Electronic structures of (In,Ga)As/GaAs quantum dot molecules made of dots with dissimilar sizes

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Using single-particle pseudopotential and many-particle configuration interaction methods, we compare various physical quantities of (In,Ga)As/GaAs quantum dot molecules (QDMs) made of dissimilar dots (hetero-QDMs) with QDMs made of identical dots (homopolar QDMs). The calculations show that the electronic structures of hetero-QDMs and homo-QDMs differ significantly at large inter-dot distance. In particular (i) Unlike those of homo-QDMs, the single-particle molecular orbitals of hetero-QDMs convert to dot localized orbitals at large inter-dot distance. (ii) Consequently, in a hetero-QMD the bonding-antibonding splitting of molecular orbitals at large inter-dot distance is significantly larger than the electron hopping energy whereas for homo-QDM, the bonding-antibonding splitting is very similar to the hopping energy. (iii) The asymmetry of the QDM increases significantly the double occupation for the two-electron ground states, and therefore lowers the degree of entanglement of the two electrons.

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I. INTRODUCTION

Vertically coupled quantum dots\textsuperscript{3,4} obtained via epitaxial growth provide a potential scheme for scalable nano-structures for quantum computing. In this scheme, two coupled quantum dots are used as a basic logic gate, via the entanglement of one exciton\textsuperscript{5} or two electronic spins.\textsuperscript{6} This proposal for gate operations, requires knowledge of the detailed physical properties of the “quantum gate” made of two quantum dots. Significant progress has been recently made\textsuperscript{7} using quantum dot molecules made of very large (\textasciitilde 500 - 1000 \textmu m) electrostatically confined dots. The limit of large quantum confinement, however, requires working with (200 \texttimes 30 \textmu m) self-assembled QDMs. So far, most experiments on self-assembled QDM are optical\textsuperscript{8} and most theories are based on continuum models, such as effective mass approximations.\textsuperscript{9} These simple models ignore or drastically simplify important real material properties such as strain, atomistic symmetries and crystal structural effects, band coupling etc. Recent studies\textsuperscript{10} show that simplification of such important effects may lead to qualitative changes in fundamental physics of the QDMs.

Previously, we have studied homopolar QDMs made of two identical quantum dots, using single-particle pseudopotential method and many-particle configuration interaction method.\textsuperscript{10,11} We have studied electron localization, double occupation rate and two-electron entanglement using a new formula for measuring the degree of entanglement formula for two indistinguishable fermions. We found that even geometrically identical dots in the QDMs lead to electronic asymmetry due to the strain effects. However, experimentally it is hard to control the shape, size and compositions of individual dots within the QDMs, so in practice, the QDMs are never made of identical dots. Actually, the top dots are tend to be larger than the bottom dots due to the strain effects.\textsuperscript{12} Indeed, the measured difference in exciton energy due to dot-size difference is about 4 meV\textsuperscript{10} for two vertically coupled dots that are 20 nm apart. Sometimes, the two dots are intentionally grown different so that they can be addressed separately.\textsuperscript{11} To provide quantitative comparison to experiments, considering the effects that asymmetry of quantum dots within the molecule, we studied the QDMs made of (In,Ga)As/GaAs quantum dots of different sizes (hetero-polar QDM).

In this paper, we study systematically the electronic properties of hetero-QDMs, including their single-particle molecular orbitals, many-particle states, double occupation and entanglement of two-electrons, and compare them to those of homo-QDMs. We found that while at short inter-dot distance, the electronic properties of hetero-QDM and homo-QDM are similar, they differ significantly at large inter-dot distance. This difference may have substantial impact in implementation of quantum gates.

II. METHODS

Figure \textsuperscript{1} shows the geometry of a hetero-QDM, consisting a pair of 3 nm tall InAs dots in the shape of truncated cones, grown on two-dimensional InAs wetting layers, embedded in a GaAs matrix. The inter-dot separation \(d\) is defined as the distance between the wetting layers of top and bottom dots. We choose the base diameter of top dots (labeled as \(\alpha\)) to be 20 nm, and that of the the bottom dots (labeled as \(\beta\)) to be 19 nm, mimicking to the fact that experimentally the top dots are slightly larger than the bottom dots.\textsuperscript{13,14} The composition of the dots vary from In\(_{0.5}\)Ga\(_{0.5}\)As at their bases to pure InAs at their top, as determined in Ref.\textsuperscript{3}.

We de-
note the dot molecules made of dissimilar dots $\alpha$ and $\beta$ as $M_{\alpha\beta}$. We also constructed the homo-QDM, consisting a pair of quantum dots $\gamma$, which have the average sizes, and the same alloy compositions of dots $\alpha$ and $\beta$ in the heteropolar dot molecule. We denote the homo-QDM as $M_{\gamma\gamma}$.

