Quantum interference from remotely trapped ions

S Gerber$^1$, D Rotter$^1$, M Henrich$^1$, R Blatt$^{1,2}$, F Rohde$^3$, C Schuck$^3$, M Almendros$^3$, R Gehr$^3$, F Dubin$^{3,4}$
and J Eschner$^3$

$^1$ Institute for Experimental Physics, University of Innsbruck, Technikerstr. 25, A-6020 Innsbruck, Austria
$^2$ Institute for Quantum Optics and Quantum Information of the Austrian Academy of Sciences, Innsbruck, Austria
$^3$ ICFO-Institut de Ciências Fotòniques, Mediterranean Technology Park, Av del Canal Olímpic, E-08860 Castelldefels, Spain
E-mail: francois.dubin@icfo.es

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Abstract. We observe quantum interference of photons emitted by two continuously laser-excited single ions, independently trapped in distinct vacuum vessels. High contrast two-photon interference is observed in two experiments with different ion species, Ca$^+$ and Ba$^+$. Our experimental findings are quantitatively reproduced by Bloch equation calculations. In particular, we show that the coherence of the individual resonance fluorescence light field is determined from the observed interference.

$^4$ Author to whom any correspondence should be addressed.
1. Introduction

Interfacing static quantum bit registers and quantum communication channels is at the focus of current research efforts aiming at the realization of quantum networks. For implementing static qubits, trapped ions are known to be promising candidates as they provide optimum conditions for quantum information processing: they offer long coherence times for information storage [1], are easily manipulated coherently and their Coulomb interaction can be utilized to realize quantum logic gates [2, 3]. Single photons, on the other hand, are well suited for transmitting quantum information over long distances [4]. Hence, ion traps may constitute the nodes of a quantum network, while communication between remote nodes may be achieved through photonic channels transmitting quantum states and entanglement [5].

Several experiments have recently addressed the realization of an atom–photon qubit interface and its basic components. Studies were performed with various systems, including atom–cavity devices [6]–[11], atomic ensembles [12]–[14] and single-trapped atoms or ions [15]–[19]. With these latter systems, the reported experimental progress comprises: single-photon interference between two ions in the same trap and continuously excited by a laser [20, 21], interference between photons emitted by two atoms in distinct traps under pulsed-laser excitation [15], two-photon interference in single-ion fluorescence under continuous excitation [17], and finally two-photon interference between two ions confined in remote traps excited by short laser pulses [18]. The latter experiments have very recently led to the demonstration of remote entanglement between the two independently trapped ions [19, 22].

Entanglement between two remotely trapped atomic systems can be established through single- or two-photon interference. In the first case, two indistinguishable scattering paths of a single-photon interfere, such that its detection leads to entanglement of the atoms [23, 24]. In the latter case, coincident detection of two photons, each being entangled with one atom [16], projects the two atoms into an entangled state [25]. A detailed comparison of the two methods is found in [26].

As originally shown by Hong, Ou and Mandel [27], two indistinguishable photons impinging simultaneously on two input ports of a beam splitter show photon coalescence, i.e. both photons will leave the beam splitter in the same output mode. Indistinguishability of the input photons is then marked by a vanishing coincidence rate between the two output ports,
corresponding to the generation of two-photon states. The contrast of two-photon interference is reduced, however, when the spatial or temporal overlap of the incident photon wave packets is only partial, such that the photons become distinguishable [11].

In the experiments reported here, we study the conditions to achieve entanglement between distant atoms with two promising systems, Ba\(^+\) and Ca\(^+\) ions. We observe and characterize quantum interference between photons emitted by two ions of the same species which are independently trapped in different vacuum chambers, approximately one meter apart. Under continuous laser excitation, resonance fluorescence photons are collected in single-mode optical fibers and overlapped at two input ports of a beam splitter. At the output ports, correlations among photon detection events are evaluated. Controlling the polarization at the beam splitter inputs, we observe high-contrast two-photon interference. Recorded correlations allow us to quantify the coherence of the single-ion resonance fluorescence under continuous excitation.

