Peculiar many-body effects revealed in the spectroscopy of highly charged quantum dots

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Coulomb interactions between electrons lead to the observed multiplet structure and breakdown of the Aufbau principle for atomic d and f shells¹. Nevertheless, these effects can disappear in extended systems. For instance, the multiplet structure of atomic carbon is not a feature of graphite or diamond. A quantum dot is an extended system containing ~106 atoms for which electron-electron interactions do survive and the interplay between the Coulomb energy, J, and the quantization energy, ΔE , is crucial to Coulomb blockade²⁻⁵. We have discovered consequences of Coulomb interactions in selfassembled quantum dots by interpreting experimental spectra with an atomistic calculation. The Coulomb effects, evident in the photon emission process, are tunable in situ by controlling the quantum dot charge from +6e to -6e. The same dot shows two regimes: $J \leq \Delta E$ for electron charging yet $J \simeq \Delta E$ for hole charging. We find a breakdown of the Aufbau principle for holes; clear proof of non-perturbative hole-hole interactions; promotion-demotion processes in the final state of the emission process, effects first predicted a decade ago6; and pronounced configuration hybridizations in the initial state. The level of charge control and the energy scales result in Coulomb effects with no obvious analogues in atomic physics.

We use InAs/GaAs dots (bound s,p and d orbitals) in tunnel contact with an electron reservoir that allows controlled charging^{4,7}. Figure 1 shows optical charging of quantum dot A. There are abrupt changes in the photoluminescence at particular voltages signifying charging events. The charge in each plateau is known by identifying the singly charged exciton, X^{1-} , which has no fine structure⁸ and a large extent in gate voltage, $V_{\rm g}$ (ref. 7). Figure 1 demonstrates charging in discrete steps from X^{6+} up to X^{6-} .

Underpinning the experiment is Coulomb blockade: the configuration of electrons with lowest energy is established by a tunnelling interaction with the electron reservoir^{4,7}. For positive charging at large negative voltages, holes accumulate in the quantum dot from the optical excitation. At each voltage, a certain number of holes must accumulate before it becomes energetically favourable for an electron to tunnel into the dot from the Fermi sea, at which point recombination yields a photon. The number of

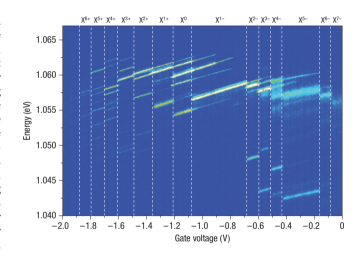


Figure 1 Charging diagram of single InAs/GaAs quantum dot A. Photoluminescence energy plotted against gate voltage, V_g . The photoluminescence intensity is represented with a colour scale, with blue (white) corresponding to 0 (2000) counts on the detector. The vertical lines mark the charging events.

holes required increases step-wise with increasing negative voltage. The transitions between the positively charged excitons are abrupt in voltage because the device does not rely on slow hole tunnelling, exploiting instead fast electron tunnelling.

We have calculated the optical properties of an InAs dot. In the single-particle step, the crystal potential is a superposition of atomic screened potentials v_{α} of atom type α at each relaxed atomic site $\mathbf{R}_{\alpha n}$, where n is the lattice site index: $V(\mathbf{r}) = \sum_{\alpha n} v_{\alpha} (\mathbf{r} - \mathbf{R}_{\alpha n})$. The pseudopotentials are adjusted so that the quasiparticle energies fit experimental data on bulk InAs and GaAs (ref. 10) thus representing valence electron correlations. The Schrödinger equation is solved in a basis set consisting of strained Bloch functions of the underlying bulk semiconductor 10. This approach

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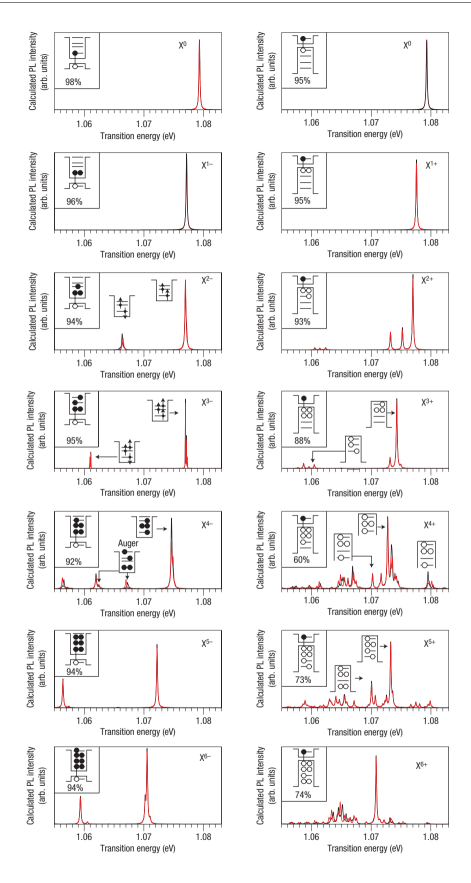


