Invited Article

Increasing the photon collection rate from a single NV center with a silver mirror

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Abstract
In the pursuit of realizing quantum optical networks, a large variety of different approaches have been studied to achieve a single photon source on-demand. The common goal for these approaches is to harvest all the emission from a quantum emitter into a single spatial optical mode while maintaining a high signal-to-noise ratio. In this work, we use a single nitrogen vacancy center in diamond as a quantum emitter operating at ambient conditions and we demonstrate an increased photon count rate up to a factor of 1.76 by placing a silver mirror fabricated on the end facet of an optical fiber near the emitter.

Keywords: NV center, diamond, mirror, collection rate, single photon source

1. Introduction
In recent years, the negatively charged nitrogen vacancy (NV) center in diamond has become one of the most attractive solid state quantum emitters due its robustness, stable emission and optically detected magnetic resonance (ODMR). By exploiting the quality of the NV center, several fundamental phenomena like spin-spin entanglement and detection of quantum coherent evolution of spin states have been demonstrated [1–4]. This enables the construction of a basic quantum spin register which is an important step towards the construction of a large scale quantum network. Furthermore, due to its atomic confinement and its long spin coherence time under ambient conditions the single NV center constitutes an excellent magnetic field sensor both in terms of sensitivity and spatial resolution. Besides the spin coherence time, the sensitivity of an NV center as a magnetic field sensor is ultimately limited by the amount of detected photons [5–7] and thus a high optical collection rate is of high importance.

Different methods have been proposed to increase the photon collection rate from an NV center. These approaches include, among others, the placement of the NV center in a nano-cavity directly processed into the diamond host material [8], micro-pillar structures [9, 10], and solid immersion lenses [11]. In a standard bulk diamond host material with an index of refraction of $n_D = 2.42$, most of the light emitted by an NV center remains inside the diamond due to total internal reflection. This limitation could be overcome by placing detectors on the side of the diamond sample [12], where a 100-times increased signal could be obtained compared to the standard approach with a confocal microscope. Another approach to increase the photon collection rate is to place the emitter at or close to the field maximum of a waveguide mode, such that emission predominantly occurs into the guided mode. This has been shown experimentally for both dielectric [13, 14] and plasmonic [15–17] waveguides.

Another approach to increase the photon collection rate is to place a metallic mirror in close vicinity of the emitter [18]. This modifies the angular emission pattern of the emitter and changes the spontaneous decay rate, as originally
demonstrated for Eu⁺ ions [19, 20] and very recently for single NV centers [21].

In this article we report on an increase of the photon count rate from three different single NV centers by placing a silver mirror fabricated on the end-facet of an optical fiber in the vicinity of the emitter. Although the silver mirror is fluorescing under intense laser light illumination we show that a high signal-to-noise ratio (SNR) can be obtained with this approach.

The article is structured as follows. In the first part, we introduce our experimental methods concerning our measurement setup, sample and mirror preparation and measurement approach and we present our measurements demonstrating increased photon collection rates. In the second part, we discuss the two contributions to an increased photon collection rate; the change of the local density of states in the vicinity of the mirror, and the increase of the total collection rate due to reflection of the NV center fluorescence from the mirror and into the mode of collection. We introduce a model for each contribution and compare the net increase of photon collection predicted by these with the experimentally measured value. Finally, we summarize our findings.

2. Experimental methods

Our sample was prepared on a plasma cleaned fused quartz substrate with a thickness of 0.17 mm optimized for a standard oil immersion microscope objective. After a plasma cleaning process, we spin-coated nano-diamonds with diameters < 50 nm (MSY 0-0.05, Microdiamant AG) on the substrate. The diamond solution was diluted and ultrasound cleaned prior to deposition. The density of the nano-diamonds was chosen to be sufficiently low such that optically active nano-diamonds containing single NV centers could be addressed individually and characterized by our homemade built scanning fluorescence confocal microscope. The silver mirror was prepared on the end-facet of a cleaved optical fiber having a diameter of 125 µm. The cleaved end facet was silver-coated with a thickness > 200 nm by electron beam evaporation of silver (Alcatel SCM 600) leaving a typical surface roughness of RMS < 2 nm.