2. Experimental apparatus

To realize the experiments, we developed a new assembly which integrates a linear Paul trap and two high-numerical-aperture laser objectives (custom HALO lenses designed by Linos). The diffraction-limited lenses are mounted inside the vacuum vessels on opposite sides of the trap; their positions are individually controlled by three-axes piezo-mechanical actuators and allow us to collect up to 8% of the total fluorescence (4% with each lens). The principal components of the experimental setup are described in the following.

2.1. HALO lenses

Each custom-designed objective for collecting the fluorescent light covers a numerical aperture of \(\text{NA} = 0.4\) with a focal length of 25 mm, thus collecting 4% of the total solid angle. The objectives are designed to be diffraction limited over a wide spectral range from ultraviolet to near infrared. The back focal length varies between 13.7 mm (at 400 nm) and 15.1 mm (at 870 nm), leaving several millimeters of space between the radio frequency electrodes of the ion trap and the front surface of the objective. These consist of four lenses which are all anti-reflection coated for the respective wavelengths (barium: 493 nm, 650 nm, 1762 nm; calcium: 397 nm, 422 nm, 850–866 nm). The calculated transmission of each HALO objective is for calcium: 98.8% (850–866 nm), 87.5% (at 729 nm), 97.2% (at 422 nm) and 95.7% (at 397 nm); for barium 97% (at 650 nm) and 98% (at 493 nm). The objectives are mounted on vacuum compatible slip-stick piezo translation stages (Attocube \(xyz\)-positioner, ANPxyz 100). These exhibit a travel of 5 mm and 400 nm step size that allows us to maximize the collection efficiency of the emitted resonance fluorescence and to address single ions individually when loading the trap with a string of ions.

The objectives have been characterized by various methods: for the first batch of objectives, of which one is used in the barium experimental setup, the manufacturer characterized the wavefront aberrations with a Zygo interferometer to be smaller than \(\lambda/10\) at 493 nm. Interference experiments proved this precision by obtaining single-ion self-interference with a 72% visibility [20]. The second batch of objectives was characterized using a wavefront sensor which showed a root mean square aberration of 0.073\(\lambda\) at 670 nm for three consecutive HALO lenses, measured over 25% of the aperture as depicted in figure 1.
Figure 1. Characterization of the high-numerical aperture lenses using a wavefront sensor. A 670 nm laser is coupled through a fiber. The fiber output is collimated by a HALO lens, and coupled through the setup consisting of the two opposing HALO lenses and the ion trap. The again collimated beam is sent to a wavefront sensor after passing the three HALO lenses. The beam covers 25% of the exit aperture of the lenses. Left: Gaussian beam profile on the wavefront sensor with a beam waist of 2.3 mm. Right: wavefront aberration of the beam giving an RMS value of $0.049 \mu m = 0.073 \lambda$ (at 670 nm).

2.2. Linear trap design

In the following experiments, we use a macroscopic linear trap, identical to the one described in [28].

2.3. Experimental setup for barium ions

The schematic experimental setup, located in Innsbruck, and the relevant level scheme of $^{138}$Ba$^+$ are shown in figure 2. One single Ba$^+$ ion is loaded into a ring trap (ion 1), a second ion into a linear trap (ion 2) using photoionization with laser light near 413 nm [29]. The ions are continuously driven and laser-cooled by the same narrow-band tunable lasers at 493 nm (green) and 650 nm (red) exciting the $|S_{1/2}\rangle \rightarrow |P_{1/2}\rangle$ and $|P_{1/2}\rangle \rightarrow |D_{3/2}\rangle$ transitions, respectively. After Doppler cooling, the ions are left in a thermal motional state in the Lamb–Dicke regime [30]. Laser parameters as well as magnetic fields at the location of the ions are calibrated by recording excitation spectra [31].

