Efficient excitation of dye molecules for single photon generation

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Efficient excitation of dye molecules for single photon generation

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Abstract
A reliable single photon source is required for many aspects of quantum technology. Organic molecules are attractive for this application because they can have high quantum yield and can be photostable, even at room temperature. To generate a photon with high probability, a laser must excite the molecule efficiently. We develop a simple model for that efficiency and discuss how to optimise it. We demonstrate the validity of our model through experiments on a single dibenzoterrylene (DBT) molecule in an anthracene crystal. We show that the excitation probability cannot exceed 75% at room temperature, but can increase to over 99% if the sample is cooled to liquid nitrogen temperature. The possibility of high photon generation efficiency with only modest cooling is a significant step towards a reliable photon source that is simple and practical.

1. Introduction

Many applications have need of a fast, reliable source of individual photons on demand [1]. These range from quantum sensing [2] through quantum communication [3] to full-scale photonic quantum computing [4]. The standard way to obtain single photons is to make a pair by parametric down conversion or spontaneous four-wave mixing, and use the detection of one photon to herald the presence of the other. The pair production occurs at random times, meaning the source has to run at a low rate to ensure only one pair is created at a time, so the simultaneous production of, say, 10 pairs from 10 separate sources is not a practical possibility [5]. There has been some progress in multiplexing a large number of heralded photon sources with suitable delays, but this approach still presents technological challenges [6–9]. A promising solution to this problem is to excite a single quantum system with a trigger pulse, then collect the photon that is spontaneously emitted. This is sometimes called a source of photons on demand. An isolated atom provides a particularly well defined quantum system [10, 11], but the need for ultra-high vacuum and some sort of trap makes this difficult to scale up in practical devices. Consequently, there is great interest in solid-state alternatives [12] such as quantum dots [13–16], defects in diamond [17, 18], impurities in other solids [19], and our system of choice—single organic molecules [20–22]. If the photons are required to interfere with each other, they should be identical and then the source needs to be cooled to very low temperature to suppress spectral broadening due to thermal phonons. However, there are a number of applications where broadband single photons can still be useful, such as quantum imaging [23], quantum communication [3], or other applications requiring a number-squeezed light source.

In this paper we consider the use of a single dye molecule as a source of individual photons, produced by spontaneous emission after laser light excites the molecule. In particular, we investigate the use of polycyclic aromatic hydrocarbons (PAH) because these can have a radiative yield very close to 1 [24, 25]. Using a simple rate model to describe the steady, excited-state population induced by the laser, we discuss how to optimise that efficiency and discuss how to maximise it. We demonstrate the validity of our model through experiments on a single dibenzoterrylene (DBT) molecule in an anthracene crystal, and we show that high photon generation efficiency can be achieved when the sample is cooled to liquid nitrogen temperature. This is a significant step in making a fast, reliable photon source that is simple and practical.
increased to as continuous-wave excitation.

With strong excitation, such that

\[ \Gamma_z \text{ increased to fit our room temperature data.} \]

Here \( R_n \) is the spontaneous decay rate of the population \( p_n \). The sum goes up to the highest vibrational level labeled by \( n_{\text{max}} \).

After sufficient time, the populations relax to a steady state, where \( \frac{dp_n}{dt} = 0 \) and therefore

\[ p_e = \frac{\sum_n R_n}{R_0 + \sum_n R_n + \Gamma_0}. \]

With strong excitation, such that \( R_0 + \sum_n R_n \gg \Gamma_0 \), \( p_e \) saturates to the value

\[ p_{e,\infty} = \left( 1 + \frac{R_0}{\sum_n R_n} \right)^{-1}. \]

If the only significant excitation is on the ZPL, \( p_e \) saturates to \( \frac{1}{2} \). If instead the excitation is entirely to states \( S_{1,n} \), then \( R_0 = 0 \) and \( p_e \) saturates to 1. This reproduces the well-known result that the steady state of a driven, damped 3-level system can have a population inversion, whereas the damped 2-level system cannot.

