Shaping the Ultrafast Temporal Correlations of Thermal-Like Photons

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We show that the temporal correlations between two light beams arising from a broadband thermal-like source can be controlled in the femtosecond regime. Specifically, by introducing spectral phase-only masks in the path of one of the beams, we show that the timing and strength of the photon correlations can be programmed on demand. This example demonstrates that the interbeam second-order coherence function propagates as a phase-sensitive ultrafast wave packet in the path towards the detectors, and is thus, susceptible to be modified by acting on just one of the beams. For quite some time, it has been thought that this could only happen with sources showing time-energy entanglement. Our work shows that such a property is due to the existence of a certain type of correlation, but not necessarily the entanglement.

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Introduction.—The statistical properties of broadband light sources play a key role in understanding the physics behind their emission mechanisms. The theory of optical coherence provides a convenient mathematical framework to account for these stochastic effects by means of a hierarchy of correlation (or coherence) functions [1,2]. The first-order coherence function quantifies the electric field correlations of the source and is used to describe the field superposition effects that appear in the most common interferometers. In the temporal domain, the interference effects of broadband light sources can easily be controlled on demand, thanks to the availability of programmable pulse shapers [3]. Conversely, the measurement of the first-order correlation function yields invaluable information about the interaction of light with matter, and it lies at the heart of some of the most popular bioimaging tools such as optical coherence tomography [4] or spectral endoscopy [5], just to name a few.

The second-order correlation function [1,2] provides a unique fingerprint of the fundamental properties of any radiation source. For example, for thermal-like light sources, this quantity explains why it is twice more probable to detect two photons in coincidence when single-photon detectors are located equidistantly from the source than in any other situation. This effect was first observed in the spatial domain by the groundbreaking experiments of Hanbury Brown and Twiss (HBT) [6]. They showed that the coherence area was related to the angular size of the radiation source [7]. Interestingly, a similar effect also appears in the temporal domain [8], i.e., for thermal-like sources, the photons appear correlated for time instants shorter than the inverse of the bandwidth of the source. On the contrary, for fermions [9] and photons showing sub-Poisson statistics [10], the effect is just the opposite.

The study of higher-order correlations of different types of light sources has been an active area of research for more than six decades and it continues attracting attention due to its relevance in fundamental science [11,12]. However, the measurement of second- and higher-order coherence functions of broadband light sources is especially challenging since, due to their large bandwidths, ultrafast (typically in the femtosecond regime) detection schemes are needed [13]. In fact, it was not until very recently that the second-order coherence properties of true thermal photons were experimentally observed [8].

In this Letter, we demonstrate the control, at ultrafast scales, of the second-order coherence properties of thermal-like photons. The key enabling point is that the intensity correlations of two incoherent light beams originating in the same thermal source are sensitive to the spectral phase difference between the two beams, an effect pointed out in Ref. [14]. Therefore, the temporal correlations of the photons can be tailored on demand by placing a programmable spectral phase-only filter in the path of just one of the beams. This effect is quite intriguing, since unlike with spectral amplitude masks [15], spectral-phase filters do not alter the hierarchy of correlation functions of the individual light beam on which they act [16].

Theory.—The theory behind this experiment can be readily understood in the framework of statistical optics as the propagation of the correlation of the source’s intensity fluctuations [17]. Alternatively, a full quantum formalism might be used instead [18]. Let us consider the scheme of Fig. 1(a). An incoherent broadband light source with...
power spectrum \(S(\omega)\) and obeying Gaussian statistics is split into two arms. The effect of propagation in individual arms can mathematically be accounted for by a linear time-invariant filter, with complex optical transfer function \(H_i(\omega)\), where \(i = u, l\) refer to the upper or lower paths. Upon propagation, both light beams are jointly detected by an ultrafast detector and correlated in intensity with each other, leading to a quantity proportional to the second-order coherence function

\[
G^{(2)}(t_1, t_2) = \langle I_u(t_1)I_l(t_2) \rangle. \tag{1}
\]

