacings are considerably smaller than those of electrons. The first confined hole state is found to be about 190 meV above the VBM of bulk GaAs, for the pure, \(h/b=2.5/20\) nm dot (Fig. 2). Unlike the case for electrons, the energy spacing between hole \(s\) and \(p\) levels depends strongly on the height of the dots,\(^{40}\) while being relatively insensitive to the base size of the dots.

(b) **Shell definition:** The well-defined \(s\), \(p\), \(d\) shell-structure for electrons does not exist for holes (Fig. 2), as the \(p_{1−2}, p_{1−3}, d_{1−2}, d_{1−3}\) splitting are much larger than those for electrons.\(^{41}\) The hole \(p\)-level splittings are also shown in Fig. 3 for different dot heights [Fig. 3(a)] and bases sizes [Fig. 3(b)]. For the smallest dots, \(h/b=2.5/25\) nm, the \(p\) splitting is about 11 meV (Table I), more than three times the value for electrons. This splitting is about half of the hole \(s-p\) energy spacing. Note that the pseudopotential calculated \(p-p\) splitting is much larger than 1.3 meV given by the \(k\)-\(p\) method (which includes piezoelectric effect).\(^{42}\) For taller dots, the \(p-p\) splittings are even larger. As a consequence, the \(p_2\) levels are energetically very close to the \(d_1\) levels, leading, as we will see below, to a nontrivial charging pattern that breaks Hund’s rule and the AFubau principle.\(^{18}\)

(c) **Wave function characters:** An analysis of the wave function show that these levels have somewhat mixed \(S\), \(P\), or \(D\) characters. For example, for a lens-shaped InAs/GaAs QD of height=3.5 nm and base=25 nm, the “\(s\) level” has 92% \(S\) character, and the two “\(p\) levels,” \(p_1\) and \(p_2\), have 86% \(P\) character, respectively (see Table II of Ref. 31). Therefore, we can still label the single-particle levels as \(s\), \(p_1\), \(p_2\), etc. These single-particle levels, \(s\), \(p_1\), \(p_2\), do not have pure HH character either, being instead 91%, 86%, and 92% HH-like, respectively. As the aspect ratio height/base increases, the mixture of angular momentum and HH and LH characters becomes stronger. For example, as shown in Ref. 43, for a InGaAs/GaAs alloy dot, with 25.2 nm in base and 2 nm in height, \(s\) level has 90% \(S\) character, while both \(p_1\) and \(p_2\) levels have 84% \(P\) character. When the height of the dot increases to 7.5 nm (with fixed base size), the leading angular momentum characters for these three levels are 84% \(S\), 78% \(P\), and 75% \(P\), respectively. Similarly, for the flat dot(2 nm in height), the mixture of LH state character is about 4%–9%, but increases to 11%–17% for the tall dot (7.5 nm in height).

**B. Coulomb integrals:** Atomistic versus 2D-EMA description

Another piece of information that decides the addition energies [Eq. (2)] is the Coulomb integral between the par-
We list the pseudopotential-calculated Coulomb energies of $s$ orbitals $J_{ss}$ for electrons in the first row of Table II and for holes in the first row of Table III. For electrons, $J_{ss}^{ee} \approx 22–25$ meV, and for holes $J_{ss}^{hh} \approx 20–27$ meV for typical dots. These numbers can be directly compared with experimental value from electron/hole charging experiments, since $J_{ss} = \Delta(1,2)$ [Eq. (12)]. The typical experimental values of $J_{ss}$ for electrons is about $4, 16, 44$ 19–27 meV and