The sample-fiber setup is depicted in figure 1. NV centers were excited with a linearly polarized continuous wave laser of wavelength λ₀ = 532 nm through an immersion oil objective with a numerical aperture NA = 1.4. After the excitation through the same objective, the fluorescence light was band-pass filtered with high transmission in the range between 647–785 nm and detected by two avalanche photodiodes (APDs) in a Hanbury, Brown and Twiss configuration [22]. Our APDs have dark count rates of 100 counts s⁻¹ and 500 counts s⁻¹, respectively, and, if required, we analyzed the temporal correlations between the two APDs using a time-to-amplitude converter.

After the sample was prepared, the silver coated fiber mirror was introduced by mounting the fiber on a xyz-piezo stage above the substrate as indicated in figure 1. To controllably approach the mirror we imaged the substrate plane with the pump laser by introducing a camera in one of the APD channels. Then, the fiber could be recognized as a second reflecting plane when lowered towards the substrate. Finally, the fiber was aligned in order to cancel out any visible intensity gradient in the reflecting plane to align the fiber end-facet parallel to the substrate plane.

To clarify the data acquisition, we shortly explain the measurement approach. The pump was focused on a chosen NV center. The fiber was then scanned in the z-direction in steps of 20 nm for a given range. Next, the pump beam was moved in the xy-direction to an area without any emitters. An equivalent second scan was finally performed yielding a background measurement.

2.1. Characterization of NV centers

Before introducing the silver mirror we selected and characterized three nano-diamonds containing single NV centers. For simplicity, these NV centers are referred to as NVa, NVb, and NVc throughout the manuscript. The emission of single photons from the NV centers was verified by measuring the second-order correlation function g²(τ) for NVb together with a fit based on the model of a three level system yielding g²(0) = 0.16. With the same approach we find g²(0) = 0.24 and g²(0) = 0.37 for NVa and NVc, respectively, proving that all NV centers are single photon emitters as g²(0) < 0.5. The pump power dependent count rate of NVb is shown in figure 2(a) we plot as an example the measurement result of g²(τ) for NVb together with a fit based on the model of a three level system yielding g²(0) = 0.16. With the same approach we find g²(0) = 0.24 and g²(0) = 0.37 for NVa and NVc, respectively, proving that all NV centers are single photon emitters as g²(0) < 0.5. The pump power dependent count rate of NVb is shown in figure 2(a) we plot as an example the measurement result of g²(τ) for NVb together with a fit based on the model of a three level system yielding g²(0) = 0.16. With the same approach we find g²(0) = 0.24 and g²(0) = 0.37 for NVa and NVc, respectively, proving that all NV centers are single photon emitters as g²(0) < 0.5. The pump power dependent count rate of NVb is shown in figure 2(a) we plot as an example the measurement result of g²(τ) for NVb together with a fit based on the model of a three level system yielding g²(0) = 0.16. With the same approach we find g²(0) = 0.24 and g²(0) = 0.37 for NVa and NVc, respectively, proving that all NV centers are single photon emitters as g²(0) < 0.5.
the spectrum of NVb which shows the zero phonon line around 637 nm and the phonon side band peak of about 700 nm, which is characteristic for the negatively charged NV center at room temperature. Finally, all measurements reported on in the following were carried out for polarizations of the pump laser yielding the maximum count rate.

2.2. Characterization of the silver mirror

Evaporated silver is known to fluoresce under intense excitation with laser light having a wavelength of $\lambda_0 = 532$ nm [23, 24]. Hence, the silver mirror will serve as a source of background noise when brought within a few $\mu$m from an NV center. Additionally, fluorescence originating from the substrate and the immersion oil will contribute to the background noise. These contributions are small compared to the fluorescence from the silver for the mirror-substrate distances considered here and will thus be included when we address the ‘silver fluorescence’, ‘mirror fluorescence’ or the ‘background’. As mentioned previously, we obtained a background measurement for each signal measure. The recorded count rate resulting from silver fluorescence is presented in

Figure 2. (a) Measured second order correlation function of NVb, obtained for a pump power of 480 $\mu$W. (b) Count rate as function of pump power measured for NVb without a mirror. A fit with a saturation function (red straight line) yields a saturation pump power $P_{sat} = 224$ $\mu$W. Inset: spectrum of NVb.