From the perspective of building a triggered photon source, we want a trigger pulse to produce a steady state with the largest \( p_e \), it can, and to do so quickly enough that the resulting spontaneous emission delivers a photon with high probability and well-defined timing. This argues for excitation with a sub-nanosecond pulse, and with a low rate \( R_0 \). It should be noted that pulsed excitation with a pulse length much longer than the coherence time in these systems (~20 fs) achieves the same \( p_{e,\infty} \) as continuous-wave (cw) excitation, and these are the type we consider. As we now show, that requires excitation on the blue side of the ZPL, together with a low enough transverse relaxation rate.

Figure 1. Dispersed emission spectrum of a DBT molecule. Black dots: spectrum measured at room temperature. Blue line: analytical spectrum (see equation (5)) constructed to reproduce the low-temperature emission spectrum in figure 1(c) of Trebbia et al. Red line: the same analytical spectrum, but with \( \Gamma_z \) increased to fit our room temperature data. Inset: molecular structure of DBT and generic energy level diagram showing pumping on the zero-phonon line (ZPL, red double-ended arrow), to a higher vibrational level (blue up arrow), fast non-radiative decays (black curvy arrows) and decays to higher vibrational levels of the ground state (maroon down arrows).

2. Modelling the molecular excitation

The diagram inset in figure 1 illustrates the energy levels of a suitable molecule. The ground electronic state is a singlet \( S_0 \), whose vibrational sublevels are designated \( S_{0,n} \) being a label for the vibrational excitations. The first excited singlet, \( S_1 \), has sublevels \( S_{1,n} \). The optical excitation \( S_{0,0} \rightarrow S_{1,0} \) is followed by stimulated decay back to \( S_{0,0} \) and by spontaneous decay in a few ns to the \( S_{0,n} \) levels. Any population in \( S_{0,n} \) relaxes in a few ps to \( S_{0,0} \). Alternatively, the molecule may be excited from \( S_{0,0} \) to \( S_{1,n} \). From here the molecule relaxes (again in ps) to the \( S_{1,0} \) state, from which it decays spontaneously to the \( S_{0,n} \) levels. It is also possible for \( S_1 \) to decay to the lowest-lying triplet state, but for efficient fluorophores this decay branch is weak and therefore we neglect it in the present discussion.

We can reasonably require that the rate for excitation be much slower than the (GHz) rate for relaxation of the vibrational excitations. Then an excitation to \( S_{1,n>0} \) is followed promptly by an incoherent transfer of population to the state \( S_{1,0} \), whilst the subsequent decay to \( S_{0,0} \) is immediately followed by relaxation to \( S_{0,0} \). Writing the populations of \( S_{1,0} \) and \( S_{0,0} \) as \( p_s \) and \( p_e \) respectively, it is then a good approximation to consider that \( p_s + p_e = 1 \) and the excited state population obeys the rate equation

\[ \frac{dp_s}{dt} = p_s \sum_{n=0}^{n_{\text{max}}} R_n - p_s (R_0 + \Gamma_1). \] (1)

Here \( R_0 \) is the stimulated transition rate on the transition \( S_{0,0} \leftrightarrow S_{1,0} \), and \( \Gamma_1 \) is the spontaneous decay rate of the population \( p_e \). The sum goes up to the highest vibrational level labeled by \( n_{\text{max}} \).

If instead the excitation is entirely to states \( S_{1,n} \), then \( R_0 = 0 \) and \( p_e \) saturates to 1. This reproduces the well-known result that the steady state of a driven, damped 3-level system can have a population inversion, whereas the damped 2-level system cannot.

From the perspective of building a triggered photon source, we want a trigger pulse to produce a steady state with the largest \( p_e \), it can, and to do so quickly enough that the resulting spontaneous emission delivers a photon with high probability and well-defined timing. This argues for excitation with a sub-nanosecond pulse, and with a low rate \( R_0 \). It should be noted that pulsed excitation with a pulse length much longer than the coherence time in these systems (~20 fs) achieves the same \( p_{e,\infty} \) as continuous-wave (cw) excitation, and these are the type we consider. As we now show, that requires excitation on the blue side of the ZPL, together with a low enough transverse relaxation rate.
The excitation rates $R_n$ in equation (3) are found by solving the optical Bloch equations in the steady state:

$$R_n = \frac{1}{2} \Omega_n^2 \frac{\Gamma_2}{\delta_n^2 + \Gamma_2^2},$$

where $\Omega_n$, $\delta_n$, and $\Gamma_2$ are respectively the Rabi frequency, laser (angular) detuning from resonance, and transverse relaxation rate for the transition $S_{0n} \leftrightarrow S_{1n}$ [29, 30]. The Rabi frequency is

$$\Omega_n = \frac{1}{\hbar} |\vec{d}_n \cdot \vec{E}|,$$

where $\vec{d}_n = \langle S_{0n}|\vec{d}|S_{1n}\rangle$ is the dipole transition matrix element and $\vec{E}$ is the electric field driving the transition. Since all the transitions are driven by the same component, $E_{\parallel}$, of the light field, we have

$$R_n = \frac{I}{\epsilon_0 \hbar c} d_n^2 \frac{\Gamma_2}{\delta_n^2 + \Gamma_2^2},$$

where the effective plane-wave intensity at the molecules, defined as $I = \frac{1}{4} \epsilon_0 E_{\parallel}^2 c$, is proportional to the laser light intensity, $I_{\text{ext}}$, incident on the sample. Equation (6) allows us to evaluate equation (3) once we know the dipole matrix elements, the detunings and the transverse relaxation rates. In the following sections, we test this model in the specific case of a dibenzoterrylene (DBT) molecule in anthracene, but it could be equally well applied to other organic molecules with known dephasing rates and vibrational levels (see supplementary information available online at stacks.iop.org/JPCO/2/115027/mmedia).

### 2.1. A specific case: dibenzoterrylene in anthracene

Many PAH molecules are fluorescent in both the visible and DBT, with the formula $C_{18}H_{12}$ and the structure shown inset in figure 1, is no exception. Pure DBT appears green as it absorbs both blue and red light, but it is the red absorption from 700-800 nm, that is of interest here. Anthracene makes an excellent host because it is transparent at 700-800 nm, and all its electronically excited levels lie above the $S_1$ levels of DBT, making it impossible for the DBT to exchange electronic excitation with its host [26, 30-33]. Moreover, DBT is highly photostable at room temperature when it is embedded in a crystal of anthracene [34, 35].

When DBT decays from the $S_{1n}$ level, its fluorescence spectrum has Stokes sidebands due to the transitions $S_{1,0} \rightarrow S_{0n}$, whose transition moments are $\vec{d}_n = \langle S_{0n}|\vec{d}|S_{1n}\rangle$. Supplementary information lists the frequencies $\omega_n$ and relative intensities $I_n$ of these lines, taken from the low-temperature fluorescence spectrum in figure 1(c) of Trebbia et al [28]. From these we deduce the relative strengths of the transition moments: $\vec{d}_n^2 \propto \eta_n/\omega_n$. In these types of molecule, excitation of the optically active electron has very little influence on the vibrating bonds, and therefore the vibrational frequencies and wavefunctions are similar in the upper and lower electronic states [36]. We therefore make the reasonable approximation that the Franck-Condon factors are the same for the excitation $S_{0n} \rightarrow S_{1n}$ and the decay $S_{1n} \rightarrow S_{0n}$, and hence that $d_n$ is equal to $\vec{d}_n$. In this way, we use the measured fluorescence spectrum to give us a set of (relative) values for the $d_n$ to be used in equation (6):

$$d_n^2 \approx \vec{d}_n^2 \propto \eta_n/\omega_n.$$

We turn now to the damping rates to be used in equations (2) and (6). Previous measurements [30] give the population decay rate $\Gamma_{1n}/(2\pi) \approx 40$ MHz, corresponding to a spontaneous decay lifetime of 4 ns, and this remains the same whether at room temperature or at low temperature [37]. The damping rates $\Gamma_2$ in equation (6) describe relaxation of the optical coherences $|S_{1n}\rangle \langle S_{0n}|$. At 4 K, $\Gamma_2$ is small enough that one can excite a single $S_{1n}$ level and observe fluorescence proportional to that $S_{1n}$ alone, with a spectral width (FWHM) of 2 $\Gamma_2$. Using the strong $n = 7$ line 8.7 THz to the blue of the ZPL (see figure 1), we have measured a value $\Gamma_2 = 23(3)$ GHz (see supplementary information), corresponding to a decay time for the vibrationally excited population of ~4 ps. For simplicity we assume that all the $S_{1, n>0}$ states have similar vibrational relaxation times (the exact values are not critical for our purpose here).