Here, the symbol \(\langle \rangle\) denotes ensemble averaging and \(I_u(l)(t)\) is the optical intensity at the upper (lower) arm. It can be calculated as \(I_u(l)(t) = |E_u(l)(t)|^2\), where

\[
E_u(l)(t) = \int d\omega \tilde{E}_0(\omega)H_u(l)(\omega) \exp(-i\omega t). \tag{2}
\]

This equation accounts for the propagation of the different realizations of the instantaneous electric field of the source, \(E_0(\omega)\), with \(\tilde{E}_0(\omega)\) denoting the Fourier transform of \(E_0(t)\). The detected intensity, \(I_u(l)(t)\), is a random function with ultrafast temporal variations. As is well known, these fluctuations typically occur on the scale on the order of the inverse of the bandwidth of the source [19]. The second-order correlation function can be calculated by substituting Eq. (2) into Eq. (1), leading to

\[
G^{(2)}(\tau = t_2 - t_1) = I_u I_l + |\Delta(\tau)|^2. \tag{3}
\]

Here,

\[
I_u(l) = \int d\omega S(\omega)|H_u(l)(\omega)|^2 \tag{4}
\]

is the energy carried in the upper (lower) arm and

\[
\Delta(\tau) = \int d\omega S(\omega)H_u^*(\omega)H_l(\omega) \exp(-i\omega \tau) \tag{5}
\]

is the term accounting for the correlation of the intensity fluctuations. To achieve the above results, we have made explicit use of the assumptions that (i) the light source is statistically stationary and (ii) it obeys Gaussian statistics. Condition (i) implies that the source is spectrally uncorrelated, \(\langle \tilde{E}_u(\omega)\tilde{E}_l(\omega') \rangle = S(\omega)\delta(\omega - \omega')\), while condition (ii) yields the Gaussian moment theorem that

\[
\langle E_u(t_1)E_u(t_2)E_l(t_3)E_l(t_4) \rangle = \langle E_u(t_1)E_l(t_3) \rangle \langle E_u(t_2)E_l(t_4) \rangle + \langle E_u(t_1)E_l(t_3) \rangle \langle E_u(t_2)E_l(t_4) \rangle [1].
\]

It is important to note that when no filters are applied, \(H_u(l)(\omega) = 1\), Eqs. (3)–(5) reduce to the well-known result for thermal-like light beams

\[
G^{(2)}(\tau) = I_0^2 + |\Gamma(\tau)|^2. \tag{6}
\]

with \(I_0 = \Gamma(\tau = 0) = 1\), \(\Gamma(\tau) = \int d\omega S(\omega) \exp(-i\omega \tau)\) being the mutual coherence function of the optical source. As one can observe, at \(\tau = 0\), the function \(G^{(2)}(0) = 2I_0^2\), whereas at time separations \(\tau\) much longer than the coherence time of the source, \(\tau \gg \tau_c\), we have \(\Gamma(\tau) = 0\) and \(G^{(2)}(\tau \gg \tau_c) \approx I_0^2\). The factor of 2 in the ratio of the correlation function to the background, \(g^{(2)}(0) = G^{(2)}(0)/G^{(2)}(\tau \gg \tau_c)\), clearly observed in the experiments of Boitier et al. [12], can thus, be accounted for within a statistical optics framework [13].

Obviously, by placing a narrow band filter, the coherence time of the source is increased and thus, likewise is the temporal window in which the photons appear to be correlated [13,19]. However, Eq. (5) indicates that the second-order coherence function has a more intricate dependence with the spectral phase of the filters placed in the beam paths. This is in radical contrast to Eq. (4), which indicates that the mean value of the ultrafast intensity in the individual beams actually does not depend on the spectral...
phase. In fact, the hierarchy of correlation functions of a thermal-like beam remains unaffected by a spectral phase-only filter [16]. Equations (3)–(5) provide a more general picture of the interbeam temporal correlations, by considering nonequal paths in the propagation towards the detectors, revealing a more subtle spectral phase dependence.

