Direct photonic coupling of a semiconductor quantum dot and a trapped ion

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Coupling individual quantum systems lies at the heart of building scalable quantum networks. Here, we report the first direct photonic coupling between a semiconductor quantum dot and a trapped ion and we demonstrate that single photons generated by a quantum dot controllably change the internal state of an Yb+ ion. We ameliorate the effect of the sixty-fold mismatch of the radiative linewidths with coherent photon generation and a high-finesse fiber-based optical cavity enhancing the coupling between the single photon and the ion. The transfer of information presented here via the classical correlations between the σ+-projection of the quantum-dot spin and the internal state of the ion provides a promising step towards quantum state-transfer in a hybrid photonic network.

Single atoms and ions are among the key players in the realization of elementary quantum information processing protocols [1]. High-fidelity state preparation and readout paired with long coherence times of internal and external degrees of freedom have enabled the implementation of small quantum processing units [2–4]. In recent years, optically active spin qubits in the solid state [5], such as semiconductor quantum dots (QDs) [6] and impurity centers in diamond [7], have emerged as complementary systems. Albeit having shorter coherence times, these systems offer ultrafast quantum control via larger electrical dipole moments [8] and on-chip integration [9–13] without the need for a continuously operating trap architecture. While prototype photonic networks of identical constituents, such as single atoms [14,15], or spins in diamond [16,17] have been demonstrated, the concept of hybrid quantum networks has recently been proposed as an exciting alternative [18–20]. Initial progress towards hybrid systems includes single-photon sources coupled to atomic vapours [21,22] and superconducting qubits to solid-state spin ensembles [23,24]. These specific experiments rely on large ensembles and, in some cases, spatial proximity to within the coherence length of the interaction. The formation of a modular network where fundamentally differing individual quantum systems communicate over long distances is an important goal, which so far has remained elusive.

Here, we report the first direct photonic coupling between a semiconductor QD spin and a trapped ion and demonstrate that single photons from a QD change the internal state of an Yb+ ion efficiently, despite a significant mismatch in the optical properties of the two systems. To achieve this we link the atomic and solid-state nodes with single photons transmitted through an optical fiber [Fig. 1(a)].

The atomic node consists of a single 174Yb+ ion in a radio-frequency (RF) Paul trap located inside a recently developed fiber-based high-finesse Fabry-Perot cavity [25]. The miniature RF-Paul trap is made of two very fine tungsten needles at 100 µm distance giving rise to trap frequencies in the range of 2π × 1–3 MHz. Ion fluorescence at 369 nm is collected by an in-vacuo objective (NA = 0.27) with 2% collection efficiency and guided onto a photomultiplier tube with 14% efficiency.

The cavity [25] is made from two single mode fibers (125 µm diameter) where a negative lens is machined on each tip (radii of curvature −300 ± 50 µm). After the machining process, the fibers are coated with a high reflectivity dielectric coating (asymmetric coating, T = 10 ppm and 100 ppm) resulting in a cavity finesse of F = 20000. The length of the cavity is 170 ± 10 µm and the mode waist is about 6.1 µm. The ion interacts with a single mode of the optical cavity through the 3D[3/2]1/2−2D[3/2]1/2 transition at 935 nm [Fig. 1(b)] in the intermediate coupling regime with cavity quantum electrodynamics (cQED) parameters (g,κ,γ) = 2π × (1.6, 25, 2.1) MHz. Here, g denotes the coupling strength between the ion and the cavity mode, κ the decay rate of the cavity field, and γ the decay rate of the atomic dipole moment. Figure 1(c) (right) displays the corresponding absorption spectrum for this transition. The cavity-modified decay probability from the 3D[3/2]1/2 to the D3/2 state is given by the bare probability of 2% plus the Purcell factor of 2C0/(2C0 + 1) where we have used the cooperativity C0 = g2/(2κγ), nominally 2.4(5)% in the experiment.

