Single pairs of time-bin-entangled photons

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Time-bin-entangled photons are ideal for long-distance quantum communication via optical fibers. Here we present a source where, even at high creation rates, each excitation pulse generates, at most, one time-bin-entangled pair. This is important for the accuracy and security of quantum communication. Our site-controlled quantum dot generates single polarization-entangled photon pairs, which are then converted, without loss of entanglement strength, into single time-bin-entangled photon pairs.

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I. INTRODUCTION

Entanglement of flying qubits is a fundamental principle of quantum information and communication [1,2], and is at the basis of quantum communication protocols such as quantum key distribution [3], quantum teleportation [4], and quantum secure direct communication [5]. Commonly used qubits are polarization-entangled photons generated by spontaneous parametric down-conversion [6]. There are three important limitations to this approach.

First, in optical fibers, polarization encoding is prone to thermal and mechanical disturbances, which affect the fiber’s birefringence and thereby also the polarization of the out-coupled photons. This phenomenon, called polarization-mode dispersion, is a major problem for quantum communication in real-world implementations [7,8]. A solution is time-bin entanglement [9,10]: quantum information encoded in the arrival time of photons. Time-bin entanglement is robust [11,12] and has enabled distribution of entangled photons over 300 km of optical fiber [13].

Second, parametric down-conversion is a random process and follows Poissonian statistics. If \( p \) is the probability for a pump pulse to create a pair of entangled photons, there is a probability of order \( p^2 \) to create two or more pairs of entangled photons from the same pump pulse. The generation of multiple pairs makes it unclear which photon is entangled with which, and thus reduces the accuracy and security of the quantum communication. A possible solution is to reduce the pair-creation rate, so that \( p^2 \) becomes very small. Of course, the data transmission rate is then reduced as well. A more rigorous solution, which has been shown to work for polarization entanglement (but not yet for time-bin entanglement), is to generate single pairs of entangled photons by exciting a biexciton (XX) in a semiconductor quantum dot [14–22]. A XX is an excited state consisting of two electrons and two holes. Recombination of one electron and one hole occurs under emission of one photon (the XX photon) and brings the quantum dot to the exciton (X) state, which then decays further to the ground state, again emitting one photon (the X photon). If the excitation laser pulse is longer than the lifetime of the XX (typically around 1 ns), reexcitation and multiple-pair emission can occur. In contrast, if the excitation laser pulse is much shorter than the XX lifetime, each laser pulse can create not more than one photon pair.

Third, there is the problem of scalability. A great challenge is to make large arrays of identical entangled photon sources for use in quantum information applications. Quantum dots offer scalable, nanosize sources of entangled photons. Quantum-dot sources are energy efficient, in the sense that much less optical pump power is required per entangled photon pair as compared to down-conversion sources. Some of us recently reported an array of site-controlled pyramidal InGaAs_{1−x}N_{x} quantum dots, with areas where up to 15% of the quantum dots emit polarization-entangled photons [22,23].

Is it possible to combine the aforementioned solutions and generate single pairs of time-bin-entangled photons? In 2005, Simon and Poizat suggested a method [24], but the lack of a suitable metastable state in the quantum dot, essential in their proposal, has prevented implementation. A recent approach by Jayakumar et al. [25] is based on a quantum dot, but still allows for a probability of order \( p^2 \) to create two photon pairs from the same incoming laser pulse because they split their pulse into two.

Here, we demonstrate a source of single time-bin-entangled photon pairs, using a site-controlled quantum dot. We overcome the need of a metastable state and avoid the multiphoton emission probability \( p^2 \) that is inherent to all previous implementations of time-bin-entanglement generation. Our source opens up possibilities for the transfer of spin-photon entanglement [26–29] over long distances, hyperentanglement [30], quantum dense coding [31], and deterministic entanglement purification [32], and could be developed further for integration in compact and scalable quantum information devices.