From each ion, and using the previously described HALO lenses, we collect 4% of the green resonance fluorescence which is mode-matched into single mode optical fibers. Photons are then guided to the entrance ports of a 50/50 beam splitter labeled by $I_1$ and $I_2$ corresponding to...
Figure 2. Sketch of the barium experimental setup (see text for details). Resonance fluorescence photons emitted by each ion are overlapped on a 50/50 beam splitter (GT stands for Glan–Thompson polarizers). Subsequently, the second-order correlation among photon detections is evaluated. Single photon arrival times are monitored with a time to amplitude converter (TAC) with 100 ps resolution. The inset shows the relevant electronic levels of $^{138}\text{Ba}^+$. 

to ion 1 and ion 2, respectively (see figure 2). The two output ports are denoted by $I_3$ and $I_4$. Note that the polarization of the two fiber outputs are controlled with Glan-Thompson polarizers to cancel out polarization fluctuations of the fiber transmission. The polarization of photons from ion 1 is set horizontal, while the polarization of photons from the second ion is varied. Finally, the beam splitter output ports are again coupled into single mode fibers to guarantee optimum spatial mode overlap. Correlations are obtained by subsequently monitoring and correlating the arrival times of photons at the two PMTs (Hamamatsu, H7421P-40 SEL with a specified minimum quantum efficiency of 40% at 493 nm) in a Hanbury–Brown and Twiss setup with sub-nanosecond time resolution. The visibility of the interferometer is measured to be 97% by sending laser light through the input fibers.

2.4. Experimental setup for calcium ions

The schematic of the setup for the $^{40}\text{Ca}^+$ experiment, located in Barcelona, and the corresponding level scheme are shown in figure 3. The setup consists of two linear Paul traps mounted in two vacuum vessels separated by about one meter. In each of the two traps a single $^{40}\text{Ca}^+$ ion is loaded. For this purpose a novel two-step photoionization method has been developed [32], which involves a single-pass frequency-doubling source of 423 nm light and a high-power LED at 389 nm. The ions are continuously excited and cooled on the $|S_{1/2}\rangle \rightarrow |P_{1/2}\rangle$ electronic transition by a laser at 397 nm. A second laser at 866 nm resonant with the $|P_{1/2}\rangle \rightarrow |D_{3/2}\rangle$ transition prevents optical pumping into the metastable $|D_{3/2}\rangle$ state. Both lasers are locked via transfer cavities to a reference laser, which itself is stabilized to an atomic cesium line via Doppler-free absorption spectroscopy [33]. Excitation parameters, e.g. laser intensities and frequencies, are set to maximize the count rate of the fluorescence light while maintaining sufficient laser cooling.
Figure 3. Sketch of the calcium experimental setup (see text for details). Fluorescence photons are coupled into the two inputs of a single mode fiber beam splitter. Stable polarization is then ensured using fiber polarization controllers (successive loops) and optionally a polarizing beamsplitter (PBS) for control. Second-order correlations are computed by recording arrival times at the detectors (PMT) using single-photon counting electronics. The overall time resolution of the detection setup is 1.5 ns. The inset shows the relevant electronic levels of a $^{40}\text{Ca}^+$ ion.

Fluorescence photons from both ions are collected using the previously described HALO lenses and then coupled into the two input ports of a single-mode fiber beam splitter. A polarizer before the input coupler projects onto horizontal polarization. Thereafter, fiber polarization controllers give full and independent control over the polarizations arriving at the two beam splitter inputs. The use of a fiber beam splitter instead of a free-space setup ensures optimum spatial overlap of the photon wave-packets at the beam splitter.

At the outputs of the fiber beam splitter, two photomultipliers (Hamamatsu, H7360-01) with specified $\eta \approx 0.25$ quantum efficiency detect arriving photons. Their arrival times are recorded with picosecond resolution (PicoHarp 300 photon counting electronics). The transit time spread of the photomultipliers sets the overall time resolution in this experiment to about 1.5 ns.

3. Theoretical description

In this part, we present a basic theoretical description to quantify two-photon interference for our experimental configuration. As depicted in figures 2 and 3, we label the two input ports of the beam splitter as $I_1$ and $I_2$. The corresponding fields, $\hat{E}_1$ and $\hat{E}_2$ are written in an $(\vec{e}_x, \vec{e}_y)$ polarization basis. In the following, fluorescence photons emitted on the $|P_{1/2}\rangle$ to $|S_{1/2}\rangle$ electronic transition are expressed in terms of Pauli lowering operators, $\hat{\sigma}^-_1$ and $\hat{\sigma}^-_2$, for ions 1 and 2, respectively. These are associated with the creation of a single photon. Without any loss of generality, we assume that $\hat{\sigma}^-_1$ transforms the field into $x$ polarization such that

$$\hat{E}_1(t) = \hat{\sigma}^-_1(t) \vec{e}_x,$$

$$\hat{E}_2(t) = \hat{\sigma}^-_2(t)(\cos \phi \vec{e}_x + \sin \phi \vec{e}_y),$$

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where $\vec{e}_x$ and $\vec{e}_y$ denote unit vectors in the x- and y-directions, respectively. In the previous expressions, let us note that the source-free part of the input fields has been neglected since noise contributions play a minor role in our experiments [34].

