We report photon correlation measurements that allow us to observe unique signatures of biexcitons in a single self-assembled InAs quantum dot. Photon correlation measurements of biexciton emission exhibit both bunching and antibunching under continuous-wave excitation while only antibunching is observed under pulsed excitation. Cross-correlation between biexciton and single-exciton peaks reveal highly asymmetric features, demonstrating that biexciton and exciton emissions have strong correlations due to cascaded emission. The anticipated correlation between the polarization of exciton and biexciton emissions however, is absent under our excitation conditions. Photon correlation measurements also provide evidence for the identification of the charged exciton emission.

78.67.Hc, 42.50.Dv, 78.55.-m, 78.55.Cr

It is by now widely accepted that various quantum dot (QD) structures exhibit features in transport \cite{12} or optical spectroscopy \cite{13,14} that indicate full three-dimensional confinement of carriers. Identification of QDs as artificial atoms has been strengthened by the recent observation of strong photon antibunching in single-exciton emission \cite{13,14}, which is the typical signature of an anharmonic quantum system: after a photon is emitted from a single two-level (anharmonic) emitter, the system is necessarily in the radiatively inactive ground state and a second photon cannot be emitted immediately after the first one. Even though the coherence properties of QD single-exciton emission closely follow those of atoms, the overall spectral features of single QDs are significantly more complicated. Since the size of QDs is roughly two-orders of magnitude larger than those of atoms, multiparticle excitations give rise to emission peaks with energies comparable to that of a single-exciton. Of primary importance in QD spectroscopy is the biexciton state, which corresponds to a doubly-excited QD with completely filled lowest electron and hole energy levels. When the biexciton state decays by radiative recombination, the final-state is a single-exciton state and the generated photon is shifted as compared to the single-exciton emission due to Coulomb interaction between the carriers. Biexciton emission in QD spectroscopy has been traditionally identified using the (quadratic) pump-power dependence of the corresponding peak.

In this letter, we demonstrate that photon correlation measurements provide a powerful tool for characterizing the multixciton spectral features of QDs. Our measurements provide a strong support for the identification of a biexciton emission peak, by demonstrating its strong correlations with the subsequent single-exciton emission. We observe that biexciton intensity autocorrelation exhibits bunching together with antibunching or only antibunching under continuous-wave (cw) excitation depending on the excitation level. In contrast, we find strong antibunching under pulsed excitation. The large difference between the levels of antibunching under continuous-wave and pulsed excitations points out to the importance of excitation mechanism and the role of free carriers in QD physics. The lack of polarization correlation between biexciton and single-exciton emissions indicates that spin dephasing is likely to play a key role under non-resonant excitation. We also observe that a third emission peak in QD spectra exhibits strong correlations with both exciton and biexciton fluorescence: we argue that these correlation signatures suggest the identification of this additional line as a charged exciton emission.

Our self-assembled InAs QDs were grown by molecular beam epitaxy (MBE) using the partially covered island technique \cite{9}. Growth resulted in typically lens shaped QDs with a base diameter of 40-50 nm and a height of 3 nm, having their single-excitonic emissions between 925 nm and 975 nm in the spectrum. In our sample, the QDs were embedded in the center of a 200 nm thick GaAs microdisk structure located above a 0.5 μm thick Al0.65Ga0.35As post. The diameter of the disks was 5 μm and the average number of QDs within the disks was less than 1. Details of the microdisk processing can be found elsewhere \cite{15}. Our experimental setup consisted of a combination of a low-temperature diffraction-limited scanning optical microscope and a Hanbury Brown and Twiss (HBT) \cite{10} setup for photon correlation measurements. The QD sample was mounted in a He-flow cryostat and cooled to 4-7 K. The cryostat was moved by a computer-controlled translation stage, thus allowing for scanning across the sample. The QDs were optically excited either with a continuous wave diode laser (operating at 785 nm), a continuous wave Ti:Sa laser (operating at...
760 nm) or a mode-locked Ti:Sa laser (82 MHz, 250 fs, operating at 790 nm), generating electron-hole pairs in the GaAs barrier layer which are subsequently captured by the QDs within a short timescale (< 35 ps) [11]. The same microscope objective (NA = 0.55) mounted outside the cryostat was used to both excite the QDs (diffraction limit : ∼ 1.7 μm) and collect the emitted fluorescence light. The collected light was spectrally filtered either by a 0.5 m monochromator or narrowband interference filters (FWHM=0.5 or 1 nm) before being detected in the HBT setup, consisting of a 50/50 beamsplitter and two single-photon-counting avalanche photodiodes. The APDs were connected to the start and stop inputs of a time to amplitude converter (TAC). The TAC output was stored in a multichannel analyzer (MCA) to yield the number of photon pairs n(τ) with arrival time separation of τ = t_{start} − t_{stop}. An electronic delay was introduced into the stop channel in order to measure the photon correlation for negative time delays. The time resolution of the HBT setup was 420 ps. Under our experimental conditions with detection efficiency of about 0.1%, the measured spectra we deduced lifetimes of 3.6 ns, 3.7 ns and 2.6 ns for the X1, X2, and XX emissions respectively. The resultant ratio of \( t_{XX}/t_{X1} = 1.4 \) is consistent with exciton and biexciton lifetime measurements performed on CdSe/ZnSe QDs [14]. We also note that, due to the different photonic environment created by the microdisk, the single-exciton lifetime we observe (3.6 ns) is larger than typical lifetimes measured (~1 ns) in bulk QD samples [7].

