Tailored photon-pair generation in optical fibers

Offir Cohen, Jeff S. Lundeen, Brian J. Smith, Graciana Puentes, Peter J. Mosley, and Ian A. Walmsley
Clarendon Laboratory, University of Oxford, Parks Road, Oxford, OX1 3PU, UK
(Dated: April 14, 2009)

We experimentally control the spectral structure of photon pairs created via spontaneous four-wave mixing in microstructured fibers. By fabricating fibers with designed dispersion, one can manipulate the photons’ wavelengths, joint spectrum, and, thus, entanglement. As an example, we produce photon-pairs with no spectral correlations, allowing direct heralding of single photons in pure-state wave packets without filtering. We achieve an experimental purity of $85.9 \pm 1.6\%$, while theoretical analysis and preliminary tests suggest $94.5\%$ purity is possible with a much longer fiber.

PACS numbers: 42.50.Dv, 42.50.-p, 42.65.-k

Optical quantum technologies such as photonic quantum computing [1], quantum cryptography [2], and quantum metrology [3] take advantage of entanglement between photons. Conventionally these technologies are based on qubit encoding, which uses a discrete variable (e.g. polarization). Entanglement can also exist between continuous variables (CVs), such as the momentum, time, position, or frequency of photons. Here we focus on frequency entanglement, which can be used to make timing measurements [4] and clock synchronization [5] more precise, and cancel the detrimental effect of dispersion in interferometers [7]. The alternative to qubit encoding, CV encoding, is a particularly powerful tool that increases the information capacity of each photon. This capacity can, for example, improve the security of quantum cryptography [8] and simplify quantum computing algorithms [9]. All of these applications require different degrees and types of entanglement, and therefore, it is important to understand how to controllably create and manipulate these qualities [6, 10, 11, 12].

Equally important, although it has received less attention, is the difficult task of eliminating entanglement entirely, which can play a deleterious role in the production of heralded pure-state single photons. In photonic quantum computing, photon state impurity increases the error rate of logic gates by limiting the visibility of the Hong-Ou-Mandel (HOM) [13] interference driving these gates [1]. Processes such as spontaneous parametric down-conversion (SPDC) and spontaneous four-wave mixing (SFWM) produce photons with a small probability, but always in pairs. In “heralding,” the detection of a photon in one beam from these sources indicates the presence of its twin and, thus, projects the other beam into a single photon state. Unfortunately, the frequencies and transverse momenta of photon pairs produced in SPDC or SFWM are typically entangled, arising from correlations due to energy and momentum conservation constraints [11] [14]. In this case, heralding projects the other photon into an impure (i.e. “mixed”) quantum state [15]. Although this entanglement can be removed (asymptotically) by spectral and spatial filtering, doing so seriously degrades source performance in terms of production rate and heralding efficiency [16]. Generating and eliminating entanglement in photon-pair production can be seen as opposite sides of the same goal: the control of the emitted joint frequency-momentum quantum state. Here we present evidence that through the flexibility of SFWM in fiber one can reach both objectives.

In SPDC, several approaches have been developed to control, directly at the point of production, the frequency state of the emitted photon pairs. However with bulk nonlinear materials, one is limited to two control parameters: pumping geometry [6, 12] and natural material dispersion [16]. This limits our ability to tailor photon-pair frequency states. Moreover, there has been little success controlling the spatial state of the emitted photons from bulk materials, resulting in poor coupling to waveguides such as fibers.

Optical fiber sources based on SFWM are now one of the brightest photon-pair sources even when filtered and offer significant advantages over bulk material SPDC. The photons, generated in single spatial modes, are ideal for mating with waveguides in integrated optical circuits, a promising platform for scalable photonic quantum computing [18]. Only recently has the full potential for control of the joint frequency state of photon pairs produced by SFWM in optical fiber been brought to light [19], and it has yet to be fully exploited experimentally [20]. A crucial difference from SPDC is that one can fabricate a photonic crystal fiber (PCF) to satisfy a wide range of dispersion requirements [21] for SFWM. This new control parameter more than compensates for loss of pump geometry control that is crucial for spectral state engineering in bulk materials [6, 12]. Thus, SFWM in optical fiber allows expanded control of the spectral structure of the produced photon-pair states compared to SPDC.

