Absolute emission rates of spontaneous parametric down-conversion into single transverse Gaussian modes

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(Received 15 January 2008; published 28 April 2008)

We provide an estimate of the absolute values of the emission rate of photon pairs produced by spontaneous parametric down-conversion in a bulk crystal when all interacting fields are in single transverse Gaussian modes. Both collinear and noncollinear configurations are covered, and we arrive at a fully analytical expression for the collinear case. Our results agree reasonably well with values found in typical experiments, which allows this model to be used for understanding the dependency on the relevant experimental parameters.

DOI: 10.1103/PhysRevA.77.043834

PACS number(s): 42.65.Lm, 42.50.-p

I. INTRODUCTION

For the last two decades, spontaneous parametric downconversion (SPDC) has been the workhorse process for the generation of correlated photon pairs. These can easily be cast into maximally entangled states, which are useful, e.g., for violating Bell inequalities [1], or for the investigation of other fundamental aspects of quantum mechanics. Progress in SPDC-based photon pair sources has allowed the emitted pairs from such sources to play a leading role in demonstration of quantum-information techniques, and made its way to almost practical applications like quantum key distribution.

A number of studies have established the basic understanding of SPDC [2–4], based on energy conservation, and momentum conservation when participating light fields are treated as plane waves [5]. Many of the more recent applications often necessitate significant manipulation and transport of the photon pairs; this can be achieved in a convenient way by guiding the light in single-mode optical fibers. The basic idea of modeling SPDC in this regime is to map the optical modes propagating in the fibers into freely propagating modes of the electromagnetic field in the nonlinear conversion material, where they interact with a pump field. These freely propagating spatial modes can be described in good approximation by paraxial Gaussian beams, and any optimization strategy will involve some sort of mode matching of such interacting beams.

Previous studies of SPDC light coupled into single-mode fibers have focused on optimizing the coupling efficiency, defined as the ratio of photon pairs to single photons that are observed, because this is a quantity that can be measured easily in an experiment. This quantity is important for developing loophole-free tests of Bell's inequality [6], heralded single-photon sources [7–9], or simply sources of high pair brightness [10].

So far, theoretical work in this area has focused mostly on such secondary parameters, and no closed expression for the absolute rate of photon pairs was available for typical experimental configurations. This made it difficult to estimate whether a particular experimental source implementation could be improved with respect to a particular figure of merit, be it total rate or spectral brightness.

In this paper, we try to address this problem and derive an expression for the absolute rate of SPDC emission from a bulk crystal into Gaussian modes. The work connects to earlier investigations of absolute SPDC rates with beams of finite diameter by Kleinman and Klyshko [3,4]. It was found there that the overall rate of pair production is independent of the spot size of the pump beam [3], and that the conversion efficiency of pump photons into correlated pairs integrated over all emission directions [4] is on the order of 10^{-8} /mm for a typical nonlinear material. The restriction on specific spatial modes defined by single-mode optical fibers in the more recent applications, however, made it difficult to relate their results directly to experiments. Our description applies to both type-I and type-II phase-matching conditions, and covers collinear and noncollinear geometries important for the generation of polarization-entangled photon pairs [11,12].

II. MODEL

The basic process of SPDC can be understood as the spontaneous decay of a photon from a pump field into two daughter photons propagating in two—possibly different—target modes, with the process being mediated by a material with a nonlinear optical susceptibility. The physical implementation of SPDC utilizes the lowest order of the nonlinear susceptibility tensor $\chi^{(2)}$ in an appropriate material. While energy conservation allows the decay process to take place in many target modes, phase-matching requirements need to be engineered to allow conversion to take place into any particular pair of directions.