The single-particle energy levels and wavefunctions of $M_{\alpha\beta}$ and $M_{\gamma\gamma}$ are obtained by solving the Schrödinger equations in a pseudopotential scheme,

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

where the total electron-ion potential $V_{\text{ion}}(\mathbf{r})$ is a superposition of local, screened atomic pseudopotentials $v_\alpha(\mathbf{r})$, and a nonlocal spin-orbit potential $V_{\text{so}}$ i.e., $V_{\text{ion}}(\mathbf{r}) = \sum_{n,\alpha} v_\alpha(\mathbf{r} - \mathbf{R}_{n,\alpha}) + V_{\text{so}}$. The atomic position $\{\mathbf{R}_{n,\alpha}\}$ is obtained from minimizing the total bond-bending and bond-stretching energy using the Valence Force Field (VFF) model.\footnote{12, 13} The atomic pseudopotentials $v_\alpha$ ($\alpha=$In, Ga, As) are fitted to the physically important quantities of bulk InAs and GaAs, including band energies, band-offsets, effective masses, deformation potentials and alloy bowing parameters, etc.\footnote{14} Because for electrons the spin-orbit coupling is extremely small in the InAs/GaAs quantum dots, we ignored this effect. In general, including the spin-orbit coupling effect will introduce mixture of different total spin states. Equation (1) is solved in the basis of $\{\phi_{mn\lambda}(\mathbf{k})\}$ of Bloch orbitals of band index $m$ and wave vector $\mathbf{k}$ of material $\lambda (=\text{InAs, GaAs})$, strained uniformly to strain $\varepsilon$ following Ref.\footnote{15}

The Hamiltonian of interacting electrons can be written as,

$$
H = \sum_{i\sigma} c_i^* \hat{\psi}_{i\sigma} \hat{\psi}_{i\sigma} + \frac{1}{2} \sum_{ijkl \sigma \sigma'} \Gamma_{ijkl}^{\sigma \sigma'} \hat{\psi}_{i\sigma}^\dagger \hat{\psi}_{j\sigma'}^\dagger \hat{\psi}_{k\sigma'} \hat{\psi}_{l\sigma},
$$

where $\hat{\psi}_{i\sigma}(\mathbf{r}) = c_{i\sigma} \psi_{i\sigma}(\mathbf{r})$ is the field operator, whereas $c_{i\sigma}$ is a fermion operator. $\psi_i = \sigma_u, \sigma_g; \pi_u, \pi_g$ are the single-particle eigenfunctions of the $i$-th molecular orbital, and $\sigma, \sigma' = 1, 2$ are spin indices. The $\Gamma_{ijkl}^{\sigma \sigma'}$ are the Coulomb integrals between molecular orbitals $\psi_i, \psi_j, \psi_k$ and $\psi_l$.

$$
\Gamma_{ijkl}^{\sigma \sigma'} = \int d\mathbf{r} d\mathbf{r}' \frac{\psi_i^\dagger(\mathbf{r}) \psi_j^\dagger(\mathbf{r}') \psi_k(\mathbf{r}') \psi_l(\mathbf{r})}{\epsilon(\mathbf{r} - \mathbf{r}')|\mathbf{r} - \mathbf{r}'|}. \quad (3)
$$

The $J_{ij} = \Gamma_{ij}^{\sigma \sigma'}$ and $K_{ij} = \Gamma_{ij}^{\sigma' \sigma}$ are diagonal Coulomb and exchange integrals respectively. The remaining terms are called off-diagonal or scattering terms. All Coulomb integrals are calculated numerically from atomistic wavefunctions.\footnote{16} We use a phenomenological, position-dependent dielectric function $\epsilon(\mathbf{r} - \mathbf{r}')$ to screen the electron-electron interaction.\footnote{17} The many-particle problem of Eq. (2) is solved via the CI method, by expanding the $N$-electron wavefunction in a set of Slater determinants, $|\Psi_{e_1, e_2, \ldots, e_N}\rangle = c_{e_1}^\dagger c_{e_2}^\dagger \cdots c_{e_N}^\dagger |\Phi_0\rangle$, where $c_{e_i}^\dagger$ creates an electron in the state $e_i$. The $\nu$-th many-particle wavefunction is then the linear combinations of the determinants,

$$
|\Psi_{\nu}\rangle = \sum_{e_1, e_2, \ldots, e_N} A_{\nu}(e_1, e_2, \ldots, e_N) |\Phi_{e_1, e_2, \ldots, e_N}\rangle. \quad (4)
$$

For the two-electron problems, our calculations include all possible Slater determinants of six confined molecular orbitals.