For an ideal 50/50 beamsplitter the transmission ($t$) and reflection ($r$) coefficients obey $-i \cdot t = r = \frac{1}{\sqrt{2}}$ [35]. Therefore, field operators at the output arms read

\begin{align}
\hat{E}_3(t) &= \frac{1}{\sqrt{2}} \{[\hat{\sigma}_1(t)e^{i\psi} + i \cos \phi \hat{\sigma}_2^-(t)]\vec{e}_x + [i \sin \phi \hat{\sigma}_2^-(t)]\vec{e}_y\}, \\
\hat{E}_4(t) &= \frac{1}{\sqrt{2}} \{[\hat{\sigma}_1(t)e^{i\psi} + \cos \phi \hat{\sigma}_2(t)]\vec{e}_x + [\sin \phi \hat{\sigma}_2(t)]\vec{e}_y\},
\end{align}

(2)

where $\psi$ represents a random phase between the fields emitted by the two ions. In fact, our experimental setup does not ensure sub-wavelength mechanical stability which imposes the inclusion of $\psi$. The overall correlation function can then be expressed as $G_{\text{tot}}^{(2)}(t, t + \tau) \propto \sum_{(i,j)=[x,y]} \{\hat{E}_3^\dagger(t)\hat{E}_4^\dagger(t + \tau)\hat{E}_4(t + \tau)\hat{E}_3(t)\}$ where $\hat{E}_{(3,4),x}^\dagger$ corresponds to the part of the field $\hat{E}_{(3,4)}^\dagger$ polarized along $x$. In the following, let us assume that the emission properties of ions 1 and 2 are identical. After averaging over all possible values of $\psi$, such that position-dependent non-classical correlations are excluded [36, 37], one finds

\begin{align}
G_{\text{tot}}^{(2)}(t, t + \tau, \phi) &\propto \frac{1}{2} \{[\hat{\sigma}_1^+(t)\hat{\sigma}_1^-(t + \tau)\hat{\sigma}_1^+(t + \tau)\hat{\sigma}_1^-(t)] + [\hat{\sigma}_2^+(t)\hat{\sigma}_2^- (t + \tau)\hat{\sigma}_2^+(t + \tau)\hat{\sigma}_2^-(t)] \nonumber \\
&\quad - 2 \cos^2(\phi)\langle \hat{\sigma}_1^+(t)\hat{\sigma}_1^-(t + \tau)\rangle\langle \hat{\sigma}_2^+(t + \tau)\hat{\sigma}_2^-(t) \rangle \\
&\quad + 2\langle \hat{\sigma}_1^+(t + \tau)\hat{\sigma}_1^-(t + \tau)\rangle\langle \hat{\sigma}_2^+(t)\hat{\sigma}_2^-(t) \rangle \}.
\end{align}

(3)

The first two terms in equation (3) represent second-order correlations between photons both emitted by the same ion, ions 1 and 2 for the first and second term, respectively. The third term shows the interference between two photons emitted by different ions. It can be rewritten as a product of the individual first-order correlation functions, $G^{(1)}$, as $-2 \cos^2(\phi)[G^{(1)}(t, t + \tau)G^{(2)}(t, t + \tau)^*]$. This term reflects the degree of indistinguishability of photons at the input ports of the beam splitter and consequently vanishes for $\phi = \pi/2$, i.e. for orthogonal polarizations. The fourth term represents the level of random correlations between photons emitted by different ions. This contribution reads as the product of the mean number of photons emitted by each ion, which we denote $\langle n \rangle^2$. Under the assumption of identical emission properties for ions 1 and 2, we can set $G_{1}^{(2)}(t, t + \tau) = G_{2}^{(2)}(t, t + \tau) = G^{(2)}(t, t + \tau)$ and $G_1^{(1)}(t, t + \tau) = G_2^{(1)}(t, t + \tau) = G^{(1)}(t, t + \tau)$ and obtain

\begin{align}
G_{\text{tot}}^{(2)}(t, t + \tau, \phi) &= \frac{1}{2} \{[G^{(2)}(t, t + \tau) - \cos^2(\phi)]G^{(1)}(t, t + \tau)\}^2 + \langle n \rangle^2].
\end{align}