II. POLARIZATION–TIME-BIN INTERFACE

In our experiment, a site-controlled semiconductor quantum dot generates single photon pairs in the polarization-
entangled state $|\langle HH \rangle + e^{i\phi} |VV\rangle\rangle / \sqrt{2}$, where $H$ ($V$) stands for horizontal (vertical) polarization and $\phi$ is a phase angle. The probability of multiple-pair emission from one excitation pulse is strongly suppressed. The polarization-entangled state is then converted into the time-bin-entangled state $|\langle ee \rangle + e^{i(\phi+\sigma)} |ll\rangle\rangle / \sqrt{2}$, where $e$ ($l$) stands for the early (late) time bin. This conversion occurs in a polarization-time-bin interface, consisting of an unbalanced Mach-Zehnder interferometer with polarizing beam splitters (PBSs) and a polarizer (Fig. 1). The horizontal (vertical) term of the two-photon wave function takes the short (long) path, and is thus converted into the early (late) term of the time-bin-entangled wave function. The path-length difference determines the separation between the two time bins, 4.3 ns in our case, and also the phase $\sigma$. The polarizer at 45° behind the second PBS erases all polarization entanglement, leaving only the time-bin entanglement. In a quantum communication network, where messages are encoded in the polarization bases, this interface can realize the transfer of quantum information to the time-bins. The resulting time-bin-entangled qubits could be faithfully transmitted through an optical fiber suffering from mechanical vibrations or thermal instability. The 50% intensity loss at the polarizer could, in principle, be avoided by operating a fast Pockels cell in front of the polarizer. This Pockels cell should then switch between the early time bin and the late time bin, and, by polarization rotation, ensure that the early and late parts of the wave function obtain the same polarization. Other realizations of conversion between polarization and time-bin entanglement (with spontaneous parametric down-conversion) are described in Refs. [33–36].

III. XX-X RADIATIVE CASCADE

A. Pyramidal quantum dot

In the present experiment, the single pairs of polarization-entangled photons are produced in one pyramidal quantum dot [37]. Our sample consists of an array of pyramids [Fig. 2(a)], where each pyramid contains a single embedded quantum dot. Figure 2(b) shows a schematic sketch of the internal structure of a pyramid, consisting of several epitaxial layers. The pyramids were grown by metal-organic vapor phase epitaxy in 7.5-μm-pitch tetrahedral recesses etched in (111)B-oriented GaAs. Details of the growth method are described in Ref. [22].

B. Excitation

We excited one pyramidal quantum dot with 639 nm, 100 ps, 80 MHz laser pulses. The diameter of the excitation spot was 1 μm, much smaller than the 7.5 μm distance between the pyramids, so that we could easily capture the optical emission from just one quantum dot. The emission spectrum [Fig. 2(c)] shows clear XX, X, and trion (X$^-$) emission lines. In our experiment, we spectrally selected the XX and X emission. Weaker emission lines on the sides [visible in Fig. 2(c)] appear only under strong excitation of the quantum dot, when a significant fraction of the pulses creates more than two electron-hole pairs in the quantum dot. In such case, charge carriers at higher energy levels, by electrostatic interaction, shift the emission energy of the charges at the lowest energy levels. By spectrally rejecting the side emission lines, we made sure that for one excitation pulse, only one XX photon and one X photon were measured. During all measurements, the sample was maintained at a constant temperature of 5.1 K in a closed-cycle cryostat. Except when stated otherwise, we used an excitation power of 150 nW at the sample. At this power, the average number of created electron-hole pairs per excitation pulse was 0.5.

C. Characterization of the XX-X radiative cascade

We observed that the XX line has a quadratic dependence on power and the X line has a nearly linear dependence up to saturation [Fig. 2(d)]. Time-resolved photoluminescence measurements show a XX lifetime of $0.72 \pm 0.03$ ns and an X lifetime of $1.25 \pm 0.04$ ns. Both XX and X emission are unpolarized. We measured a fine-structure splitting of $S = 0.6 \pm 0.2$ μeV [Fig. 2(e)], corresponding to a precession period of the X spin state of $h/S = 7.3 \pm 1.9$ ns, where $h$ is Planck’s constant [18]. Since this period is much longer than the X lifetime, spin precession has only a small influence on the correlations between the polarization of the XX and X photons. This small value for the fine-structure splitting, which is a special feature of our pyramidal quantum dots [22], enables measurement of quantum entanglement without the need for strict temporal postselecion and is thus of great importance for practical implementations of quantum communication.

IV. SINGLE PAIRS OF POLARIZATION-ENTANGLED PHOTONS

A. Single-photon correlation measurements

Single-photon time-resolved correlation measurements were performed by splitting the emission into two arms, each containing a spectrometer and an avalanche photodiode (APD). The outputs of the two APDs were connected to the time tagging module, which registers the differences in arrival time between signals from both APDs. To determine the sub-
Poisonian statistics, Hanbury-Brown Twiss autocorrelation measurements were performed by selecting with both spectrometers the X emission or the XX emission. Time-resolved cross-correlation measurements, to study the emission from the XX-X radiative cascade, were performed by selecting with one spectrometer the XX emission and with the other the X emission.