The physical model of the three interacting optical modes is depicted in Fig. 1. We treat the pump beam and the target modes for the down-converted light as propagating paraxial beams with a Gaussian transverse profile. The beams overlap within a nonlinear optical crystal of finite length l, with surfaces normal to the propagation direction of the pump beam. Pump and target modes propagate in one plane, but need not be parallel. We further assume that the three interacting modes overlap in a region without a significant variation of the transverse profile along their respective propagation di-

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FIG. 1. (Color online) Schematic of the down-conversion model considered in this paper. Pump, signal, and idler beams are treated as paraxial beams with a Gaussian transverse mode. The x axis is perpendicular to the plane of the diagram. Coordinate systems of the signal and idler $(x_{s,i}, y_{s,i}, z_{s,i})$ are tilted by angles θ_s , θ_i with respect to the coordinate system of the pump.

rections. This is a reasonable assumption for typical Gaussian beam parameters and conversion crystal lengths used in experiments [10].

We follow the tradition of referring to the target modes as signal (index s) and idler (index i) and choose coordinate systems where the $z_{s,i,p}$ directions are parallel to the main propagation direction for each mode s, i, p (p refers to the pump mode). The spatial mode function of the electrical field for each of the modes can be written as

$$g(\mathbf{r}) = e^{ikz}U(x,y) = e^{ikz}e^{-(x^2+y^2)/W^2},$$
(1)

where k denotes the z component of the corresponding wave vector, W the Gaussian beam waist parameter, and x, y, zrefer to the corresponding coordinate system for each mode. To simplify the overlap calculations, we use normalization constants α for the envelope functions U(x, y) such that

$$\alpha^2 \int dx \, dy |U(x,y)|^2 = 1 \tag{2}$$

in their corresponding coordinate systems, which implies

$$\alpha_{p,s,i} = \sqrt{\frac{2}{\pi W_{p,s,i}^2}}.$$
(3)

We note that the spatial mode function $g(\mathbf{r})$ satisfies Maxwell's equations only approximately. For the calculations presented below, however, this poses no problem. Furthermore, the dispersion relation connected with this mode function has the confinement correction

$$\omega^2 = c^2 \left(k^2 + \frac{2}{W^2} \right).$$
 (4)

Again, for practical beam diameters W of about 100 wavelengths considered in this paper, this correction term is small enough to be safely neglected.

A. Pump mode

The pump mode is aligned with the main coordinate system x, y, z, and treated as a classical monochromatic field of amplitude E_p^0 . We further assume (as is customary) that we have no significant depletion of the pump in the downconversion crystal. The electrical field of the pump can thus be written as

$$\mathbf{E}_{p}(\mathbf{r},t) = \frac{1}{2} [\mathbf{E}_{p}^{(+)}(\mathbf{r},t) + \mathbf{E}_{p}^{(-)}(\mathbf{r},t)] = \frac{1}{2} [E_{p}^{0} \mathbf{e}_{p} g_{p}(\mathbf{r}) e^{-i\omega_{p}t} + \text{c.c.}],$$
(5)

with a polarization vector \mathbf{e}_p , and a corresponding angular frequency ω_p . Using the normalization expression (2), we can connect the electrical field amplitude E_p^0 with the optical power *P* in the pump beam:

$$|E_p^0|^2 = \alpha_p^2 \frac{2P}{\epsilon_0 n_p c},\tag{6}$$

with the refractive index n_p for the pump field, the electrical field constant ϵ_0 , and the speed of light c in vacuum.

B. Target modes

First, we need to take care of the possibly noncollinear propagation of the target modes with respect to the pump. By introducing target mode angles $\theta_{s,i}$, and using an orientation as indicated in Fig. 1, we express the spatial coordinates of the target modes in terms of the main coordinates x, y, z:

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$$\begin{pmatrix} x_{s,i} \\ y_{s,i} \\ z_{s,i} \end{pmatrix} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \cos \theta_{s,i} & \pm \sin \theta_{s,i} \\ 0 & \mp \sin \theta_{s,i} & \cos \theta_{s,i} \end{pmatrix} \begin{pmatrix} x \\ y \\ z \end{pmatrix}.$$
 (7)