Figure 3. (a) Fluorescence map of an electron beam evaporated silver mirror as a function of mirror position and pump power relative to the saturation power $P_{sat} = 224$ $\mu$W of NVb. (b) Silver mirror fluorescence normalized and averaged with respect to the pump power. Oscillations with a period corresponding to 532/2 nm are observed. The green curve is shown as a guide to the eye and has a period of 266 nm. Inset: Spectrum of an electron-beam evaporated silver mirror obtained at a pump power of 480 $\mu$W and located roughly 500 nm from the substrate. The two peaks at 549 nm and 807 nm are outside the detection band of our setup.
Figure 3(a) as a function of mirror position (to be discussed) and pump power. Assuming that the silver fluorescence is spatially uniform and constant in time, we obtain the background signal important for characterizations of the NV centers, as reported later in the manuscript. From this measurement it is obvious that the largest mirror fluorescence with a count rate of $14 \text{kCts s}^{-1}$ is obtained for small substrate-mirror separations and at high pump power of $\approx 480 \mu\text{W}$, see figure 3(a). Due to Fresnel reflections on the glass-air interface, a standing wave of the pump laser builds up between the silver mirror and the glass substrate causing periodic changes of the background signal. This is revealed by the plot presented in figure 3(b), where for each pump power the signal was normalized to its maximum value and averaged over all powers. As expected, the oscillation period corresponds to half the wavelength of the pump laser. In the inset of figure 3(b), we plot the spectrum of the silver fluorescence and we clearly see that the silver fluorescence behaves spectrally as a very broadband and uniform noise source within our detection band.

2.3. NV centers in the vicinity of a silver mirror

Next, we focus on the count rate from the NV centers in the vicinity of a silver mirror. For each NV center the count rate is recorded as a function of mirror-substrate separation $D$ for different pump powers. The background corrected count rate obtained from all three NV centers as function of mirror position is presented in figure 4 for a pump power of $80 \mu\text{W}$ (a) and $640 \mu\text{W}$ (b). The maximum oscillation peaks are labeled so they occur for a mirror position of 0 nm. We have chosen to crop the data for mirror positions smaller than $-200 \text{nm}$ since the mirror at this point is touching the substrate which is recognized by a sudden anharmonic behavior of the fluorescence with decreasing mirror position in each mirror scan measurement.

For the pump power $P = 80 \mu\text{W}$ and substrate-mirror separations in the range 1000–4000 nm, significant count rate oscillations are visible, with periods corresponding to (or close to) half the pump laser wavelength $532/2 \text{ nm}$. This occurs due to the standing field created from interference between the pump beam approaching the mirror and the reflected pump beam propagating in the opposite direction. In contrast to this, when $P > P_{\text{sat}}$ (see figure 4(b)) oscillations of the count rate corresponding to $532/2 \text{ nm}$ are barely visible since in this range the NV center level populations are almost independent on small changes of the pump power.

All measurements presented in figure 4 show that for substrate-mirror separations below 1000 nm, strong count rate oscillations occur. For $P > P_{\text{sat}}$, the period of these oscillations dominate the pump associated oscillations and appear to be approximately 350 nm corresponding to the maximum of the phonon broadened sideband peak in the NV centers emission spectrum. Taking the maximum count rate in the high excitation power limit relative to the count rate without a mirror, enhancements by a factor of $1.44 \pm 0.040$, $1.76 \pm 0.045$, and $1.57 \pm 0.036$ are observed for NVa, NVb, and NVc, respectively.

To investigate the underlying dependencies more thoroughly, we scan for NVb the excitation power in steps of $100 \mu\text{W}$ and record the count rate as a function of substrate-mirror separation. The result of this background corrected measurement is presented as a color map in figure 5(a) and the background corrected result normalized to the counts without mirror (see figure 2(b)) is shown in figure 5(b). The largest enhancement is obtained for small NV center-mirror separations yielding a factor of around four at a low pump power and a factor of around 1.75 for a high pump power. In accordance with the measurements shown in figure 4, for $P \geq P_{\text{sat}}$ the oscillations corresponding to a $532/2 \text{ nm}$ period...
decrease and the signal oscillates with longer periods corresponding to the half maximum of the NV center spectrum. When measuring certain parameters, one cannot rely on a subsequent subtraction of background fluorescence and hence the SNR is most relevant. The SNR is obtained as the ratio between the background corrected count rate (see figure 5(a) for NVb) and the count rate from the silver mirror (see figure 3(a)). It is plotted in figure 6 for NVb. At a low pump power the highest SNR is measured with a value of 40, while for high pump powers the SNR decreases to around 12 due to a largely increased background fluorescence of the silver mirror.