Experiment.—The experimental setup is depicted in Fig. 1(b). The optical source is the amplified spontaneous emission radiation of an erbium-doped fiber amplifier. It has $\sim 200$ mW average power and 2.15 THz bandwidth (measured at full width at half maximum). The thermal-like behavior of this kind of source has been demonstrated in different experiments [12,20]. The light is polarized and split into two different paths, similarly to the HBT setup of Ref. [6] (Fig. 2). In one of the arms, a commercially available pulse shaper (Finisar Waveshaper 1000S) is located. This device can program $H_s(\omega)$ in both amplitude and phase with 10 GHz resolution within the whole communications C band (1527.4–1567.4 nm). We carefully measured the time of flight through the upper arm and then prepared a fiber link of 7.238 m for the lower arm so that the timing difference between the two arms was located within the range of the motorized stage of the cross correlator. The optical power spectra at different arms are shown in Fig. 1(c). As can be seen, $H_s(\omega)$ introduces a slight cutoff at longer wavelengths. This effect can easily be taken into account through Eqs. (2)–(5) and is included in the subsequent simulations. Afterwards, the light beams coming from both arms are sent to a homemade background-free noncollinear intensity cross correlator. This machine has the capability to measure the intensity cross correlation [Eq. (3)] with femtosecond accuracy [21].

The fiber link in the lower arm is placed to adjust the timing. However, it introduces residual dispersion, i.e., $H_l(\omega) = \exp(i\Phi_{2\omega} \omega^2/2)$, where the angular frequencies are referred to baseband for simplicity. This problem can be circumvented since, according to Eq. (5), this residual dispersion can be compensated by programming the same dispersion [14] with the pulse shaper ($\sim -0.0215$ ps$^2$, in this case). The resulting compressed cross correlation is displayed in Fig. 2, normalized with respect to the background (solid blue line). It shows a peak with a width of 351 fs and $g^{(2)}(0) = 1.91$, very close to the ideal value of 2. To make sure that we have achieved the shortest possible cross correlation, we compare the result with the expected one taking into account Eqs. (3)–(5) (dash-dotted red line). There is an excellent agreement between the two curves, indicating that the second-order coherence function is indeed "transform-limited". To illustrate the phase-sensitive nature of the cross correlation, we simply reverse the sign of the programmed dispersion with the shaper. The ensuing cross correlation width increases to 494 fs and the peak drops significantly (dotted green curve). In the following, we will add the above required dispersion amount to the different spectral phases implemented with the pulse shaper.

The results shown in Fig. 2 demonstrate that the temporal correlations of two light beams generated by the same thermal source, and thus lacking entanglement, can be tailored via remote control of the dispersion that sees one of the light beams, similarly to the effect observed with entangled two-photon states, such as the ones produced by means of spontaneous parametric down conversion [22,23]. We should remark that this does not mean that the light beams considered here share the same characteristics with the entangled two-photon states. However, both light sources have the required correlations that enable its observation.

In a further step, we illustrate that the photon temporal correlations can be controlled on demand by spectral phase shaping. Concretely, we implement the following spectral phase $H_s(\omega) = \exp[i\eta \sin(\omega T)]$, with $\eta$ and $T$ real independent parameters to be programmed with the pulse shaper. Let us show first what we should expect, by introducing this expression into Eq. (5). Since the residual dispersions are compensated we can safely consider $H_l(\omega) = 1$, thus we get

$$G^{(2)}(\tau) = I_0^2 + |\Delta(\tau)|^2, \quad (7)$$

with

$$\Delta(\tau) = \Gamma(\tau) \otimes \sum_n J_n(\eta) \delta(\tau - nT). \quad (8)$$