Fig. 1 shows the power dependent photoluminescence (PL) spectra of the single QD that we analyze. At low pump powers, the single-exciton emission peak (X1) dominates the spectrum. At higher pump powers, we observe that two other peaks become dominant: among these, the lower energy one (XX) has an energy (red) shift of 3.5 meV from X1 and its intensity has a quadratic dependence on pump power; these are typical signatures for biexciton emission in self-assembled InAs QDs. The third peak (X2) is red-shifted from the single-exciton peak by about 500 μeV. All three emission peaks are resolution limited at 70 μeV, and none of them is polarized. To ensure that X1 originates from a single QD exciton emission, we have carried out photon auto-correlation measurements where both APDs were illuminated by X1 emission: under cw and pulsed excitation, X1 emission was found to exhibit perfect antibunching (Fig. 1 inset) and single-photon source operation [12,13], respectively. We have also performed time correlated single photon counting experiments on X1, X2, and XX emissions to measure their lifetimes [14]. Those measurements were performed in the very low excitation regime where the decay times of the resulting spectra were determined by the lifetimes of the corresponding emissions [14]. From the measured spectra we deduced lifetimes of 3.6 ns, 3.7 ns and 2.6 ns for the X1, X2, and XX emissions respectively. The resultant ratio of \( t_{X1}/t_{XX} = 1.4 \) is consistent with exciton and biexciton lifetime measurements performed on CdSe/ZnSe QDs [14]. We also note that, due to the different photonic environment created by the microdisk, the single-exciton lifetime we observe (3.6 ns) is larger than typical lifetimes measured (~1 ns) in bulk QD samples [7].
will also exhibit simple antibunching, as has been observed for single-exciton emission. Figures 2(a) and 2(b) show $g^{(2)}(\tau)$ for the XX emission under cw excitation, at pump powers corresponding to X1 emission intensities that are 0.7 and 1.0 of the (exciton) saturation level, respectively. Both curves, obtained using a 0.5 nm interference filter, exhibit antibunching ($g^{(2)}(0) = 0.95$ in Fig. 2(a), $g^{(2)}(0) = 0.6$ in Fig. 2(b)) with similar decay times of 1 ns. The curve in Fig. 2(a) also exhibits bunching ($g^{(2)} = 1.4$) that decays with a decay time of 3.5 ns. Bunching here originates from the fact that the detection of a photon at the biexciton transition results in the projection of the QD wave-function onto the single-exciton-state X1. When the average occupancy of X1 in steady-state is lower than unity, post-measurement-state has higher occupancy in the single-exciton-state than pre-measurement-state, and is more likely to result in re-excitation of the biexciton state. An analysis of the QD dynamics using 3-level rate equations indicates that $g^{(2)}(\tau)$ should indeed exhibit bunching that decays in a timescale determined by the single-exciton lifetime of 3.6 ns which is in agreement with the experimental result (Fig. 2(a)). This analysis also predicts that antibunching at $\tau = 0$ should turn into bunching in a timescale determined by the biexciton lifetime in the low excitation regime.

We could observe strong bunching but no antibunching in biexciton auto-correlation measurements when we use a 1 nm interference filter, indicating the importance of additional broadband radiation at biexciton energy that appears to be correlated with exciton emission. While the pump laser wavelength has a strong effect on the observability of biexciton antibunching, pump intensity plays no significant role in determining the recovery time of the antibunching dip. On the other hand, the stronger biexciton antibunching under pulsed excitation suggests that the free carriers could still have an adverse effect (Fig. 2(c)): we remark that under pulsed excitation, free-carriers recombine in a time-scale that is much faster than the biexciton radiative recombination time, and therefore their influence on biexciton dynamics is expected to be minimal. The antibunching dip in Fig. 2(a) is also affected by the fact that due to the presence of bunching, the correlation function recovers to a value exceeding unity within a biexciton lifetime; the effect of time resolution is therefore more pronounced in the vicinity of zero time delay compared to the excitation regime when no bunching is observed (Fig. 2(b)).

