

shell emission is recovered with an energy significantly lowered in comparison with the emission line of a single exciton due to exchange interaction (bandgap renormalization).

The calculations are in good agreement with the experimental observations. As the emission in quantum-dot lasers originates from recombination of excitonic complexes, these calculations should be applicable to the electronic structure of such lasers. The demonstration of 'atomic spectroscopy' in a solid-state system might also open up possibilities of using quantum dots for quantum information processing²⁶. Quantum dots offer flexibility in designing and controlling their interaction with optical pulses. Optically driven coupled quantum dots might lead to the realization of quantum gates, a building block of a quantum computer. A major obstacle to realizing an optically driven quantum computer in semiconductors is the fast dephasing of interband polarization due to carrier-carrier scattering. The hidden symmetries in excitonic artificial atoms imply that the four-particle correlations in the hamiltonian—which cause the dephasing—cancel. Therefore the realization of hidden symmetries may prove useful for the construction of decoherence-free (as far as carrier-carrier interactions are concerned) quantum dynamics. □

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Optical emission from a charge-tunable quantum ring

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Quantum dots or rings are artificial nanometre-sized clusters that confine electrons in all three directions. They can be fabricated in a semiconductor system by embedding an island of low-bandgap material in a sea of material with a higher bandgap. Quantum dots are often referred to as artificial atoms because, when filled sequentially with electrons, the charging energies are pronounced for particular electron numbers^{1–3}; this is analogous to Hund's rules in atomic physics. But semiconductors also have a valence band with strong optical transitions to the conduction band. These transitions are the basis for the application of quantum dots as laser emitters⁴, storage devices^{5–7} and fluorescence markers⁸. Here we report how the optical emission (photoluminescence) of a single quantum ring changes as electrons are added one-by-one. We find that the emission energy changes abruptly whenever an electron is added to the artificial atom, and that the sizes of the jumps reveal a shell structure.

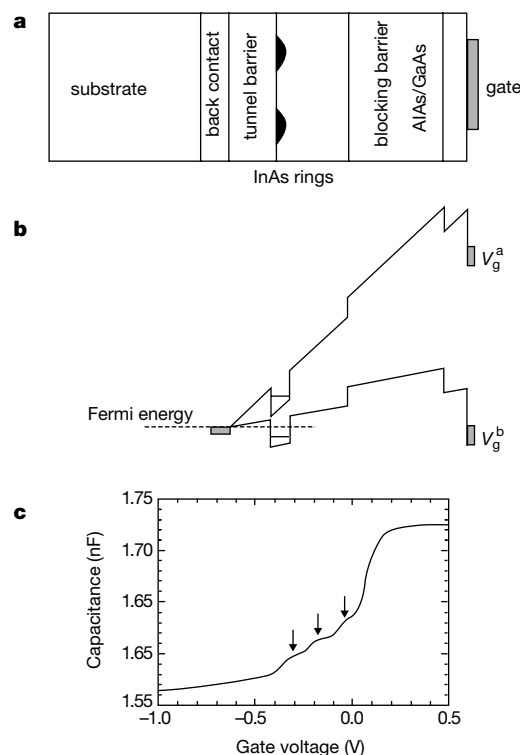


Figure 1 The semiconductor device for charging quantum rings with electrons. **a**, The layer structure. The tunnel barrier is 25 nm thick, and the separation between back contact and surface is 175 nm, so that a change in gate voltage ΔV_g implies a change in electrostatic potential (in eV) of $\Delta V_g/7$. **b**, The band diagram at two different voltages, V_g^a and V_g^b . **c**, The capacitance of the device used for the optical experiments (gate area 1.8 mm²) at 4.2 K. The arrows mark three charging peaks.