The solid-state node consists of an Indium Arsenide (InAs) QD in a Schottky diode placed inside a 4.2-K magneto-optical bath cryostat [Fig. 1(a)] giving access to neutral and negatively charged QD configurations, as well as electric and magnetic field tuning of the optical transitions [27]. The samples contain a distributed Bragg reflector formed from alternating GaAs/AlGaAs layers.
below the QD layer to increase the collection efficiency around 920-960 nm. A superhemispherical zirconia solid immersion lens (Weierstrass geometry) mounted on the top surface of the sample is used to further increase the photon collection and improve spatial resolution. Resonant optical excitation and collection from single QDs is achieved using a confocal microscope with a 90:10 beam-splitter. We collect both QD fluorescence and laser scattering via a 0.5 NA aspheric lens. Linearly polarized excitation, and cross-polarized detection allows us to suppress the laser scattering by a factor of 10^5. For an excitation intensity of $I = I_{\text{sat}}$, where $I_{\text{sat}}$ denotes the measured laser intensity for which the steady-state QD excited state population is 1/4, the signal-to-laser background ratio is 70:1 (20:1) in single- (two-) laser experiments. Pulsed laser excitation is realized using acousto-optic modulators. Figure 1(c) (left) shows the absorption spectrum for the $|0\rangle \rightarrow |+1\rangle$ transition illustrated in Fig. 1(b). The inset displays the full emission spectrum consisting of the zero-phonon line and the spectrally broad phonon sideband.

The resonantly generated QD photons are coupled into a 50-m long optical fiber and transmitted to the atomic node located 25 meters away. The overall transmission probability of $5 \times 10^{-4}$ is given by the photon-extraction efficiency from the QD sample (3.5% into the first lens) and losses in the optical link from the QD to the ion (1.4% transmission from the first lens to the cavity mirror).

We first demonstrate the excitation of the atomic node with single photons from the solid-state node. We prepare the ion in the lowest Zeeman level $|m_J = -3/2\rangle$ of the $^2D_{3/2}$ manifold by optical pumping. This state has a natural lifetime of 50 ms and absorbs only $\sigma^+$-polarized photons. We then generate a single-photon stream from the bright neutral exciton transition ($|0\rangle \rightarrow |+1\rangle$) of the QD, as illustrated in Fig. 1(b). This transition has a radiative linewidth of $\Gamma_{\text{QD}} = 2\pi \times 250(10)$ MHz, which is broadened further by spectral diffusion processes.

We drive the QD with an excitation intensity of $I = 0.5I_{\text{sat}}$ for a variable time $T$, which determines the total number of photons transmitted to the atomic node [see Fig. 2(a)]. Absorption of a QD photon transfers the ion into the $^2S_{1/2}$ electronic ground state with a probability given by the intermediate $^3D_{3/2}$ state’s cavity-modified branching ratio of 91:9. We then probe the $^2S_{1/2} \rightarrow ^2P_{1/2}$ transition of the ion, where fluorescence at 369 nm verifies a successful state change. In contrast, we infer that a photon absorption event did not take place if the ion remains in the ‘dark’ $^2D_{3/2}$ state.

Figure 2(b) displays the measured ion-state transfer probability as a function of $T$ for a fixed photon rate impinging on the fiber cavity of $\gamma_{\text{QD}} = 9(1) \times 10^4$ s$^{-1}$. The exponential saturation behavior, displayed by the solid curve,
yields a characteristic transfer time of 1.08(4) ms, which corresponds to 97(9) QD photons impinging on the cavity. This yields a single-photon absorption probability of \( p_{\text{abs}} = 1.0(2)\% \) at this excitation power. This is a conservative estimate as it includes the 13% of the QD photons that are red-detuned by a few hundred GHz due to phonon-assisted emission [30, 31], as seen in the inset of Fig. 1(c) and do not interact with the ion.

Figure 3(a) shows the dependence of the ion-state transfer probability on the spectral overlap between the QD photons and the ion transition. We tune the spectrum of QD emission in the strong excitation regime \( (I = 11I_{\text{sat}}) \) across the ion-cavity resonance and monitor the internal state of the ion [29]. The recorded state-transfer probability arises from the convolution of the QD single-photon spectrum, \( S(\omega) \), with the cavity-coupled ion absorption spectrum, \( L(\omega) \), as shown in the inset of Fig. 3(a), thus providing a measure of the spectral bandwidths of the two systems. As expected, the detuning dependence follows the Mollow-triplet signature of the QD emission spectrum. The narrow peak at zero detuning stems from the coherently scattered component as well as the residual laser due to imperfect suppression and its measured width, \( \approx 20 \text{ MHz} \), is set directly by \( L(\omega) \).

