Toward tripartite hybrid entanglement in quantum dot molecules

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Abstract

Establishing the hybrid entanglement among a growing amount of matter and photonic quantum bits is necessary for scalable quantum computation and long-distance quantum communication. Here we demonstrate that charged excitonic complexes forming in strongly correlated quantum dot molecules are able to generate tripartite hybrid entanglement under proper carrier quantization. The mixed orbitals of the molecule construct multi-level ground states with sub-meV hole tunneling energy and relatively large electron hybridization energy. We show that appropriate size and interdot spacing keeps the electron particle weakly localized, opening extra recombination channels by correlating ground-state excitons. This allows for creation of higher order entangled states. Nontrivial hole tunneling energy, renormalized by multi-particle interactions, facilitates the realization of the energy coincidence among only certain components of the molecule optical spectrum. This translates to the emergence of

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favorable spectral components in a multi-body excitonic complex which sustain principal oscillator strengths throughout the electric field-induced hole tunneling process. We particularly analyze whether the level broadening of favorable spin configurations could be manipulated to eliminate the distinguishability of photons.

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1. Introduction

Strong localization of charge carriers in quantum dots (QDs) maintains a sufficiently long coherence time of their spin quantum bits (qubits), easing the realization of quantum correlation between the carrier spin and flying photons, which is a compulsory step toward developing secure quantum communication [1]. Experimental evidence reported so far on photon (or spin) qubit entanglement in isolated QDs [2–4] is a valid manifestation of their ability in this context. Although realizing bipartite quantum entanglement has been increasingly studied in recent years, there exists no theoretical proposal on generating higher order entangled states in real QD structures. In the simplest picture, the reason lies in the anharmonic energy level spectrum of a typical QD that impedes color matching amid its absorption energies, making the excitonic transitions distinguishable [5, 6]. Since the entangled degree of freedom is mostly defined based on the polarization of a photon (or the spin state of a carrier), this color distinguishability reduces the degree of entanglement, even in a bipartite system. Current solutions offered to establish the coincidence in the energy of photons are often perturbational [3, 7–10] or architectural [11–13], allowing for the creation of up to bipartite quantum correlated states. Actualizing higher order entangled states, however, seems unlikely in a single QD due to the absence of more than two color-matching excitons \( X \) in its optical spectrum [14]. Experimental demonstration of tripartite entanglement has so far succeeded only for spin states in diamond NV centers [15] and photon states via spontaneous parametric downconversion [16].

Considering the fact that photon entanglement in QDs relies on the sequential decay of correlated excitons [6], preserving the spatial overlaps of excitonic orbitals in the quantum dot molecule (QDM) is a requisite to maintain this correlation. Creating higher order entangled states thus demands for excitons to spatially spread across the entire QDM, rather than a single QD, otherwise the excitons localized in different dots are partially correlated merely via their Coulomb interactions. This means that one constituting particle of the exciton, that is the electron or hole, has to be weakly quantized to establish a strong correlation between the two dots. In contrast to the previous experiments [17, 18], here we show that weakly localized electron orbitals with relatively large tunneling energy retain the correlation and give rise to optically active ‘indirect’ excitonic recombinations, opening new channels of entangled photon emission. The QDM in our model consists of two axially stacked In(Ga)As QDs embedded in a GaAs nanowire [19]. An axial electric field can then be exploited to drive the interdot hole tunneling, and eventually to entangle or disentangle electron–hole pairs [20, 21]. In contrast to the electron, the hole tunneling energy is noticeably renormalized by the spin–orbit interaction in III–V materials; see the supplementary information, available from stacks.iop.org/NJP/16/023019/mmedia and [22]. Moreover, the emergence of new entangled channels strongly depends on the spin fine structure of holes. Thus, a detailed quantum
mechanical treatment is required to address all of the underlying phenomena, including spin–orbit and Coulomb interactions, in a few-body populated QDM.

In this paper, we study the capability of QDMs as potential sources of higher order entanglement. We show that the configurations of multiexcitons can be deterministically controlled by a tunneling mechanism between the hybridized s-shells via an external source of electric field. We analyze the particular case of hybrid entanglement in which the spin of a carrier resting in a metastable ground state is correlated with the polarization of twin photons. To this end, we confine our model to the sequential decay of a negatively (positively) charged biexciton, $XX^-$ ($XX^+$), down to the singly charged electron (hole) state, $XX^- \rightarrow X^- \rightarrow e$ ($XX^+ \rightarrow X^+ \rightarrow h$), although the scheme can be generalized to higher orders with more complexity. We demonstrate that only a limited number of transitions are favorable in the field-dependent spectrum of QDM due to the few-body interactions and tunneling effects.

2. Basic concepts and theoretical approach

The energy level structure of a typical single QD is composed of s, p, d and higher shells with resemblance to the atomic states, where the s-shell is referred to as the ground state. In a QDM, the orbitals of each individual shell from single dots are mixed and their corresponding energy levels shift based on the coupling strength. The electron and hole s-shells of a single dot, $s_e$ and $s_h$, can host up to two interacting excitons $X$ (a biexciton $XX$); thus the formation of any higher order complex, such as the negatively charged biexciton $XX^-$ ($X + X + e$), inevitably involves the p-shells; see figure 1(a). However, p-shell carriers are not immune to phonon-mediated decay into the ground state, and the initialization and quantum manipulation of carrier spin in the s-shell is generally preferred.

The QDM structure we address is a symmetric double dot embedded inside a III–V nanowire. Analogous to any other quantum-confined structure, the starting point to study the QDM is to find its bound states, namely the molecular orbitals. Since we will be dealing with multi-particle states in the following sections, we also refer to these molecular orbitals as single-particle states. To solve for the molecular orbitals we consider the whole QDM as a confining heterostructure and diagonalize the modified Luttinger–Kohn Hamiltonian introduced in [23]. The molecular orbitals could be approximately represented by the linear combination of dot-localized orbitals beyond the thin barrier limit [24]. They construct the basis set required to form the Hilbert space of multi-particle states. An electric field applied along the nanowire axis is able to change the character of molecular orbitals into dot-localized orbitals, and thus tailor the corresponding Coulomb interactions. We solve for the single- and multi-particle states at several different axial electric fields and resolve the photoluminescence intensity of multi-particle transitions at grid points and midpoints by interpolation.

The QDM is designed such that the electron molecular orbitals are weakly quantized over the entire molecule and their orbitals negligibly deform in response to the axial electric field, preserving the multi-particle correlations and switching on additional transitions required for the cascade emission. The electron molecular orbitals remain weakly localized even after the introduction of Coulomb interactions in the multi-particle states. This is because the hole molecular orbitals are likewise delocalized at zero axial field. In contrast to the electron states, hole single particles quickly localize in response to the electric field. We will demonstrate that the variation range of Coulomb interactions versus electric field is negligible compared to the electron tunneling energy.
In section 6 we study the charged biexciton cascade in QDMs in the presence of hole tunneling. The emergence of favorable cascade transitions depends on the spin fine structure of the contributing complexes, namely $XX^-$ and $X^-$ ($XX^+$ and $X^+$), and localization of the molecular orbitals under the variable electric field. Furthermore, the color distinguishability of bright transitions relies on the hole hybridization energy, the existing Coulomb interactions and the Zeeman energy shift induced by the magnetic field used for the spin control. The field-dependent photoluminescence spectrum of the QDM is calculated for different transitions contributing in the $XX^- \rightarrow X^- \rightarrow e$ and $XX^+ \rightarrow X^+ \rightarrow h$ cascades. These spectra indicate that only a few spectral components, corresponding to both initial and final favorable spin configurations, are bright. Unfavorable spin configurations refer to those multi-particle states that the Coulomb blockade or Pauli spin blockade prevent them from forming.

The problem of hybrid entanglement between the spin and photons urges coherent control of the electron or hole spin by means of an external magnetic field [25]. Carrier spin preparation is normally accomplished through exciting the singly charged QD into the charged exciton ($X^- = X + e$ or $X^+ = X + h$) state and waiting until the spontaneous decay leaves the QD either in $|\uparrow\rangle$ or $|\downarrow\rangle$ states, while the spin state is coherently controlled through a Raman transition driven by a few picosecond pulse at its qubit resonance frequency [26, 27]. The formation of charged biexcitons, $XX^-$ and $XX^+$, from the $|\uparrow\rangle$ or $|\downarrow\rangle$ states is then feasible through a two-photon excitation process [28]. We presume single-electron states $|e\uparrow\rangle$ and $|e\downarrow\rangle$ to be nearly metastable because the $|\uparrow\rangle \Leftrightarrow |\downarrow\rangle$ spin-flip transitions are hardly induced by the hyperfine interaction of the electron spin with the nuclear spin ensemble even at small magnetic fields.
(γ_{1}|↑⟩ \leftrightarrow |↓⟩) \ll \Gamma$, but are instead regulated by the spin-flip Raman process and the intensity of the stimulating laser [26, 29].

The spectral response of QDM to the magnetic field appears in the Zeeman shift of multi-particle states and their spin coupling, thus directly influencing the color and linewidth distinguishability of transitions. Along with the Zeeman shift, the exchange interaction causes energy splittings in the spin fine structure of multi-particle states. However, by performing a detailed configuration interaction (CI) calculation, we demonstrate that the scale of Zeeman splittings rapidly exceeds the exchange splittings, thus the energy indistinguishability of transitions is primarily violated by the Zeeman effect. To obtain a clear sense of the magnetic field intensity required for spin state initialization and manipulation in QDs, we rely on earlier experiments reporting acceptable fidelities [30, 31].