The background fluorescence from the silver mirror limiting the SNR can be a result of many effects such as roughness of the cleaved fiber facet, the grain type and size for the deposited silver and silver-composites like silver-oxide and silver-sulfide forming on the silver surface in ambient air [25–27]. A careful optimization of the silver deposition can however remedy the low SNR. We tested an alternative deposition method which was a wet chemical deposition [28]. Despite that this approach provided uniform smooth silver deposition independent on surface orientation, we found that the electron beam evaporation method yields silver layers which fluoresced less and in a more stable manner under 532 nm laser illumination.

To resolve the different frequency components contributing to the count rate oscillations at various pump powers, we performed a fast Fourier transform (FFT) with respect to the substrate-mirror separation of the background corrected relative signal shown in figure 5(b). The power spectral density obtained from the FFT is presented in figure 7. Instead of plotting the spatial frequency component $k_d$ we re-scaled the axis and plotted $2/k_d$ to better illustrate the one-to-one wavelength correspondence. Two significant $2/k_d$-components are visible in figure 7: one at around 532 nm confirming the dependence on the pump laser wavelength and a second one at around 700 nm. It is clearly visible that the 532 nm dependence decreases for an increasing pump power and the contribution almost vanishes at a power of $P \gtrsim 2P_{sat}$. In

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**Figure 5.** (a) Background corrected count rate of NVb as a function of mirror position and pump power. (b) The same as in (a), but normalized to the count rate without the mirror ($D = \infty$) presented in figure 3(b).

**Figure 6.** The signal-to-noise ratio determined from the ratio between background corrected absolute counts (see figure 5(a)) and mirror fluorescence (see figure 3(a)).
contrast to this, the peak centered around $2/k_d = 700$ nm is almost independent of the pump power and clearly dominates the spectrum for $P_{\text{exc}} > 1.5 \times P_{\text{sat}}$. The spatial component around 700 nm corresponds to the maximum of the phonon broadened side band peak of the NV center’s emission spectrum (see inset in figure 2(b)) and therefore we conclude that these oscillations occur due to an enhancement of the NV center’s spontaneous emission rate in the vicinity of the mirror.

3. Modeling the collected fluorescence

First we consider the photon collection increase due to the Purcell enhancement. When a mirror is introduced in the vicinity of a dipole emitter the rate of spontaneous emission is changed due to a variation of the local density of states. The enhancement of an emitter’s spontaneous emission decay rate is generally known as the Purcell effect [31]. In the vicinity of a mirror this effect can be interpreted as a driven oscillating dipole, where the emitted field interferes with the dipole upon reflection on the mirror and thus changes the dipole transition [19, 32]. Our simulation is carried out for a dipole emitting light of $\lambda_0 = 700$ nm representing the NV center phonon sideband and recognized as a present spatial frequency in figure 7. The relative change of the spontaneous decay rate $\Gamma/\Gamma_0$, where $\Gamma_0$ is the decay rate in a homogeneous environment, versus the distance to a metallic mirror is plotted in figure 8 for three different dipole orientations with respect to the mirror plane. Here we see a significant difference in the $\Gamma/\Gamma_0$ versus mirror position behavior depending on the dipole orientation.

The intrinsic property of the NV center having two orthogonal dipole transitions spanning a plane ensures that there will always be a non-zero projection onto the mirror plane. To describe the complete relative weighting of each of these dipole orientations, one needs to take into account both the projection of the pump laser polarization onto each dipole transition as well as the projection of each dipole transition onto the mirror. This is, however, beyond the scope of this paper.