\section*{III. Basic Electronic Structures at the Single-Particle Level}

\subsection*{A. Double-dot molecular orbitals}

We first show the electronic structure of isolated dots $\alpha$ and $\beta$. The single-dot electron $s$ and $p$ levels of dots $\alpha$
and $\beta$ are shown on the right panel of Fig. 2 We see that the $s$-$p$ energy spacing of dot $\alpha$ is $\epsilon(p_\alpha) - \epsilon(s_\alpha) = 52$ meV and that of dot $\beta$ is $\epsilon(p_\beta) - \epsilon(s_\beta) = 59$ meV, compared to 54 meV of dot $\gamma$ (not shown). The energy level of $s_\beta$ is slightly ($\sim 6$ meV) higher than $s_\alpha$, because dot $\beta$ is smaller than dot $\alpha$ and therefore has larger confinement. The $p$ levels of all dots have a small energy splitting due to the underlying atomistic asymmetry, e.g., $\delta(\epsilon(p_\alpha) = 1$ meV. We further calculated the fundamental exciton energy of dot $\alpha$, $E_X(\alpha) = 1153$ meV, and that of dot $\beta$, $E_X(\beta) = 1159$ meV. The energy difference in exciton of dots $\alpha$ and $\beta$ is about 6 meV, in agreement with experiment. The fundamental exciton energy of the “averaged” dot $\gamma$ is $E_X(\gamma) = 1156$ meV.

When two dots $\alpha$ and $\beta$ couple, the bonding and antibonding “molecular orbitals” ensue from the single-dot orbitals. The energy levels of molecular orbitals are shown on the left panel of Fig. 2. We show the single-particle levels of molecular orbitals $\sigma_g, \sigma_u$ originating from $s$ orbitals, and $\pi_u$, and $\pi_g$ originating from $p$ orbitals. The bonding and anti-bonding splitting $\Delta_\sigma = \epsilon(\sigma_u) - \epsilon(\sigma_g)$ and $\Delta_\pi = \epsilon(\pi_g) - \epsilon(\pi_u)$ increase with the decrease of inter-dot distance, because the coupling between the top and bottom dots gets stronger. This picture is similar to what we obtained for homo-QDMs. However, there is an important difference between the homo-QDMs $M_{\gamma\gamma}$ and hetero-QDMs $M_{\alpha\beta}$: in the former case, the bonding and anti-bonding splitting $\Delta_\sigma$ and $\Delta_\pi$ decay to almost zero at large inter-dot distance, while in the later case, $\Delta_\sigma$ and $\Delta_\pi$ tend to constants ($\Delta_\sigma \sim 7$ meV, $\Delta_\pi \sim 10$ meV here), because the molecular orbitals gradually convert at large inter-dot distance to single dot energy levels, e.g. the $\sigma_g$ levels convert to top dot $s$ orbitals, and $\sigma_u$ convert to bottom dot $s$ orbitals, therefore the energy splitting between the first and second molecular states at large distances is approximately the energy difference between $s$ orbitals of the top and bottom dots, i.e., $\Delta_\sigma \sim \epsilon(s_\beta) - \epsilon(s_\alpha) \neq 0$ for $M_{\alpha\beta}$.

Figure 2 shows that at inter-dot distance $d = 10$ nm, the molecular orbital levels are about 25 meV higher than the isolated dot levels, although the direct electronic coupling between two dots is much smaller than this quantity. This energy shift results from the long range strain effects experienced by one dot due to the presence of the second dots. This effect is missed in EMA-type model calculations which ignore strain effects.

**B. Single dot-localized orbitals**

The above discussions pertain to the basis of double-dot molecular orbitals. An alternative way to study QDMs is to use a dot-localized basis. We have demonstrated that dot-localized orbitals can be a useful tool to analyze the QDM physics, including the electron double occupation, and two-electron entanglement.

Dot-localized orbitals $\chi_\eta$ can be obtained from a unitary rotation of molecular orbitals, i.e.,

$$\chi_\eta = \sum_{i=1}^{N} U_{\eta,i} \psi_i ,$$  

where, $\psi_i$ is the $i$-th molecular orbital, and $U$ is a unitary matrix, i.e., $U^\dagger U = I$. We choose the unitary matrices $U$ that maximize the total orbital self-Coulomb energy $\eta$. The procedure of finding $U$ is described in the Appendix B of Ref.9. As we will show below these dot-localized orbitals $\chi_\eta$ have the advantage of being only weakly dependent to the inter-dot coupling. This invariance may provide simplified pictures for qualitatively understanding of the QDM physics.