Here, $\otimes$ denotes the convolution operation and $J_n$ are the Bessel functions of the first kind. The above equation indicates that, due to the effect of the spectral filter, there appear several temporal locations at which the normalized second-order coherence function shows a peak. These locations are determined through the parameter $T$, which in turn can be programmed by the pulse shaper. Furthermore, the

![FIG. 2 (color online). Measured normalized second-order correlation function $g^{(2)}(\tau)$ when the spectral phase is corrected with the pulse shaper (continuous blue line), compared with the expected $g^{(2)}(\tau)$ trace (dash-dotted red line). When the programmed phase is reversed in sign, the value $g^{(2)}(0)$ drops and the cross correlation broadens (dotted green).](image-url)
The pulse shaper is programmed with the spectral phase
second accuracy because it is programmed as a phase enco-

The results of this experiment are presented in Fig. 3. The
temporal location of the peaks can be changed with femto-
second accuracy because it is programmed as a phase enco-
ded in the spectral domain, as shown in Figs. 3(a) and 3(b).
In Fig. 3(c), we display the effect of changing the strength
parameter \( \eta \). As expected from the theoretical calculation
taking into account Eq. (3), the probability of detecting
coincident photons at \( \tau = 0 \) increases with respect to the
previous case, where \( \eta = 2 \) rad. The above example shows
that the temporal correlations of the two beams that propa-
gate through different paths can be tailored while leaving
completely unaltered the hierarchy of the correlations of the
individual light beams. In other words, despite acting on
one of the beams, the effect only manifests when perform-
ing intensity cross correlations.

Summary.—We have shown that the ultrafast photon
temporal correlations (second-order coherence function)
of two thermal-like light beams which traverse different
paths depend on the spectral phase difference of the filters
located in their paths. With the aid of a programmable
pulse shaper, we have demonstrated that the strength and
the time instant at which the second-order coherence
function shows a peak can be controlled on demand in a
subpicosecond temporal scale. This physical phenomenon
reveals that the second-order coherence properties of two
optical beams originating from the same thermal-like
source propagate as an ultrashort phase-sensitive wave
packet, whose characteristics can be affected while leaving
unaltered the hierarchy of correlation functions of the

FIG. 3 (color online). Shaping of the photon temporal corre-
lations. Simulation (left column) and measurement (right col-
mum) of the normalized second-order correlation function \( g^{(2)}(\tau) \)
when the pulse shaper is programmed with the spectral phase
\( \exp[i \eta \sin(\omega T)] \). The relevant parameters appear in the figure.

strength of these peaks can be independently controlled
by a suitable choice of the strength parameter \( \eta \).

Discussion.—The possibility of tailoring the temporal
correlations of light beams on demand is a necessary
ingredient for the exploration and development of quantum-inspired classical technologies, i.e., applications
with highly beneficial capabilities that can be understood
with the laws of classical physics to emulate effects first
revealed in the realm of quantum mechanics [25]. This is
the case, for instance, of two-photon lithography [26,27] and
remote dispersion and temporal modulation cancellation
[18,22,28,29]. In all these cases, classical and quantum
sources share the same enabling characteristics (certain
frequency correlations) that allow us the sought-after obser-
vation. Importantly, the classical source generally shows a
higher brightness and easier implementation than its quan-
tum counterpart, facilitating the realization of the system.

One paradigmatic example is quantum optical coherence
tomography [30], a counterpart quantum version of the
usual optical coherence tomography (OCT) technique,
which makes use of broadband entangled photon pairs to
eliminate the disturbing decrease of sensitivity in OCT that
usually accompanies the presence of chromatic dispersion.
Notwithstanding, correlations between light beams, and
not entanglement, is the key enabling factor behind disper-
sion compensation in OCT [31]. Therefore, a beneficial
effect first put forward in a quantum context, could also be
implemented with a more convenient, i.e., with a higher
intensity, light source.

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[1] L. Mandel and E. Wolf, Optical Coherence and Quantum
Optics (Cambridge University Press, Cambridge, England,
1995).