B. Single-photon statistics

Hanbury-Brown Twiss autocorrelation measurements gave $g^{(2)}(0) = 0.13$ as the maximum for the central peak for X photons [Fig. 3(a)] and $g^{(2)}(0) = 0.22$ for XX photons [Fig. 3(b)], while XX-X cross-correlation measurements gave $g^{(2)}(0) = 3.5$ [Fig. 3(c)]. These correlation results show that the quantum dot is a sub-Poissonian source, i.e., a single-photon source, of photon pairs from the XX-X cascade. Note that for a Poissonian source, a down-conversion source, for example, the central peak in an autocorrelation measurement, is as high as the neighboring peaks.

The measured nonzero value for $g^{(2)}(0)$ in the autocorrelation measurements can be explained by reexcitation. Some biexcitons and excitons decay within the 100-ps-long excitation pulse, so that the quantum dot can be excited again by a pump photon, leading to emission of more than one XX photon or X photon. Purer single-photon statistics can thus be obtained by reducing the excitation pulse duration. We therefore also took Hanbury-Brown Twiss autocorrelation measurements to study the emission from the XX-X radiative cascade.

FIG. 2. (Color online) Pyramidal quantum-dot emitter. (a) Scanning electron microscopy image of the sample, containing an array of pyramids with a 7.5 μm pitch. Each pyramid contains a quantum dot. Scale bar: 30 μm. Tilt angle: 60°. (b) Sketch of the internal epitaxial layer structure of a pyramid. The quantum dot is marked in red. (c) Emission spectrum of one quantum dot at two excitation powers. Indicated are the exciton (X), biexciton (XX), and trion (X*) emission lines. (d) The XX (red) and X (black) emission intensities vs excitation power. Fits to power functions show a quadratic dependence (power 2.0) for the XX emission and a nearly linear dependence (power 1.1) for the X emission. (e) Energy separation between XX and X vs polarization angle. This measurement determines the fine-structure splitting: the energy splitting between the two X spin states is 0.6 ± 0.2 μeV.

FIG. 3. Single-photon time-resolved correlation measurements. (a) X-X autocorrelation (start: X, stop: X), (b) XX-XX autocorrelation (start: XX, stop: XX), and (c) XX-X cross correlation (start: XX, stop: X) with 639 nm, 100 ps, 80 MHz excitation pulses at 150 nW. (d) X-X autocorrelation, (e) XX-XX autocorrelation, and (f) XX-X cross correlation with 750 nm, 3 ps, 80 MHz excitation pulses at 150 nW.
To determine the full quantum state of the photon pair, a standard quantum state tomography was performed, following the method described by James et al. [38]. We performed 16 time-resolved cross-correlation measurements, with one spectrometer selecting the XX emission and the other the X emission, with the following polarization selections: $HH, HV, HD, HL, VH, VV, VD, VL, DH, DV, DD, DL, LH, LV, LD$, and $LL$. The polarization selections were made using quarter-wave plates, half-wave plates, and polarizers. Additional half-wave plates were used to ensure that the same polarization always enters the spectrometers, and thus to avoid any effect of polarization sensitivity of the spectrometers on the results. For each measurement, we used an integration time of 300 s.

The observed density matrix [Figs. 4(b) and 4(c)] shows a polarization-entangled state with a concurrence of 0.54 ± 0.03, where a positive value indicates quantum entanglement [39]. The measured coincidence numbers, on which this density matrix is based, are given in the Appendix. The fidelity to the maximally entangled state ($|HH⟩ + |VV⟩)/√2$ is $0.722 ± 0.006$, where 0.5 is the classical limit. The fidelity to $(|HH⟩ + e^{0.144iπ}|VV⟩)/√2$, also a maximally entangled state, is $0.758 ± 0.006$. It has been observed before that the highest fidelity in quantum dots is found with respect to a state $(|HH⟩ + e^{iϕ}|VV⟩)/√2$ with a small phase $ϕ$ [21]. This phase could be attributed to precession of the $X$ spin state, or to a birefringence in the sample or in the setup. In our calculations, we used a time window of 3 ns, which is 2.4 times the $X$ lifetime, thus including more than 90% of the correlation counts. For the calculation of the density matrix, we used a maximum-likelihood estimation, following Ref. [38].