At room temperature the optical coherence is so rapidly dephased by the interaction with thermal phonons that $\Gamma_2$ is a thousand time larger and the absorption lines are no longer resolved. Indeed, we are unable to measure the width of the room-temperature absorption spectrum directly because the excitation laser does not scan far enough. Instead, we measure the dispersed fluorescence spectrum while exciting continuously at 730 nm, with the result shown by the black dots in figure 1. We use this spectrum to determine the room-temperature value of $\Gamma_2$, as follows.

The blue curve in figure 1 shows the narrow lines of the dispersed fluorescence spectrum, $S_{flu}$, at 2 K,

$$S_{flu} = \sum_n \eta_n \frac{\Gamma_2^2}{(\omega - \omega_n)^2 + \Gamma_2^2},$$
in which $\omega$ is the frequency of the emitted light and $\Gamma^2_{2n}$ are the relaxation rates of the coherences $|S_{1,n}\rangle\langle S_{0,n}|$. We have taken $\Gamma_{20} = \Gamma_{10}/2$ and $\Gamma_{2n-1,0}/(2\pi) = 23$ GHz on the assumption that all the vibrational excitations relax at roughly the same rate. The broad spectrum (red line) is obtained by adding a further rate $\Gamma_{2n}^2$ to these low temperature rates in order to model the dephasing of the optical dipole by thermal phonons. On fitting this curve to the data we obtain the value $\Gamma_{2n}^2/(2\pi) = 8.2$ THz. We expect the coherence $|S_{1,n}\rangle\langle S_{0,0}|$ to suffer the same thermal dephasing and therefore we conclude that $\Gamma_{2n}/(2\pi) = 8.2$ THz at room temperature.

With these values of $d^2_{2n}$, $\Gamma_{2n}$, and $\Gamma_{10}$, we can substitute equation (6) into equation (2) to give the excited state population $p_e$ prepared by the trigger pulse. Figure 2(a) is a plot of $p_e$ at room temperature against the intensity of the pulse, for several blue detunings $\Delta\nu$ of the laser from the ZPL. As anticipated in equation (3), $p_e$ saturates at various levels $p_{e,\infty}$, the lowest being 0.55 when $\Delta\nu = 0$ and the highest being 0.75 when $\Delta\nu = 40$ THz. At 50 THz detuning $p_{e,\infty}$ decreases again because there are only 23 vibrationally excited states in the model, and indeed no more have been observed experimentally. Thus, it seems likely that $p_{e,\infty} = 0.75$ is the highest achievable at room temperature. Figure 2(b) shows the same information, but plotted against detuning for several intensities, indicated by the colour of the lines.

3. DBT Molecule at room temperature

We now describe experiments to test this theory on a single DBT molecule at room temperature.

3.1. Experimental setup

Individual DBT molecules were observed using the confocal microscope shown schematically in figure 3. The molecule is excited either by a cw tuneable titanium sapphire laser (Coherent MBR) or by a pulsed diode laser centred at 781 nm (Picoquant). The laser light is coupled to the table through a polarisation maintaining fibre (PMF), and is collimated using an aspheric lens (AL). A half wave plate (HWP) and polarising beam splitter (PBS) clean the linear polarisation of the fibre output, then a second HWP rotates the linear polarisation to lie along the optical transition dipole of the DBT molecule. A neutral density filter (ND) controls the pump power and a filter (Filter) removes any light of unwanted wavelength coming from the laser or from fluorescence of optical components up to this point. This is a 780 ± 6 nm band pass, a 760 ± 6 nm band pass, or a 750 nm short pass, depending on the chosen laser wavelength. After passing through the dichroic mirror (DM) and beam sampler (BS) (used to monitor the forward power), the angle of the beam is scanned by two galvo mirrors. The beam then passes through lenses, L1 (focal length $f_1 = 75$ mm) and L2 ($f_2 = 250$ mm), in a 4f’ arrangement, where the scanning mirrors lie at $2f_1'$ + $2f_2'$ from the back focal plane of the microscope objective (Obj, a Nikon PlanApo 100x, 0.9NA). This arrangement allows the galvo mirrors to scan the focal spot on the sample, while always filling the aperture of the objective to produce a Gaussian spot of ~720 nm FWHM. The DBT-doped anthracene crystal, shown inset in figure 4(a), is grown by co-sublimation [27], then placed on a glass coverslip and protected by a thin, spin-coated layer of polyvinyl alcohol (PVA). This sample is held in place using a vacuum chuck. Red-shifted fluorescence from the DBT molecules travels back to the dichroic mirror, which reflects it through an 800 nm long pass filter (LPF, to block scattered excitation light), and into a multimode fiber (MMF). A 50:50 fiber splitter delivers the light to two avalanche photodiodes (Perkin Elmer, APD) so that the second-order correlation function, $g^{(2)}(\tau)$, can be measured.
3.2. Single molecule images