Cross-correlation measurements complement the identification of the biexciton emission: since X1 population is enhanced as a result of the detection of an XX photon, strong correlations between the X1 and XX emissions can be expected. Fig. 3 shows such a photon cross-correlation measurement, obtained by illuminating the start APD by the X1 emission and stop-APD by the XX emission. The depicted quantity here is $g^{(2)}(\tau) = \langle(I_{XX}(t)I_{X1}(t+\tau) - \langle I_{XX}(t) \rangle \langle I_{X1}(t) \rangle)/\langle I_{X1}(t) \rangle \langle I_{XX}(t) \rangle)$, where $I_{X1}(t)$ and $I_{XX}(t)$ are the intensities of the X1 and XX emissions, respectively. Remarkable features of this cross-correlation include strong antibunching for $\tau > 0$ and strong bunching for $\tau < 0$ with a close to resolution-limited transition between the two regimes. For $\tau > 0$, suppression of a joint X1 and XX event arises from the fact that following the detection of an X1 photon, which projects the QD onto its ground-state, detection of an XX photon is very unlikely. Strong bunching for $\tau < 0$ follows from the fact that detection of an XX photon projects the QD onto the X1 state, as discussed earlier. Asymmetry in cross-correlation measurements under pulsed excitation have been recently reported [10]. We claim that the signature depicted in Fig. 3 proves that the XX emission arises from the decay of the biexciton state into the single-exciton state. The strong antibunching in cross-correlation is yet another indication that the additional broadband radiation is correlated with the X1 emission.
increases the relative intensity of charged exciton emission. Presence of carbon in these samples is well known, making it likely that a QD is p-doped.

Finally, it has been predicted that the radiative decay of a single QD biexciton state will result in polarization-entangled-state generation \[17\]. To observe such polarization correlations, we have measured the polarization dependence of the X1-XX cross-correlation. Under cw-excitation, we have seen no evidence for polarization correlations. We believe that spin-decoherence that has been observed to occur in nanosecond timescales for these QDs under non-resonant excitation is responsible for the lack of polarization-correlation \[18\].

In summary, we have used photon auto- and cross-correlation measurements to identify dominant spectral features of a single QD, and characterize the recombination dynamics under various excitation conditions. Given the difficulty of accurate theoretical calculations and the richness of the QD spectra which differs significantly from one QD to another, we believe that the techniques described here will be invaluable in understanding individual QDs. Further experiments under different excitation conditions are needed to understand the polarization correlations and eventually for the generation of entangled-photon states.

This work was supported by David Packard Foundation and an Army Research Office grant. C.B. acknowledges support from the Deutsche Forschungsgemeinschaft.

**FIG. 4.** Cross-correlation measurements under cw diode laser excitation at 785 nm. (a) The X2 fluorescence is sent to the start APD while XX is sent to the stop APD. The absence of bunching demonstrates that XX emission does not populate the X2 state. (b) The X2 emission is sent to the start APD and the X1 emission to the stop APD. The antibunching dip shows that both transitions stem from the same QD. The asymmetry in the dip indicates a faster recovery for the X1 state after X2 detection.

Having identified the two principal lines in QD spectrum, the next natural question is whether photon correlation spectroscopy can tell us anything about the origin of the X2 emission. The cross-correlation between the X2 and XX emissions only shows antibunching (Fig. 4(a)), indicating that while those emissions arise from the same QD, the radiative decay of the biexciton state does not populate the X2 state. From the pump power dependent PL spectra (Fig. 1), it can be seen that the X2 emission has a stronger pump power dependence than X1 but saturates earlier than the XX line. This may already suggest an identification of X2 as a charged-exciton (trion) line. To provide further evidence, we have carried out cross-correlation measurements between the X2 and X1 emissions \(g^{(2)}(\tau)\) where the start and stop APDs were illuminated by the X1 and X2 lines, respectively. The resulting X1-X2 cross-correlation function (Fig. 4(b)) clearly shows asymmetric antibunching with \(g^{(2)}(0) = 0.3\), which proves once again that the two lines originate from the same QD. The asymmetry with fast recovery for \(\tau > 0\) is expected if X2 arises from a charged exciton: the post-measurement state of charged-exciton emission is a singly-charged QD. We would expect single-charge injection into the QD to be much faster than triple charge injection, which in turn determines the recovery-time for \(\tau < 0\). Given that the X2 emission of this QD is stronger than we typically see in other QDs, we could envision the presence of an acceptor or donor impurity that
lak, S. Charbonneau, D. Leonard, R. Leon, P. M. Petroff, and J. L. Merz, Phys. Rev. B 54, 11548 (1996).
[12] P. Michler, A. Kiraz, C. Becher, W.V. Schoenfeld, P.M. Petroff, Lidong Zhang, E. Hu, and A. Imamoglu, Science 290, 2282 (2000).
[13] C. Santori, M. Pelton, G. Solomon, Y. Dale, and Y. Yamamoto, Phys. Rev. Lett. 86, 1502 (2001).
[14] D.V. O’Connor and D. Phillips, *Time Correlated Single Photon Counting* (Academic Press, London, 1984).
[15] G. Bacher, R. Weigand, J. Seufert, V.D. Kulakovskii, N.A. Gippius, A. Forchel, K. Leonardi, and D. Hommel, Phys. Rev. Lett. 83, 4417 (1999).
[16] I. Robert et al. (to be published)
[17] O. Benson, C. Santori, M. Pelton, and Y. Yamamoto, Phys. Rev. Lett. 84, 2513 (2000).
[18] M. Paillard, X. Marie, P. Renucci, T. Amand, A. Jbeli, and J.M. Gérard, Phys. Rev. Lett. 86, 1634 (2001).