Comparing with figure 4(b), where $P > 2P_{\text{sat}}$ for all three NV centers, we find indications that all of them show similarities with the horizontally oriented emitter recognized by clear oscillations growing in amplitude for an approaching mirror within 1 $\mu$m and a maximum decay rate terminating in a significant drop when approaching the mirror-emitter distance $D = 0$.

Assuming that the maximum fluorescence peak in figure 4(b) indeed corresponds to the maximum decay rate peak in figure 8 we should experimentally obtain a fluorescence increase of about 1.4 contributed by the Purcell enhancement at an emitter-mirror separation of about 200 nm.

We now consider the increase of photon collection due to reflection of the NV center fluorescence from the mirror and into the mode of collection which we for simplicity call the geometrical factor. When considering a dipole emitter resting near the surface of a substrate, optical near field effects will determine the angular emission pattern of the dipole [33–35]. The angular emission pattern then depends on the refractive indices of the substrate and the emitter’s host material as well as the distance and the orientation of the dipole with respect to the interface. For multilayer structures the dipole emission pattern can be modeled by a Fourier integration technique of transmitted and reflected plane waves [36–38]. Exploiting this technique, we simulate the emission pattern of a dipole

![Figure 7](image1.png)

**Figure 7.** Power spectral density (PSD) of spatial components obtained via a fast Fourier transform with respect to the mirror separation. We plot $2/k_d$ instead of $k_d$ in order to directly illustrate the underlying spatial component.

![Figure 8](image2.png)

**Figure 8.** Relative spontaneous decay rate of a dipole emitter in the vicinity of a silver mirror for different dipole orientations with respect to the mirror surface. The emission wavelength of the dipole was chosen to be $\lambda_0 = 700$ nm as it corresponds to the peak emission wavelength of an NV center at room temperature. The quantum efficiency of the emitter was assumed to be one in these calculations.

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located in air 25 nm above a silica substrate with an index of refraction $n_{SiO_2} = 1.46$, as depicted in figure 9(a). We again choose the dipole transition frequency to correspond to a wavelength of $\lambda_0 = 700 \text{ nm}$ according to the spatial component in figure 7 corresponding to the phonon side band peak of the NV center. Vertical (horizontal) orientation refers to perpendicular (parallel) alignment of the dipole axis to the plane of the substrate. The calculated emission pattern for this case is presented in figure 9(b). It is clear that independent of the dipole orientation a fraction of the emitted light will escape into the upper half space and thus can not be collected by a microscope objective placed below the glass substrate. With an index-matched high NA microscope objective of NA = 1.4 light within a half angle of 73.5° can be collected which yields a total collection efficiency of 0.79 (0.83) for a horizontally (vertically) aligned dipole. If a silver mirror with $n_{Ag}(\lambda_0 = 700 \text{ nm}) = 0.16761 + 4.2867i$ is positioned 200 nm above the emitter, the dipole pattern drastically changes, as illustrated by the plot in figure 9(c). The total collection efficiency then increases to 0.99 (0.96) for a horizontally (vertically) aligned dipole. This corresponds to a geometrical factor in the photon collection being in the range 1.16–1.25 depending on the dipole orientation.

Note that the simulation only takes angular dependence into account. This means that light originating from elsewhere in the $xy$-plane relative to the emitter as a result of multiple scattering between the mirror and the substrate is pictured as stemming from the $xy$-coordinate of the emitter. But since the lens diameter of the objective $D_{\text{len}} \gg D + d$, $d$ being the emitter-substrate distance shown in figure 9(a), this phenomenon will not affect our collection and the emission pattern of figure 9(c) is representative.

Considering both the Purcell factor and the geometrical factor, an increase of the total photon count rate up to a factor of 1.75 is expected for a horizontally aligned dipole separated by approximately $D = 200 \text{ nm}$ from the mirror.

4. Discussion

Multiplying the geometrical factor and the Purcell factor yields a total factor ranging from 1.62 to 1.75 depending on the dipole orientation. The measured collection enhancement of $1.76 \pm 0.045$ for NVb fits well this range suggesting that the mirror has reached a distance of approximately 200 nm from the emitter. Since the same mirror, and thus the same mirror alignment, was used for all three NV centers, all of them are probed for the same mirror distances outlining the maximum oscillation peak nearest to the mirror as depicted in figure 4. The collection enhancement factors of NVa and NVc are not represented in the 1.62–1.75 interval. Since the models presented only consider single dipole emitters we note that this simplification of the NV center can contribute to the deviation. Furthermore, effects stemming from the unknown individual diamond host geometry are not considered.