The nanometre-sized clusters for our experiments were made by self-assembly. We initially grew InAs quantum dots on GaAs by exploiting the strain-driven change from two-dimensional to three-dimensional growth which occurs after a coverage of 1.5 monolayers of InAs. A pause was introduced in the overgrowth with GaAs during which In migrated, causing the dots to become ring-like^{9,10}. The rings have an *s*-like ground state and a *p*-like excited state, as for dots, so that the general features of shell filling are the same for both dots and rings. The Aharonov–Bohm effect, an interference of electron waves in a quantum ring, is not relevant here as we do not apply a magnetic field to generate a phase difference between clockwise and anticlockwise paths around the ring.

We were faced with two experimental challenges. The first was to design a structure to load quantum rings sequentially with electrons. Our concept is to embed the InAs rings between highly doped GaAs and a surface gate electrode¹¹, as shown in Fig. 1a. At large, negative gate voltages (V_g^a in Fig. 1b), the ring level lies well above the Fermi energy and is unoccupied. At a more positive V_g , the ring level is resonant with the Fermi energy and electrons tunnel into and out of the ring. A further increase in V_g (V_g^b in Fig. 1b) traps one electron in the ring. We can monitor the tunnelling by using the capacitance between gate and back contact^{12,13}. Three charging peaks can be made out in the capacitance of a large ensemble of rings (Fig. 1c). The rise in capacitance at $V_g = 0.1$ V corresponds to tunnelling into the thin InAs layer between the rings (the wetting layer).

The second challenge was to measure a single ring. This is important as otherwise the 20-meV inhomogeneous broadening in the photoluminescence (PL) obscures the shifts expected on charging. We measured PL with a low-temperature confocal microscope which has a diffraction-limited spatial resolution of 610 nm at a wavelength of 950 nm. Our ring densities are $\sim 5 \times 10^9 \text{ cm}^{-2}$, implying that several tens of rings lie in the focus. Individual rings can be selected through their emission energy: we looked for rings in the low-energy tail of the spectral distribution. In order to investigate rings closer to the peak of the ensemble distribution, we increased the spatial resolution by defining 300-nm-sized holes in an aluminium film on the sample surface. Even in this case, we always detected emission from several rings. However, as each ring has unique charging voltages, the V_g dependence of the PL allows us to determine which PL lines belong to which ring. We present here data from a sample without a metal mask as the signal/noise ratio is superior.

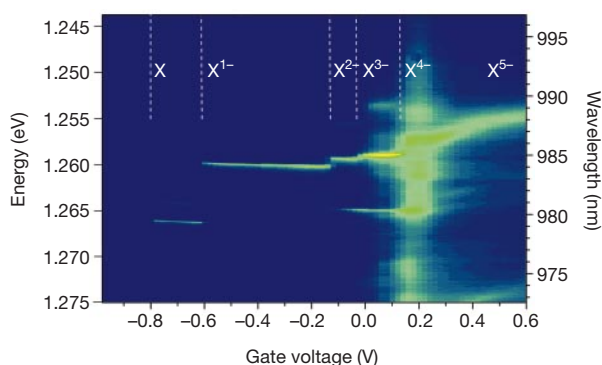


Figure 2 A colour-scale plot of the photoluminescence (PL) versus gate voltage at 4.2 K. Here dark blue, green and yellow correspond to low, medium and high signals, respectively. PL was excited by generating electron–hole pairs in the wetting layer with a laser diode emitting at 822 nm wavelength. The pump power was kept below 1 μW in order to populate the rings with no more than one exciton at a given time. The PL was dispersed with a 0.25-m grating spectrometer and detected with a cooled CCD camera, a set-up with spectral resolution 0.2 nm. The main features are from a single ring. The PL at 1.265 eV (980 nm), however, pronounced around 0.0 V, comes from a second ring.