A viable solution for erasing these sub-meV energy splittings is to employ a high-efficiency downconversion technique. The method consists of cross-correlating the emitted photons from QDs with a few-picosecond pulse in a nonlinear medium such as a periodically poled lithium niobate (PPLN) waveguide [32]. The time resolution of pulses determines the arrival time of single photons and therefore broadens their linewidths (the lifetime of an exciton is commonly beyond 0.5 ns \( \gg \) few ps). This idea could be extended to an arbitrary stream of anti-bunched photons, including entangled photons. Figure 1(c) schematically shows how the portion of fully entangled photons being delivered by the source can be increased through the linewidth broadening. Twin photons \( P_1 \) and \( P_2 \) have an initial linewidth equivalent to \( \Gamma \) and their color mismatch is \( \delta \). Assuming that \( \delta \) remains constant, the overlap region, which implicitly represents the fraction of quantum correlated photons, grows by enhancing the linewidth \( \Gamma \rightarrow \Gamma' \). To quantitatively measure the energy coincidence of photons, we utilize the concept of concurrence developed for entanglement in both regular and time-reordering schemes [33–35]. In the spin fine structure of a typical QD, concurrence is generally a function of energy mismatches and linewidths of excitonic levels. We calculate the evolution of concurrence versus the external magnetic field intensity and photon linewidth. The results reveal that a sufficiently high concurrence, as compared to the previous experimental results [36, 37], could be reached by manipulating the linewidths properly. It is worth noting that the QDM with strongly correlated excitons can be exploited to generate higher order entangled photon states or hybrid spin–photon states, and the current work serves as a prototypical study on the application of QDMs in creating tripartite Greenberger–Horne–Zeilinger and W-states [38].

3. Molecular orbitals in a nanowire double quantum dot

The double QD we study consists of two In_{0.5}Ga_{0.5}As insertions with the ideal \( D_\infty h \) symmetry embedded inside a [001]-oriented GaAs core–shell nanowire; see figure 2(a). The QD diameter and its vertical aspect ratio, \( a_h = h_D/D_0 \), are chosen as 20 nm and 0.25, respectively. This ratio leads to comparable mutual interactions between electrons and holes, thus the multi-particle states accumulate less correlation energy [13]. The interdot spacing \( D_s \) determines the wave function symmetry of the hole particle ground state \( |h_0⟩ \) and the sign of its tunneling matrix element \( t_h \) to hop up to the first excited state \( |h_1⟩ \). Our numerical calculations at the single-particle level demonstrate that the \( |h_0⟩ \) symmetry reversal, as well as the sign inversion of the tunneling matrix element [39] (\( t_h \) becoming negative), occur at \( D_s \approx 1.8 \) nm. The dominant heavy hole-like part of the ground-state envelope function \( \psi_{J=3/2, J_z=3/2}^h \).
Figure 2. (a) Schematic of QDM embedded in a [001]-oriented core–shell nanowire. $D_s$: the interdot spacing between the left ($L$) and right ($R$) single dots. $h_D$: height of the single QD and $D_D$: diameter of the single QD. Although the structure of the QDM is assumed to be symmetric, the strain field relaxes away from the substrate and becomes axially asymmetric. (b), (c) Electron and hole molecular energies versus the interdot spacing. The first and second electron excited states are energetically inseparable for $D_s < 3$ nm as the whole QDM acts similar to a single dot for electrons at the thin barrier limit. The highlighted region in (c) specifies where the hole ground state is bonding (axially symmetric). The hole tunneling matrix element becomes negative at $D_s = 1.8$ nm where the hole ground state and first excited states anticross. (d), (e) Envelope function of the hole ground state $\phi_{h0}$ plotted along the nanowire axis at $D_s = 1$ and 2 nm showing its bonding and antibonding nature versus $D_s$. The highlighted region shows where the hole ground state is symmetric.

(see supplementary information, available from stacks.iop.org/NJP/16/023019/mmedia) is illustrated in figures 2(d)–(e), where its parity flips moving from $D_s = 1$ to 2 nm.

The molecular ground and first excited states of the hole, $|h_0\rangle$ and $|h_1\rangle$, are hybridizations of the two dot-localized s-shell states, $|s_{h,R}\rangle$ and $|s_{h,L}\rangle$. These two mixtures are highly correlated at small double-dot spacing, where the $|s_{h,R}\rangle$ and $|s_{h,L}\rangle$ orbitals spread uniformly over the QDM, and become $|s_{h,R}\rangle$-like or $|s_{h,L}\rangle$-like when the spacing is sufficiently increased. An analogous situation occurs for the molecular ground state $|e_0\rangle$ and first excited state $|e_1\rangle$ of the electron in terms of losing the correlation versus $D_s$. Figures 2(b) and (c) illustrate the evolution of the corresponding electron and hole energies, $E_{e0}$, $E_{e1}$, $E_{h0}$ and $E_{h1}$ against the interdot spacing. $4 < D_s < 6$ nm can be seen to be a reasonable range for the interdot distance for the following reasons: (i) hole hybridization energy is suppressed within this range since the tunneling component caused by the spin–orbit interaction weakly exceeds the diagonal tunneling matrix element (the definitions of these tunneling components are explained in the supplementary information (see the supplementary information, available from stacks.iop.org/NJP/16/023019/mmedia)). (ii) The electron first excited state (predominantly s-like) is noticeably coupled to the second excited state (predominantly p-like) given $D_s < 3$ nm.
Figure 3. (a), (b) S- and p-shell hole molecular orbitals versus the electric field indicated in kV cm\(^{-1}\). Hole orbitals become localized inside single dots at \(E_z = \pm 3\) kV cm\(^{-1}\), thus their wave function could be approximated by the dot-localized states. Applying an axial electric field reduces the symmetry character of orbitals from \(\sim D_{2d}\) down to \(C_{2v}\). (c) Electron molecular orbitals. Owing to the small effective mass of the electron, its molecular orbitals reshape reluctantly in response to \(E_z\). The electron wave function remains as the linear mixture of dot-localized states within \(-5 < E_z < 5\) kV cm\(^{-1}\). The \([001]\) axis is slightly rotated with respect to the left panel (a and b) in order to illustrate the diffusion of the electron orbital inside the GaAs barrier (\(t_e > t_h\)). (d) Electron (four states, 2s and 2p) and hole (seven states, 2s, 4p and 1d) energies as a function of the axial electric field.

as a result of their trivial energy spacing; this leads to a pronounced correlation between the two molecular states \(|e_1\rangle\) and \(|e_2\rangle\) for \(D_s < 3\) nm. (iii) \(D_s\) could be increased limitedly because the hole tunneling matrix element rapidly vanishes. Hereinafter and in order to observe the field-induced spectrum of the molecule at a fixed interdot spacing we set \(D_s\) equal to 5 nm.

4. Molecular orbitals versus the axial electric field

In the next step, an axial electric field \(E_z (z \equiv [001])\) is applied in order to tailor the absorption energies of the QDM. Owing to the \(D_{\infty h}\) symmetry of single QDs, the orbital densities are quite similar once \(E_z = 0\), neglecting trivial asymmetries caused by the strain-induced fields; see figures 3(a)–(c). Both electron and hole molecular orbitals exhibit approximately \(D_{2d}\) symmetry,
spreading throughout the molecule where each orbital lobe is \( C_{2v} \)-symmetric. Immediately upon applying the axial electric field, the heavy hole with highest energy \( |h_0\rangle = |h^2_i\rangle \) moves toward lower potential energy in QD\(_R\), and effectively evacuates QD\(_L\) at \( E_z \approx \pm 3 \text{kV cm}^{-1} \) acquiring \( C_{2v} \) symmetry; \( |h_0\rangle = |s_{h,R}\rangle \). The very same situation occurs for \( |h_1\rangle = |h^2_i\rangle \) under negative comparable electric fields. The Galium intermixing in the QD composition facilitates both hybridization and interdot diffusion of the electron and hole. As implied from its dispersion, the electron orbital is stiff against electric field variations, otherwise it would gain considerable kinetic energy \([13]; \) see figure 3(c). Thus \( |e_0\rangle = |e^1_i\rangle \) and \( |e_1\rangle = |e^2_i\rangle \) orbitals sustain their spread out over both QDs maintaining the interdot coupling, \( |e_0\rangle = |s_{e,L}\rangle + |s_{e,R}\rangle \) and \( |e_1\rangle = |s_{e,L}\rangle - |s_{e,R}\rangle \).

S-shell electron and hole energies undergo an anticrossing in proximity to \( E_z = 0; \) see figure 3(d). In accordance with trivial orbital deformations, the s-shell electron energies experience relatively small variations compared to the s-shell hole energies. Along with the relatively large energy spacing between \( |e_0\rangle \) and \( |e_1\rangle \), this leads to the predominant hole contribution in any conversion between the excitonic configurations at low electric fields (\( |E_z| \leq 1.5 \text{kV cm}^{-1} \)). Owing to the strain-induced potentials, electrons and holes do not necessarily localize symmetrically in response to negative and positive electric fields, prohibiting a perfectly symmetric evolution of spectral features versus the axial electric field.