To arrive at a rate of photon pairs generated via SPDC, we use the quantized field operators. We do that by introducing a quantization length L in the propagation direction for clarity in the counting of modes, and postulate periodic boundary conditions; later we will drop this requirement. Following the notation in Eq. (5), the electrical field operators take the form

$$\hat{\mathbf{E}}_{s,i} = \frac{1}{2} [\hat{\mathbf{E}}_{s,i}^{(+)}(\mathbf{r},t) + \hat{\mathbf{E}}_{s,i}^{(-)}(\mathbf{r},t)]$$
$$= \frac{i}{2} \sum_{k_{s,i}} \sqrt{\frac{2\hbar\omega_{s,i}}{n_{s,i}^2\epsilon_0}} \frac{\alpha_{s,i}}{\sqrt{L}} \mathbf{e}_{s,i} g_{s,i}(\mathbf{r}) e^{-i\omega_{s,i}t} \hat{a}_{k_{s,i}} + \text{H.c.} \quad (8)$$

Here, $\mathbf{e}_{s,i}$ indicate the polarization vectors, and $n_{s,i}$ and $\omega_{s,i}$ the corresponding refractive indices and angular frequencies of the target modes. The modes are indexed by the scalar moduli $k_{s,i}$, and the corresponding full wave vectors in the pump coordinates are given by

$$\mathbf{k}_{s,i} = k_{s,i} (\mp \sin \theta_{s,i} \mathbf{e}_{y} + \cos \theta_{s,i} \mathbf{e}_{z}). \tag{9}$$

The longitudinal wave vector components serve as a complete discrete mode index $k_{s,i}=2\pi m_{s,i}/L$ with integer numbers $m_{s,i}$. The coefficients before the raising and lowering operators in Eq. (8) are chosen such that the free field Hamiltonian \hat{H}_0 for the target modes takes the usual form

$$\hat{H}_{0} = \sum_{k_{s,i}} \hbar \omega_{s,i} \left(\hat{a}_{k_{s,i}}^{\dagger} \hat{a}_{k_{s,i}} + \frac{1}{2} \right).$$
(10)

C. Interaction Hamiltonian

The SPDC process is enabled by a nonlinear optical material whose presence is described by the Hamiltonian \hat{H}_{I} , written in the interaction picture with time dependence of the raising and lowering operators [13]:

$$H_{I} = -\frac{2\epsilon_{0}\chi^{(2)}}{8} \int_{-\infty}^{\infty} dx \, dy \int_{-l/2}^{l/2} dz \, \mathbf{E}_{p}^{(+)} \hat{\mathbf{E}}_{s}^{(-)} \hat{\mathbf{E}}_{i}^{(-)} + \text{H.c.}$$
$$= d \int_{-\infty}^{\infty} dx \, dy \int_{-l/2}^{l/2} dz \sum_{k_{s},k_{i}} \frac{\hbar \sqrt{\omega_{i}\omega_{s}}}{n_{s}n_{i}} \frac{\alpha_{s}\alpha_{i}E_{p}^{0}}{L}$$
$$\times e^{-i\Delta\omega t} g_{p}(\mathbf{r}) g_{s}^{*}(\mathbf{r}) g_{i}^{*}(\mathbf{r}) \hat{a}_{k_{s}}^{\dagger}(t) \hat{a}_{k_{i}}^{\dagger}(t) + \text{H.c.}$$
(11)

We have assumed a crystal of infinite transverse (x, y) extent, which is justified when the beam diameters are much smaller than the crystal dimensions. We introduce a frequency mismatch $\Delta \omega = \omega_p - \omega_s - \omega_i$. The effective nonlinearity *d* captures the contraction of the nonlinear susceptibility tensor with the corresponding polarization vectors $(2d=\mathbf{e}_p\chi^{(2)}:\mathbf{e}_s\mathbf{e}_i)$ [13]. With this notation, the type of phase-matching condition (type I or II) is reflected in an appropriate effective nonlinearity *d*.