(4)

At steady state ($t \rightarrow \infty$), the normalized correlation function is given by $g^{(2)}(\tau, \phi) = G_{\text{tot}}^{(2)}(\tau, \phi)/G_{\text{tot}}^{(2)}(\tau \rightarrow \infty, \phi)$, and it follows that

\begin{align}
g^{(2)}(\tau, \phi) = \frac{1}{2} g^{(2)}(\tau) + \frac{1}{2} [1 - \cos^2(\phi)] g^{(1)}(\tau)^2].
\end{align}

(5)

In equation (5), we note that the two-photon interference contribution (last term) has an amplitude which is given by the individual coherence properties of the interfering fields, even for $\phi = 0$, i.e. for identical polarization. Indeed, $|g^{(1)}|$ continuously decreases, starting from $|g^{(1)}(0)| = 1$, with a time constant equal to the coherence time of the resonance fluorescence. Hence, for $\phi$ set to 0, the coincidence rate vanishes due to both the discrete character of photon...
emission by a single-ion \((g^{(2)}(0) = 0\), anti-bunching) and the complete photon coalescence at the beam splitter. On the other hand, for \(\tau \neq 0\), the individual coherence of interfering fields is revealed by the \(g^{(2)}\) function.

4. Experimental results

Figure 4 presents the second-order correlation function, \(g^{(2)}_{\text{tot}}\), when photons emitted by the two ions are made fully distinguishable by setting \(\phi = 90\). The top (bottom) panel of figure 4 shows the results obtained with the barium (calcium) experiment. In both cases, the coincidence rate is such that \(g^{(2)}_{\text{tot}}(0, 90) \approx 0.5\), as expected from equation (5). Indeed, for input photons with orthogonal polarization, the last term in equation (5) reduces to \(1/2\) independent on \(\tau\). This reflects that among all possible correlations, one half occurs between photons emitted by different ions. These are here rendered distinguishable by their orthogonal polarization such that interference is absent. In figure 4, the result of our theoretical model (equation (5)) is also presented. It is obtained by solving the Bloch equations for our experimental parameters, including eight relevant Zeeman electronic levels for the ion internal states while neglecting motional degrees of freedom [31]. There is a good agreement for both measurements. We also present in the inset of figure 4 the normalized second-order correlation function of a single-ion, \(g^{(2)}\). It is obtained by blocking one fiber entrance and it shows almost ideal anti-bunching \((g^{(2)}(0) = 0.03\) with no background subtraction) and an optical nutation, \(g^{(2)}(13\,\text{ns}) = 2.9\).

Figure 5 shows experimental results obtained for \(\phi = 0\) such that photons impinging at the two input ports of the beam splitter are indistinguishable. As expected, the normalized number of coincidences strongly drops around \(\tau = 0\) which signals two-photon interference [17]. From the measured values of \(g^{(2)}_{\text{tot}}(0, 90)\) and \(g^{(2)}_{\text{tot}}(0, 0)\), we deduce that the two-photon interference contrast reaches \(89(2)\%\) and \(80(10)\%\) for the barium and calcium experiments, respectively. In the latter case, the contrast is limited by the time resolution of the photodetectors. Contrasts are furthermore derived from raw experimental data, without correcting for accidental correlations induced by stray light photons or dark-counts of the photodetectors.

As previously mentioned, individual coherence properties of interfering fields modify the overall correlation function for \(\phi = 0\). First, for \(\tau = 0\), incident photons are detected simultaneously such that these are fully indistinguishable and therefore the amplitude of two-photon interference is maximal. By contrast, for \(\tau \neq 0\) the degree of indistinguishability of the input photons is characterized by the temporal overlap of their wave packets. The length of these wave packets, in fact, corresponds to the photon coherence time which then governs the contrast of the interference [11]. This property is reflected by the last term of equation (5) which is also represented by a dashed–dotted line in figure 5. It continuously increases before saturating to \(1/2\) for \(|\tau| \approx 10\,\text{ns}\). This reveals that the coherence time of the resonance fluorescence does not exceed \(20\,\text{ns}\) in our experiments. This indicates that incoherent scattering is dominant, far longer coherence times would otherwise be observed. As a consequence, let us note that the amplitude of the optical nutation is reduced by \(\approx 5\%\) compared with the non-interfering case \((\phi = 90)\).