Figure 2 Calculated photoluminescence (PL) spectra for different exciton charges for the dot with circular base. The dot is a cylindrical InAs dot with 25 nm diameter and 2 nm height sitting on two monolayers of wetting layer. Red denotes [110] polarization, black [110] polarization calculated for 5 K. The insets in the upper-left corner of each panel show the leading configuration of the initial states along with its weight in per cent. For some selected photoluminescence lines, the leading final-state configuration is shown.

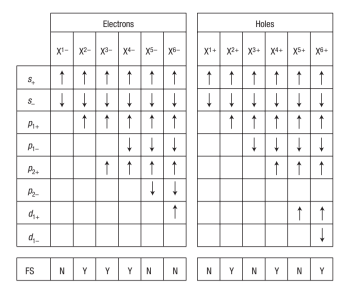


Figure 3 Calculated leading configurations for charged excitons. Left: one hole and *N* electrons; right: one electron and *N* holes. For the positively (negatively) charged states, the single electron (hole) in the initial state is in the lowest electron (hole) state. The last line indicates whether fine-structure effects are predicted (Y) or not (N).

captures the multiband, intervalley and spin-orbit interactions and also includes first- and second-order piezoelectric effects¹¹. All of the valence single-particle levels of the dots are represented, for example, for an N-atom dot there are 2N occupied singleparticle levels ($N \sim 10^6$). These levels have mixed angular momenta, mixed heavy hole-light hole and complex nodal structure, much like the states of a giant molecule containing a comparable number of atoms. This is different to effective-mass models^{6,12} where the orbitals are hydrogenic-like. In the many-body step, we use a configuration interaction approach where all of the Slater determinants constructed from 12 electron and 12 hole singleparticle states (counting spin) interact. The interaction consists of Coulomb and exchange integrals calculated from the atomistic single-particle wavefunctions. The screening function for these integrals is calculated following the microscopic model of Resta¹³ and exhibits a smooth transition from unscreened at short range to screened at long range. This naturally includes both longand short-range exchange¹⁴. Using atomistic orbitals to calculate many-particle interactions is different to most calculations of correlation in quantum nanostructures where the basis of singleparticle states is taken from effective-mass models. That the interelectronic integrals computed from effective-mass wavefunctions are very different to those computed from atomistic wavefunctions can be dramatically seen by the different ensuing symmetries of the charged exciton ground states¹⁵. Our method has provided a quantitative understanding of the unusual X²⁻ and X²⁺ fine structure16.

Calculations for a cylindrical dot are shown in Fig. 2 where the dominant initial-state configuration is given for each charge state with its weight. The final-state configurations are shown for selected transitions. We interpret the experimental results with the model without fitting any of the nanostructure parameters. This approach is limited only by the imperfect knowledge of the real dots' morphology. The experiment does not measure directly the composition of the ground-state configuration. Nevertheless, information can be deduced on the basis of the sensitivity of the photoluminescence spectrum to the initial-state configuration,

closed and open shells having very different optical signatures; and to the initial-state spin through fine structure, an electronhole exchange interaction which leads to polarization-dependent energy splittings⁸. Only an unpaired hole (electron) in the positively (negatively) charged initial state leads to fine structure allowing an experimental discrimination between zero and finite spin. We find an excellent correspondence between our measured and calculated photoluminescence spectra, allowing us to uncover a number of physical effects.