<table>
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<tr>
<th>Height (nm)</th>
<th>2D-EMA</th>
<th>1.5</th>
<th>2.5</th>
<th>3.5</th>
<th>5.0</th>
<th>7.0</th>
</tr>
</thead>
<tbody>
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<td>22.6</td>
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<td>0.83</td>
<td>0.84</td>
<td>0.85</td>
<td>0.86</td>
</tr>
<tr>
<td>$J_{pd}/J_{ss}$</td>
<td>0.59</td>
<td>0.67</td>
<td>0.73</td>
<td>0.75</td>
<td>0.76</td>
<td>0.77</td>
</tr>
<tr>
<td>$J_{dd}/J_{ss}$</td>
<td>0.69</td>
<td>0.63</td>
<td>0.72</td>
<td>0.75</td>
<td>0.75</td>
<td>0.77</td>
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<tr>
<td>$J_{pp}/J_{ss}$</td>
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<td>0.84</td>
<td>0.84</td>
<td>0.86</td>
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<tr>
<td>$J_{pd}/J_{ss}$</td>
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<td>0.61</td>
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<tr>
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<tr>
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<td>0.23</td>
<td>0.22</td>
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</tr>
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<td>0.80</td>
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<td>0.70</td>
<td>0.70</td>
<td>0.87</td>
<td>0.73</td>
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<td>$J_{pd}/J_{ss}$</td>
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<td>0.80</td>
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<td>0.12</td>
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<td>0.12</td>
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for holes\textsuperscript{11,16,45} 20–25 meV. We plot in Fig. 4 $J_{ss}$ for electrons and holes versus height for base $b=25$ nm dots. For flat dots, the electron-electron Coulomb energy $J_{ee}$ is smaller than that of holes $J_{hh}^{(hh)}$. However, $J_{ee}$ is larger than $J_{hh}^{(hh)}$ for taller dots. The crossover is at about 2.5 nm for the $b=25$ nm dots. Note that in our calculation, the two nearly degenerate electron $p$ orbitals $p_1$ and $p_2$, are spatially almost orthogonal to each other. However, in the simple parabolic 2D-EMA model,\textsuperscript{16} the two degenerate $p$ orbitals $p_+ = p_x + i p_y$ and $p_- = p_x - i p_y$ have same spatial function differing only by a phase factor. As a result, the exchange interaction between $p_1$ and $p_2$ is much smaller than that of $p_+$ and $p_-$. Furthermore, the simple 2D-EMA model predicts $J_{p,p_+} = J_{p,p_-}$, which is not true in the atomistic description, where $J_{p, p_2}$ is much smaller than $J_{p, p_1}$ and $J_{p, p_2}$.

In a parabolic 2D-EMA model, there is only one free parameter in calculating the Coulomb integrals for each type of carriers: The effective length $l_p$ for electrons and $l_p$ for holes. This leads to simple geometric relationships as illustrated in Eq. (17). These relations are listed in the second column of Tables II and III. However, as noted in Sec. II C, a detailed analysis of atomistic wave functions\textsuperscript{31,43} show that they are not of pure conduction band character for electrons or HH, LH characters for holes; nor do they have pure $s$, $p$ angular momentum characters as predicted by 2D-EMA model. Since inter-valley, inter-band coupling and the underlying atomistic symmetries are also ignored in the 2D-EMA model, we might expect that the simple relations between the Coulomb integrals of Eq. (17) must be somehow different in the atomistic approach relative to those predicted by the EMA, even for rather flat lens-shaped dots having parabolic-like energy level pattern, $\epsilon_p - \epsilon_0 \approx \epsilon_p - \epsilon_0$, as chosen here.

The ratios $J_{a/b}/J_{ss}$ and $K_{a/b}/J_{ss}$, derived from atomistic calculations are compared with the parabolic 2D-EMA model relationships in Table II for electrons and in Table III for holes for dots of different aspect ratios. To visualize the differences, we also plot the relative error for Coulomb energies $(J_{a/b}/J_{ss})_\text{atm} - (J_{a/b}/J_{ss})_\text{EMA}$ and for exchange energies $(K_{a/b}/J_{ss})_\text{atm} - (K_{a/b}/J_{ss})_\text{EMA}$ in Fig. 5(a) for electrons and in Fig. 5(b) for holes, where “atm” means atomistic results and “EMA” means the 2D-EMA model results. For flat lens-shaped dots which have a parabolic-like level spectrum, we see qualitative agreement between the atomistic calculations and the 2D-EMA model. For such flat dots, the errors are generally within 20% of $J_{ss}$. For taller dots, the agreement is worse, as can be seen from Fig. 5. For even taller QDs (e.g., height/base > 5/25 nm), the large biaxial strain will develop a “hole trap” at the InAs/GaAs interface, which lead to the hole localization on the interface of the dots\textsuperscript{31,43} In these cases, where the hole (envelope) wave functions are totally different from those predicted by 2D-EMA model, the relationship between Coulomb and exchange energies are, accordingly, very different from those of 2D-EMA model.