The microscope image in figure 4(a) shows fluorescence from isolated DBT molecules over a 20 μm square area of the anthracene crystal. We selected a single molecule, illuminated it with the 781 nm pulsed laser, and monitored the time Δt between excitation and the detection of photons to build up the histogram shown in figure 4(b). Fitting the decay curve to a single exponential decay $Ae^{-1/\Gamma_1}$, shown as a red line, we measured the excited state lifetime $1/\Gamma_1 = 3.53^{(2)}$ ns.

Changing to ~70 μW of cw laser light at 780 nm, and using both APD detectors, we measured $g^{(2)}(\tau)$, plotted in figure 4(c). This shows a strong dip going down to 0.33(4) at $\tau = 0$. With the high dephasing rate at room temperature, the $S_{10}$ level is populated incoherently, and therefore the ideal second order correlation function takes the simple form [30]

$$g^{(2)}(\tau) = 1 - \frac{1}{N_{\text{eff}}} e^{-\left(1+51\tau_0/\tau\right)} ,$$

where $N_{\text{eff}}$ is the effective number of molecules contributing to the signal, $S = I/I_{\text{sat}}$ is the saturation parameter, and $\tau$ is the time delay between the first and second detection event. The blue dashed curve in figure 4(c) is a fit to this function with $1/\Gamma_1$ fixed at our measured value. That works well in the wings but has a discrepancy near $\tau = 0$ because of the 455 ps (rms) timing jitter of our detection system. When we convolve equation (9) with a Gaussian to model timing jitter of the detectors.

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**Figure 3.** Schematic diagram of the confocal microscope. Dark green beam indicates the pump light, red is the fluorescence and grey is the white light used for imaging. PMF: polarization maintaining fibre; AL: aspheric lens fibre coupler; HWP: half-wave plate; PBS: polarizing beam splitter; ND: neutral density filter; DM: dichroic mirror; BS: beam sampler; PM: power meter; FMM: flip-mount mirror; WL: white light source; L1: first lens; L2: second lens; Obj: microscope objective lens; CCD: charge-coupled device camera; LPF: 800 nm long pass filter; MMF 50:50: multimode fibre beam splitter; APD: avalanche photodiode.

**Figure 4.** (a) Confocal microscope image showing fluorescence from individual DBT molecules in an anthracene crystal. The scale bar is 5 μm. Inset: white light image of the crystal, which is ~0.5 mm across. (b) Histogram of time delays, $\Delta t$, between pulsed excitation of a single molecule and subsequent detection of a fluorescence photon. The bin size is 26.7 ps. Red line: single exponential fit to the data, giving excited-state lifetime 3.53(2) ns. (c) Black dots: measured $g^{(2)}(\tau)$ for the same molecule, with grey shading indicating 1σ Poissonian uncertainty. Blue dashed line: exponential fit to the data. Red curve: fit using equation (9) convolved with a Gaussian to model timing jitter of the detectors.
accidental coincidences. After correcting for that \([33]\), we conclude that \(N_{\text{eff}} = 1.02(4)\) and hence that we are indeed looking at a single molecule.