Most of the scaling aspects of the parametric downconversion process connected with the geometry of the interaction are determined by the overlap integral $\Phi(\Delta k)$ of the three mode functions $g_{p,s,i}(\mathbf{r})$ in the crystal:

$$\Phi(\Delta \mathbf{k}) = \int dz \int dy \, dx \, g_p(\mathbf{r}) g_s^*(\mathbf{r}) g_i^*(\mathbf{r})$$
$$= \int dz \int dy \, dx \, e^{i\Delta \mathbf{k} \cdot \mathbf{r}} U_p(\mathbf{r}) U_s(\mathbf{r}) U_i(\mathbf{r}). \quad (12)$$

In this expression, $\Delta \mathbf{k} = \mathbf{k}_p - \mathbf{k}_s - \mathbf{k}_i$ describes the wave vector mismatch. Since pump and target modes are defined in the *y*-*z* plane, there are no wave vector components in the *x* direction and hence $\Delta k_x = 0$. Carrying out the integration in the transverse directions (*x*, *y*) we arrive at

$$\Phi(\Delta \mathbf{k}) = \frac{\pi}{\sqrt{AC}} e^{-\Delta k_y^2/4C} \int dz \ e^{-Hz^2 + izK},$$
 (13)

with the abbreviations

$$A = \frac{1}{W_p^2} + \frac{1}{W_s^2} + \frac{1}{W_i^2},$$
 (14)

$$C = \frac{1}{W_p^2} + \frac{\cos^2 \theta_s}{W_s^2} + \frac{\cos^2 \theta_i}{W_i^2},$$
 (15)

$$D = \frac{\sin 2\theta_s}{W_s^2} - \frac{\sin 2\theta_i}{W_i^2},\tag{16}$$

$$F = \frac{\sin^2 \theta_s}{W_s^2} + \frac{\sin^2 \theta_i}{W_i^2},\tag{17}$$

$$H = F - \frac{D^2}{4C},\tag{18}$$

$$K = \Delta k_y \frac{D}{2C} + \Delta k_z.$$
(19)

The exponential term before the residual integral in Eq. (13) represents the approximate transverse wave vector mismatch. This term can be ignored only if one of the beams is infinitely large $(W_{p,s,i} \rightarrow \infty)$, or if there is perfect transverse phase matching.

The residual integral along z in Eq. (13) can be rewritten in a form that allows for a physical interpretation. We introduce Φ_{z} , where

$$\Phi_{z} \coloneqq \int_{-l/2}^{l/2} dz \ e^{-Hz^{2} + izK} \tag{20}$$

$$=l\int_{0}^{1}du \ e^{-\Xi^{2}u^{2}}\cos(\Delta\varphi u).$$
(21)

The phase mismatch is now defined as $\Delta \varphi \coloneqq Kl/2$. The argument $\Xi \coloneqq \sqrt{Hl/2}$ in the exponential can be viewed as a "walk-off" parameter due to noncollinear mode propagation. This parameter is useful for identifying a thin- and a thick-crystal regime [6]. In our model, these regimes refer to the physical boundary conditions imposed on the interaction volume by the geometry of the pump and target modes.

In the thick-crystal regime with a large walk-off parameter ($\Xi > 1$), the overlap integral Φ_z depends mostly on the characteristic beam parameters $W_{p,s,i}$ and not much on the physical boundaries of the nonlinear material. For $\Xi \rightarrow \infty$ the length of the crystal ceases to play a role altogether:

$$\Phi_z \approx l \frac{\sqrt{\pi}}{2\Xi} \operatorname{erf}(\Xi) = \sqrt{\frac{\pi}{H}} \operatorname{erf}(\Xi).$$
 (22)