At first sight, it would seem that if the values of $J_{ss}$ in the parabolic 2D-EMA model are taken to be those calculated atomistically, the Coulomb integrals predicted by the parabolic 2D-EMA model are only a few meV off from those calculated by atomistic theories. However, these small differences will change significantly the electronic phase-diagrams of carriers, as we will see below.
TABLE IV. Orbital occupations for N holes in InAs/GaAs QDs of different height/base obtained from single-particle atomistic pseudopotential and many-particle CI calculations. The orbital occupations are given by the leading configurations in CI results.

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<th>Height/Base (nm)</th>
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<th>N=3</th>
<th>N=4</th>
<th>N=5</th>
<th>N=6</th>
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<td>s²</td>
<td>s¹pₚ¹</td>
<td>s¹p₁</td>
<td>s¹p₁d₁</td>
<td>s¹p₁d²</td>
</tr>
<tr>
<td>3.5/20</td>
<td>s¹</td>
<td>s²</td>
<td>s¹pₚ¹</td>
<td>s¹p₁</td>
<td>s¹p₁d₁</td>
<td>s¹p₁d²</td>
</tr>
<tr>
<td>2.5/25</td>
<td>s¹</td>
<td>s²</td>
<td>s¹pₚ¹</td>
<td>s¹p₁</td>
<td>s¹p₁d₁</td>
<td>s¹p₁d²</td>
</tr>
<tr>
<td>3.5/25</td>
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<td>s¹p₁d²</td>
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<tr>
<td>2.5/27.5</td>
<td>s¹</td>
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<td>s¹pₚ¹</td>
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<td>s¹p₁d₁</td>
<td>s¹p₁d²</td>
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<tr>
<td>3.5/27.5</td>
<td>s¹</td>
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<td>s¹pₚ¹</td>
<td>s¹p₁</td>
<td>s¹p₁d₁</td>
<td>s¹p₁d²</td>
</tr>
</tbody>
</table>

C. Ground state configurations: Atomistic versus 2D-EMA description

1. Generic phase diagrams for 2D-EMA model

To calculate the addition energies, we first need to know the spin and orbital configurations of the few-particle ground state, which minimizes the energy E(N) of the N-particle system. For an illustration, we first use the “single configuration approximation,” i.e., retain only the Slater determinants that have the same orbital degree of freedom, and keep the exchange (spin-spin) coupling between these Slater determinants. The ground states are those configurations that have the lowest total energy. For electrons, the calculated ground state for N=1–6 are shown in Fig. 1, which shows that the electron charging pattern follows Hund’s rule. The ground state spin and orbital configurations for holes are listed in Table IV for N=1–6 holes. In contrast to electrons, the hole charging patterns show complicated behaviors that defy the Hund’s rule and the Aufbau’s principle for most of the cases of N=5, 6 holes.18

To understand the differences between the ground-state configurations of electrons and holes and the driving forces for these differences, we developed a general phase-diagram approach18 that classifies the many-particle configurations for electrons and holes in quantum dots in terms of simple electronic and geometric parameters. To do so, we calculate for each particle number N, the configuration which minimizes the total-energy under the single-configuration approximation at different p₁-p₂ splitting (δp₁,p₂ = εp₂ - εp₁) and p₂-d₁ energy spacing (δp₂,d₁ = εd₁ - εp₂) using Coulomb integrals J and K in units of Jˢ. This approach gives a phase diagram as a function of the parameters (N; δp₁,p₂, δp₂,d₁) in units of Jˢ, which yields for N=4 two electronic phases:

\[ \Sigma = (s¹s¹)(p¹)(p¹), \]
\[ \Delta = (s¹s¹)(p¹)(p¹), \]

Here, we have adopted a spectroscopic notation for a system with cylindrical symmetry. For N=5, there are three possible phases:

\[ 4Δ = (s¹s¹)(p¹)(p¹)(d¹), \]
\[ 2Δ = (s¹s¹)(p¹)(p¹)(d¹), \]