FIG. 4. (Color online) Polarization entanglement. (a) Measurements of correlations between the polarization of the $XX$ photon and the polarization of the $X$ photon. The first letter represents the polarization of the $XX$ photon and the second letter represents the polarization of the $X$ photon, where $H, V, D, A, L$, and $R$ stand for horizontal, vertical, diagonal, antidiagonal, left-handed, and right-handed polarization, respectively. (b) Real and (c) imaginary parts of the measured density matrix of the polarization quantum state of the two photons from the $XX$-X cascade. The positive matrix elements are orange and the negative matrix elements are green.
The fidelity $F(\rho, \psi)$ of a density matrix $\rho$ to a pure state $|\psi\rangle$ is calculated from $F = \langle \psi | \rho | \psi \rangle$.

V. SINGLE PAIRS OF TIME-BIN-ENTANGLLED PHOTONS

A. Time-bin quantum state tomography

To obtain single time-bin-entangled photon pairs, we converted the single polarization-entangled photon pairs from the pyramidal quantum dot using our polarization–time-bin interface (Fig. 1). As already mentioned, the polarizer in this interface eliminates all polarization entanglement. We analyzed the time-bin quantum state by time-bin quantum state tomography [35,40]. The photons have to be sent again through an unbalanced interferometer, so that the early and late terms of the two-photon wave function overlap with each other. Essentially, we used the method of Bussières et al. [35], where PBSs in the interferometer enable the time-bin quantum state tomography by employing wave plates and polarizers in a similar way as in polarization quantum state tomography. For the time-bin quantum state tomography, we used an integration time of 1800 s for each measurement.

The essence of this form of time-bin quantum state tomography is that the time-bin entanglement is converted back into polarization entanglement. In our experiment, this conversion back into polarization entanglement was established by two elements in our setup. First, the time-bin-entangled photons were sent back through the unbalanced Mach–Zehnder interferometer, as indicated by the orange paths in Fig. 1. Second, we measured with our correlation electronics not just the time difference between the XX photon and the X photon, as in the regular polarization quantum state tomography, but we measured the timing of the XX photon and the X photon with respect to the trigger signal from our pulsed excitation laser. Based on their arrival times with respect to the laser pulse, the detected photons were discriminated into three categories: photons that traveled twice the short path in the interferometer, photons that traveled once the short path and once the long path (here it is fundamentally uncertain whether they first traveled the short path and then the long path or first the long path and then the short path), and photons that traveled twice the long path [Fig. 5(a)]. The photon pairs traveling twice the short (long) path were, with certainty, in the $|ee\rangle$ ($|ll\rangle$) state and are therefore, as a result of our measurement, not time-bin entangled. The pairs of photons that traveled once the short path and once the long path were entangled and therefore we temporally postselected those photons, again using a time window of 3 ns. We measured the correlations in the polarizations of the postselected pairs. Here, a measured $V$ ($H$) polarization means that the photon was in the early (late) state. Likewise, a measured $D$ polarization corresponds to $(|l\rangle + |e\rangle)/\sqrt{2}$. $A$ corresponds to $(|l\rangle - |e\rangle)/\sqrt{2}$, $L$ to $(|l\rangle - i|e\rangle)/\sqrt{2}$, and $R$ to $(|l\rangle + i|e\rangle)/\sqrt{2}$. Thus, time-bin entanglement can be measured by measuring polarization entanglement. The postselection and the concomitant loss of intensity could, in principle, be avoided by polarization rotation with a fast Pockels cell.

B. Time-bin density matrix

The resulting density matrix is shown in Figs. 5(b) and 5(c). The measured coincidence numbers, on which this density matrix is based, are given in the Appendix. The concurrence is $0.58 \pm 0.07$, which demonstrates time-bin entanglement. The fidelity to $(|ee\rangle + e^{0.672s}|ll\rangle)/\sqrt{2}$ is $0.74 \pm 0.02$. These values are, within experimental error, the same as the values we had obtained for polarization entanglement under the same excitation conditions, showing that the conversion takes place without loss of entanglement strength. As a result of traveling twice through the interferometer, the phase $\chi$ between the two components of the wave function is different from the phase that was measured with the polarization quantum state tomography.

VI. DISCUSSION

Compared to Poissonian parametric down-conversion sources of time-bin-entangled photons, our pyramidal quantum dot source has several advantages. First, the relative rate of pulses where more than one pair is emitted is strongly reduced, as shown by the antibunching data [Fig. 3]. Second, our type of sample offers the potential of scalability. The pyramidal quantum dot used in this study is just one among tens of thousands of very similar position-controlled quantum
dots on the sample, as, indeed, site-controlled pyramidal quantum dots have demonstrated to be the highest uniformity quantum-dot system to date [41,42]. Current research efforts in quantum-dot growth will result in fabrication techniques with even higher control of shape, composition, and uniformity. The goal is to make arrays where all quantum dots emit strongly entangled photons in nearly identical quantum states. Third, the generation efficiency (in terms of power) is much higher than for nonlinear crystals.

