PAPER

Revealing superradiant emission in the single-to-bulk transition of quantum emitters in nanodiamond agglomerates

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Abstract

Individual quantum emitters form a fundamental building block for emerging quantum technologies. Collective effects of emitters, such as superradiance, might improve the performance of applications even further. When scaling materials to larger sizes, however, the optical density of states is modified by the surrounding material, and the collective coupling in small domains might be covered by transitions to bulk properties due to the presence of multiple collectively emitting domains, which inhomogeneously add. Here, we probe the optical properties of nitrogen vacancy centers in agglomerates of nanodiamonds. We quantify the transition from individual emitters to bulk emission by fluorescence lifetime measurements, and find a transition to occur on a length scale of ∼ 3 wavelengths around the emitter. While our lifetime measurements are consistent with superradiant decay, the second-order correlation function, which is a standard measure to reveal collective properties, fails to probe collective effects for our case of an ensemble of collectively contributing domains to the emission. Therefore, we propose and apply a new measure to trace collective effects based on the intensity fluctuations of the emitted light. Our work points toward systematically studying collective effects in a scalable solid-state quantum system, and using them for quantum optical applications in agglomerates of highly-doped nanodiamonds.

1. Introduction

Micro- and nano-scale objects have become increasingly relevant for technological applications [1, 2]. The length scale of a material platform is often essential to understand its properties. When its size increases, a transition occurs at a characteristic length scale to establish macroscopic bulk properties. Prominent examples are the structure of nano-gold complexes [3], aqueous solutions of gold and silver colloids [4], or the emergence of nanoscopic aqueous droplets of acid formed within a superfluid helium cluster [5]. Impurities immersed in such a material will experience the bulk properties also beyond a particular length scale, which for optically active impurities can be of the order of the wavelength of light. For multiple impurities, an additional transition occurs, beyond which the impurities act collectively and develop properties that are not present in individual nano-particles, making collective effects attractive for possible applications. Such collective effects can lead to various physical phenomena, such as the well-known superradiance predicted by Dicke [6]. An ensemble of emitters emit energy collectively, yielding insight into quantum mechanical processes through a macroscopic effect [7, 8]. Collective emission has been observed in gases [9], quantum dots [10–12], van-der-Waals heterostructures [13], and ultracold atoms [14], for example. Recently, superradiance has been observed in optically active nitrogen-vacancy (NV) centers in diamond in the optical [15] and microwave regime [16]. An important question for future applications, such as quantum metrology [17–20], light harvesting [21, 22] or low-threshold lasing [23], is if the signature of collective effects prevails when the system size crosses the transition to bulk properties.
In this respect, the negatively charged NV center in diamond is a promising technological platform. It is a widely investigated material system [24], specifically for emerging quantum technology such as spin-magnetometry [25], quantum information processing [26], and non-classical light sources [27]. This point defect is well-known as a room-temperature single-photon source [28] and can be produced as bulk- and nanodiamond [29, 30] and prepared as individual emitters or assembled as large, dense ensembles. Likewise, these defect centers change their (quantum) optical properties, such as optical decay rate and collective emission, as a function of the size of the diamond host crystal and the distance between individual defect centers. On the one hand, NV centers in nanodiamonds are subject to a reduction of their optical decay rate in comparison to bulk diamond due to the increased surface-volume ratio and thereby reduced optical density of states (DOS) [27, 31]. On the other hand, nanodiamond samples with high NV-concentration and a particle size above 100 nm show collective emission, i.e. Dicke superradiance [6, 15] which can be established even in the presence of strong dephasing [32]. These collective effects are accompanied by a super-Poissonian photon distribution leading to an increase in the second-order correlation at zero time delay $g^{(2)}(0) > 1$, the so-called photon bunching. These properties make NV centers in nanodiamonds well-suited to study parallel transitions to bulk properties and collective response.

A profound size effect on the optical properties concerning lifetime reduction towards the bulk-lifetime in agglomerates of small nanocrystals with a size of 5 nm and 55 nm was shown but not investigated in detail [33]. Further, such increased optical emission and photon bunching in the nanosecond regime of NV ensembles in single nanodiamonds has been interpreted as superradiance in reference [15].

We use agglomerates of nanodiamond crystals as an intermediate state between bulk diamond and individual nanodiamonds, where poly-crystalline quasi-2D sections form on a glass substrate. Each doped nanodiamond ($n_{NV} \approx 10$ ppm) comprises >1000 NV centers, which can be concentrated at sub-wavelength scales. Additionally, the size of agglomerates can be controlled, rendering them an ideal testbed to systematically study the optical properties of quantum emitters for growing system sizes.