(1) Anomalous ground states. We determine the ground-state configurations of charged excitons by minimizing the many-body energy as a function of the distribution of electrons and holes in the single-particle levels. Figure 3 shows that a system consisting of one hole and two, three, five, six, seven electrons and a system consisting of one electron and two, three, four, five holes follows the Aufbau principle whereas the others, X^{3-} , X^{5+} and X^{6+} , violate the Aufbau principle in that one low-energy level remains empty whereas a higher-energy level is filled. The calculations for X³⁻ predict an open-shell initial configuration (Fig. 2) that recombines into two final states split by 16 meV through electron-electron exchange. The initial state violates Aufbau in that the p_{1-} level remains empty whereas the higher-energy p_{2+} is filled: for dots of small height, the loss in single-particle energy on promoting $\varepsilon(p_{1-})$ to $\varepsilon(p_{2+})$ is more than compensated by the gain in exchange energy associated with forming a higher total spin¹⁷. Experimentally, the dot A X³⁻ photoluminescence is dominated by the open-shell lines split by 14 meV (Fig. 4), in excellent agreement with the theoretical prediction; the two closed-shell lines (green circles in Fig. 4) can be made out and arise from thermal occupation of the higher-energy initial state. Furthermore, the upper X³⁻ photoluminescence line has a clear fine structure, as predicted. For dot B (see Supplementary Information, Fig. S1), the openshell and closed-shell X3- lines have similar intensities—the two configurations are close to degenerate in this case; for dot C (see Supplementary Information, Fig. S2), the closed-shell X3photoluminescence lines dominate. These observations point to a slight decrease in rotational symmetry from dot A to B to C.

Theory predicts a very simple spectrum for X^{5-} that originates from a strongly dominant closed-shell initial configuration with no fine structure. In a calculated elongated dot, the lower X^{5-} peak weakens and a new lower-energy peak emerges (see Supplementary Information, Fig. S3). Experimentally, there is clear shell-filling for X^{5-} : dots A, B and C all have a large- V_g plateau extent and an absence of fine structure. Dot A shows one lower photoluminescence peak, consistent with the theory for the symmetric dot; dots B and C show two lower peaks, consistent with the theory for the elongated dot.

The open-shell ground state of X^{2+} predicted by the theory is confirmed by experiment. A polarization dependence in the transitions is found in both theory and experiment. Theoretically, X³⁺ has a closed-shell ground state with no polarization dependence in the photoluminescence. Likewise, the experimental photoluminescence has no measurable polarization dependence. In the experiment, the principal X³⁺ photoluminescence peak is slightly blue-shifted relative to the principal X2+ peak, closer to the theoretical prediction for the closed-shell X³⁺ (red-shift of 3 meV) than the open-shell X³⁺ (blue-shift of 6 meV). The lines in the experimental X³⁺ photoluminescence above the principal peak from dot A are not reproduced in the theory but these features are of lower relative intensity from dots B and C. The openshell X4+ configuration with fine-structure splitting is confirmed experimentally through the strong polarization dependence of the transitions. X⁵⁺ has a most fascinating behaviour. In contrast to X^{5-} , X^{5+} is predicted to violate Aufbau in that the first d state is occupied before the second p state is filled. Furthermore, we

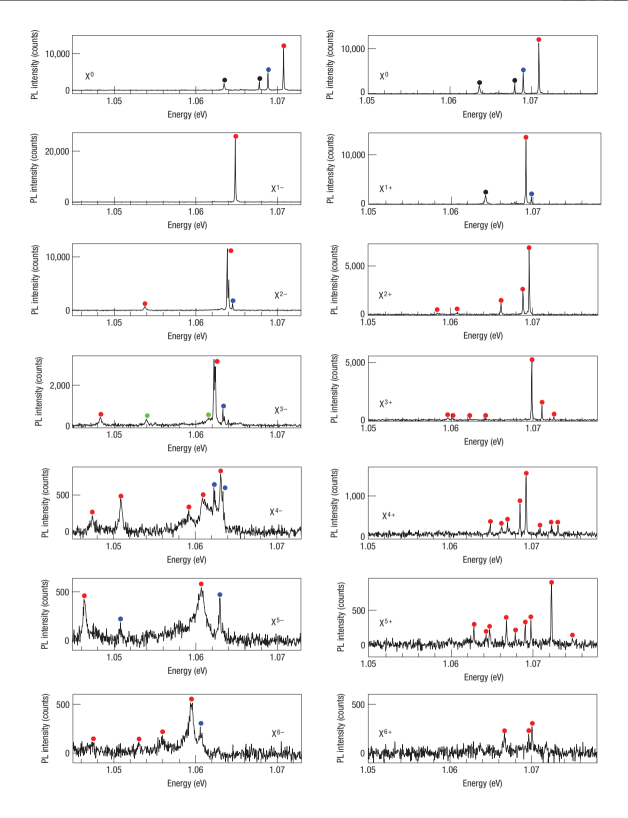


Figure 4 Measured photoluminescence spectra from dot A for different exciton charges. Photoluminescence intensity (840 s integration time) is plotted against energy for V_0 corresponding to the centre of each charging plateau. Individual peaks are labelled: red circle denotes a photoluminescence peak from the particular exciton with charge n; blue circle emission from exciton with charge n-1; black circle biexciton-related emission; green circle emission from an excited initial-state configuration.