We recorded the PL as a function of V_g . The data are represented as a colour-scale plot in Fig. 2. At $V_g \sim -0.7$ V there is a single, sharp peak which is the emission from a single ring. At -0.6 V the PL energy decreases abruptly. The jump in energy occurs because an additional electron becomes trapped in the ring. This voltage, -0.6 V, is more negative than the first charging voltage of the ensemble average (Fig. 1c), and this correlates with the particularly low emission energy of the ring in Fig. 2. At higher V_g , further steps can be made out in the PL. In each case, a jump arises when an additional electron is added to the ring. The first jump at -0.6 V, 6.0 meV in energy, represents the binding energy of a singly charged exciton (X^{1-}). The second jump represents the energy needed to add an additional electron to the X^{1-} to form X^{2-} , and so on. The first jump in energy is large, the second small, the third also small, and the fourth reasonably large. This is an optical manifestation of Hund's rules for electron charging. Whenever an electron can be added with the benefit of exchange energy, as is the case for the second and third steps, the charging energy is small. Whenever an electron is added to complete a sub-shell, the charging energy is large.

On moving from X^{1-} to X^{2-} , a satellite appears on the low-energy side of the main peak. The satellite can be just made out in the colour-scale plot, and is a clear feature in the individual spectra, a few of which are shown in Fig. 3. We can be confident that the two emission lines come from the same ring because the satellite

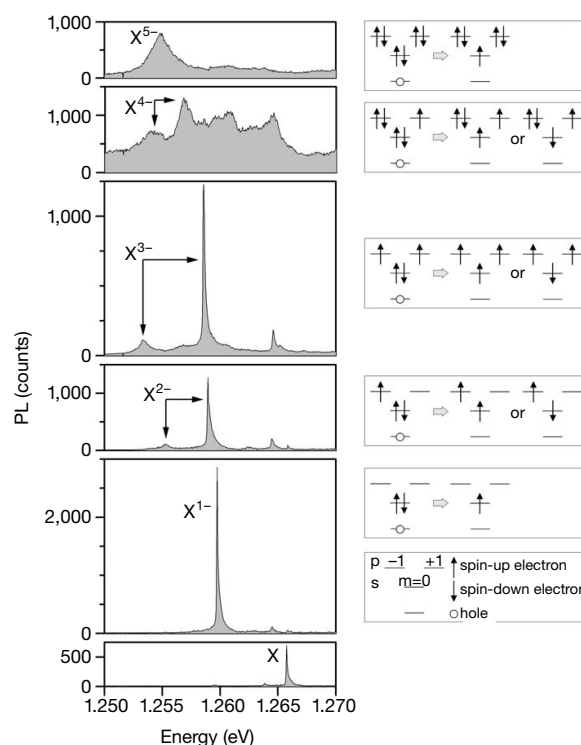


Figure 3 Photoluminescence from a single quantum ring for different charge states. Spectra are shown at $V_g = -0.76, -0.16, -0.10, 0.40, 0.22$ and 0.50 V, corresponding to emission from the $X, X^{1-}, X^{2-}, X^{3-}, X^{4-}$ and X^{5-} excitons, respectively. The PL red-shifts on charging, and a satellite occurs on the low-energy side of the main peak for X^{2-} and X^{3-} . Note that the other, weaker emissions (for instance on the high-energy side of the main X^{2-} PL) belong to another ring, and that the asymmetry in the sharp lines is an artefact of the spectrometer. The broad peak between the X^{3-} main peak and the satellite may be emission from an excited initial state which recombines before it can relax. The level diagrams show the emission processes from initial state to possible final states. We consider *s* and *p* states in the conduction band with angular momenta of 0 and ± 1 , and the *s* state in the valence band. In the experiment, both electrons and holes are unpolarized. To avoid redundancy in the figure, we sketch cases with electrons preferentially polarized up and an unpolarized hole.

emerges exactly when the PL changes from X^{1-} to X^{2-} (see also Fig. 4). Also, the main line weakens at the transition but the total intensity stays constant to within 10%. We claim that the X^{2-} has two lines because there are two different final states¹⁴, as shown in the level diagrams in Fig. 3. For X^{2-} , the two possible final states have either parallel spins (a triplet state) or antiparallel spins (a singlet state). In exact analogy to the excited states of the helium atom, these two states are separated by twice the exchange energy, $2X_{sp}$, and have degeneracies of 3 and 1, respectively. We measure the splitting to be 3.6 meV, that is $X_{sp} = 1.8$ meV. The intensity ratio is $7 \pm 2 : 1$, not the 3 : 1 expected from the degeneracies. This could be due to the electron–hole exchange interaction which tends to align the spins in the initial state, favouring the triplet final state^{15,16}.