The thin-crystal regime refers to a small walk-off parameter, $\Xi \ll 1$, so that the characteristic beam parameters have almost no influence on Φ_z . In particular, this applies for collinear arrangement of all modes ($\theta_i = \theta_s = 0$), where $\Xi = 0$. In this case, $K = \Delta k_z$, and

$$\Phi_z = l \operatorname{sinc}(\Delta \varphi). \tag{23}$$

This reveals the well-known influence of the longitudinal phase mismatch on the down-conversion spectral properties [3]. Figure 2 shows the overlap contribution Φ_z/l as a function of the phase mismatch $\Delta\varphi$ for various walk-off parameters Ξ . We note that, as Ξ becomes large, the spectrum becomes Gaussian-like. If Ξ is identified as the degree of (non)collinearity, it suggests that the SPDC spectral profile far from collinear emission can be very broad. In other words, for the same beam parameters $W_{s,i,p}$, the bandwidth of collinear emission will be narrower than for noncollinear emission.

D. Spectral emission rate

To obtain absolute emission rates, we make use of Fermi's golden rule as an expression for the transition rate $R(k_s)$ between the initial vacuum state $|i\rangle = |0_{k_s}, 0_{k_i}\rangle$ and a final state $|f\rangle = \hat{a}_{k_s}^{\dagger} \hat{a}_{k_i}^{\dagger} |0_{k_s}, 0_{k_i}\rangle$ with the mode pair k_s, k_i populated with



FIG. 2. (Color online) Longitudinal overlap function Φ_z/l as a function of the total phase mismatch $\Delta \varphi = Kl/2$ over the crystal for various walk-off parameters Ξ . For Ξ =0, the typical sinc-shaped spectral distribution is revealed, whereas for large walk-off parameters $\Xi > 1$ the phase-matching condition is determined by the overlap region formed by pump and target modes, and develops into a Gaussian distribution.

one photon each. Fermi's rule applies for asymptotic scattering rates, so the relation between k_s and k_i is fixed by energy conservation:

$$\Delta \omega = \omega_p - k_s \frac{c}{n_s} - k_i \frac{c}{n_i} = 0.$$
⁽²⁴⁾

We first evaluate the transition rate $R(k_s)$ to a *fixed signal* target mode k_s . The density of states ρ per unit of energy $\Delta E = \hbar \Delta \omega$ is extracted out of a quasicontinuum of states for the mode k_i :

$$\rho(\Delta E) = \frac{\Delta m}{\Delta k_i} \frac{\partial k_i}{\partial (\hbar \Delta \omega)} = \frac{L}{2\pi \hbar c} \frac{n_i}{\hbar c},$$
(25)

where $\Delta m / \Delta k_i = L/2\pi$ denotes the number of modes per unit of wave vector component k_i .

With the transition matrix element expressed in terms of the overlap integral $\Phi(\Delta \mathbf{k})$,

$$\langle f | \hat{H}_{I} | i \rangle = d \frac{\hbar \sqrt{\omega_{s} \omega_{i}}}{n_{s} n_{i}} \frac{\alpha_{s} \alpha_{i}}{L} E_{p}^{0} \Phi(\Delta \mathbf{k}), \qquad (26)$$

the transition rate is then given by

$$R(k_s) = \frac{2\pi}{\hbar} |\langle f | \hat{H}_I | i \rangle|^2 \rho(\Delta E)$$
(27)

$$= |d\alpha_s \alpha_i E_p^0 \Phi(\Delta \mathbf{k})|^2 \frac{\omega_s \omega_i}{n_s^2 n_i c L}.$$
 (28)

The spectral emission rate per unit of angular frequency ω_s is obtained by multiplying $R(k_s)$ with the number of modes k_s in a unit interval of ω_s , which is $Ln_s/2\pi c$. We finally arrive at

$$\frac{dR(\omega_s)}{d\omega_s} = \left(\frac{d\alpha_s \alpha_i E_p^0 \Phi(\Delta \mathbf{k})}{c}\right)^2 \frac{\omega_s \omega_i}{2\pi n_s n_i}.$$
 (29)

At this point, the earlier introduced quantization length L has vanished, as expected.