The solid curve gives the absorption spectrum calculated from the optical Bloch equations for the QD emission...
and ion-cavity absorption [29]. The highest probability of single-photon absorption for coherently scattered QD photons occurs at the exact cavity-ion resonance frequency, in stark contrast to the spectrally mismatched incoherent counterpart.

Figure 3(b) displays the dependence of the ion-state transfer probability on the QD excitation laser intensity. We see a pronounced efficiency increase in the low excitation regime, where the dominant contribution to the QD photon spectrum is inherited from the continuous-wave (cw) excitation laser field [31]. We measure a maximum value of 1.2(2)% at \( I = 0.1I_{\text{sat}} \), which corresponds to 1.4(2)% after accounting for the above-mentioned phonon-assisted emission. This value is comparable to the 1.8(2)% measured independently for cw laser of equivalent intensity, which represents the upper limit on ion-photon interaction observed with this cavity. In the high-excitation regime \( (I \gg I_{\text{sat}}) \), where the main contribution to resonance fluorescence is incoherent, the absorption probability reduces by an order of magnitude, consistent with the theoretical prediction [solid curve in Fig. 3(b)]. The laser-like absorption probability demonstrates that coherent scattering can provide an efficient interface for systems presenting a strong radiative linewidth mismatch. Quantum-network protocols based on coherent scattering [32] are inherently probabilistic owing to the sub-unity photon generation rate in this regime. That said, our results still predict an overall 20% higher efficiency of ion-state transfer over a deterministic generation scheme even when the coherent photon-generation probability is 10%.

As a prerequisite for quantum-state transfer, we demonstrate classical communication between our solid-state and atomic nodes such that the internal state of the ion and the projection of the QD spin are correlated. First, we switch to a negatively charged QD under 0.7-T magnetic field in Faraday configuration. This provides optical access to the spin projection of the QD states via the Zeeman splitting of the ground and excited states [Fig. 4(a)]. The \(|\uparrow\rangle-|\downarrow\downarrow\rangle\) transition is tuned on resonance with the ion transition, while the \(|\downarrow\rangle-|\uparrow\downarrow\rangle\) transition is off-resonant by 20 GHz. Second, we prepare the desired spin mixture through optical pumping by driving the \(\sigma^-\) transition with a pulse of variable duration, \(\tau\), and the \(\sigma^+\) transition with a 600-ns probe pulse. This alternating-pulses protocol provides a \(\sigma_z\) projection of the electron spin ranging from \(p_\uparrow = 0.072(2)\) to \(p_\uparrow = 0.81(1)\) [29]. State-preparation and photon-generation steps are alternated with a repetition rate of 670 kHz during the QD-ion interaction time of 700 \(\mu\)s. Once again, the ion is prepared in the \(m_J = -3\) Zeeman state of the \(2D_{3/2}\) manifold and absorbs \(\sigma^+\)-polarized photons leading to a state transfer to \(2S_{1/2}\).

Figure 4(b) presents the theoretically expected as well as the measured correlation between the QD spin state and the internal state of the ion. The dashed curve indicates the ideal correlation for our state-transfer experiments, while the solid curve represents the expected correlation calibrated for the presence of residual laser background. This sequence maps the \(\sigma_z\)-spin component of the QD to the internal state of the ion within an average uncertainty of 3.8%. Our results show that an arbitrary QD spin projection is reproduced faithfully on the atomic node in the form of \(2S_{1/2}\) internal-state projection.

The optical interface demonstrated here links two quantum systems with significantly different optical characteristics via the exchange of single photons. By coherent photon generation and cQED techniques we have achieved direct coupling between these systems with an efficiency that surpasses limitations set by their intrinsic properties. Our work can be extended to achieve faith-
ful quantum-state transfer and distant entanglement between a QD and an ion. The hyperfine states of the trapped ion (e.g. in $^{171}$Yb$^+$) can serve as a long-term quantum memory for the QD spin qubits. A key challenge for the implementation of such a scheme is reaching sufficient coupling strength between the nodes. While the overall efficiency is limited to $5 \times 10^{-6}$ here, we note that a 20-fold improvement in QD-photons collection efficiency has been recently achieved [33], a 10-fold reduction of loss in the optical link is straightforward, and increasing the absorption probability by a factor 30 through cavity improvements is within reach. Collectively, these technical steps could provide a three orders of magnitude improvement in our current node-to-node coupling in a not too distant future.

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