SFWM can be described as the virtual absorption of two pump ($p$) photons followed by the emission of a photon pair (the signal, $s$, and idler, $i$). The pump photons may originate from two distinct pulses or frequencies, an additional control parameter over SPDC, enabling further possibilities for state tailoring [19]. The general form
of the photon-pair frequency state can be expressed as

$$|\psi\rangle = \int \int d\omega_s d\omega_i \, f(\omega_s, \omega_i)|\omega_s\rangle|\omega_i\rangle,$$

(1)

where $|\omega_s\rangle|\omega_i\rangle$ is a photon-pair state with signal (idler) frequency $\omega_s$ ($\omega_i$), and $f(\omega_s, \omega_i)$ is the joint-spectral-amplitude. In this Letter, we focus on the case where both pump photons originate from the same pump pulse. Energy conservation gives rise to the dependence of $f(\omega_s, \omega_i)$ on the pump spectral amplitude and is assumed throughout, while momentum conservation centers the joint spectral amplitude at zero wave-vector mismatch, i.e. $\Delta k = 0$, with a width inversely proportional to the length of the fiber $L$ [19]. Here we consider a birefringent fiber [22] where the pump is polarization-orthogonal to the signal and idler, in which the wave-vector mismatch, $\Delta k$ is

$$\Delta k = 2k_p + \frac{2}{3} \gamma P_p + 2\Delta n \frac{\omega_p}{c} - k_s - k_i,$$

(2)

Here $k_j$, $j = (p, s, i)$, is the dispersion-dependent wave vector for the pump, signal and idler respectively, $P_p$ is the peak pump power, $\gamma$ is the fiber nonlinear coefficient, and $\Delta n$ is the birefringence. Fiber dispersion plays a central role in $\Delta k$, and thus in the joint spectral properties of the signal and idler photons created. In PCFs, the dispersion varies rapidly with frequency, allowing one to produce completely uncorrelated or highly entangled photon pairs within a small range of pump frequencies. Fiber birefringence gives an additional element of control over $\Delta k$ that has yet to be exploited experimentally, and permits one to choose the central frequencies of the photon pair states. Indeed, a key advantage of birefringent PCF fiber is that the joint spectral amplitude is completely tunable through the fiber and pump configurations. This allows the signal and idler wavelengths to be tuned within the sensitivity range of silicon-based photon-counting detectors while remaining far from the pump wavelength to eliminate contamination by Raman background, hampering several previous experiments.

To demonstrate the potential of these source engineering techniques we set out to create photon pairs in an unentangled spectral state with a joint amplitude that is factorable, i.e. $f(\omega_s, \omega_i) = f_s(\omega_s)f_i(\omega_i)$. In this situation, photon pairs are generated in only two field modes, as required for heralding pure-state single photons. To achieve this, one first fabricates or chooses a fiber in which, atypically, the group velocity of one of the generated photons is equal to that of the pump [19]. This puts tight requirements on the fiber dispersion, particularly if one wishes to choose the center wavelengths of the pump $\lambda_{p0}$ and generated photons at which this condition occurs. A signature of this group-velocity condition is that if the pump center wavelength is varied around $\lambda_{p0}$, the center wavelength of one of the generated photons remains unchanged. If the pump is at $\lambda_{p0}$ and has an appropriate bandwidth (see Ref. [19]), an unentangled spectral state will be generated.