For X^{3-} a similar argument applies; in this case the final state is analogous to the excited lithium atom. We have diagonalized the interaction matrix for three spin-1/2 electrons, and calculate a splitting of $3X_{sp}$ and an intensity ratio 2 : 1. Experimentally, the splitting is 1.44 times larger than for the X^{2-} , remarkably close to the predicted 1.5, and the satellite becomes more intense, 2 ± 0.5 times weaker than the main peak. It is surprising that a treatment of the Coulomb interactions with perturbation theory works so well. This approach is justified when the electron and hole wavefunctions extend less than the excitonic Bohr radius, 10 nm; yet here the circumference of the ring is ~ 100 nm. The X^{3-} is the most complicated case; X^{4-} is one electron short of a filled p shell and should behave like X^{2-} . In the data, the splitting for X^{4-} returns almost to the value for X^{2-} but the PL becomes very broad.

At large and negative V_g , the PL disappears; we interpret this as field ionization of the excitons. The electron tunnelling time, τ_t , decreases rapidly with increasing $|V_g|$, eventually becoming smaller than the recombination time, τ_r . At a smaller $|V_g|$ ($\tau_t > \tau_r$), the 0-electron and the 1-electron states become degenerate and electron tunnelling occurs. We detect this as a jump in the PL energy. Figure 4 shows how the high-energy peak weakens and the low-energy peak strengthens with small steps in V_g . It is puzzling that we observe both X and X^{1-} over a large range of V_g , some 20 mV, corresponding to a change in electrostatic energy of about 3 meV. This energy is much larger than the thermal energy (0.36 meV). Heating from the laser is unlikely as an explanation, as broad single electron tunnelling peaks have been observed in transport experiments on similar samples¹⁷. The 3-meV energy scale is also much larger than ring–ring interactions (1 meV) which are significantly screened by the adjacent back contact¹³. To account for this result, we propose that there are temporal fluctuations in the potential (on a timescale large compared to τ_t), caused by some sort of impurity state close to the ring. The effect is analogous to the spectral diffusion observed in the PL from single CdSe dots¹⁸. Figure 4 shows also the transition from X^{1-} to X^{2-} , and this too shows a gradual cross-over from one line to the other. Given the 3-meV fluctuations, it is at first sight surprising that we see sharp PL lines at all. The explanation is that the PL energy is very sensitive to the charge in the ring, but only weakly

sensitive to the electric field (almost flat plateaux in Fig. 2). The combination of the fluctuations with the slight V_g -dependence of the PL energy constitutes a broadening mechanism. We predict this to be about 20 μ eV for our sample, comparable in fact to the expected homogeneous linewidth¹⁹.

For highly charged excitons, X^{2-} and above, the final state after emission of a photon is an excited state. In the simplest case, the ring emits a photon and then some time later relaxes into the ground state. But if the final state relaxes very quickly, the emission will be broadened by the energetic uncertainty of the final state. Therefore, we can use the PL linewidth as a measure of the relaxation rates of the final states. For X^{2-} , the satellite is clearly broader (full-width at half-maximum, FWHM, 0.6 meV) than the main line (FWHM < 0.25 meV), implying a singlet decay time of 1.1 ps and that the singlet decays faster than the triplet. The different decay rates are a consequence of the spin: the triplet state has to flip a spin to relax, the singlet state does not. For the singlet, phonon scattering—which preserves electron spin—is the likely mechanism for the relaxation. Relaxation by phonon scattering is therefore fast on the timescale of recombination, a conclusion consistent with other experiments^{19–21}. The argument holds also for X^{3-} , which also shows a broad satellite PL (FWHM 1.2 meV).