1. Single-particle energies of dot-localized orbitals

The single-particle levels of dot-localized orbitals and the hopping (or tunneling) term between two dots can be obtained from

$$\epsilon_\eta = \langle \chi_\eta | \hat{H}_0 | \chi_\eta \rangle = \sum_i U_{\eta,i}^* \epsilon_i U_{\eta,i} ,$$  

$$t_{\eta_{12}} = \langle \chi_{\eta_1} | \hat{H}_0 | \chi_{\eta_2} \rangle = \sum_i U_{\eta_1,i}^* \epsilon_i U_{\eta_2,i} ,$$

where, $\epsilon_i$ is the single-particle energy of $i$-th molecular orbital and $\hat{H}_0 = \sum_{\sigma} \epsilon_\sigma \psi_\sigma^\dagger \psi_\sigma$ is the single-particle Hamiltonian. Figure 3 depicts the single-particle levels $e_T$ and $e_B$ of the dot-localized orbitals of both top and bottom dots, for inter-dot distances $d$ in the range from 4 nm to 10 nm. (Here, we denote the top dot $T$ and the bottom dot $B$, to distinguish them from isolated dots $\alpha$, $\beta$ and $\gamma$). $e_T$ and $e_B$ of $M_{\alpha\beta}$ are shown in the black
solid lines, and those of \( M_{\gamma\gamma} \) are shown in the red solid lines. At large \( d \), the energy difference \( e_B - e_T \sim 6 \) meV for \( M_{\alpha\beta} \), is close to the value of difference \( \epsilon(s_{\beta}) - \epsilon(s_{\alpha}) \) between \( s \) orbitals of isolated dots \( \alpha, \beta \). This energy difference gets smaller when the two dots move closer, because the energy levels of the top dot rise faster than those of bottom dots due to the strain asymmetry. For the homo-QDMs \( M_{\gamma\gamma} \), \( e_T \) and \( e_B \) are almost degenerate. The small difference (\( \sim 1 \) meV) between them is due to the strain and alloy effects. We also plot in Fig. 3 the energies of molecular orbitals \( \sigma_{\alpha} \) and \( \sigma_{\beta} \) in dashed lines for \( M_{\alpha\beta} \). As we see, for \( d > 9 \) nm, the dot-localized state \( e_B \) of \( M_{\alpha\beta} \) is almost identical to the molecular orbital \( \sigma_{\alpha} \), while \( e_T \) merges with \( \sigma_{\beta} \), indicating at large \( d \), molecular orbitals convert to dot-centered orbitals for \( M_{\alpha\beta} \).

The quantity \( 2t \) measures the coupling strength between the top and bottom dots, and directly determines the two-electron properties such as singlet-triplet splitting in the QDM. We calculate this hopping energy between the \( s \) orbitals of top and bottom dots at different inter-dot distances for both \( M_{\alpha\beta} \) and \( M_{\gamma\gamma} \) in Fig. 4. (We ignore the orbital index “s” to simplify the notation.) We find that \( 2t(M_{\alpha\beta}) \) and \( 2t(M_{\gamma\gamma}) \) are almost identical at all inter-dot distance. However, the hopping energies calculated here are much larger than what we obtained for the pure InAs/GaAs QDMs, because the alloy QDM have much smaller energy barrier between two dots than pure QDM. In general, the quantity \( 2t \) does not equal to the bonding-antibonding splitting \( \Delta_{\sigma} = \sqrt{\delta^2 + 4t^2} \), where \( \delta = \epsilon(e_T) - \epsilon(e_B) \), being the energy difference of \( s \) orbitals of the top and bottom dots. For homo-QDMs, where \( \delta/2t \ll 1 \), we have \( 2t \sim \Delta_{\sigma} \) as seen in Fig. 4. However, for hetero-QDMs, \( \Delta_{\sigma} \) may be significantly different from \( 2t \), especially at large inter-dot distances, where \( \delta/2t \gg 1 \), also illustrated in Fig. 4. Experimentally,\(^{19}\) one usually measures the bonding-antibonding splitting rather than the hopping \( 2t \). Therefore, to get the hopping energy between two dots, one need to know the energy difference \( \delta \) of two dots.