Predicting $\lambda_{p0}$ requires precise knowledge of the fiber dispersion. Accurate birefringence and dispersion data for commercially manufactured fibers are not published, making direct measurement of the fiber properties necessary. Based on preliminary modeling, we chose a 40 cm long PCF from Crystal-Fibre (model NL-1.8-750), quoted to have zero group-velocity dispersion (GVD) at 750 and 1110 nm, and $\gamma = 99$ [W km$^{-1}$] at 780 nm. We pumped along its fast axis with a mode-locked Ti:Sapphire laser (50 fs full width half maximum (FWHM), 76 MHz repetition rate). The laser bandwidth was reduced with a tunable filter to $\Delta \lambda_p = 4$ nm, which applies a square window function of width $\Delta \lambda_p$. This filter is used throughout the paper. The average power of the filtered pump was 1 mW. The signal and idler wavelengths were recorded as the pump was tuned from 765 nm to 795 nm (Fig. 1(b)). This maps out the “phasematching curve” from which the fiber dispersion and birefringence can be determined. Modeling the fiber as having a step-index-profile [23] and fitting to this data yielded a core-diameter and air-filling-fraction of 1.7507 $\mu$m and 51.1% for the fast axis, and 1.7488$\mu$m and 50.5% for the slow axis. This difference implies a birefringence of $\Delta n \approx 1.5 \times 10^{-5}$ at 785 nm. From these fiber parameters, and with $P_p = 0$, we predict a phasematching curve (Fig. 1(a)) that allows entanglement types ranging from frequency correlated to anti-correlated, all within the frequency range of a Ti:Sapphire laser pump. The experimental phasematching data shows that for $\lambda_p \approx 785$ nm, the idler wavelength is, to first order, independent of the pump.

At this point, signal and pump have the same group velocity. Although the model-based theoretical fit is quite accurate (less than 1 nm discrepancy), it pre-

![FIG. 1: (a) Theoretical phasematching curve (i.e. wavelength pairs where $\Delta k = 0$) (solid lines) for orthogonally polarized SFWM in our fiber. (b) Magnified view of boxed region in (a). Curve is fit to experimentally measured signal (bottom $\square$) and idler (top $\bigcirc$) central wavelengths as a function of pump wavelength. Vertical dashed lines indicate the zero GVD wavelengths. Points where frequency correlated ($\times$) and anti-correlated states ($\triangle$) can be created are marked. Thin sections of the curve indicate where factorable states are possible.](image-url)
dicts factorability at $\lambda_p = 783$ nm. Assuming a Gaussian pump laser spectrum with a bandwidth of 20 nm FWHM centered on $\lambda_p = 783$ nm that is subsequently filtered to $\Delta \lambda_p = 8$ nm and using our fiber model we calculate the theoretical joint spectral amplitude $f(\omega_s, \omega_i)$. The numerical Schmidt decomposition of this result [17] predicts the purity of the heralded photons to be 86%.

![FIG. 2: Fiber spectrum (inset) and polarization HOM experimental setup. Although some Raman background remains at the idler wavelength, lack of background at the signal entrance. Although some Raman background remains at the idler wavelength, lack of background at the signal entrance. The tuneable filter reduced the pump bandwidth to $\Delta \lambda_p = 8$ nm, centered on $\lambda_p = 785$ nm with 1.4 mW of average power. A half wave plate (HWP1) and PBS split the pump into clockwise and counterclockwise pulses traveling through the PCF. Achromatic HWPs (AHWP1 and AHWP2) oriented the polarizations of both pump pulses with the fast axis of the fiber and ensured the spent pump exited PBS1 back towards the laser. The photon pairs from the clockwise (H) and counterclockwise (V) directions exit from the same PBS1 port. Any remaining pump was removed with a notch filter (NF). A prism separated the signal and idler photons ($\lambda_{s(i)} = 720(860)$ nm, $\Delta \lambda_{s(i)} = 3.4(0.84)$ nm standard deviation) for both H and V. Occasionally, one H and one V photon pair were created simultaneously. This case was indicated by a silicon avalanche photodiode (APD) detection of two signal photons (APDA and APDC) behind PBS3. This heralded the existence of two idler photons, the subjects of our purity measurement. To ensure good spatial overlap the idler photons were passed through a single-mode fiber (SMF), which made their polarization elliptical. A quarter wave plate (QWP) partially restores linear polarization, leaving them with a residual ellipticity $\tan \chi$ (the ratio between the major and minor axes of the elliptical polarization). The polarization HOM interference was implemented by HWP4 and PBS2 with outcomes detected at APDB and APDD.