In case of real-time recordings, where post-subtraction of the background fluorescence is not an option, the ratio between the emitter signal and the mirror background becomes critical. This includes temporal correlations such as measuring the auto-correlation function $g^{(2)}(\tau)$ or demonstrating the Hong–Ou–Mandel effect. On the other hand, introducing a mirror for collection enhancement for ODMR measurements would provide a higher noise floor but would however simultaneously increase the contrast, thus improving the distinguishability between e.g. different spin states due to the increase in collection efficiency [5–7].

Choosing a different emitter enabling an alternative spectral filtering might increase the SNR if the primary emitter signal is confined to a narrow band in contrast to the NV center. For this purpose quantum emitters like the silicon-vacancy defects in diamond [39–41], vacancy defects in silicon carbide [42] and nano-structured quantum dots [43] can be suited for increasing the collection efficiency using a mirror.

Examples of experiments which can benefit from our photon collection enhancement method are experiments with opto-mechanical membranes [8, 44], and ODMR detection schemes [5, 6, 45]. For a high density ensemble of NV centers it can be a challenge to reach the saturation pump power which can approach 1 W [6]. In this case one would both benefit from the cavity effect of the pump and the combined geometrical factor and Purcell factor which we for our system demonstrated to be a four-fold increase in photon collection, see figure 5(b). Regardless whether a saturation power can be
reached a significant increase in photon collection can be made for the purpose of improving sensitivity of ODMR.

5. Summary

In summary, we have demonstrated a method for increasing the photon collection rate consisting of a silver coated standard optical fiber. The collection enhancement was demonstrated for three different NV centers where we found count rate increases of $1.44 \pm 0.040$, $1.76 \pm 0.045$, and $1.57 \pm 0.036$. We thoroughly investigated the pump power dependence for one of the NV centers finding a SNR of 12 when saturating the NV center. We introduced two theoretical models which when combined predicted a maximum enhancement of 1.75. We finally discussed our method as a simple way to increase the collection rate for experiments with quantum emitters.