Since the electrons are weakly localized in our electric field range, the correct representation is to assign \( |P\rangle = |s_{e,L}\rangle + |s_{e,R}\rangle \) and \( |N\rangle = |s_{e,L}\rangle - |s_{e,R}\rangle \) to the associated molecular states inward and away from the resonance. In the case of holes, however, we refer to the molecular states as \( |R\rangle = |s_{h,R}\rangle \) and \( |L\rangle = |s_{h,L}\rangle \) away from the resonance as their orbital could be well approximated by the dot-localized states there. To simplify the nomenclature, we agree to use the \( |R\rangle \) and \( |L\rangle \) notation even for holes close to the anticrossing; however, we are implicitly aware of the fact that \( |R\rangle \approx |s_{h,L}\rangle - |s_{h,R}\rangle \) and \( |L\rangle \approx |s_{h,R}\rangle + |s_{h,L}\rangle \) at this point.

5. Charged biexciton cascades in quantum dot molecules: role of hole tunneling in the energy coincidence of transitions

Since generating multi-partite correlated states in our scheme relies on the cascade recombination of charged multi-particle complexes, namely \( XX^- \) and \( XX^+ \), inspecting the contributing transitions of the QDM in the presence of hole tunneling sheds light on the essential properties we seek in the energy of emitted photons. To this end, we consider the exemplary cascades sketched in figure 4 and calculate the transition energies of the diagram. We focus on the cross-entanglement scheme where the hole tunneling energy plays a substantial role. In this scheme, the energy coincidence between \( XX^+_i \rightarrow X_i^- \) and \( X_i^+ \rightarrow e \) (or \( XX^+_i \rightarrow X_i^+ \) and \( X_i^- \rightarrow h \) \( i \neq j \) transitions must be established; \( i \) \( j \) and \( k \) sweep over possible spin configurations of \( X^- \) and \( X^- \). In the diagram shown in figure 4, the \( XX^- \) spin configurations are \( |e_{\sigma_1}e_{\sigma_2}h_{\sigma_1}h_{\sigma_2}\rangle = |P_1P_1N_1R_1L_\emptyset\rangle \) and \( |P_1P_2N_1R_1L_\emptyset\rangle \), where one electron molecular state is fully occupied and other states are singly filled up. Upon two inequivalent \( |P_1R_\emptyset\rangle \) and \( |P_1L_\emptyset\rangle \) recombinations starting from \( |P_1P_1N_1R_1L_\emptyset\rangle \) (\( |P_1P_1N_1R_1L_\emptyset\rangle \)) and \( |P_1N_1R_\emptyset\rangle \) (\( |P_1N_1R_\emptyset\rangle \)) emerge, whose energy spacing, labeled by \( \Delta X^+_i = \delta_{\text{exc}} X^+_i = \delta_{\text{exc}} X^+_i \) in figure 4, lies in the hole tunneling matrix element along with the direct and exchange Coulomb interactions. The absorption energy detuning between the \( \sigma_{1i}^+ \)-photon in path \( \mathcal{P}_1 \) and the \( \sigma_{4i}^+ \)-photon in path \( \mathcal{P}_4 \) under zero magnetic
Figure 4. Cascade decays initiated from $|P_1 P_1 N_1 R_0 L_0\rangle$ and $|P_1 P_1 N_1 R_0 L_0\rangle$ negative biexcitons down to intermediate negative trion and electron states, producing right $\sigma^+$ and left $\sigma^-$ polarized photons. The axial electric field is assumed to be nonzero, $E_z \neq 0$, squeezing hole orbitals into dot-localized states. Weak localization of the electron is schematically represented by its orbital filling the whole QDM. $B_x = 0$, thus spin configurations are still not Zeeman-shifted. Spin–photon pair entanglement could be established between each two paths of the left and right panels (subscripts denote the path numbers). Due to the spin–orbit interaction, degeneracies in the charged configurations are lifted (i.e. nonzero $\delta_{\text{exc}}^X$) even at zero magnetic field.

\[ \Delta_{X}^{\alpha} = \Delta_{\text{exc}}^{\alpha} \]  

represents the energy spacing of intermediate trions ruled by the hole tunneling matrix element and Coulomb interactions.

Field, $\delta_{\sigma^+} = (\approx \delta_{\sigma^-})$, neglecting the difference between exchange interactions, is given by

\[ \delta_{\sigma^+} = 2t_h + J_{hh} - J_{ee} - \Delta_{\text{exc}}^X - \Delta_{\text{exc}}^X - \Delta_{\text{exc}}^X - \Delta_{\text{exc}}^X, \]  

(1)

where $t_h = (E_{X}^R - E_{X}^L)/2$ represents the hole tunneling matrix element, $J_{ab}$ stands for the mutual direct Coulomb interaction between particles $a$ and $b$ in the molecular states A and B, and $\Delta_{\text{cor}} (\Delta_{\text{exc}})$ denote the correlation (exchange) energy stored in the complex $\alpha$. Superscript AB denotes the potential of a single particle resting in molecular state A being felt by another single particle localized in state B. In order to obtain the above equation, the total energies of the initial and final complexes in each transition are derived and subtracted. Equation (1) holds once the two paths (here $P_1$ and $P_4$) do not share identical initial (negative biexciton) and ground (electron) states.

As implied by equation (1), $\delta_{\sigma^+}^X$ can be suppressed upon comparable attractive ($J_{\text{exc}}^{AB}$) and repulsive ($J_{\text{exc}}^{AB}$) Coulomb interactions assuming $t_h$, $\Delta_{\text{cor}}$, $\Delta_{\text{exc}}^{\alpha} \ll J_{ab}^{AB}$ [40]. Fulfilling the condition $J_{hh}^{LR} + J_{ee}^{NN} = J_{eh}^{NR} + J_{eh}^{NL}$ then requires a relative similarity between the electron molecular states as well as left- and right-localized hole states. The necessity of this resemblance further unfolds when considering a single QD: $XX^-$ forms once a biexciton fills up the s-shell and a single electron resides in the p-shell. Since transitions between the s and p shells are ideally forbidden $M_{sp} \ll M_{ss}, M_{pp}$ ($M_{AB}$ stands for the oscillator strength of transition between the states A and B), either (a) one recombination within each path is optically unfavorable or (b) the final metastable state varies between the s and p shell, hindering the energy coincidence. In the QDM proposed here, this coincidence is likely to occur only when the two decaying paths...
share the same ground state, \( |N_{+}⟩ \) or \( |P_{+}⟩ \), as the electron tunneling matrix element exceeds few meVs.

All charged configurations in figure 4 further split after introducing a magnetic field in the Voigt geometry (transverse to the primary quantization axis, \( B_{x} \neq 0 \)). The induced energy splitting \( \Delta_{B}^{a_{i},a_{j}} \) between charged configurations \( a_{i} \) and \( a_{j} \) then relies on their Zeeman mixing renormalized by the few-body correlations. The order of this splitting under an adequately intense magnetic field (\( \gtrsim 200 \mu eV \)) is large enough compared to the dephasing linewidth of the \( \sigma^{+} \) (or \( \sigma^{-} \)) transition to impair the certainty in the phase of spin–photon pair wave function. Applying weak magnetic fields, on the other hand, could not maintain enough fidelity of the few-photon-based spin readout [30]. Note that in addition to the recombinations shown in figure 4, dark excitons \( (e_{1}h_{0} \text{ or } e_{1}h_{0}) \) become optically active in the Voigt geometry and give rise to extra cascade combinations; this is further detailed in section 6. The above criteria could be safely applied to other existing \( XX^{-} \rightarrow X^{-} \rightarrow e \) cascades and also to \( XX^{+} \rightarrow X^{+} \rightarrow h \) cascades.

Accounting only for the bright transitions, the spin–photon pair entangled states from paths \( \mathcal{P}_{1,4} \) and \( \mathcal{P}_{2,3} \) are

\[
1/\sqrt{2}(|\sigma^{+}(\omega_{1})\sigma^{+}(\omega_{4})⟩ \otimes |\downarrow⟩ + |\sigma^{-}(\omega_{4})\sigma^{-}(\omega_{1})⟩ \otimes |\uparrow⟩)
\]

and

\[
1/\sqrt{2}(|\sigma^{-}(\omega_{2})\sigma^{-}(\omega_{3})⟩ \otimes |\downarrow⟩ + |\sigma^{+}(\omega_{3})\sigma^{+}(\omega_{2})⟩ \otimes |\uparrow⟩),
\]

respectively, where \( \hbar \omega_{1}^{\pm} = \hbar \omega_{1}^{\pm} \pm \delta^{\sigma_{1},4} + \mathcal{O}_{1}(\delta_{exc}, \Delta_{B}) = \hbar \omega_{1}^{\pm} + \mathcal{O}_{2}(\delta_{exc}, \Delta_{B}) = \hbar \omega_{2}^{\pm} + \mathcal{O}_{3}(\delta_{exc}, \Delta_{B}) \), and \( \mathcal{O}(\delta_{exc}, \Delta_{B}) \) represents the energy correction due to the exchange and Zeeman splittings. As offered by figure 3(d), the hole anticrossing and thus the negative trion cascades, \( \Delta_{-}^{X} \) and \( \Delta_{-}^{X} \), are in the order of 0.9 meV (\( t_{0} < 0.45 \text{ meV} \)) at \( D_{z} = 5 \text{ nm} \). This small tunneling energy allows the hole to readily commute between the dots and render higher energy resolutions in the detuning \( \delta^{\sigma_{1},4} \) adjustment.