2. Coulomb integrals of dot-localized orbitals

The Coulomb integrals in the dot-localized basis can be obtained from Coulomb integrals of molecular orbitals as follows,

\[
\Gamma_{\eta_{\alpha},\eta_{\beta}} = \sum_{i,j,k,l} U_{\eta_{i},i}^* U_{\eta_{j},j} U_{\eta_{k},k} U_{\eta_{l},l} \Gamma_{i,j}^{i,j}, \tag{8}
\]

where \( \Gamma_{i,j}^{i,j} \) are the Coulomb integrals in the molecular basis. The direct Coulomb integrals \( J_{TT}, J_{BB} \) and \( J_{TB} \) for \( M_{\alpha\beta} \) are shown in Fig. 5. The Coulomb integrals \( J_{TT} \sim J_{\alpha\alpha} = 21.4 \) meV and \( J_{BB} \sim J_{\beta\beta} = 22.3 \) meV, are almost constants at all inter-dot distances, suggesting that the dot-localized orbitals are approximately unchanged for different inter-dot distance \( d \). \( J_{\beta\beta} > J_{\alpha\alpha} \), as dot \( \beta \) is smaller than dot \( \alpha \). The inter-dot Coulomb interaction \( J_{TB} \) decay slowly as \( 1/d \). The exchange energies (not shown) between the top and bottom electrons is orders of magnitude smaller than the hopping energy, and therefore can be ignored in practice. For the homo-QDM \( M_{\gamma\gamma} \), we found that on-site Coulomb energies \( J_{TT} \sim J_{BB} \), both are very close to the average values of \( J_{TT} \) and \( J_{BB} \) of \( M_{\alpha\beta} \). The inter-dot Coulomb energies \( J_{TB} \) of \( M_{\alpha\beta} \) and \( M_{\gamma\gamma} \) are also extremely close.

FIG. 4: (Color online) The inter-dot hopping energy \( 2t \) (solid lines) of hetero-QDM \( M_{\alpha\beta} \) and homo-QDM \( M_{\gamma\gamma} \). We also show the bonding-antibonding splitting \( \Delta_{\sigma} \) of \( M_{\alpha\beta} \) and \( M_{\gamma\gamma} \).

FIG. 5: The Coulomb energies of dot-localized orbitals of hetero-QDM \( M_{\alpha\beta} \). \( J_{TT} \) and \( J_{BB} \) are the \( s \) orbital self-Coulomb energies of top and bottom dots respectively, whereas \( J_{TB} \) are the Coulomb energies between \( s \) orbitals of the top and the bottom dots.
FIG. 6: (Color online) Two-electron states for (a) hetero-QDM $M_{\alpha\beta}$ and (b) homo-QDM $M_{\gamma\gamma}$, including the singlet $^1\Sigma_u^+$, $^3\Sigma_u^+$, $^3\Pi_g$ states and the 3-fold degenerated triplet states $^3\Sigma_u$ as well as two 3-fold degenerated triplet states $^3\Pi_u$.

IV. TWO ELECTRONS IN THE DOT MOLECULE

A. Many-body energy states

The two-electron-in-a-QDM problem is of special interest, as it is the prototype of quantum gate using QDMs. We calculate the two-electron energy levels by the configuration interaction method using Slater determinants constructed from confined molecular orbitals $\sigma_g$, $\sigma_u$, and $\pi_u$, $\pi_g$, which give 66 configurations in total. The two-electron energies $\Sigma$ and $^3\Pi_u$ for hetero-QDMs $M_{\alpha\beta}$ are plotted in Fig. 6(a). To compare with homo-QDMs, we show the two-electron states of $M_{\gamma\gamma}$ in Fig. 6(b). The energy levels of $M_{\alpha\beta}$ are similar to those of $M_{\gamma\gamma}$, in the following way: (i) The order of the CI levels is unchanged, particularly the ground states are still the singlet states $^1\Sigma_u^{(a)}$ at all inter-dot distances; (ii) The trend of each CI level vs. inter-dot distance $d$ is similar to what we obtained for $M_{\gamma\gamma}$. There are also some differences between the hetero-QDMs $M_{\alpha\beta}$ and homo-QDMs $M_{\gamma\gamma}$, especially at larger inter-dot distances. For example, in the homopolar QDMs, the $^1\Sigma_u$ state is almost degenerate with $^1\Sigma_g^{(b)}$ at large inter-dot distance, while in $M_{\alpha\beta}$, $^1\Sigma_g^{(b)}$ is about 13 meV higher than $^1\Sigma_u$ at $d=10$ nm. At large $d$, $^1\Sigma_u$ and $^1\Sigma_g^{(b)}$ correspond to the states that two electrons localized on the same dots. The energy difference between $^1\Sigma_g^{(b)}$ and $^1\Sigma_u$ is due to the size difference of dots $\alpha$ and $\beta$.