References

[1] Dutt M V G, Childress L, Jiang L, Togan E, Maze J, Jelezko F, Zibrov A S, Hemmer P R and Lukin M D 2007 Science 316 1312–6
[2] Drea A, Spinicelli P, Maze J R, Roch J-F and Jacques1 V 2013 Phys. Rev. Lett. 110 060502
[3] Dolde F, Jakobi I, Naydenov B, Zhao N, Pezzagna S, Trautmann C, Meijer J, Neumann P, Jelezko F and Wrachtrup J 2013 Nan. Phys. 9 139–43
[4] Pfaff W et al 2014 Science 345 532
[5] Acosta V M, Jensen K, Santori C, Budker D and Beausoleil R G 2013 Phys. Rev. Lett. 110 213605
[6] Jensen K, Acosta V M, Jarmola A and Budker D 2013 Phys. Rev. B 87 014115
[7] Maletinsky P, Hong S, Grinolds M S, Hausmann B, Lukin M D, Walsworth R L, Loncar M and Yacoby A 2012 Nat. Nanotechnology 7 320–4
[8] Faraon A, Santori C, Huang Z, Acosta V M and Beausoleil R G 2012 Phys. Rev. Lett. 109 033604
[9] Hausmann B J M, Khana M, Zhang Y, Babinec T M, Martinick K, McCutcheon A, Hemmer P R and Loncar M 2010 Diam. Relat. Mater. 19 621–9
[10] Choy T J, Hausmann B J M, Babinec T M, Bulu I, Khan M, Maletinsky P, Yacoby A and Loncar M 2011 Nat. Photonics 5 738–43
[11] Hadden J P, Harrison J P, Stanley-Clarke A C, Marseglia L, Ho Y-L D, Patton B R, O’Brien J L and Rarity J G 2010 Appl. Phys. Lett. 97 241901
[12] le Sage D, Pflam L M, Bar-Gill N, Belthangady C, Lukin M D, Yacoby A and Walsworth R L 2012 Phys. Rev. B 85 121202(R)
[13] Schröder T, Fujiwara M, Noda T, Zhao H-Q, Benson O and Takeuchi S 2012 Opt. Express 20 10490
[14] Schröder T, Schell A W, Kewes G, Barth M, Aichele T and Benson O 2011 SPIE 7943 794312-1
[15] Akimov A V et al 2007 Nature 450 402
[16] Kolesov R, Groitz B, Balasubramanian G, Stihrl1 R J, Nicolet A A L, Hemmer P R, Jelezko F and Wrachtrup J 2009 Nat. Phys. 5 470–4
[17] Huck A, Kumar J S, Shakoor A and Andersen U L 2011 Phys. Rev. Lett. 106 096801
[18] Lee K G, Chen X W, Eghlidi H, Kukura P, Lettow R, Renn A, Sandoghdar V and Götzinger S 2011 Nat. Photonics 8 166–9
[19] Drexhage K H 1970 J. Lumin. 1 693–701
[20] Amos R M and Barnes W L 1997 Phys. Rev. B 11 557249
[21] Mohtashami A and Koenderink F 2013 New J. Phys. 15 043017
[22] Hanbury Brown R and Twiss R Q 1956 Nature 178 1046–8
[23] Peyser L A, Vinson A E, Bartko A P and Dickson R M 2001 Science 291 103–6
[24] Pfaff W, Vos A and Hanson R 2013 J. Appl. Phys. 113 024310
[25] Pettersson L A A and Snyder P G 1995 Thin Solid Films 270 69–72
[26] Burge D K, Bennett J M, Peck R L and Bennett H E 1969 Surf. Sci. 16 303–20
[27] Bennett J M, Stanford J L and Ashley E J 1970 J. Opt. Soc. Am. 60 224–32
[28] Malureanu R, Alabastru A, Cheng W, Kiyani R, Chichkov B, Andryieuski A and Lavrinenko A 2011 Appl. Phys. A 103 749–53
[29] Wolters J, Sadzak N, Schell A W, Schröder T and Benson O 2013 Phys. Rev. Lett. 110 027401
[30] Beha K, Batalov A, Manson N B, Bratschitsch R and Leitenstorfer A 2012 Phys. Rev. Lett. 109 077404
[31] Purcell E M 1946 Phys. Rev. 69 681
[32] Chance P R, Prock A and Silbey R 1978 Molecular fluorescence and energy transfer near interfaces (Advances in Chemical Physics vol 37) ed I Prigogine and S A Rice (Hoboken NJ: John Wiley and Sons)
[33] Lukosz W and Kunz R E 1977 Light emission by magnetic and electric dipoles close to a plane interface: I. Total radiated power J. Opt. Soc. Am. 67 12
[34] Lukosz W and Kunz R E 1977 Light emission by magnetic and electric dipoles close to a plane dielectric interface: II. Radiation patterns of perpendicular oriented dipoles J. Opt. Soc. Am. 67 12
[35] Lukosz W 1979 J. Opt. Soc. Am. 69 11
[36] Neyts K A 1998 J. Opt. Soc. Am. A 15 962–71
[37] Chen X-W, Choy W C H and He S 2007 J. Disp. Technol. 3 110–7
[38] Chen X-W, Götzinger S and Sandoghdar V 2011 Opt. Lett. 36 3545–7
[39] Neu E, Agio M and Becher C 2012 Opt. Express 20 19956
[40] Neu E, Steinmetz D, Riedrich-Meiller J, Gsell S, Fischer M, Schreck M and Becher C 2011 New J. Phys. 13 025012
[41] Wang C, Kurtisfeier C, Weinfurter H and Burchard B 2006 J. Phys. B: At. Mol. Opt. Phys. 39 37–41
[42] Castelletto S, Johnson B C, Ivády V, Stavrias N, Umeda T, Gali A and Ohshima T 2014 Nat. Mater. 13 151–6
[43] Shields A J 2007 Semiconductor quantum light sources Nat. Photonics 1 215–23
[44] Ovartchaiyapong P, Lee K W, Myers B A and Jayich A C B 2014 Nat. Commun. 5 4429
[45] Budker D and Romalis M 2007 Nat. Phys. 3 227–34