The four-fold coincidence (detection of two signal and two idler photons) probability as a function of the polarization rotation $\theta$ induced by HWP4 is

$$P_4(\theta) = \frac{1}{2} [ (1 - p) + (1 + p) \cos^2(2\chi) \cos^2(2\theta) ],$$

where $p = Tr[\rho_V \rho_H]$ depends on both the indistinguishability and purity of the photons, and $\rho_{H(V)}$ is the density matrix of the heralded horizontal (vertical) photon. In the case $\rho_H = \rho_V$, i.e. the photons are identical, $p$ is the purity of the photons. Thus, from a fit of $P_4(\theta)$ we can find a lower bound on the purity.

![FIG. 3: Measured counts in 100s for the polarization HOM experiment. (a) Singles counts of APDs A (•), B (○), C (□) and D (△). (b) Two-fold coincidence counts $R_{AB}$ (•), $R_{CD}$ (○), $R_{AD}$ (□) and $R_{BC}$ (△). (c) Raw four-fold coincidence counts. (d) Normalized four-fold coincidence counts with theoretical fit corresponding to purity 82.1 ± 1.6%. Error bars represent propagated errors assuming Poissonian count statistics.

The raw count data as a function of $\theta$ is shown in Fig. 3. The singles rates oscillate due to laboratory climate control 12-minute cycling and is independent of $\theta$. Fig. 3(d) displays the four-fold counts normalized to account for these fluctuations according to

$$P_4(\theta) = \frac{R_{ABCD} \left( 1 + \cos^2(2\chi) \cos^2(2\theta) \right) \times r \times d}{2 \left( [R_{AB} \times R_{CD}] + [R_{AD} \times R_{BC}] \right)}.$$
where $R_{ABCD}$ is the four-fold coincidence rate, $R_{XY}$ ($X, Y \in (A, B, C, D)$) is a two-fold coincidence rate, $d$ is the counting time for each data point, and $r$ is the laser repetition rate. Assuming identical photons, the fit of Eq. (3) to this normalized data (leaving $\chi$ free) yields an idler photon purity of $82.1 \pm 1.6\%$. The role of the pump time dependence and peak power in SFWM is largely untested. Varying the pump power from 0.2 to 1.4 mW, we find the signal and idler central wavelengths change by only 0.5 nm. Moreover, we see no purity variation for power less than 1 mW. Above this power the observed purity decrease may be caused by higher-order processes leading to more than one photon pair being produced per pump pulse, as well as self and cross phase modulation.

The measured purity in our experiment is mainly limited by the fiber length available to us. Our model predicts that the purity will approach unity as $L$ becomes large [19] for $L = 100$ m the theoretical purity is 98.5%. Unfortunately, current commercial fibers of this length are prohibitively expensive. Alternatively, one could choose a fiber with a larger difference between pump and signal group velocities [19].

To confirm this prediction, we repeated the HOM experiment using a longer fiber, $L = 1$ m, of the same type (see Fig. 4). The two fibers, which were obtained at different times, had factorable points $\lambda_p = 786$ nm, and bandwidth $\Delta \lambda_p = 6$ nm, we measured a purity increase of 4% to 85.9 ± 1.6%, while the purity predicted by the Schmidt decomposition method increased by 4% to 90%. Both test fibers exhibited an experimental purity, as determined by a fit to Eq. (4), of 4% less than the respective theoretical predictions, suggesting a 94.5% experimental purity might be possible with a 100 m fiber.

We have demonstrated the capability to tailor photon-pair states through SFWM in PCF by producing highly-pure heralded single photons without any spectral filtering, a significant source of loss in standard sources. Eliminating this loss is critical because current photonic quantum computing circuitry loses by post selection, which is not feasible for scalable protocols. Our experiment validates recent theoretical results showing that with careful control of dispersion in a fiber, SFWM can generate a wide variety of entangled or, in our case, unentangled states [19]. PCFs provide the flexibility to produce tailored joint spectral states from the visible to telecommunications spectral regions. However, in order to facilitate concurrent use of multiple heralded photon sources, future fabrication of these fibers should address the issue of uniformity between and within fiber samples.

This work was supported by the EPSRC through the QIP IRC, by the EC under QAP funded by IST directorate, and by the Royal Society.

Note added: Since this submission, it has come to our attention that similar results have been obtained by another research group [25].

* Electronic address: o.cohen1@physics.ox.ac.uk

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