3.3. Scattering rate at room temperature

Having characterised this particular molecule, we investigated the saturation of its scattering rate by analysing the confocal microscope images. The intensity distribution of the laser spot was a slightly elliptical Gaussian with principal axes along the two axes of the scan and with standard deviations \(\sigma_x\) and \(\sigma_y\). In equation (2), the rates \(R_p\) are all proportional to the laser intensity, so a molecule at position \((x_0, y_0)\), illuminated by a spot centred on \((x, y)\), gives a signal

\[
R(x, y) = R_{\infty} \frac{S_0}{S_0 + \exp \left[ \frac{1}{2} \left( \frac{x-x_0}{\sigma_x} \right)^2 + \frac{1}{2} \left( \frac{y-y_0}{\sigma_y} \right)^2 \right]},
\]

where \(R_{\infty}\) is the fully saturated rate of photons detected from the molecule and \(S_0 = I_0/I_{\text{ext}}\) is the saturation parameter when the spot is centred on the molecule. When \(S_0 \ll 1\) this gives an image that reproduces the intensity distribution of the laser spot, but as the molecule saturates, the image becomes wider and flatter.

With the excitation laser frequency set at a blue detuning \(\Delta \nu\) from the ZPL, we took microscope images over a range of intensities and made a global fit of equation (10) to these images, to obtain a value for \(R_{\infty}\) at that detuning (see also supplementary information). This was done with the same single molecule for 8 different values of \(\Delta \nu\) with the results plotted in figure 5(a). It is straightforward to convert these to values of \(R_{\infty}\) through the relation \(R_{\infty} = \alpha R_{\infty,0} \Gamma I_0\), where \(\alpha\) is the total efficiency for collection and detection. On fitting equation (3) to the data in figure 5(b), with \(\alpha\) as the only free parameter, we find the result \(\alpha = 1.03(2) \times 10^{-3}\) for this particular molecule.\(^1\) The correspondence between our data and the theory gives us confidence that equations (3) and (6) provide a correct description, and therefore that one cannot expect to achieve more than 75% steady-state excitation at room temperature.

The same data also provide information on the variation of \(I_{\text{sat}}\) with detuning, as plotted by the data points in figure 5(b). Figure 2 shows that \(I_{\text{sat}}\) at room temperature should increase monotonically with detuning, but this is not the behaviour in figure 5(b). The reason is that we measure (and plot) the intensity \(I_{\text{sat}}\) incident on the surface of the sample, whereas \(I\) in the theory refers to the intensity at the site of the DBT molecule itself. The ratio of these two is frequency dependent because the sample has several parallel dielectric interfaces making a stack of etalons. We know that the anthracene crystal on the glass substrate is more than 5 \(\mu\)m thick and that the PVA spin-coated top layer is 100 nm thick. A simple model with these two layers reproduces the structure seen in figure 5(b) when we set the anthracene thickness to be 7 \(\mu\)m, as shown by the blue curve. If we also include the 120 \(\mu\)m-thick glass substrate, the additional fringes provide an explanation for the scatter of the data points around the line in figure 5(b) (see supplementary information).

\(^1\) Note, however, that the value of \(\alpha\) varies by as much as 50% from one molecule to another. We think this is due to variation in the depth of the molecule within the crystal.
4. DBT molecule at 77 K

The room-temperature limit of $p_{\infty} \approx 0.75$ is not high enough to make a good triggered photon source, but a significant improvement is possible if $\Gamma 2\nu$ can be reduced. Figure 6(a) shows $p_{\infty}$ as a function of the blue detuning of the trigger pulse, calculated from equation (3) for five values of $\Gamma 2\nu$. The 8.2 THz curve is the same as the red curve in figure 5, peaking at $p_{\infty} = 0.75$ near a detuning of $35 \text{–} 40$ THz. That peak is improved by reducing the dephasing rate, reaching $p_{\infty} = 0.99$ when $\Gamma 2\nu/(2\pi) = 0.7$ THz. The spectral structure in $p_{\infty}$ becomes increasingly evident as $\Gamma 2\nu$ is reduced because the resonances become narrower, with the dips in $p_{\infty}$ corresponding to the regions between resonances, where the effect of the off-resonant $R_0$ is more significant.