For N=6, we find four phases,

\[ 5Σ = (s¹s¹)(p¹)(p¹)(d¹)(d¹), \]
\[ 3Π = (s¹s¹)(p¹)(p¹)(d¹), \]
\[ 1Σ = (s¹s¹)(p¹)(p¹)(p¹), \]
\[ 1Σ* = (s¹s¹)(p¹)(p¹)(d¹). \]

We first apply this approach to a parabolic 2D-EMA model. We relax the restrictions in the parabolic 2D-EMA model of degenerate shells (εp₁ = εp₂, εd₁ = εd₂ = εd₃) and of equidistant shells (εp - εd = εd - εp) and allow δp₁,p₂ and δp₂,d₁ to vary. The resulting phase diagrams are shown in Fig. 6 for N=4–6.

2. Ground states for specific dots in the 2D-EMA model

To decide which of these phases is a ground state for a given dot, we need to know in Fig. 6(a) the actual value of δp₁,p₂/Jˢ and δp₂,d₁/Jˢ for this realistic dot. For electrons in self-assembled dots, the single-particle energy spacing is usually more than twice the Coulomb energy,4,16 i.e., δp₁,p₂ > 2Jˢ. For holes, δp₂,d₁ = 1.17Jˢ was determined from recent experiments11,13 and δp₁,p₂ = 0 is assumed in 2D-EMA model. This places in Fig. 6(a), for both electrons and holes, phases 3Σ, 2Π, and 1Σ as ground states for N=4–6, respectively, where the lens represents a lens-shaped self-assembled QD, and labels “c” and “h” inside the lens are symbols for electron and hole, respectively. As a comparison, we also show the electron ground-state phases for an electrostatic dot (∼50 nm) represented by a circle.

3. Generic phase diagrams for atomic approach

We now use atomistic Coulomb and exchange energies J and K, listed in Tables II and III to recalculate the phase diagram (N; δp₁,p₂, δp₂,d₁) for electrons and holes in Figs. 6(b) and 6(c), respectively. We see that the phase boundaries change dramatically for both electrons and holes.18 For example, for N=6 electrons, phase 3Σ* disappears [Fig. 6(b)], while phase 5Σ disappear for N=5 holes.
4. Ground states for specific dots in the atomistic approach

We next use $\delta_p$, $\delta_d$, and $J_{ss}$ from Tables I and II to decide the ground states for electrons in self-assembled InAs/GaAs QDs. For example, for a InAs/GaAs dot with 2.5 nm in height and 20 nm in base, we have $\delta_p/J_{ss} \sim 0.1$ and $\delta_d/J_{ss} \sim 3$. These parameters give the ground-state phase $^1 \Pi$ for 5 electrons, and the ground-state phase $^3 \Sigma$ for 6 electrons. As we see, even though the phase boundaries calculated from atomistic theories are very different from those calculated from 2D-EMA model, the ground-state configurations are remain the same as those predicted by the parabolic 2D-EMA model. This is because in the phase diagrams, the coordinates of the electrons in self-assembled dots are far away from other competing phases. For $N=4$ electrons, the high spin state phase $^3 \Sigma$ (Hund’s rule) and low spin state $^1 \Delta$ are relatively close in the phase diagram. The intrinsic $p$-level splittings in the lens-shaped dots are about $1-4$ meV (Table I), which are smaller than the electron-electron exchange energies (5 mV), and therefore the ground state is the high-spin phase $^3 \Sigma$. However, if the dot shape is elongated, adding additional $p$-level splittings, the ground state could be low-spin phase $^1 \Delta$. Actually there are some experimental evidences showing that the ground state of 4 electrons could be a low spin state.6,46