It must be said that down-conversion sources still reach better entanglement fidelities. Yet, quantum-dot sources show rapid progress towards purer (polarization-)entangled states [43–47]. A recent approach is to create polarization-entangled photons in a [111]-grown quantum dot embedded in a nanowire [47,48], where the cylindric symmetry reduces the fine-structure splitting [49]. The nanowire shape additionally offers high brightness [50] and coherent emission [47], as well as a directional Gaussian emission profile [51].

Possible applications of single pairs of time-bin-entangled photons include quantum communication via fiber, where, thanks to the time-bin encoding, the two-photon quantum state is not affected by fiber instabilities. A grating could be used to split the XX beam from the X beam. After transmission through the fibers, it can be useful to convert the time-bin entanglement back into polarization entanglement, as is done in our setup, because the photon polarization can be easily manipulated. Other possible applications involve hyperentanglement (simultaneous polarization and time-bin entanglement), which could be obtained by removing the polarizer behind the interferometer in Fig. 1. Schuck et al. [31] have shown that it is possible to use time-bin entanglement, in addition to polarization entanglement, for a more complete Bell-state analysis and quantum dense coding. Finally, by expanding the setup to a form like the one proposed by Sheng and Zhou [32], it should be possible to realize deterministic polarization-entanglement purification with single-photon pairs.

VII. CONCLUSION

We generated single time-bin-entangled photon pairs from a pyramidal quantum dot. The strength of entanglement is maintained in our polarization–time-bin interface. Our source of single time-bin-entangled photons could be used in a quantum communication scheme, where entanglement is preserved in optical fibers suffering from mechanical or thermal vibrations. If desired, the entanglement can be converted back into the polarization bases, so that the photons could be further processed by widely available polarization-sensitive optical components. Recently, generation of entangled photons from a semiconductor quantum dot at a telecom wavelength has been realized [52]. By combination with our technique, and with resonant excitation [25,45] and electrical injection [20], one could make an electrically driven on-demand source of single time-bin-entangled photons at a telecom wavelength.

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APPENDIX

The measured numbers of coincidence counts for the polarization quantum state tomography are given in Table I. The selected time window was 3 ns. The first letter indicates the XX polarization and the second letter indicates the X polarization. For this measurement, the photons were sent directly to the tomography setup, without going through the interferometer. Calculation based on these results gives the following density matrix in the basis \([|HH\rangle,|HV\rangle,|VH\rangle,|VV\rangle\):

\[
\begin{pmatrix}
0.405 & 0.019 + 0.022i & -0.012 - 0.014i & 0.333 - 0.158i \\
0.019 - 0.022i & 0.110 & 0.008 - 0.056i & -0.053 - 0.011i \\
-0.012 + 0.014i & 0.008 + 0.056i & 0.112 & -0.020 - 0.028i \\
0.333 + 0.158i & -0.053 + 0.011i & -0.020 + 0.028i & 0.373
\end{pmatrix}.
\]

This matrix is given in Figs. 4(b) and 4(c).

The measured numbers of coincidence counts for the time-bin quantum state tomography are given in Table II. Again a time window of 3 ns was selected and again the first (second) letter indicates the XX (X) polarization. The photons were sent...
through the polarization–time-bin interface to create time-bin entanglement, and again through the interferometer for the time-bin tomography measurement. In this measurement, $H$ corresponds to $I$, $V$ to $e$, $D$ to $(l + e)/\sqrt{2}$, and $L$ corresponds to $(l - ie)/\sqrt{2}$. The calculation based on the results in Table II gives the following density matrix in the basis $\{|ll\rangle, |le\rangle, |el\rangle, |ee\rangle\}$:

$$
\begin{pmatrix}
0.303 & 0.071 - 0.067i & 0.054 + 0.069i & -0.178 + 0.297i \\
0.071 + 0.067i & 0.106 & 0.060 + 0.057i & -0.063 + 0.037i \\
0.054 - 0.069i & 0.060 - 0.057i & 0.108 & 0.071 + 0.048i \\
-0.178 - 0.297i & 0.063 + 0.037i & 0.063 - 0.037i & 0.483 \\
\end{pmatrix}.
$$

(A2)

This matrix is given in Figs. 5(b) and 5(c).