6. Photoluminescence spectra of charged complexes under an electric field

In this section, we present the results of CI calculations, incorporating the Coulomb direct and exchange interactions in a universal few-body Hamiltonian constructed from the single particle orbitals (see the supplementary information, available from stacks.iop.org/NJP/16/023019/mmedia). Table 1 shows the different classes of excitonic complexes studied here. The case that electron molecular orbitals were strongly localized in different dots: \( |P⟩ = |s_{e,R}⟩ \) and \( |N⟩ = |s_{e,L}⟩ \). The s-shell transitions then could be categorized as direct and indirect as shown in table 1. The indirect transitions are switched off in such a QDM hosting weakly correlated excitons. In a relatively symmetric QDM with weakly localized electrons, however, they have the chance to recombine with either left- or right-localized holes depending on the total energy of the final complex or particle. This will be detailed in the following sections.

6.1. Negatively charged exciton, \( X^{-} \rightarrow e \) transition

Neglecting the spin degree of freedom, 12 transitions could occur between the negative exciton and the single electron levels (\( 6X^{-} \times 2e \)). Among these, certain tunneling-assisted transitions
Table 1. Labels and number of s-shell configurations of multi-particle complexes studied here. All four complexes $X^-$, $X^+$, $XX^-$ and $XX^+$ have six spin-excluded components free of exchange interactions. The last two columns show the number of direct, $e_1 N_R h_\uparrow$ or $e_1 N_R h_\downarrow$ and indirect, $e_1 N_{P R} R_\perp h_\perp$ or $e_1 N_{P R} R_\parallel h_\parallel$, bright excitons existing in each complex.

| Complex          | Label | s-shell configurations (no spin) | s-shell bright channels (spin included) |
|------------------|-------|---------------------------------|----------------------------------------|
| N-trion ($X^-$)  | eeh   | 6 $X^- \rightarrow e$           | Total 24, Direct 12, Indirect 12       |
| P-trion ($X^+$)  | hhe   | 6 $X^+ \rightarrow h$           | Total 24, Direct 12, Indirect 12       |
| N-biexciton ($XX^-$) | eehh  | 6 $XX^- \rightarrow X^-$         | Total 72, Direct 36, Indirect 36       |
| P-biexciton ($XX^+$) | hhhee | 6 $XX^+ \rightarrow X^+$          | Total 72, Direct 36, Indirect 36       |

can be ruled out depending on how the single particles redistribute their orbitals according to the Coulomb interactions. The formation mechanism of $X^-$ consists of initializing one electron spin in the ground state followed by a photoexcitation creating an electron–hole pair. The favorable configuration for the two electrons is to occupy different molecular states $|P\rangle$ and $|N\rangle$ as a consequence of repulsive interaction. $|P\rangle$ and $|N\rangle$, however, remain as a strong superposition of $|s_{e,L}\rangle$ and $|s_{e,R}\rangle$ in our electric field range. The hole particle may exist either in the left or in the right QD, whether it is energetically favorable or not. For the sake of simplicity, hereafter, we represent $|s_{e,L}\rangle$ and $|s_{e,R}\rangle$ by $|L\rangle$ and $|R\rangle$, respectively.

Figure 5 illustrates the photoluminescence intensity of $X^-_{ee}$ ($\equiv X^- \rightarrow e$) transitions versus the axial electric field and the associated energies. The transition from the initial configuration $i$ to the final configuration $f$ is represented by $i_f$ and all spectral components are plotted with reference energy located at the QDM center. A relatively large energy spacing between $|P\rangle$ and $|L\rangle$ translates to two families of transitions appearing in the spectrum. Among all the features, four components around 1.155 and 1.17 eV exhibit bright photoluminescence intensity. (a) Bright transitions: $X^-_{ee}|e\rangle \equiv PN_{NR}, PNL_{NL}, PNR_{P} \text{ and } PNL_{P}$ for $E_\perp < 0.5 \text{ kV cm}^{-1}$. Note that in this representation, particles flip their localization ($R \Rightarrow L$) once the electric field is reversed, with the exception of those particles undergoing an anticrossing. We observe no pronounced distinction between the $M_{NNR}$ ($M_{PL}$) and $M_{NLL}$ ($M_{PR}$) oscillator strengths. This primarily relates to the conformity of electron configurations in the two initial states of these transitions bearing a hole anticrossing, and also to the extension of electron orbitals spreading over the entire molecule: $PN_{NR}$ and $PNL_{N}$ transitions are coupled sharing the same final state ($|N\rangle$ for $E_\perp < 0.5 \text{ kV cm}^{-1}$) and $\langle P|R\rangle \approx \langle P|L\rangle$ remains unaffected upon heavy hole tunneling.

The anticrossing energy $\Delta_{PR-PL} \approx 0.8 \text{ meV}$ at $E_\perp = 0.5 \text{ kV cm}^{-1}$ is comparable to twice the heavy hole tunneling matrix element $\sim 0.86 \text{ meV}$ (see the supplementary information, available from stacks.iop.org/NJP/16/023019/mmedia) but has been slightly renormalized by the Coulomb correlations accumulated in $X^-$. (b) Semi-bright transitions, including $NNL_{NR}$, $NRR_{NR}$, $PPL_{P}$ and $PR_{P}$ ($E_\perp > 0.5 \text{ kV cm}^{-1}$), have two electrons with the same molecular state in their initial trion. The photoexcited $X^-_{ee}$ state is then unfavorable to form, thus the oscillator strength drops significantly regardless of whether there are recombining particles resting in the same dot or not. (c) Dark transitions, such as $PR_{NR}$, exist where one electron post-tunneling is involved after recombination (not shown here). We attribute the asymmetry evident in the trion features mostly to the modest asymmetry of the electron and hole orbitals in response to oppositely oriented axial fields.
Figure 5. Left and right panels: electron and hole configurations in the bright and semi-bright spectral components of the negative trion $X^-$. Center: photoluminescence intensity of $X^-$ bright and semi-bright components plotted against the axial electric field and $X^- \rightarrow e$ transition energy. Recombining particles are colored blue. The averaged energies associated with different components are highlighted by dashed lines: blue, favorable bright configurations and white, semi-bright configurations. Squares show the region of trion-bound hole anticrossing: $\Delta_{NR-NL}^X \approx 0.8 \text{meV}$ comparable to the magnitude of hole anticrossing gap; the subscript shows the recombining excitons in each anticrossing.

6.2. Negatively charged biexciton, $XX^- \rightarrow X^-$ transition

In contrast to $X^-$, favorable $XX^-$ configurations accommodate two holes separated in different dots while one electron molecular state is fully and the other one is singly populated. Figure 6 shows the two transition families experiencing the hole anticrossings around 1.158 and 1.175 eV. Analogous to $X^-$, the bright components comprise the following recombinations: $XX^-\{eehh\} = PNNR_{PR}P_{NL}$, $PPNR_{PL}P_{NL}$, $PPNR_{PL}P_{RR}$ and $PPNR_{PL}P_{PL}$. A part of the semi-bright components originate from those configurations having two holes in the same dot, that is $XX^-\{eeeh\} = eeeR_{PR}R_{PR}$ or $eeeL_{RR}L_{RR}$, where the unlabeled electron could choose both $|P\rangle$ and $|N\rangle$ molecular states. The rest form once the fully populated electron molecular orbital of the trion state reshapes along the direction of the electric field, rather than in the opposite direction. Some features, including bright components, are subjected to sizeable energy shifts up to 8 meV as a function of the electric field. All $XX^-$ configurations comprising separated holes rise in energy once the electric field is suppressed since $J_{hh}^{LR}$ increases. This increase returns to the significant enhancement of the $\langle h_0|\hat{h}_1 \rangle$ spatial overlap despite the quick decrease in the $|R\rangle$ and $|L\rangle$ orbital densities. In those components where the two holes populate the same QD, $J_{hh}^{LL,RR}$ drops at smaller electric fields because both $\langle h_0,\hat{h}_1 \rangle (\langle h_1,\hat{h}_1 \rangle)$ and the orbital densities diminish.

We focus on the two lower-energy bright $XX^- \rightarrow X^-$ transitions previously illustrated in figure 4, $PPNR_{PL}P_{RR}$ and $PPNR_{PL}P_{PL}$. Assuming exactly identical initial configurations (spin included), the two transitions are tunnel coupled through their final trion states and separated by $\Delta_h^{X^-} = \Delta_N^{X^-}{NR-NL}$ splitting at $E_z = 0.5 \text{kV cm}^{-1}$. The highlighted energy lines
in figure 6, however, merely indicate the average energy of each component regardless of its spin fine structure. Comparing figures 5 and 6, we notice that $PPNL_{PNL}$ and $PN_{NL}$ transitions almost energetically coincide at $E_z = -5 \text{kV cm}^{-1}$ (marked by the yellow dotted line) where $PPNL_{PNL}$ and $PN_{NL}$ coincide. Neglecting the correlation energies in equation (1), $J_{hh}$ is subjected to $\sim 5 \text{meV (50.25–46.25 meV)}$ reduction in magnitude due to the $h_0 - h_1$ spatial splitting caused by a negative electric field equivalent to $5 \text{kV cm}^{-1}$. $J_{ee}$ remains almost unchanged (48.5 meV), while $J_{eh}$ and $J_{eh}$ each undergo a $-0.4$ and a 1.6 meV shift. The size of the QDM and the interdot spacing allow for less confinement along the angular momentum quantization axis $z$ compared to the previously reported single or stacked QDs \cite{21, 41}. The moderate axial confinement then leads to the smooth variation of $h-h$ interaction while $e-h$ and $e-e$ interactions stay roughly fixed. Although $|h_0\rangle$ and $|h_1\rangle$ molecular orbitals are spatially separated at such fields, their excitons are correlated via the bound electrons.