An obvious feature in Figs 2 and 3 is the marked broadening and increase in intensity in the emission at $V_g \approx 0.1$ V. This looks similar to the broad emission of highly excited dots²². We suggest that the PL is broad at 0.1 V because relaxation of the final state always proceeds quickly, regardless of spin. A new relaxation channel must open at 0.1 V. We know from the capacitance that at 0.1 V the wetting layer begins to fill, and the new channel is therefore an Auger process: electrons in the quantum ring lose energy by promoting an electron in the wetting layer to an unoccupied state at higher energy²³. Auger processes can swap the spins of the two participating electrons. In fact, spin-swapping processes are most likely²⁴. This interpretation is supported by the nature of the PL at higher positive voltages, where the PL to some extent narrows again. The ring electrons can only interact with electrons in the continuum a few meV away from the Fermi energy, so that once the Fermi energy is very high, the Auger rate will decrease. The broadening at 0.1 V is accompanied by a rapid increase in overall intensity. This is partly caused by an increase in capture efficiency, but other factors may be involved. $\square$

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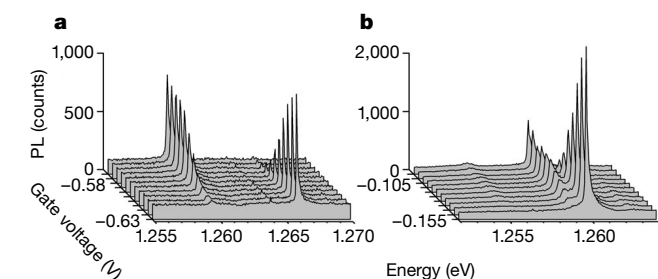


Figure 4 PL of a single quantum ring in the transition from one charge state to the next. **a**, X to X^{1-} ; **b**, X^{1-} to X^{2-} . Panel **b** also shows the emergence of the satellite at 1.255 eV.

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Gigantic optical nonlinearity in one-dimensional Mott–Hubbard insulators

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The realization of all-optical switching, modulating and computing devices is an important goal in modern optical technology. Nonlinear optical materials with large third-order nonlinear susceptibilities ($\chi^{(3)}$) are indispensable for such devices, because the magnitude of this quantity dominates the device performance. A key strategy in the development of new materials with large nonlinear susceptibilities is the exploration of quasi-one-dimensional systems^{1,2}, or ‘quantum wires’—the quantum confinement of electron–hole motion in one-dimensional space can enhance $\chi^{(3)}$. Two types of chemically synthesized quantum wires have been extensively studied: the band insulators of silicon polymers, and Peierls insulators of π -conjugated polymers and platinum halides. In these systems, $\chi^{(3)}$ values of 10^{-12} to 10^{-7} e.s.u. (electrostatic system of units) have been reported^{3–7}. Here we demonstrate an anomalous enhancement of the third-order nonlinear susceptibility in a different category of quantum wires: one-dimensional Mott insulators of $3d$ transition-metal oxides and halides. By analysing the electroreflectance spectra of these com-

pounds, we measure $\chi^{(3)}$ values in the range 10^{-8} to 10^{-5} e.s.u. The anomalous enhancement results from a large dipole moment between the lowest two excited states of these systems.