predict that the hole charging sequence is perturbed by the presence of the electron: without the electron, the second p state is not occupied at all¹⁵. Curiously, the predicted initial configuration

is open shell, yet the photoluminescence is almost unpolarized, both in the experiment and in the theory, signifying a zero-spinstate coupling of the unpaired holes. Small fine-structure effects are still present in the theoretical results, originating from the admixed (27%) configurations in the initial state; these effects are beyond the experimental resolution. Theoretically, the signature of the open-shell X^{5+} is the presence of a multitude of peaks with comparable intensity, whereas the closed-shell X^{5+} configuration has one strong peak accompanied by many very weak transitions. Experimentally, there are several strong photoluminescence lines, strongly supporting the open-shell configuration. The non-Aufbau filling of hole states continues for X^{6+} where p_2 is left half empty. A polarized experimental spectrum (dots A and B) with a few peaks agrees with the theoretical prediction but the X^{6+} photoluminescence is very weak.

- (2) Non-perturbative Coulomb interactions. A perturbation treatment of the Coulomb interactions predicts a blue-shifted X^{1+} on the basis of a red-shifted X^{1-} (ref. 18). Indeed, our calculated Coulomb energies¹⁷ $J_{hh} = 25.9 \, \text{meV} > |J_{eh}| = 25.3 \, \text{meV} > J_{ee} = 24.9 \, \text{meV}$ lead to a $J_{hh} J_{eh} = 0.6 \, \text{meV}$ blue-shift of X^{1+} with respect to X^0 . However, this effect is countered by the non-perturbative mixing of the $h_s^2 e_s^1$ configuration with other configurations, a mixing that produces an overall red-shift of X^{1+} , a clear feature in both experiment (Fig. 4) and theory (Fig. 2). This simple fact proves that hole–hole Coulomb interactions cannot be treated perturbatively.
- (3) Exchange-split final states. In general, charged exciton states with an open-shell initial configuration decay into final states split by electron–electron or hole–hole exchange. For instance, X^{2-} , X^{3-} , X^{4-} and X^{6-} each have two photoluminescence lines arising from two possible final spin states, Fig. 2. The open-shell X^{2+} decays into three states, not two as for X^{2-} . The two holes in the X^{2+} final state are predominantly heavy hole in character and therefore follow the rules for spin-3/2 particles: there is a two-fold degenerate state with J=3 and two singlets with J=0 and J=2. The X^{2+} initial state is split by the electron–hole exchange interaction and has total angular momentum f=2 and f=1 resulting in three transitions: $f=1 \rightarrow J=0$, $f=1 \rightarrow J=2$ and $f=2 \rightarrow J=3$. For all exciton charges, the exchange splittings in experiment and theory are in perfect correspondence with good quantitative agreement.
- (4) Correlation-induced 'promotion-demotion' exciton lines. The X⁴⁻ photoluminescence consists of more than the two peaks expected from electron exchange. This observation heralds a new consequence of Coulomb interactions. When the number of carriers after recombination exceeds two, a final configuration can be generated by demoting one particle to a lower level (for example, p to s) and promoting another to a higher level (for example, pto $d)^{6,19}$. These Auger-like configurations are themselves optically dark but admixture with bright configurations has a powerful effect on the photoluminescence spectrum. For X⁴⁻, we find two further photoluminescence lines related to promotion-demotion where the electron d orbitals are occupied in the final states even though the d orbitals are empty in the initial state (Fig. 2). The original effective-mass model with parabolic confinement predicts six bright photoluminescence lines for X4- (ref. 6) but this is artificially high owing to the high degeneracies in the single-particle states. Our calculations predict instead four X4- bright lines. The inner two have significant promotion-demotion character and would not appear without the *d* shell. The energies of the innermost transitions depend on the symmetry—the two peaks move further apart in an elongated dot (see Supplementary Information, Fig. S3). In the experiment, Fig. 4 shows two weak peaks at low energy, a strong peak at high energy and two further peaks intermediate in energy with strange line shapes. Analysis of dots B and C reveals in each case three peaks at low energy again with some peculiar structure around the peak at high energy. Our conclusion is that the experiment matches the theory well but there are further peaks at high energy in the measured photoluminescence, arising perhaps

from an excited initial-state configuration. X^{5-} has a closed initial shell and does not have exchange-split final states. Nevertheless, both the measured and calculated X^{5-} spectra exhibit a splitting into two dominant lines, a direct consequence of promotion–demotion. The large calculated mixing of Auger configurations in the final states (40% and 60% in the high and low peaks, respectively) gives rise to the large $\simeq 15$ meV splitting of the two peaks. For the elongated dot, theory predicts that the lower-energy peak splits into two (see Supplementary Information, Fig. S3). This is exactly what we observe for dots B and C (see Supplementary Information, Figs S1, S2).