6.3. Positively charged complexes, $X^+ \rightarrow h$ and $XX^+ \rightarrow X^+$ transition

Analogous to the negative trion, formation of a positive trion consists of spin initialization and photoexcitation, but the hole spin is coherently controlled here. Favorable configurations form once the initial hole and the photogenerated hole populate different molecular states. This is implied by figure 7(a) where the bright features correspond to separated holes: $X^+\langle hhe\rangle_{h[h]} = LR_{NL}, LR_{NR} (E_z > 0), RL_{NR}$ and $RL_{NL} (E_z < 0)$. Regardless of the number of transitions, small energy corrections and the spectral redshift, we seek band bendings akin to the $XX^- \rightarrow X^-$
transitions since the energy evolution of each component versus the electric field is primarily attributed to the changes in $J_{hh}^{LR}$ ($J_{hh}^{LL}$ or $J_{hh}^{RR}$ in semi-bright features). In contrast to $XX^-$ and $X^+$, $XX^+$ contains three interacting hole particles, complicating the photoluminescence spectrum. The highly favorable configurations develop with segregated electrons; however, the $e-e$ repulsion does not severely influence the formation of pair electrons in our axial field range, because the energy difference between $P_L - P_U$ (or $N_L - N_U$) and $P_U - N_L$ repulsions is insignificant. Transitions in figure 7(b) can be divided into two classes based on their final trion state: (a) components in which the hole particles in the final state live in the same dot. These transitions are more responsive toward $h-h$ repulsion compared to (b) components with segregated holes in their final trion, thus bearing a higher energy shift at larger fields. No features from the $XX^+ \rightarrow X^+$ and $X^+ \rightarrow h$ spectra coincide due to the relatively large hybridization energy of the extra electron living in $XX^+$. This signifies that multi-partite energy matching via both negatively and positively charged complexes is, in general, feasible in QDMs where electron and hole tunneling matrix elements are tantamount [42].

In the final part of this section, let us concisely address the case of the [001] nanowire replaced with its [111] counterpart. The strain-driven asymmetry of single particle states seen above, and also the asymmetry of transition energies versus the axial electric field are more pronounced in [111]-oriented double QDs as a result of the built-in piezoelectric field [43]. This internal axial piezoelectric field separates $|h_0\rangle$ and $|h_1\rangle$ molecular orbitals even at $E_z \approx 0$, further splitting the locations of electron and hole anticrossings at low electric fields. The advantage of [111]-oriented quantization, however, is that orbitals save their $C_{3v}$ symmetry irrespective of

Figure 7. Photoluminescence intensity depicted for (a) positive trion $X^+$ and (b) positive biexciton $XX^+$ against the axial electric field and transition energies. Bright components in the $X^+$ spectrum form with spatially separated holes. The $XX^+ \rightarrow X^+$ spectrum is, on average, 20 meV blueshifted with respect to the $X^+ \rightarrow h$ bright transitions due to the electron hybridization energy. Bright components of $XX^+$ include initial configurations with separated electrons and final configurations $X^+$ with separated holes: $LLNP_{LRN}$, $LRRNP_{LRN}$, $LLRP_{LRP}$ and $LRRNP_{LRP}$. Semi-bright transitions in $XX^+ \rightarrow X^+$ comprise either two neighboring electrons in the initial state or two neighboring holes in the final state. The energy splitting between the $RLNP_L$ and $RLNP_R$ ($LRNP_L$ and $LRNP_R$) components at small fields primarily correspond to hole anticrossing in their final states.
the electric field amplitude. As mentioned above, the symmetry conversion from $D_{2d}$ down to $C_{2v}$ upon applying the axial electric field on [001] QDMs lifts the bright exciton degeneracies. In contrast, the elevated $C_{3v}$ symmetry predicted in [111] QDMs is immune to the symmetry-breaking Stark effect.

7. Analytical Hamiltonian: effect of multi-particle interactions on the spin fine structures

In this section, we develop a simplified version of the $XX^-$ and $X^-$ Hamiltonians to comprehend the effect of Coulomb interactions on the spin-conserved tunnelings. Providing an analytical model helps us to visualize the behavior of the $XX^-$ and $X^-$ spin fine structure versus the electric field, without getting involved with the details of the CI method. We confine our study to the favorable configurations of $XX^-$ and $X^-$ shown in figure 4. For simplicity, we consider the $e–h$ exchange interactions of single particles residing in different molecular states to be equal, i.e. $\delta_{\text{exc}}^{N_L} = \delta_{\text{exc}}^{PR} = \delta_{\text{exc}}^{NR} = \delta_{\text{exc}}^{PL} = \delta_{\text{exc}}^{eh}$. This assumption is valid within our range of electric fields where electron and hole molecular orbitals consistently keep their overlap integral and associated mean field potentials rather unchanged. The $4 \times 4$ few-body Hamiltonian describing the nondegenerate bright configurations of the negative trion $X_{\text{eh}}=P_{Nh}$ could be expanded on basis states $N_i P_j L_{\parallel}, N_i P_j R_{\parallel}, N_i P_j L_{\parallel}$ and $N_i P_j R_{\parallel}$:

$$
\mathcal{H}_{X^-}^{\text{B}} = 
\begin{pmatrix}
\zeta_L^{X^-} & -t_h & 0 & 0 \\
-t_h & \zeta_L^{X^-} + \Delta J_X^+ - \zeta_E & 0 & 0 \\
0 & 0 & \zeta_L^{X^-} + 2\delta_{\text{exc}} & -t_h \\
0 & 0 & -t_h & \zeta_L^{X^-} + \Delta J_X^+ - \zeta_E & 2\delta_{\text{exc}}
\end{pmatrix}.
$$

(4)

Here, $\zeta_L^{X^-}$ is the total energy of $N_i P_j L_{\parallel}$, $\zeta_E$ signifies the energy shift caused by the electric field when the reference energy is located at the center of the left QD, $\zeta_L = \bar{\sigma} E_z (D_s + h_D)$, $\bar{\sigma}$ is the unit charge and $\Delta J_X^+ = J_{\text{PR}}^+ + J_{\text{PL}}^+ - J_{\text{NR}}^+ - J_{\text{eh}}^+$. The slight difference between s-shell molecular orbitals leads to nonzero $\Delta J_X^+$. The mean value of the $e–h$ exchange energy $\delta_{\text{exc}}^{eh}$ is calculated by averaging over $\langle \psi_{\uparrow} \psi_{\downarrow} | C | \psi_{\downarrow} \psi_{\downarrow} \rangle - \langle \psi_{\uparrow} \psi_{\downarrow} | C | \psi_{\downarrow} \psi_{\uparrow} \rangle$ versus $E_z$, where $C$ is the Coulomb interaction operator.

We notice that the bright states $N_i P_j L_{\parallel} (\equiv N_i P_j L_{\parallel}) \equiv N_i P_j L_{\parallel}$ and $N_i P_j R_{\parallel} (\equiv N_i P_j R_{\parallel}) \equiv N_i P_j R_{\parallel}$ are fourfold degenerate, while $N_i P_j L_{\parallel} (\equiv N_i P_j L_{\parallel})$ and $N_i P_j R_{\parallel} (\equiv N_i P_j R_{\parallel})$ are twofold degenerate as long as $\delta_{\text{exc}}^{NL} = \delta_{\text{exc}}^{PR} = \delta_{\text{exc}}^{NR} = \delta_{\text{exc}}^{PL}$ and the trivial anisotropic exchange interactions are discarded. These assumptions are not necessarily valid in a strained QDM as could be revealed by our CI calculations, but facilitate the semi-quantitative description of the exchange couplings. The total energy spectrum of $X_{\text{eh}}=P_{Nh}$ as a function of the axial electric field is plotted in figure 8(a), where the direct Coulomb interactions are artificially discarded, $\Delta J_X^+ = 0$, and figure 8(b) where the states are Coulomb correlated, $\Delta J_X^+ \neq 0$. The quadruplets labeled by $N_i P_j L_{\parallel}$ and $N_i P_j R_{\parallel}$ (analogous to the doublets labeled by $N_i P_j L_{\parallel}$ and $N_i P_j R_{\parallel}$) evolve to resonance where the anticrossing caused by the hole tunneling appears, $\Delta \delta_{\text{exc}}^{eh} = 2\sqrt{t_h^2 + (2\delta_{\text{exc}}^{eh})^2} \approx 892 \mu eV$ having $2\delta_{\text{exc}}^{eh} = 122$ and $t_h = 430 \mu eV$. Relatively trivial variations of $\Delta J_X^+$, shown in the inset of figure 8(b), are a consequence of
Figure 8. (a) Energy level diagram of the irreducible trion basis set plotted with respect to the reference energy $\zeta_{X^-}$ as a function of the axial electric field once the direct Coulomb interactions are switched off. $N_1P_1L_R$ and $N_1P_1R_R$ ($N_1P_1L_L$ and $N_1P_1R_L$) are tunnel coupled, $\Delta J_{X^-} \simeq 892, 2\delta_{exc}^{eh} = 122 \mu eV$. (b) $\Delta J_{X} \neq 0$ (inset), but the spin fine structure exhibits a similar pattern to (a) because the attractive interactions are comparable and evolve smoothly as a function of the electric field. (c) Energy state diagram of the negative biexciton basis set calculated with respect to $\zeta_{XX^-}$ having the Coulomb interactions discarded. $\Delta J_{XX^-} \simeq 1.24 \text{meV}$, $N_1P_1P_1L_R$ and $N_1P_1P_1L_L$ are each separated from the $N_1P_1P_1L_R R_R-N_1P_1P_1L_L R_R$ doublet by $2\delta_{exc}^{eh}$ ($\zeta_{exc}^{XX^-} = 4\delta_{exc}^{eh}$). The inset illustrates the situation where the $\delta_{exc}^{RR}$ and $\delta_{exc}^{PL}$ interactions are assumed to be weak. The violet lines show the new energies of the quadruplet with $N_1P_1P_1L_R$ and $N_1P_1P_1L_L$ crossing. (d) Coulomb interactions are switched on, $\Delta J_{1^{XX^-}} \neq \zeta_{L}^{XX^-}$ and $\Delta J_{2^{XX^-}} \neq \zeta_{L}^{XX^-}$. A noticeable change in $\Delta J_{1^{XX^-}} \simeq J_{hh}^{LR} - J_{hh}^{LL}$ decouples the quadruplet from the $N_1P_1P_1L_R R_R$ and $N_1P_1P_1L_L R_R$ singlets. Inset: evolution of attractive and repulsive interactions as a function of the axial electric field.

electron delocalization and comparable direct and indirect $e-h$ interactions, thus $\Delta h_{X^-}$ is not noticeably renormalized.