In order to ascertain what level of cooling is required to reach $\Gamma 2\nu \lesssim 0.7$ THz, we placed a DBT-doped anthracene crystal in a closed-cycle cryostat (Montana Cryostation) and cooled it to the base temperature of 3.5 K. We imaged a single molecule, and measured the width of its $S_{0,0} \rightarrow S_{1,0}$ scattering resonance over a range of excitation intensities (see supplementary information). This gave a natural width of 38(2) MHz, corresponding to $\Gamma 2\nu/(2\pi) = 19(1)$ MHz and a lifetime of 4.2(2) ns. We then increased the temperature to 77 K, pumped the molecule on the strong $n = 7$ transition 8.7 THz to the blue of the ZPL, and dispersed the fluorescence on a single-photon sensitive spectrometer (Andor Shamrock with a Newton EMCCD). This gave the spectrum shown by the data points in figure 6(b), after subtracting a background of light scattered by the anthracene. The main features of this spectrum are well reproduced by the red line, which is a plot of equation (8), taking $\Gamma 2\nu/(2\pi) = 0.7$ THz. On the low-frequency side of the ZPL there is a small shoulder. The residuals plotted in the lower panel show that this is a sideband centred at a detuning of 2 THz. In section II of their paper [30], Grandi et al identified a possible local phonon mode of this system at an energy corresponding to 40 K (∼1 THz), so we surmise that the sideband here corresponds to a two-phonon excitation [38] of that mode.

Having established that liquid nitrogen temperature should be cold enough to achieve exceedingly high values of $p_{\infty}$, we put this to the test by recording the saturation of the scattering rate, as shown in figure 7. The black data points show the photon count rate in the detector as a function of the power incident on the sample, with the excitation laser tuned to the ZPL. The red line is a fit to these points of the function $R_{\infty} S/(1 + S)$, giving $R_{\infty} = 275(2)$ kcounts/s. The grey data points show the photon count rates when the excitation is tuned 14.2 THz to the blue of the ZPL, where the blue line is a fit to the same function as before, giving $R_{\infty} = 510(4)$ kcounts/s. We expect from equation (3) when $\Gamma 2\nu/(2\pi) = 0.7$ THz that $p_{\infty} = 0.50$ when pumping on the ZPL. Using this and comparing the two saturated count rates we calculate that $p_{\infty} = 0.93(1)$ for the blue detuned case, in agreement with the value 0.94 predicted from equation (3).

These results give us confidence that photons can be triggered with high efficiency using DBT in anthracene once it is cooled to liquid nitrogen temperature.

5. Conclusion and perspective

We have modelled the efficiency of triggering photon emission from a single PAH molecule, taking into account the thermal dephasing of the optical dipole. We find that the trigger pulse can produce a steady excited-state...
population $p_e$ close to 100%, but only when the stimulated emission on the ZPL is negligible. This requires the excitation laser frequency to be far detuned to the blue of the ZPL in comparison with the dephasing rate $\Gamma$.

In the case of DBT in anthracene we measure at room temperature that $\Gamma/(2\pi) = 8.2$ THz, and hence conclude that the photon generation efficiency cannot exceed 75%. We have confirmed this by isolating a single DBT molecule and investigating the saturation of the scattering rate for several frequencies of the pump laser.

By contrast, we expect that this efficiency can exceed 99% if $\Gamma/(2\pi)$ can be reduced to 0.7 THz. We have shown experimentally that this ambition can be achieved simply by cooling the sample to liquid nitrogen temperature. That is a significant step towards a fast, reliable, triggered single photon source because it is much more straightforward to operate a practical device with liquid nitrogen than with liquid helium. To our knowledge, no other types of solid state source can offer such high photon generation efficiency at 77 K.

However, other PAH molecules, most notably terrylene [25], have shown high emission rates even at room temperature. As terrylene has a lower branching ratio to the ZPL than DBT, it can reach a higher $p_e$, up to 96% at room temperature and may well be able to exceed $p_e > 99%$ with only modest cooling (see supplementary information). In order to make the best use of an emitter with such high excitation probability, it is also necessary to collect the photons efficiently, and we have not addressed that challenge here. However, it has already been demonstrated that light from a dipole source can be collected very efficiently by coupling it to a resonant optical structure [13–16, 18, 25, 32, 33], and this is what we plan to do next.

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See supplementary information for supporting content. The data from this paper is available at [39].

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