The singlet $^1\Sigma_u^{(a)}$ and triplet states $^3\Sigma$ can be used as two qubit states in quantum computing. In a proposed quantum SWAP gate, the gate operation time $\tau \sim 1/J_{S-T}$, where $J_{S-T}$ being the singlet-triplet energy splitting. The singlet-triplet splitting of $M_{\alpha\beta}$ is shown in Fig. 7 on a semi-log plot. We see that it decay approximately exponentially with the inter-dot distance. We also show in Fig. 7 the singlet-triplet splitting of the homo-QDM $M_{\gamma\gamma}$. We found that the $J_{S-T}$ of homo-QDM $M_{\gamma\gamma}$ is slightly smaller than the $J_{S-T}$ of hetero-QDM $M_{\alpha\beta}$, though the hopping energies of $M_{\alpha\beta}$ and $M_{\gamma\gamma}$ are almost identical. In the hetero-QDM case, the singlet wavefunction has more weight on the lower energy dot and therefore lowers the singlet energy and increases the singlet-triplet splitting.

B. Double occupation of one of the dots in a QDM

Double occupation means that two electrons occupy the same dot in a QDM. If the double occupation rate is high, the quantum gate operation may fail. The double occupation rate also reflects the localization properties of electrons in the QDM. If the double occupation rate is zero, each dot has one electron, whereas double occupation rate of 1 means that two electrons are always localize on a single dot. When the double occupation rate is 0.5, two electrons are delocalized between two dots. The double occupation can be conveniently analyzed in the dot-localized basis by transforming the CI equations to the dot-localized basis. In the simplest case, we consider only the “s” orbital for each dot, which give six config-
Coulomb integrals, which are much smaller than the hopping. We denote both on the top dots, or both on the bottom dots, or one on the top dots, or both on the bottom dots. We denote both on the top dots, or both on the bottom dots, or one on the top dots, or both on the bottom dots. The Hamiltonian in this basis set

\[
H = \begin{pmatrix}
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0
\end{pmatrix} 
\]

occupying the dot \(p = (T \text{ or } B)\) at the same time, i.e.,

\[
Q_{\nu}^{(v)} = \sum_{\sigma, \sigma'} P_{\sigma}(|\chi_{I, p}^{\sigma}, \chi_{I', p}^{\sigma'}|),
\]

where \(P_{\sigma}(C)\) is the weight of the configuration \(C\) in the many-body wave functions of state \(\nu\). The total probability of two electrons being on the same dot is then \(Q_{\nu}^{(v)} = Q_{\nu}^{(v)} + Q_{\nu}^{BB}\) for the \(\nu\)-th state.

We plot \(Q_{\nu}^{(v)}\) for state \(1\Sigma_{(a)}\) for \(M_{\alpha\beta}\) in Fig. 8(a) and for \(M_{\gamma\gamma}\) in Fig. 8(b). We also performed calculations on a “symmetrized” model QDM \(M_{\alpha'\alpha'}\) by setting \(e_T' = e_B' = (e_T + e_B)/2\) and \(J_{TT}' = J_{BB}' = (J_T + J_B)/2\) of \(M_{\alpha\beta}\) in Eq. (9). \(M_{\alpha'\alpha'}\) represents an ideal homo-QDM, without the asymmetry caused by strain, size and alloy composition effects. When compare the double occupation of the hetero- and homo-QDMs, we see that (i) For both types of QDMs, \(Q_{\nu}^{(v)} \sim 0.5\) at \(d \sim 4.5\) nm, meaning that two electrons are delocalized on two dots. For both QDMs, \(Q_{\nu}^{(v)}\) decays monotonically with the inter-dot distance, and at \(d \sim 10\) nm, \(Q_{\nu}^{(v)} \sim 0\), meaning that the two electrons are about each localized on one of the two dots.

On the other hand, the double occupation of individual dot \(Q_{TT}\) and \(Q_{BB}\) differ substantially for homo-QDMs and hetero-QDMs:

(ii) For the homo-QDM \(M_{\alpha'\alpha'}\), \(Q_{BB} = Q_{TT}\) and decay monotonically with the inter-dot distances. \(Q_{BB}\) and \(Q_{TT}\) of \(M_{\gamma\gamma}\) have similar features, although \(Q_{BB}\) is slightly different from \(Q_{TT}\) due to the strain and alloy effects. This feature is also seen in the homo-QDM made of pure InAs/GaAs dots. In the hetero-QDMs \(M_{\alpha\beta}\), \(Q_{TT}\) behaves very differently from \(Q_{BB}\) because the effective single-particle energy \(e_T < e_B\). Whereas \(Q_{BB}\) decays monotonically with the inter-dot distance, \(Q_{TT}\) has a maximum at \(d \sim 7\) nm. The reason is that at \(d \sim 4.5\) nm, the hopping energy \(2t\) is much larger than \(e_B - e_T\), therefore the electrons can overcome the energy barrier between the top and bottom dots and distribute evenly between two dots, leading to \(Q_{TT} \sim Q_{BB}\). At larger \(d\), \(2t \ll e_B - e_T\), and the electrons would prefer to localize on the top dots, leading to \(Q_{TT} \gg Q_{BB}\). Therefore, even when the total double occupation rate drops down, \(Q_{TT}\) still increases and reaches the maximum at \(d = 7\) nm. For \(d > 7\) nm, \(Q_{TT}\) decays as \(Q_{\nu}^{(v)}\) decays.