Finally, we present in figure 6 the number of coincidence counts as a function of the polarization angle, \(\phi\). Thereby the degree of distinguishability of interfering photons, and thus the amplitude of the two-photon interference, is smoothly tuned. From equation (5), one expects a \(\frac{1}{2}\left[1 - \cos^2 \phi\right] = \frac{1}{2} \sin^2 \phi\) dependence. The latter is represented by a solid line in figure 6, in good agreement with our experimental findings.
Figure 4. Normalized second-order correlation function for orthogonal polarization between $I_1$ and $I_2$ ($\phi = 90$, i.e. the non-interfering case). Top: experimental results obtained in the barium experiment. Data points are obtained after an accumulation of 30 min. Inset: normalized second-order correlation function when one fiber output port is blocked. Bottom: experimental results obtained in the calcium experiment. Data are obtained after 2.5 h of accumulation. In both panels, the result of our theoretical model (equation (5)) is displayed by the solid line. It is obtained by considering our experimental parameters, e.g. laser intensities and detuning, from a full eight-level Bloch equation model. The dashed and dash-dotted lines show the contributions of the first and second term of equation (5), respectively. Experimental data are all presented with 1 ns resolution, without background subtraction, including the variance obtained from shot noise (Poisson statistics at all times $\tau$).
Figure 5. Normalized second-order correlation function for indistinguishable photons impinging at the input ports of the beam splitter ($\phi = 0$). The top panel presents the results of the barium experiment while the bottom panel shows the experimental results obtained in the calcium experiment. As in figure 4, the result of our theoretical model (equation (5)) is displayed by the solid line, the dashed and dash-dotted lines showing the contributions of the first and second term of equation (5), respectively. Experimental data are again presented with 1 ns resolution, without background subtraction, including the variance obtained from shot noise (Poisson statistics). The two-photon interference contrast deduced from these measurements reaches 89(2) and 80(10)% for the barium and calcium experiments, respectively.
Figure 6. Number of coincidences $g_{\text{tot}}^{(2)}(0, \phi)$ for varying polarization angles, $\phi \in \{0, 26, 47, 64, 90\}$. The solid line shows the $\frac{1}{2} \sin^2(\phi)$ dependence of the coincidence rate at $\tau = 0$ as given by equation (5). Experimental results are obtained with the barium experiment.

5. Summary

In summary, we have observed high contrast interference between resonance fluorescence photons emitted by remotely trapped ions. Our experiments report studies with two species, namely barium and calcium. For both of them, large photon detection efficiency is reached by means of high numerical aperture HALO-lenses placed inside the vacuum vessels, close to the traps, with adjustable positioning by $xyz$-piezo actuators. For barium, we reach a detection count rate in a single mode as high as 25–30 kcps s$^{-1}$ while for calcium, we obtain 10–15 kcps s$^{-1}$. This discrepancy arises mainly from different fiber coupling efficiencies but also from different photodetector quantum efficiencies at 493 and 397 nm for barium and calcium, respectively. Furthermore, let us note that we have followed slightly different experimental approaches to demonstrate indistinguishability of resonance fluorescence photons. In the calcium experiment, we use an integrated fiber beam splitter together with passive polarization control, while in the barium case interference is achieved in free space. In the former, the contrast of the two-photon coalescence is somewhat reduced compared to the latter. To the best of our knowledge, this difference arises from the transit time spread of the photodetectors used in the calcium experiment. Apart from these technical differences, both species are equally suited for these kinds of experiments and their application for, e.g. atom–photon interfaces. Finally, both experimental results are quantitatively reproduced by eight-level Bloch equations. This allows us to quantify the coherence of resonance fluorescence photons by means of two-photon interference, in close analogy with the original work of Hong, Ou and Mandel [27].
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