The negative biexciton includes two hole particles, thus the spin-conserved tunneling of the hole is restricted by the Pauli exclusion principle. This spin blockade leads to six $XX_{eehh}^{xx-} = N_1P_1P_1hh$ doublets considering the twofold spin degeneracy of each electron. In order to resolve the corresponding energy state diagram, we consider the basis states
\[ N_1 P_1 L_1 R_1 (\equiv N_1 P_1 L_1 R_1), \quad N_1 P_1 L_1 R_1 (\equiv N_1 P_1 L_1 R_1), \quad N_1 P_1 L_1 L_2 R_1 (\equiv N_1 P_1 L_1 L_2 R_1), \quad N_1 P_1 L_1 R_1 (\equiv N_1 P_1 L_1 R_1), \quad N_1 P_1 L_1 L_2 R_1 (\equiv N_1 P_1 L_1 L_2 R_1), \quad N_1 P_1 L_1 R_1 (\equiv N_1 P_1 L_1 R_1), \quad N_1 P_1 L_1 L_2 R_1 (\equiv N_1 P_1 L_1 R_1), \quad N_1 P_1 L_1 L_2 R_1 (\equiv N_1 P_1 L_1 L_2 R_1), \quad \text{and} \quad N_1 P_1 L_1 L_2 L_3 R_1 (\equiv N_1 P_1 L_1 L_2 L_3 R_1), \]

then construct the 6 × 6 few-body Hamiltonian \( \mathcal{H}_B^{XX} \) as

\[
\begin{pmatrix}
\Delta J_1^{XX} - \zeta_{E_1} & 0 & 0 & -t_h & -t_h \\
0 & \Delta J_1^{XX} - \zeta_{E_1} & 0 & 0 & -t_h \\
0 & 0 & \Delta J_1^{XX} - \zeta_{E_1} & 0 & 0 \\
-t_h & -t_h & 0 & \Delta J_2^{XX} - \zeta_{E_1} & 0 \\
-t_h & -t_h & 0 & 0 & \zeta_{E_1}^{XX} \\
\end{pmatrix}, \tag{5}
\]

where \( \zeta_{E_1}^{XX} \) is the total energy of \( N_1 P_1 L_1 L_2 R_1 \). Assuming that the direct and indirect interactions are comparable, \( \Delta J_1^{XX} = \zeta_{E_1}^{XX} \) and \( \Delta J_2^{XX} = \zeta_{E_1}^{XX} \), we observe the singlet states \( N_1 P_1 L_1 L_2 L_3 R_1 \) undergoing an anticrossing with an energy gap equivalent to

\[
\Delta E_{XX} = 2\sqrt{2t_h^2 + (2\delta_{exc})^2} \approx 1.24 \text{ meV} \; \text{; see figure 8(c). The two singlets, however, first couple to} \quad N_1 P_1 L_1 L_2 R_1 \quad \text{or} \quad N_1 P_1 L_1 L_2 R_1 \quad \text{rather than undergoing a direct anticrossing since the two-hole tunneling is prohibited in our model. The triplet states} \quad N_1 P_1 L_1 L_2 R_1 \quad \text{and} \quad N_1 P_1 L_1 R_1 \quad \text{are separated by} \quad \zeta_{XX}^{ee} = 4\delta_{exc} \quad \text{and pass through the anticrossing region without tunnel coupling to the singlet states, because the spin state is conserved during the tunneling. The remaining states,} \quad N_1 P_1 L_1 L_2 R_1 \quad \text{and} \quad N_1 P_1 L_1 R_1 \quad \text{form a singlet–triplet pair in proximity to the anticrossing. Their degeneracy seen in figure 8(c) returns back to the assumption of all exchange interactions being equivalent. Beyond the thin barrier limit,} \delta_{exc} \quad \text{exchange interaction is negligible, thus} \quad N_1 P_1 L_1 L_2 R_1 \quad \text{and} \quad N_1 P_1 L_1 R_1 \quad \text{split by} \quad \zeta_{XX}^{ee} = 2\delta_{exc} \quad \text{as illustrated in the inset of figure 8(c).}
\]

In figure 8(d), the energy state diagram of \( XX^{ee}hh = N_1 P_1 P_1 hh \) is depicted against the electric field, taking the Coulomb interactions into account (\( \Delta J_1^{XX} \neq \zeta_{E_1}^{XX} \) and \( \Delta J_2^{XX} \neq \zeta_{E_1}^{XX} \)). \( h-h \) repulsive interactions significantly change as a function of axial localization, leading to large variations in \( J_{hh}^{LL}, J_{hh}^{LR}, J_{hh}^{RL}, \text{ and } \Delta J_1^{XX} \) versus the electric field as shown in the inset. The Coulomb splitting drastically renormalizes the energy levels of the quadruplet \( N_1 P_1 L_1 L_2 R_1, N_1 P_1 L_1 L_2 R_1, N_1 P_1 L_1 L_2 R_1, N_1 P_1 L_1 L_2 R_1, \text{ and } \), and decouples the hole spin singlets, \( N_1 P_1 L_1 L_2 R_1 \), from \( N_1 P_1 L_1 L_2 R_1 \), in the vicinity of the anticrossing gap. The splitting can be suppressed by intermixing the dot-barrier materials and thus delocalizing the \( |h_0 \rangle \) and \( |h_1 \rangle \) molecular orbitals. Since the negative trion states are rather insensitive to Coulomb interactions, the anticrossings visible in the \( N_1 P_1 P_1 hh \text{ transitions are mainly due to the tunnelcoupling of final states. Extending the same principles to positively charged complexes, the spin fine structure of the positive trion } X^+ \text{ should exhibit a similar pattern to } XX^- \text{ as long as the hole tunneling is concerned. The presence of three holes in the positive biexciton } XX^+, \text{ however, further complicates the effective Hamiltonian. We do not study the total energy of } XX^+ \text{ here, but we predict observing trivial decouplings because the } h-h \text{ repulsions effectively cancel each other out, i.e.} J_{hh}^{LL} + 2J_{hh}^{RL} - J_{hh}^{RR} - 2J_{hh}^{LR} \lesssim 2t_h.}
8. Cascade transitions under a transverse magnetic field

The CI method incorporates both isotropic and anisotropic e−h exchange interactions 
\[ \delta_{\text{exc}}^e = -\beta_{\text{exc}}^e \sigma_e \cdot J_h, \]
where \( \beta_{\text{exc}}^e \) is the coupling coefficient and \( J_h \) is the hole angular momentum operator. \( \beta_{\text{exc}}^e \) coefficients of mutual e−h exchange interactions are calculated using
\[ \langle \psi_{\sigma_e}^e | \chi_{\sigma_h}^h | \psi_{\sigma_e}^e \rangle \]
where \( V_{\sigma_e,\sigma_h}^e \) is the mean field potential caused by the mixed orbital \( \langle \psi_{\sigma_e}^e | \psi_{\sigma_e}^h \rangle \) [44]. Upon applying a transverse (nonaxial) magnetic field in the Voigt geometry, the magnetic field-induced coupling terms between the bright and dark excitons appear, enhancing the energy splittings and providing access to the desirable spin states. The off-diagonal mixing matrix elements \( \langle \alpha_i | \mathcal{H}_{ij}^B | \alpha_j \rangle \) between two arbitrary configurations \( \alpha_i \) and \( \alpha_j \) is given by
\[ \mathcal{H}_{B}^{ij} = \frac{i n_x}{2} g_{\text{eff}} B \cdot \sigma_{ij}, \]
where \( g_{\text{eff}} \) stands for the effective transverse g-factor at a particular axial electric field and \( \sigma_{ij} \) is the spin flip operator associated with configurations \( \alpha_i \) and \( \alpha_j \). The dark-bright mixing gives rise to a nonzero many-body oscillator strength \( \delta_{\text{osc}} \) of the dark states, introducing extra optically active decay in proximity to the unperturbed transitions. The activation of dark transitions may lead to the emergence of unwanted spectral features and several \( \Lambda \) systems [45], comprising metastable electron and hole ground states with unwanted spin configurations. Therefore, they shall be eliminated from the spectral window before being subjected to any additional post-processing.