Since the discovery of high-temperature superconductivity, the dynamics of charge and spin under the influence of strong electron correlation have been extensively investigated in doped and undoped copper oxides⁸. Among them, Sr_2CuO_3 (whose crystal structure is shown in Fig. 1a) is a prototype one-dimensional (1D) Mott insulator with quantum spin liquid⁹. One-dimensional Cu–O chains are composed of CuO_4 quadrilateral structures with shared corner oxygens along the b axis, in which the Cu ion is divalent (spin quantum number $S = 1/2$) and one unpaired electron exists in the $d_{x^2-y^2}$ orbital. In the Cu–O chain, a 1D electronic state is formed by the overlap of the p_x, p_y orbitals of O and the $d_{x^2-y^2}$ orbitals of Cu, as shown in Fig. 1c. Because of large on-site Coulomb repulsion energy U acting on the Cu ions, the Mott–Hubbard gap is opened in the Cu $3d$ band. The occupied O $2p$ band is located between the Cu $3d$ upper Hubbard (UH) band and the Cu $3d$ lower Hubbard (LH) band, as schematically illustrated in Fig. 1e. A strong charge transfer (CT) transition from the O $2p$ valence band to the Cu $3d$ UH band occurs around 1.75 eV, as observed in the spectrum of the imaginary part of dielectric constant (ϵ_2) shown by the broken line in Fig. 2.

Other prototypical 1D Mott insulators investigated here are the halogen (X)-bridged Ni compounds^{10,11}, $[\text{Ni}(\text{chxn})_2\text{X}]Y_2$, (Y, counter ion; chxn, cyclohexanediamine) whose general crystal structure is shown in Fig. 1b. The Ni^{3+} ions and X^- ions are arranged alternately along the b axis, forming a purely 1D electronic state composed of the p_z orbitals of X and the d_{z^2} orbitals of Ni (see Fig. 1d). Four N atoms of amino groups in two chxn molecules coordinate a Ni^{3+} ion in a plane normal to the chain axis b , and

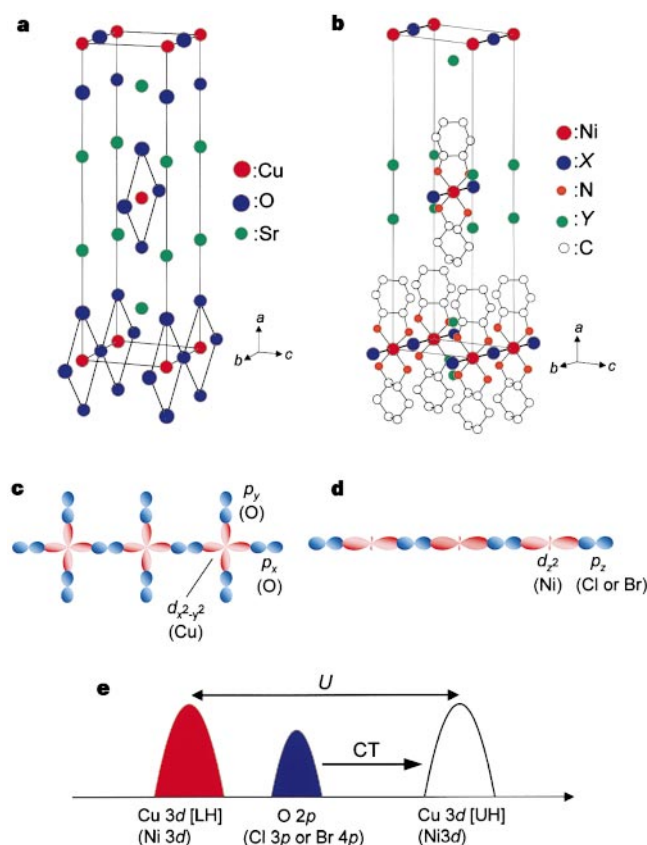


Figure 1 Crystal and electronic structures of the 1D Mott insulators. **a**, Structure of Sr_2CuO_3 . **b**, Structure of $[\text{Ni}(\text{chxn})_2\text{X}]Y_2$. **c**, Configuration of Cu $3d_{x^2-y^2}$ (red) and O $2p_x, 2p_y$ (blue) orbitals in Sr_2CuO_3 . **d**, Configuration of Ni $3d_{z^2}$ (red) and X $2p_z$ (blue) orbitals in $[\text{Ni}(\text{chxn})_2\text{X}]Y_2$. **e**, Schematic electronic structure of the 1D Mott insulators. In **b**, hydrogen atoms are omitted.