In contrast to electrons, holes have $\Delta \epsilon_h \geq J_{ss}$, and large $p_1-p_2$ splitting, small $p_2-d_1$ splitting (Fig. 2), which place the holes in a different region in the phase diagrams than electrons, where there are more competing phases. For example, for the dot with base $b=20$ nm, and height $h=2.5$ nm, we have $\delta_p/J_{ss} \sim 0.4$ and $\delta_d/J_{ss} \sim 0.17$. These relationships give, for this dot, the ground states $^1 \Delta$, $^2 \Delta$, and $^1 \Sigma^*$ for $N=4$, 5, and 6 holes, respectively, showing a nontrivial hole charging pattern that breaks the Aufbau principle.18 We also list the ground state configurations for dots of different geometries in Table IV. As we see, in most of the cases, the ground states are still $^3 \Sigma$, $^2 \Delta$, and $^1 \Sigma^*$ as shown in Fig. 6(c). However, there are some exceptions for very tall dots or very flat dots. For very flat dots (e.g., the $h/b=2.5/27.5$ nm dot), the $p_1-p_2$ energy splitting (5.8 meV) is much smaller than the $p_2-d_1$ energy splitting (9.4 meV). These parameters place the ground state of $N=5$ holes in phase $^1 \Pi$ and that of 6 holes in phase $^3 \Pi$ of Fig. 6(c). For very tall dots (e.g., the $h/b=3.5/20$ nm dot), the interfacial hole localization changes the hole-hole Coulomb integrals dramatically, and thus change the phase boundaries in Fig. 6(c). We find the ground states of the dot with $h/b=3.5/20$ nm are $^2 \Pi$ and $^1 \Sigma^*$ for 5 and 6 holes, respectively.

5. Effects of configuration interaction

If we use CI instead of single-configuration approximation, the ground states are superpositions of different configurations, but the leading CI configurations have a significant weight, being 79%, 71%, and 64% for 4, 5, and 6 holes, respectively, in the $h/b=2.5/20$ nm dot. Since the weights of leading configurations are significantly larger than other configurations, we are justified in using leading configurations to represent graphically the ground states. It is worth noting...
that for very large electrostatically confined dots, where single-particle energy spacing $\delta E_{ss} \ll 1$ (down-left corner of the phase diagrams of Fig. 6), the ground state mixes large number of configurations, which have no significant leading configurations and are, therefore, in strongly correlated states that are not discussed here.

D. Calculated charging and addition energies

Once we determined the ground state configurations, we can calculate the total energies using Eq. (4). We calculate the ground state total energies for up to 6 electrons/holes for each dot. For electrons, in the CI approach we used 6 single-particle electron levels $s, p_1, p_2, d_1, d_2, d_3$ to construct all possible Slater determinants, while for holes, we used 8 single-particle hole levels. The total number of Slater determinants for 6 electrons is 924. For 6 holes, the total number of determinants is 8008. We plot in Fig. 7 the CI total energies for the ground state of 6 carriers versus number of single-particle states included in the CI expansions. The total energies converge to about 1 meV if 6 single-particle states are used for electrons and 8 states are used for holes.

The charging energies and addition energies are calculated using Eqs. (1) and (2). The addition energies calculated from CI calculations give results that are about 1 meV different from single-configuration results for electrons, and 1–3 meV difference for holes. The addition energies are summarized in Table V. Experimentally, the charging energies of electrons and holes in self-assembled QDs are usually measured via capacitance spectroscopy, using gated structures. The (In,Ga)As/GaAs dots used in the electron charging experiments were roughly estimated to be 7 nm in height and 20 nm in base in Refs. 5 and 6. The measured $\Delta_e(1,2)$ ~ 23 meV in Ref. 5 and ~21.5 meV in Ref. 6, which, as shown in Sec. II equals roughly $F_{ee}$, the electron-electron Coulomb interaction of $s$ orbitals. The charging energies between $s$ and $p$ levels, $\Delta_e(2,3)$ was estimated to be 57.0 meV in Ref. 6. The average addition energies between $p$-states are estimated to be 18 meV in Ref. 5, and 14 meV in Ref. 6. Furthermore, in Ref. 6, $\Delta_e(4,5)$ is almost twice as large as $\Delta_e(3,4)$ and $\Delta_e(5,6)$, which might be related to the

![CI total energy (N=6) vs basis set](image)

**FIG. 7.** CI total-energy convergency test for (a) six-electron and (b) six-hole states with increasing number of single-particle orbitals in the bases. The test is done for a pure InAs/GaAs dot, with base $b$ = 25 nm, and height $h$ = 3.5 nm.