(5) Configuration mixing in the initial states. For the highly positively charged excitons, the likelihood of finding several symmetry-compatible many-body configurations is large. We obtain theoretically such a strong initial-state mixing especially for X4+ and X5+ where the majority configuration constitutes only 60% and 73%, respectively, of the initial state. The minority configurations open new recombination pathways (see also Supplementary Information, Fig. S4). Especially intriguing are photoluminescence transitions in both the experiment and the theory for X⁴⁺ and X⁵⁺ a few meV above the principal peak. An admixed configuration in the initial state with non-Aufbau character is responsible for these photoluminescence lines. For X^{4+} , it is the configuration where the fifth hole occupies the d and not the p_2 level. This opens pathways to a final configuration with an empty p_2 state, as shown in Fig. 2. Together with holehole exchange and promotion-demotion effects, this leads to very complex X4+ and X5+ photoluminescence spectra. The general feature of a principal line at high energy with a multitude of lines in a bandwidth of \sim 10 meV is a feature in both experiment and theory. There is clearly a very strong dot dependence: in the theory, elongating the dot induces a considerable change to the X4+ and X5+ spectra. Similarly, dots A, B, C show marked differences for X⁴⁺ and X⁵⁺. A more detailed understanding of X⁴⁺ and X⁵⁺ will probably be possible only with a more detailed picture of the real dot morphology.

METHODS

The quantum dots are self-assembled by molecular beam epitaxy and are separated by 17 nm of intrinsic GaAs from an n $^+$ GaAs layer. The dots are capped with 10 nm of undoped GaAs followed by a 105 nm AlAs/GaAs superlattice. Ohmic contacts are prepared to the back contact, the earth, after which a 5-nm-thick NiCr Schottky barrier is evaporated onto the sample surface. Voltage $V_{\rm g}$ is applied to the NiCr Schottky contact. Photoluminescence is excited non-resonantly at 850 nm, collected with a low-temperature confocal microscope (5 K) and detected with a spectrometer–InGaAs photodiode array set-up with a spectral resolution of 120 $\mu \rm eV$.

Some photoluminescence lines extend in voltage beyond the host plateau. It is important to make an assignment of the photoluminescence lines to a particular exciton charge. We find that some photoluminescence lines extend beyond the plateau to more positive $V_{\rm g}$ but not to more negative $V_{\rm g}$. (This is the consequence of a cascade: X^{n-} emission leaves behind n electrons, which then capture a hole and emit as $X^{(n-1)-}$ before the true ground-state configuration, X^{n-} , is re-established.) In addition, the photoluminescence intensity from a particular exciton decreases abruptly with increasing $V_{\rm g}$ once the exciton no longer forms the ground state. We can determine the $V_{\rm g}$ boundaries of each photoluminescence line to \sim 10 mV, enabling us to assign each line to a particular charge without ambiguity, Fig. 4. In Fig. 1 there are some weak biexciton-related features but these can be spotted easily through their dependence on pump power.

Tunnelling does not shift the photoluminescence lines appreciably: even at the plateau edges, energy shifts in the photoluminescence through a tunnel hybridization cannot be made out in Fig. 1. However, the photoluminescence energies depend on the electric field, a Stark effect, and we find that the corresponding dipole moment and polarizability depend on charge, particularly for the positively charged excitons⁹. To facilitate comparison with



our theory, we extrapolate in Fig. 4 the photoluminescence spectra to zero electric field. This is crucial as it reveals a red-shifted X^{1+} , not a blue-shifted X^{1+} as a cursory inspection of Fig. 1 might suggest. None of the splittings in the experimental photoluminescence depend on electric field demonstrating the validity of this method.

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Author contributions

M.E. carried out the experimental work under the supervision of R.J.W. and K.K; G.B. carried out the theoretical work in A.Z.'s group; A.B. fabricated the heterostructure in P.M.P.'s group. M.E., G.B., K.K., A.Z. and R.J.W. worked jointly on the interpretation of the results.

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