E. Total emission rate

We now can determine the total pair generation rate by integrating the spectral rate density over all frequencies ω_s . Assuming that the overlap $\Phi(\Delta \mathbf{k})$ is only nonvanishing over a small range of frequencies ω_s , the total pair generation rate can be written as

$$R_T = \left(\frac{d\alpha_s \alpha_i E_p^0}{c}\right)^2 \frac{\omega_s \omega_i}{2\pi n_s n_i} \int d\omega_s |\Phi(\Delta \mathbf{k})|^2.$$
(30)

The dependency of $\Phi(\Delta \mathbf{k})$ on ω_s can be quite involved, as in the noncollinear case ($\theta_{i,s} \neq 0$) both $\Delta k_y(\omega_s)$ and $\Delta k_z(\omega_s)$ must be considered. However, the alignment criteria for most experimental setups assume perfect longitudinal phase matching to arrive at the collection angles for degenerate down-conversion [10,11]. In these collection directions, the target mode angles $\theta_{s,i}$ are equal. Furthermore, the typical experiments use identical collection mode diameters for signal and idler, $W_s = W_i$ [10]. Under these two conditions, the phase mismatch $\Delta \varphi$ depends only on Δk_z .

We now consider the exponential term for the overlap Φ in Eq. (13) that contains Δk_y . For experiments where light centered on the degenerate wavelengths with a small bandwidth is collected (≈ 2 nm on either side of the center [10]), we will assume perfect transverse phase matching. A complete treatment with nonzero transverse phase mismatch requires a numerical procedure [14].

With perfect transverse phase matching, we can carry out the integration in Eq. (30) by reparametrizing the frequencies of the signal and idler about the degenerate SPDC frequency: $\omega_s = \frac{\omega_p}{2} - \delta_{\omega}$ and $\omega_i = \frac{\omega_p}{2} + \delta_{\omega}$. We approximate $\omega_s \omega_i \approx \frac{\omega_p^2}{4}$ by ignoring terms $O(\delta_w^2)$. From energy conservation equation (24) and the phase matching condition

$$\Delta k_z = n_s \omega_s \cos \theta_s + n_i \omega_i \cos \theta_i - n_p \omega_p, \qquad (31)$$

we obtain a dispersion relation between $d\omega_s$ and $d(\Delta k_z)$:

$$d(\Delta k_z) = \frac{(n_i \cos \theta_i - n_s \cos \theta_s)}{c} d\omega_s.$$
(32)

The emission rate can now be integrated over the longitudinal wave vector mismatch Δk_z ,

$$R_T = \frac{d^2(\alpha_i \alpha_s E_p^0)^2 \omega_p^2}{4cn_s n_i (2\pi)(n_i \cos \theta_i - n_s \cos \theta_s)} \int |\Phi(\Delta \mathbf{k})|^2 d(\Delta k_z).$$
(33)

Effectively, this is the pair emission rate for all allowed wavelengths in the direction defined by our paraxial beams. If we recall that the pump has a Gaussian envelope, and choose all beam characteristics to be equal $(W_p = W_s = W_i)$, the rate R_T finally can be written as



FIG. 3. Variation of the spectral integral S with walk-off parameter Ξ . Since the absolute emission rate is proportional to S, the largest absolute rate is obtained in the thin-crystal limit with Ξ =0.

$$R_T = \frac{4d^2 P l \omega_p^2}{3 \pi n_p n_s n_i \epsilon_0 c^2 (\pi W_p^2) (1 + \cos \theta_i^2 + \cos \theta_s^2)} \\ \times \frac{1}{(n_i \cos \theta_i - n_s \cos \theta_s)} S,$$
(34)

with the spectral integral $S := \int |\frac{\Phi_z(\Delta \mathbf{k}_z)}{l}|^2 d(\Delta k_z l/2)$. The absolute emission rate is proportional to *S*, which has a dependence on the value of the walk-off parameter Ξ as shown in Fig. 3. The spectral integral assumes its largest value $S = \pi$ in the thin-crystal limit. In this limit, closed form expressions for the spectral and total rates are