**TABLE V.** Summary of atomistically calculated electron addition energies $\Delta_e$ and hole addition energies $\Delta_h$ (in meV) of In$_{1-x}$Ga$_x$As/GaAs quantum dots for various dot heights, base sizes and Ga compositions. The experimental addition energies for electrons are extracted from Ref. 6, and the addition energies for holes are taken from Ref. 11.

| Height (nm) | 2.5 | 3.5 | 2.5 | 3.5 | 3.5 | 3.5 | 3.5 | 2.5 | 35 |
| Base (nm) | 20 | 20 | 25 | 25 | 25 | 25 | 25 | 27.5 | 27.5 |
| Ga comp. | 0 | 0 | 0 | 0 | 0.15 | 0.3 | 0.5 | 0 | 0 |
| $\Delta_e(1,2)$ | 26.3 | 24.6 | 23.3 | 21.8 | 20.5 | 18.5 | 15.2 | 22.1 | 20.5 | 21.5 |
| $\Delta_e(2,3)$ | 89.5 | 84.4 | 71.8 | 67.6 | 61.2 | 53.2 | 40.3 | 65.2 | 60.8 | $\sim$57 |
| $\Delta_e(3,4)$ | 19.7 | 20.3 | 16.7 | 16.5 | 15.9 | 12.7 | 10.6 | 15.4 | 15.5 | 11.4 |
| $\Delta_e(4,5)$ | 21.5 | 19.5 | 20.1 | 18.5 | 16.6 | 16.5 | 12.9 | 19.6 | 17.5 | 21.0 |
| $\Delta_e(5,6)$ | 19.7 | 20.3 | 16.7 | 16.5 | 15.8 | 12.4 | 10.3 | 15.4 | 15.4 | 12.2 |
| $\Delta_h(1,2)$ | 24.1 | 19.0 | 21.9 | 17.5 | 18.3 | 18.4 | 17.8 | 21.0 | 16.7 | 23.9 |
| $\Delta_h(2,3)$ | 28.7 | 21.7 | 27.2 | 21.2 | 23.0 | 23.6 | 22.6 | 26.4 | 20.6 | 34.2 |
| $\Delta_h(3,4)$ | 18.1 | 16.9 | 16.4 | 15.2 | 15.4 | 15.4 | 15.0 | 15.6 | 14.5 | 17.1 |
| $\Delta_h(4,5)$ | 26.4 | 21.6 | 25.4 | 20.8 | 23.2 | 21.7 | 19.3 | 23.8 | 20.5 | 23.2 |
| $\Delta_h(5,6)$ | 17.1 | 16.1 | 15.3 | 14.4 | 14.5 | 16.0 | 15.7 | 15.5 | 13.7 | 15.0 |
breakdown of Hund’s rule as a consequence of irregular shape of the dots. These results are listed in Table V and compared with our theoretical results. We see that the electron addition energies of $\text{H}_{20849}$In$_{1-x}$Ga$_x$/GaAs dots with height $h=3.5$ nm and base $b=25$ nm at $x=0$, 0.15, 0.3, and 0.5.

The experimental hole addition energies are taken from Ref. 11, which gives $\Delta_0(1,2) = 23.9$ meV, comparable to that of $\Delta_0(1,2)$. However, the addition energy between $s$ and $p$ orbitals, $\Delta_0(2,3) = 34.2$ meV, is significantly smaller than $\Delta_0(2,3) = 57$ meV. This result reflects that the $s$-$p$ energy spacing of holes is much smaller than that of electrons. As seen from Table V, our calculated addition energies of pure $\text{InAs/GaAs}$ dots with base $b=25$ nm and height $h=2.5$ and 3.5 nm; (b) pure $\text{InAs/GaAs}$ dots with height $h=3.5$ nm and base $b=20$, 25, and 27.5 nm; and (c) $\text{In}_{1-x}\text{Ga}_x$/GaAs dots with height $h=3.5$ nm and base $b=25$ nm at $x=0$, 0.15, 0.3, and 0.5.

FIG. 8. Electron addition energies of (a) pure InAs/GaAs dots with base $b=25$ nm and height $h=2.5$ nm and 3.5 nm; (b) pure InAs/GaAs dots with height $h=3.5$ nm and base $b=20$, 25, and 27.5 nm; and (c) $\text{In}_{1-x}\text{Ga}_x$/GaAs dots with height $h=3.5$ nm and base $b=25$ nm at $x=0$, 0.15, 0.3, and 0.5.

and $\text{H}_{20849}$$\text{H}_{20850}$In$_{1-x}$Ga$_x$/GaAs dots agree very well with this experiment.