FIG. 8: The double occupation rate of the ground-state singlet \(1\Sigma_{a}(a)\) vs. inter-dot distance for (a) hetero-QDM \(M_{\alpha\beta}\) and (b) homo-QDM \(M_{\gamma\gamma}\). Where \(t = t_{TB}\). We ignored in Eq. (9) the off-diagonal Coulomb integrals, which are much smaller than the hopping \(t\).

The calculation of the matrix elements of Eq. (9) is described in Sec. II B. The two electrons can be either both on the top dots, or both on the bottom dots, or one on the top and the other on the bottom dots. We denote by \(|\chi_{I, p}^{\sigma}, \chi_{I', p}^{\sigma'}|\) the configuration where one electron is on the \(\iota\)-th orbital of the \(p\) dot with spin \(\sigma\), and the other electron is on the \(\iota'\)-th orbital of the \(p'\) dot with spin \(\sigma'\). Then the double occupation rate \(Q_{\nu}^{(v)}\) in the many-particle state \(\nu\) is the probability of two electrons
quantum correlation function for two electrons which has similar properties as the DOE. However, the generalization of this quantum correlation function to a system that has more than two single-particle levels is complicated. We proposed a DOE measure\(^1\) for indistinguishable fermions using the Slater decompositions\(^{24,31}\) as,

\[
S = -\sum_i z_i^2 \log_2 z_i^2 ,
\]

where, \(z_i\) are Slater decomposition coefficients and \(\sum_i z_i^2 = 1\). As shown in Ref. 9, the DOE measure Eq. (11) reduces to the usual Von Neumann entropy for distinguishable particles when the two electrons are far from each other. In Refs. 25,26, a similar DOE measure was defined, which however due to a different normalization condition for \(z_i\) was used, does not reduce to the usual Von Neumann entropy even when the two electrons can be distinguished by their sites.

The DOE of Σ states calculated from Eq. (11) for the hetero-QDM \(M_{a\beta}\), the homo-QDM \(M_{\gamma\gamma}\), and the model homo-QDM \(M_{a'a'}\) are shown in Fig. 9(a), (b) and (c) respectively. All of the three QDMs have the following features: (i) \(S(\Sigma_g^{(a)})\) is close to zero (unentangled) at \(d \sim 4.5\) nm, and close to unity (fully entangled) at \(d \sim 10\) nm. (ii) \(S(\Sigma_u^{(g)})\) is almost unity (fully entangled) at all inter-dot distances. However, \(S(\Sigma_g^{(a)})\) of the homo-QDM \(M_{\gamma\gamma}\) (which is very close the \(S(\Sigma_g^{(a)})\) of \(M_{a'a'}\)) is larger than \(S(\Sigma_u^{(g)})\) of the hetero-QDM \(M_{a\beta}\), showing that the asymmetry in a QDM lowers the two-electron entanglement of the ground state singlet.

In contrast to \(S(\Sigma_g^{(a)})\) and \(S(\Sigma_u^{(g)})\), \(S(\Sigma_g^{(b)})\) and \(S(\Sigma_u^{(b)})\) are very sensitive to the asymmetry of the QDMs. In general, if the two dots have identical electronic structures (e.g., in the simple Hubbard model), \(S(\Sigma_g^{(b)}) = S(\Sigma_g^{(a)})\) and \(S(\Sigma_u^{(b)}) = S(\Sigma_u^{(a)}) = 1\) as is illustrated in Fig. 9(c) for \(M_{a'a'}\). For \(M_{\gamma\gamma}\), which is somehow asymmetric due to the strain and alloy effects, \(S(\Sigma_g^{(b)})\) is close to \(S(\Sigma_g^{(a)})\) at small \(d\), and drops down at large \(d\), whereas for \(M_{a\beta}\), \(S(\Sigma_g^{(b)})\) is different from \(S(\Sigma_g^{(a)})\) at all inter-dot distances. The slight asymmetry in \(M_{a\beta}\) also causes \(S(\Sigma_u^{(b)})\) to drop down at large \(d\), similar to \(S(\Sigma_u^{(a)})\) of \(M_{a\beta}\).