One striking feature in the energy level diagram of QDMs, as compared to single QDs, is the existence of available bright channels leading to both \( \langle e_{\uparrow} | \psi \rangle \) and \( \langle e_{\downarrow} | \psi \rangle \) electron final states. Hence, the requisite for activating dark states via a transverse magnetic field is basically eliminated, and the Faraday configuration (parallel to the strong quantization axis) could also be employed for the spin control or tuning the charged complexes. Our single particle calculations based on the gauge invariant discretization method [31] show that due to the orbital resemblance, the hole effective g-factors are comparable for the low-lying s-shells, that is \( g_{|s_h,\perp>}^x = 0.262 \) and \( g_{|s_h,\perp>}^x = 0.258 \), and smaller than the electron effective g-factors, \( g_{|s_e,n>}^x = -1.08 \) and \( g_{|s_e,\perp>}^x = -1.042 \), at \( E_z = -5 \text{ kV cm}^{-1} \). We comment that strong localization of hole orbitals under an electric field results in smaller g-factors and less Zeeman splitting of charged states. The reason lies in the fact that the g-factor can be explained as the measure of deformation the orbital undergoes in response to the external magnetic field. Once the hole orbital becomes squeezed in the real space, its Fourier transform spreads out in \( k \)-space. The hole particle then gains further kinetic energy if its orbital reshapes. The reluctance to exchange the kinetic energy prohibits the orbital from deforming by the magnetic field and the g-factor drops. Therefore, although excitons are fully entangled in the vicinity of the anticrossing, they suffer from larger Zeeman shifts under the magnetic field.

Figures 9(a) and (b) illustrate the energy spectrum of \( XX^- \) and \( X^- \) for different spin configurations in \( NPPLR \) and \( NPL \) components, respectively, as a function of the transverse magnetic field \( B_x \) at \( E_z = -5 \text{ kV cm}^{-1} \). We selected three exemplary configurations from the \( XX^- \) energy state spectrum comprising two bright recombinations with electrons localized in \( |P\rangle \). Transitions labeled as \( \sigma_{5\uparrow}^1, \sigma_{5\downarrow}^1 \) and \( \sigma_{5\downarrow}^2 \) correspond to the bright \( \langle P | R \rangle \) recombinations. The oscillator strength of \( XX^-\{NPPLR\}X^-\{NPL\} \) transitions is given in figure 9(c). The anticrossings visible at low magnetic fields correspond to the coupling to dark trion features or the bright−bright trion mixing; for example, \( \delta_{\text{osc}}^e = 11 \mu eV \) is the hole anticrossing energy between \( N_1 P_1 R_R \) and \( N_1 P_1 R_R \) states. Such anticrossing behavior changes the dominant character of Zeeman-coupled spin configurations in the course of magnetic field variations. The measure of anticrossing energy pertains to the effective g-factor of the coupled spin states.
Figure 9. (a), (e) Negative biexciton NPPLR and (b), (f) negative trion NPL and NPR energies versus the [100]-oriented magnetic field at $E_z = -5 \text{kV cm}^{-1}$. Spin configurations of $X^-$ and $XX^-$ are exchange and Zeeman coupled, and the dominant character attributed to each configuration at $B_x = 3 \text{T}$ is indicated at the right side of each panel. Configurations with all the spins flipped are approximately degenerate at $B_x = 0$. Intermediate trions configured in NPL and NPR form following $\langle P_\sigma^e | R_\sigma^h \rangle$ and $\langle P_\sigma^e | L_\sigma^h \rangle$ recombinations, respectively. Three pairs of $XX^- X$ bright transitions exhibiting large enough oscillator strengths are selected: $\sigma_1^+ (\sigma_2^-)$, $\sigma_2^+ (\sigma_3^-)$ and $\sigma_3^- (\sigma_4^+)$, as shown in (c) and (g). The subsequent $X^- e$ transitions, (d) and (h), are labeled by $\tilde{\sigma}_1^+ (\tilde{\sigma}_2^-)$, $\tilde{\sigma}_2^+ (\tilde{\sigma}_3^-)$ and $\tilde{\sigma}_3^- (\tilde{\sigma}_4^+)$ and correspond to the polarized photons shown in figure 4. The energies of final electron states $\{N_\uparrow, N_\downarrow\}$ are not shown here. Horizontal arrows in (c), (d) and (g), (h) mark dark states gaining noticeable oscillator strengths upon Zeeman mixing.
The primary character of each spin configuration at $B_x = 3$ T is indicated in figures 9(a), (b), (e) and (f).

The asymmetric character of s-shell molecular orbitals is revealed comparing figures 9(b) and (f), where localization of the trion-bound hole is slightly altered but the effective $g$-factors of trions are different. Similar to figure 9(c), the oscillator strengths of the trions are shown in figure 9(d), labeled by $\tilde{\bar{\sigma}}_\alpha^\pm$, $\bar{\sigma}_\alpha^\pm$, and $\tilde{\sigma}_\alpha^\pm$. Compared to $\sigma_{\alpha=1,3}^\pm$ ($\sigma_{\alpha}^\pm$), the trion transitions $\tilde{\bar{\sigma}}_{\alpha=1,3}^\pm$ undergo a relatively smaller energy change versus $B_x$. The maximum energy variation observed for each individual transition, however, does not exceed 200 $\mu$eV under this magnetic field range $B_x < 3$ T. The right panel of figure 9 depicts the same set of data for $XX^-\{NPPLR\}X^-\{NPR\}$ transitions: the initial $XX^-$ states are identical but the intermediate trion states emerge following the $\langle P|L \rangle$ recombination, and in addition the polarization of the emitted photon is reversed. The spectral dispersion through coupling to multiple states is considerably suppressed in both the $XX^-X^-e$ and $X^-X^-e$ spectra compared to the alternative path due to the lower number of crossings (anticrossings) seen in the NPR trion.

9. Photon pair concurrence

Full analysis of the spin–photon pair entanglement should be performed in an experimental framework or via a complete mathematical model incorporating the timing details and spin dynamics. An explicit parameter to incorporate the energy and linewidth of two arbitrary transitions in the QDM spectrum is the photon-pair concurrence. In our particular case, the tripartite concurrence could be estimated once the spin dynamics are known [46]. Here, we exclusively analyze the photon indistinguishability through calculating the concurrence $C$, which is ideally $C = 0$ for separable states and $C = 1$ for the maximally entangled states in the regular scheme, but never reaches above 0.73 in the time reordering scheme (cross-entanglement). In order to verify the entanglement between the photon pair and the third particle (electron spin), their correlations in the current spin-polarization basis as well as the rotated basis shall be measured. Previous observation of these correlations are reported for the spin-single photon entanglement [47]. In the following, we narrow down our model to the cross-entanglement scheme. The findings are then extensible to the regular scheme.

According to figure 4, the concurrence of photon pairs relies upon $\Gamma_{XX^-\rightarrow X^-}^\lambda$, $\Gamma_{X^-\rightarrow e}^\lambda$, $\lambda \in \{\sigma^+, \sigma^-\}$, and the normalized detuning $\Delta^\lambda_\alpha = [\delta^\lambda_{i,i'} + O(\delta_{\text{exc}}, \Delta_B)] / 2 \Gamma_{X^-\rightarrow e}^\lambda$, where $\Gamma_{\alpha_1\rightarrow \alpha_i}$ stands for half the spontaneous emission rate of the recombining exciton. The concurrence drops with increasing $\Gamma_{XX^-\rightarrow X^-}^\lambda / \Gamma_{X^-\rightarrow e}$ and $\Delta^\lambda_\alpha$ [35]; see supplementary information (available from stacks.iop.org/NJP/16/023019/mmedia). Both of these parameters can be optimized via the frequency conversion technique to maximize the concurrence, because the process leads to frequency and linewidth conversion of the initial $XX^-X^-$ and $X^-X^-$ photons. Assuming that the dephasing linewidth could become as narrow as 30 $\mu$eV in defect-free nanowire QDs [48], the normalized detuning drops significantly subsequent to the frequency conversion process, which is able to enhance $\Gamma$ even by one order of magnitude [47].

The above equalities relate to the photon emission rates proportional to $\langle XX^-X^-|H_{\text{em}}|X^-\rangle$ and $\langle X^-|H_{\text{em}}|e\rangle$. Here, $H_{\text{em}}$ is the coupling Hamiltonian to the optical modes $l$, $H_{\text{em}} = \sum_{l,i;\lambda} g_{l,\lambda}^i \hat{a}^\dagger_{l,\lambda} \hat{b}_i + \text{h.c.}$ where $g_{l,\lambda}^i$ is the oscillator strength of transition $c = \{XX^- \rightarrow X^-\rightarrow e\}$, $\hat{b}_\alpha^\dagger = \hat{h}_\alpha \hat{c}_\dagger$, $\hat{b}_\alpha = \hat{h}_\beta \hat{c}$, and $\hat{a}_{l,\lambda}^\dagger$ creates a photon in $l$th optical mode with polarization $\lambda$. 

20
Figure 10. (a), (d) Energy of photon pairs created across generations in cascade paths \( P_1 \) and \( P_4 \), \( P_2 \) and \( P_3 \). The respective linewidths of photons before frequency conversion are illustrated in a sense that upper transitions possess less dephasing lifetime. The frequency of each transition is labeled according to the energy diagrams shown in (c) and (f). (b) Concurrence of \( P_1 - P_4 \) cascades versus magnetic field and \( \gamma'_{\sigma_1} = 2 \). The inset shows concurrence as a function of magnetic field for \( B_x = 3 \) T. (e) Concurrence of photon pairs in \( P_2 - P_3 \) cascades; \( \gamma'_{\sigma_1} = 2 \). Inset: concurrence versus magnetic field for \( B_x = 2 \) T and 3 T.