$$\frac{d\tilde{R}(\omega_s)}{d\omega_s} = \frac{2d^2\omega_p^2 P l^2 \operatorname{sinc}^2(\Delta k_z l/2)}{9\pi n_s n_s \epsilon_0 c^3(\pi W_w^2)},$$
(35)

$$\widetilde{R}_T = \frac{4d^2 P l \omega_p^2}{9n_s n_i n_p \epsilon_0 \pi W_p^2 (n_i - n_s) c^2}.$$
(36)

F. Dependence of emission rate on beam waists

Although it is convenient to set all beam waists to be equal, this is not necessary. In fact, it can be shown that this choice does not maximize the total emission rate for a given optical pump power. Carrying out the more general derivation to arrive at an expression similar to Eq. (36), the dependency on the various beam waists W_p , W_s , and W_i (again in the thin-crystal limit) can be written as

$$R_T \propto \frac{1}{W_p^2 W_s^2 W_i^2 (1/W_p^2 + 1/W_s^2 + 1/W_i^2)^2}.$$
 (37)

To develop an alignment strategy, we may assume that the collection modes are identical $(W_s = W_i = W)$, but we reexpress the pump waist as $W_p = \gamma W$, so we obtain

$$\widetilde{R}_T \propto R_T \propto \frac{1}{W^2 (1/\gamma + 2\gamma)^2}.$$
(38)

This relationship is illustrated in Fig. 4, and exhibits a maximum of \tilde{R}_T for $\gamma = \frac{1}{\sqrt{2}}$. For $\gamma = 1$, the emission rate is about 12% lower than the maximum value. This suggests that experimental setups that are designed with equal beam waists for pump and collection modes may be further optimized,



FIG. 4. Dependence of the total pair rate \tilde{R}_T on the ratio γ between target and pump waist. The maximum emission rate can be expected at $\gamma = 1/\sqrt{2}$.

and the simple argument of maximizing a mode overlap [10] with matching beam waists does not hold.

III. PHYSICAL INTERPRETATION AND COMPARISON TO EXPERIMENTS

While R_T is proportional to the crystal length l, the spectral rate $dR(\omega_s)/d\omega_s$ is proportional to the square of a sinc function. This is in agreement with results from previous work [3]. However, our expression reveals dependencies on other factors, namely, emission geometry and pump spot size. The expression for R_T reveals that the emission rate is higher in a collinear geometry compared to a noncollinear case. This can be intuitively understood because the noncollinear case has a smaller interaction volume.

We find that the absolute rate R_T is proportional to the square of pump frequency since we reparametrize the signal and idler about the degenerate frequency, so downconversion efficiency can be improved with shorter-wavelength pumps. Both spectral and total emission rates are inversely proportional to the mode area of the beams, in contrast to previous studies, which showed that the total SPDC cannot be enhanced by focusing [3,15]. There, however, SPDC emission was not considered for specific transverse modes. The dependence of emission rates on the mode area has been reported in a previous analysis of SPDC in waveguide structures [16]. This is not surprising because the emission into paraxial beams is essentially the same problem as SPDC in waveguides, where the target modes are quantized in one dimension only. We note that our Eq. (35) is similar to the equation obtained in Ref. [16].

We should not draw the conclusion, however, that SPDC emission into single transverse modes can be arbitrarily enhanced by tight focusing. Our model is only valid in cases where the transverse profile of the beams does not vary significantly over the crystal length. For an optimization study of focus size on SPDC emission we refer the reader to Ref. [9].

For explicit comparison of Eq. (34) with experimental values, we consider our experimental setup (similar to that used in Ref. [17]). In this experiment a pump beam (beam waist $W_p=82 \ \mu$ m) at a wavelength of 351.1 nm is incident

on a 2-mm-thick β -barium borate (BBO) crystal. Two single-mode fibers are used to collect degenerate down-converted photons, which is estimated to have an external emission angle of 3.1°. The collection modes also have beam waists of $W_{s,i}$ =82 μ m.