**V. ENTANGLEMENT**

**A. Degree of entanglement for two electrons**

The degree of entanglement (DOE) is one of the most important quantities for successful quantum gate operations. For distinguishable particles such as an electron and a hole, the DOE can be calculated from the Von Neumann-entropy formulation.\(^{20,21,22,23}\) However, Von Neumann entropy formulation cannot be used directly to calculate DOE for indistinguishable particles.\(^{24,25,26,27,28,29,30}\) Schliemann et al. proposed a
be generally written as,

$$
\Psi^{(1\Sigma_g^\alpha)} = c_1 |e^+_B, e^+_B\rangle + c_2 |e^+_B, e^+_T\rangle + c_3 |e^+_T, e^+_T\rangle + c_4 |e^+_B, e^+_B\rangle ,
$$

(12)

and $|c_1|^2 + |c_2|^2 + |c_3|^2 + |c_4|^2 = 1$. Alternatively, we have

$$
\Psi^{(1\Sigma_g^\alpha)} = \sum_{i,j} \omega_{ij} |i\rangle \otimes |j\rangle ,
$$

(13)

where,

$$
\omega = \begin{pmatrix}
0 & -c_3 & 0 & -c_1 \\
c_3 & 0 & c_2 & 0 \\
0 & -c_2 & 0 & -c_4 \\
c_1 & c_3 & c_4 & 0
\end{pmatrix},
$$

(14)

and $|i\rangle , |j\rangle = |e^+_B\rangle, |e^+_T\rangle, |e^+_B\rangle, |e^+_B\rangle$. We can use Eq. (11) to calculate the DOE, where $z_1^2 = 1/2(1 - \sqrt{1 - 4(c_1c_2 - c_3c_4)^2})$ and $z_2^2 = 1/2(1 + \sqrt{1 - 4(c_1c_2 - c_3c_4)^2})$ are the eigenvalues of $\omega^4 \omega$. For a QDM with reflection symmetry, we have $c_1 = c_2$ and $c_3 = c_4$, and therefore $z_1^2 = 1/2(1 - \sqrt{1 - (1 - 4c_3^2)^2})$, and $z_2^2 = 1/2(1 + \sqrt{1 - (1 - 4c_3^2)^2})$. Using the definition of double occupation rate, $Q_{\text{tot}} = c_3^2 + c_4^2$, we have

$$
z_1^2 = 1/2(1 - \sqrt{1 - (1 - 2Q_{\text{tot}})^2}) ,
$$

$$
z_2^2 = 1/2(1 + \sqrt{1 - (1 - 2Q_{\text{tot}})^2}) .
$$

(15)

The DOE of $1\Sigma_g^\alpha$ is calculated by substituting $z_1^2$, $z_2^2$ into Eq. (11). We plot the DOE vs. double occupation rate of the above ideal model in Fig. 10 in a black solid line. We also present in the same figure, the DOE of $M_{\alpha\beta}$, $M_{\gamma\gamma}$ and $M_{\alpha'\alpha'}$ vs. double occupation rate. We found that the double occupation dependence of DOE for the homo-QDM $M_{\alpha'\alpha'}$ has perfect agreement with the analytical result, which is also true for $M_{\gamma\gamma}$ even though it has small asymmetry in the molecule due to the strain and alloy effects. We also checked the homo-QDM made of pure InAs/GaAs dots, and found the same double occupation dependence of DOE for the $1\Sigma_g^\alpha$ state, indicating this is a robust feature for homo-QDMs. However, the double occupation dependence of DOE for $M_{\alpha\beta}$ deviates from the ideal case because dots $\alpha$ and $\beta$ are different.

VI. SUMMARY

We have studied the electronic structures of quantum dot molecules made of (In,Ga)As/GaAs dots of different sizes (hetero-QDMs), and compare them to that of quantum dot molecules made of identical dots (homo-QDMs). We found that while the hetero-QDMs and homo-QDMs have relatively similar electronic structures at short inter-dot distance, they differ significantly at large inter-dot distance. (i) Unlike those of homo-QDMs, the single-particle molecular orbitals of hetero-QDMs convert to dot localized orbitals at large inter-dot distance. (ii) Consequently, the bonding-antibonding splitting of molecular orbitals is significantly larger than the electron hopping energy in a hetero-QMD at large inter-dot distance, whereas for homo-QDM, the bonding-antibonding splitting is very similar to the hopping energy. (iii) The asymmetry of the QDM will significantly increase the double occupation for the two-electron ground states, and therefore lowers the degree of entanglement of the two electrons.

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