Figures 10(a) and (d) depict the energy evolution of photons created in paths \( P_1 \) and \( P_4 \) (\( P_2 \) and \( P_3 \)) as a function of the transverse magnetic field. For the sake of simplicity, we assume that the average linewidth of upper transitions in figure 4 after frequency conversion is twice the linewidth of lower ones: 

\[
\gamma'_{\sigma} = \frac{\Gamma'_{XX^- \rightarrow X^-}}{\Gamma'_{X^- \rightarrow e}} \approx 2; \quad \Gamma'_{\sigma_1} = \Gamma'_{\sigma_4} = 2\Gamma'_{\sigma_2} = 2\Gamma'_{\sigma_3}.
\]

Although this ratio is independent of the dephasing linewidth and primarily relies on the time resolution of the laser source pumping the PPLN, our assumption does not pose any constraint to the generality of results. We further assume that the frequencies of single photons are downconverted with the same ratio, i.e. \( \omega'_i = 1.2.3.4 = \eta \omega_i = 1.2.3.4 \). The concurrence of the photon pairs in paths \( P_1 \) and \( P_4 \) is then given by [35, 49]

\[
C = \frac{4}{\pi^2} \int \int \frac{\Gamma_{\sigma_1} \Gamma_{\sigma_4} W_{\eta}(\omega_m, \omega_n) \Gamma_{\sigma_2} \Gamma_{\sigma_3}}{(\omega_m + \omega_n - \Omega_{4;XX^-} - i\Gamma'_{\sigma_1})((\omega_m + \omega_n - \Omega_{4;XX^-} + i\Gamma'_{\sigma_2})W_{\eta}(\omega_m, \omega_n))} \times \frac{d\omega_m}{\omega_m - \omega_3;X^- - i\Gamma'_{\sigma_2}} \frac{d\omega_n}{(\omega_m - \omega_4;X^- - i\Gamma'_{\sigma_1})},
\]

(8)
where $\Omega_{1,XX^e} = \omega_1' + \omega_3'$ and $\Omega_{4,XX^e} = \omega_2' + \omega_4'$; see figure 10(c). The simplest additional phase above $W_n'(\omega_m, \omega_n)$ can be a linear phase with time delay $\tau$, that is $W_n'(\omega_m, \omega_n) = \exp[i(\omega_m - \omega_n)\tau]$. Since the detuning between the two paths $\mathcal{P}_1$ and $\mathcal{P}_4$, $\Delta_\omega = |\omega_1' - \omega_4'| + |\omega_2' - \omega_1'|$, is constantly nonzero in our setup, the time delay suggested by Pathak and Hughes, $\tau_o = \text{ln}(1 + \Gamma_{XX^e \rightarrow X^-_1}^\prime)/2\Gamma_{X_1^- \rightarrow e}/\Gamma_{XX^e \rightarrow X^-_1}$, does not necessarily optimize the concurrence here. The optimum $\tau_o$, however, could be resolved empirically once the actual level broadenings are determined.

The concurrence of photon pairs versus the magnetic field and $\Gamma_{XX^e \rightarrow X^-_1}$ is plotted in figure 10(b). The magnetic field magnitude is set above 2 T where the spin initialization and readout are experimentally feasible. The transition oscillator strengths are comparable and steady in this range, permitting the detuning $\Delta_\omega$ to be the only parameter restraining the concurrence. According to the continuous increase in $|\omega_1' - \omega_4'|$ splitting, the local extremum occurs at $\Delta_{\text{min}}^\omega$ at which $\sigma_1^-$ and $\sigma_\downarrow$ coincide. The inset shows how noticeably the concurrence is improved once $\Gamma_{XX^e \rightarrow X^-_1}$ increases from 100 up to 200 $\mu$eV. Figure 10(e) depicts the same plot for the concurrence of photon states across generations in $\mathcal{P}_2$ and $\mathcal{P}_3$. Energy detunings $|\omega_4' - \omega_1'|$ and $|\omega_2' - \omega_4'|$ evolve oppositely versus the magnetic field, thus the concurrence remains insensitive toward its variations, as shown in the inset of figure 10(e).

The above example demonstrates that level broadenings up to half the photon–photon detuning energy ($\sim200\mu$eV) could improve the cascade concurrence considerably. On the other hand, exerting the magnetic field in the Faraday configuration would keep the dark states inactive and conserve the bright transition oscillator strengths undispersed by prohibiting any spin-flip mixing. Coherent control of the spin rotation may then be more favorable in Faraday configuration owing to its clean emission spectrum. We note that intermediate exchange-coupled trion states are assumed to be immune to the single spin flip, e.g. $N_1 \rightarrow N_1 P_1 L^\perp \rightarrow N_1 P_1 L^\perp$, or the cross-dephasing process, e.g. $N_1 \rightarrow N_1 P_1 L^\perp \rightarrow N_1 P_1 L^\perp$, during the excitation cycle. Principally, weak axial quantization of molecular orbitals gives rise to smaller spin relaxation times of the exciton-bound electron and hole [50, 51]. The hole molecular orbitals are squeezed at a higher electric field here, but the electron orbitals preserve a fixed volume regardless of the electric field magnitude.

Apart from the photons across generations in figure 4, entanglement between the photons within generations could also be established via the frequency conversion technique. The first emitted photons in paths $\mathcal{P}_1$ and $\mathcal{P}_3$ (or second emitted photons in $\mathcal{P}_2$ and $\mathcal{P}_4$) are in fact energetically separated only by the measure of exchange and Zeeman splittings. This implies that the charged biexciton cascades offer several path options to produce tripartite entangled states. In addition, we notice that the level of concurrence calculated here delicately depends on $\gamma_p$. Without applying the frequency conversion technique, $\gamma_p$ represents a relative parameter comparing the natural linewidths of $XX^e \rightarrow X^-_1$ and $X^-_1 \rightarrow e$ transitions. Therefore, the concurrence is essentially untouched and determined merely by the QDM structure. Upon a frequency conversion, each $\Gamma_{XX^e \rightarrow X^-_1}$ and $\Gamma_{X^-_1 \rightarrow e}$ could be manipulated individually, leading to a tunable range of $\gamma_p$. We predict that higher values of concurrence are obtainable by tailoring the $XX^e$ and $X^-_1$ level broadenings with different scales as concurrence grows monotonously once $\Gamma_{XX^e \rightarrow X^-_1}/\Gamma_{X^-_1 \rightarrow e}$ decreases [35].

Finally, we comment on how a QDM could be exploited to generate GHZ or $W$ entangled states of photons. The existence of two low-lying $s$-shells in the energy level spectrum of QDM leaves room for three correlated excitons to form a triexciton $XXX$ under proper
pumping conditions. Sequential decay of \( XXX \) down to the ground state, \( XXX \rightarrow XX \rightarrow X \rightarrow G \), create three correlated photons whose color indistinguishability could be manipulated via the frequency conversion technique. Different configurations of initial and intermediate bright states then prepare a set of decay paths comprising diverse combinations of photon polarizations \( |\sigma_{1}^{\pm} \sigma_{2}^{\pm} \sigma_{3}^{\pm} \rangle \). A two-path combination might be able to produce a GHZ state, such as \( \frac{1}{\sqrt{2}}( |\sigma_{1}^{+} \sigma_{2}^{+} \sigma_{3}^{+} \rangle + |\sigma_{1}^{-} \sigma_{2}^{-} \sigma_{3}^{-} \rangle ) \), while a three-path combination might give rise to a W-state. We notice that the color distinguishability in a spin-free state only counts on direct and exchange energies as no external magnetic field is involved. Provided that the combinations at each level, \( XXX \) or \( XX \) or \( X \), are chosen from spin-flipped configurations, the color distinguishability pertains only to exchange energies which are typically limited as compared to the Zeeman splitting.

10. Summary

We investigated the feasibility of producing tripartite spin–photon pair entanglement from charged biexcitons in QDMs. In the proposed structure, two QDs are coupled sharing their s-shell ground states with a relatively small hybridization energy of the hole particle (\( t_h < 0.5 \) meV) that facilitates switching between different configurations without the extensive cost of tunneling energies. We particularly analyzed the case of In(Ga)As stacked QDs embedded in [001]-oriented GaAs nanowires. However, the results can be generalized to other types of QDMs sustaining strongly correlated orbitals. We showed that the weak quantization of the electron opens new channels of recombination for higher order entanglement. Our calculations in the few-body framework revealed that charged biexcitons and excitons exhibit dominant spectral features as a direct consequence of existing mutual interactions. The variety of available transitions is then narrowed down to a few favorable initial and final states. Neglecting the small exchange splittings, \( XX^- \rightarrow X^- \rightarrow e \) cascades represent relative energy matching under an axial electric field, while \( XX^+ \rightarrow X^+ \rightarrow h \) channels lack any color coincidence under our range of fields. This is primarily attributed to the large electron hybridization energy. By providing an exemplary cascade of \( XX^- \rightarrow X^- \rightarrow e \) transitions, we examined the double-dot spin fine structure and Zeeman shifts under the magnetic field in the Voigt geometry. The strong quantization of the hole particle away from the anticrossing lead to its small transverse \( g \)-factor, thus charged complexes undergo less Zeeman shift. Upon the empirically approved range of magnetic field required for the spin manipulation (\( B < 3 \) T), photon energy detunings were calculated below 400 \( \mu \)eV. Sufficient photon concurrence is then achievable by manipulating the level broadenings through the frequency conversion technique. Our analysis is a demonstration of how properly sorted multi-particle states could be exploited to create multi-partite entangled states in an engineerable solid state source.

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24
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