To study trends of addition energies for the electrons and the holes, we depict the electron addition energies for different dot heights [Fig. 8(a)], bases [Fig. 8(b)] and alloy compositions [Fig. 8(c)]. Similarly, the hole addition energies are plotted in Figs. 9(a)–9(c) for dot heights, bases and alloy compositions. We see the following:

(i) Up to six carriers, the electron addition energies $\Delta(N-1,N)$ have a single peak at $(N-1)=2$. The peak is due to the single-particle energy gap between the $s$-shell and $p$-shell. On the other hand, all hole addition energies have two peaks at $(N-1)=2$ and $(N-1)=4$, respectively, where the first peak come from the single-particle energy gap between the $s$ orbital and $p_1$ orbital and the second peak is associated with the energy difference between the $p_1$ and higher energy orbitals.

FIG. 9. Hole addition energies of (a) pure InAs/GaAs dots with base $b=25$ nm and height $h=2.5$ and 3.5 nm; (b) pure InAs/GaAs dots with height $h=3.5$ nm and base $b=20$, 25, and 27.5 nm; and (c) $\text{In}_{1-x}\text{Ga}_x$/GaAs dots with height $h=3.5$ nm and base $b=25$ nm at $x=0$, 0.15, 0.3, and 0.5.
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(ii) Electron addition energies decrease with increasing height and base of the dots, and the hole addition energies share the same trend. However, the electron addition energies are more sensitive to the base of the dots and relatively insensitive to the height. In contrast, the hole addition energies are very sensitive to the heights of the dot, and relatively insensitive to the base. This dependence suggests that the electron wave functions are more two-dimensional like in the QDs, while hole wave functions are more extended in [001] direction.

(iii) Electron addition energies show a simple trend for alloyed In$_{x}$Ga$_{1-x}$As/GaAs QDs, which are consistently smaller for Ga rich dots. However, the hole addition energies of alloy dots show more complicated behaviors. The reasons of this complication are the following: (1) Alloy dots have different trends in hole single-particle energy level spacings (see Table I); (2) the ground state configurations might be different for different alloy compositions.

IV. SUMMARY

We systematically studied the electron/hole addition energy spectra using single-particle pseudopotential plus many-particle CI methods. Considering the single-particle step, we find for electrons that there is a shell structure and that the $p$-$p$ and $d$-$d$ splittings are about 1–4 meV depending on the dot geometry. For holes, the single-particle step reveals large ($5–11$ meV) $p_{1}$-$p_{2}$ splitting and absence of a well defined shell structure. Considering the $e$-$e$ and $h$-$h$ Coulomb integrals, we find that atomistically calculated ratios between various Coulomb integrals $J_{ij}$ differ by about 20% from those in the 2D-EMA model calculation. These differences lead to many-particle phase diagrams that differ significantly from those predicted by the parabolic 2D-EMA model. In particular, the “unusual” hole single-particle spectrum and Coulomb integrals lead to many-particle ground states that defy the Hund’s rule and the Aufbau principle for holes. The predicted ground-state configurations and the addition energies calculated in this pseudopotential plus CI scheme compare well with experiments.

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40 For alloyed dots, the same dependence with height of the \( s-p \) spacing is found. See Ref. 43.

41 The piezoelectric effect also contributes to the \( p \)-state splitting of the electron states, as has been shown in Ref. 21. For the splitting of the hole states, the effect is more subtle. We find for a pure InAs lens shaped dot with 25 nm base and 3.5 nm height, using piezoelectric coefficients from bulk materials, that the hole \( p-p \) splitting without piezoelectric effect is 9.5 meV and with the piezoelectric effect is 9.7 meV, using piezoelectric coefficients from bulk materials. The change (0.2 meV) due to piezoelectric effects is rather small. We therefore ignored piezoelectric effects in our calculations.


48 The electric fields applied to the dots are typically of the order of 50 kV/cm for electron charging devices (as gleaned from Refs. 4–6) and 100 kV/cm for hole charging devices (Refs. 11 and 13). We performed additional calculations to estimate the effect of this field on the addition energies and obtained changes for addition energies to be about 1 meV at 100 kV/cm. We therefore, ignored this correction in our calculations.