Specifically, for BBO the effective nonlinearity is given by $d=d_{22}\cos^2\theta_c\cos 3\phi_c$. The angle between pump wave vector and crystal optical axis is $\theta_c=49.7^\circ$, while the azimuthal angle is $\phi_c=60^\circ$, resulting in an effective nonlinearity of 9×10^{-13} m/V ($d_{22}=2.11 \times 10^{-12}$ m⁻¹ V⁻¹ according to Ref. [18]). The observed pair rate is approximately 800 pairs mW⁻¹ s⁻¹ with a pair-to-singles ratio of 0.23.

The walk-off parameter for this setup is $\Xi = 0.933$, indicating that the overlap integral is intermediate between the thin- and thick-crystal limits. The largest observable rate according to our model is $2(0.23 \times R_T) = 1100 \text{ mW}^{-1} \text{ s}^{-1}$. The additional factor of 2 is used because in experiments, the geometry is used to collect down-conversion emission in two decay paths.

The source of the discrepancy between experiment and our model is hard to identify. The assumptions used in the model make it an overestimate, primarily in the reparametrizing of signal and idler frequencies about the degenerate wavelength. Experimentally, there are several sources of uncertainty, the main one being the difficulty in establishing pump power very accurately. For example, the average observed value was arrived by measuring the power using two different power meters (a Newport Model 818-UV reported 11.7 mW while a Coherent Fieldmaster reported 9 mW). The uncertainty in pump power estimation, however, is not sufficient to make the observed result compatible with the calculated value.

According to the model, the conversion efficiency into Gaussian transverse modes for our experimental setting will be 3×10^{-12} mm⁻¹ of crystal length. Other experimentally reported rates in the literature reveal similar down-conversion efficiencies [8,10,19]. The total conversion efficiency of SPDC was found to be on the order of 3×10^{-8} mm⁻¹ sr⁻¹ by Klyshko [4] (for degenerate SPDC with a 500 nm pump wavelength). Experimentally, our collection angle is 3.3×10^{-5} sr. If we convert our units to be

comparable with Klyshko's result, we obtain an efficiency of $7 \times 10^{-8} \text{ mm}^{-1} \text{ sr}^{-1}$.

IV. CONCLUSION

In conclusion, we have presented expressions that provide absolute values for the rate and bandwidth of correlated pairs emitted in bulk crystal SPDC which are in a single Gaussian mode. These modes may be defined by the collection profile of single-mode fibers, selecting emission in a specific pair of directions. The single-mode treatment reduces the complexity in the final expression for the rate equations.

We find that the expression for absolute rates given by the model are slightly larger than experimental observations. The model may thus be treated as an idealized case for the total pair emission rate. The small difference between experimentally observed rates and predictions according to the closed expression for R_T , however, suggests that experimental setups using single-mode collection fibers (e.g., [8,10]) operate close to the optimal limit.

Substantial increase of the emission rates are to be expected from larger nonlinearities, since emission rates are proportional to d^2 . Small mode diameters are also expected to enhance emission rates, as has been convincingly reported for SPDC experiments using waveguide structures [20–22], and a similar theoretical analysis [16] for those cases. Overall spectral brightness will be improved by combining larger nonlinearities with collinear mode confinement in longer structures. Even then, however, the spectral width is still ultimately determined by the longitudinal wave vector mismatch. This indicates that very dramatic improvements (by several orders of magnitude) to the generated pair rate in a narrow bandwidth necessary for addressing atomic systems is not very likely to be expected from bulk crystal emission.

ACKNOWLEDGMENTS

The authors acknowledge financial support from ASTAR under SERC Grant No. 052 101 0043. Special thanks to a referee for initiating the discussion on the beam waist ratios in this paper.

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