Here, we probe the optical properties as the diamond host transits to bulk-like behavior, and we reveal the collective phenomena in this condensed-matter material. To this end, we investigate the dependence and scaling of (quantum) optical properties of NV-ensembles on the system’s size in a systematic study of different agglomeration states ranging from single nanodiamonds of size 100 nm to large quasi 2D agglomerates consisting of up to 10 000 nanodiamonds, as shown in figures 1(a) and (b). Experimentally, the change of agglomerate size leads to apparently contradicting results in the two main observables usually considered, fluorescence lifetime on the one hand and photon–photon correlation on the other hand. The main observation is shown in figure 1, where details for the measurements will be given later. For growing agglomerate size, the fluorescence lifetime decreases (figure 1(c)), consistent with the interpretation of superradiance. By contrast, the second-order correlation function for zero time delay shows a decreasing value for increasing agglomerate size (figure 1(d)), inconsistent with the interpretation of superradiant effects on a first glance. We trace this discrepancy to an average over different collective domains contributing to the emission. We find that, for large numbers of collective domains, the correlation signal would decrease to $g^{(2)}(0) = 1$, even in the case where superradiance can be expected. To resolve this discrepancy, we introduce a novel way to quantify collective effects via the Fano factor, quantifying number fluctuations of the photon statistics, and justify this in a theoretical model. Applying this model, we find that our observations are consistent with the interpretation of superradiant emission. We furthermore identify the relevant length scale on which bulk-like properties can be established and the scaling of collective properties in agglomerates. Our results are not restricted to NV centers but also applicable to other (diamond) color centers like silicon-vacancy or germanium-vacancy centers, or quantum dots.

## 2. Experimental system and measurement sequence

The negatively charged NV center is a paramagnetic point defect in the diamond lattice composed of a substitutional nitrogen atom (N) and an adjacent vacant lattice site (V) [24]. This defect features local energy levels in the diamond’s band-gap, which can be addressed with optical and microwave wavelengths, as depicted in figure 2. The electronic ground state and optically excited state are spin-triplet states ($m_s = 0, \pm 1$) with separation energy of 1.945 eV corresponding to a zero phonon line (ZPL) at $\lambda_{ZPL} = 637$ nm, followed by phononic bands of vibrational modes. The lifetime of the optically excited state differs for NV centers in nanodiamonds ($\tau_{ND} \approx 23–25$ ns) and bulk diamond ($\tau_{bulk} = 12–13$ ns) [27, 34–36]. Further, a second decay path via intersystem-crossing (ISC) into long-lived ($\tau_{sing} \approx 150$ ns [37]) singlet states and a subsequent second ISC back to the triplet ground state exists. The $m_s = \pm 1$-states
Figure 1. Raster scan of nanodiamonds in different agglomeration states. (a) SEM images of different agglomeration states ranging from single nanodiamonds over planar sections to an agglomerate with 3D features. The scale bar in each image has a size of 2 μm, and a scale of the emission wavelength λ_{ZPL} is shown for comparison. SEM-images were taken at the end of all measurements at the Hitachi SU8000 SEM; the same agglomerates and nanodiamonds were found using marker structures on the glass-substrate. (b) Normalized count rate measured at a 2D confocal scan as a function of x- and y-position; each measurement was normalized to the maximum count rate in the respective data set; all scans have the same scale; the red rectangle gives the region of the corresponding SEM-image. (c) Count rate weighted lifetime measurements of the different agglomerates shown in (a) and (b). A multi-exponential decay is observed, where larger agglomerates show a lifetime reduction, the zoom-in shows the initial fast decay. (d) Time-averaged second-order correlation measurements in dependence on the time window t′ with maximum signal to noise ratio of the shown agglomerates. The photon bunching reduces on a time scale smaller than the fluorescence lifetime.

preferably decay via this path, and therefore, optical pumping leads to spin polarization of the m_s = 0 ground state.

The experimental setup consists of a home-built confocal microscope, as depicted in figure 2. NV centers are optically excited by an off-resonant pulsed laser source featuring a wavelength of λ_L = 517 nm, pulse-widths of 500 ps and repetition rates of 5 MHz. The sample consists of highly doped nanodiamonds in an aqueous solution [38]. They have an average size of 100 nm and more than 1000 NV centers per single crystal. We drop cast them as received on a glass substrate and dry the sample with a contact heat plate to vaporize the water. Fluorescence radiation of NV centers is collected by the same microscope objective transmitted through a dichroic mirror and wavelength-filtered by a bandpass to transmit a small band of 637(3) nm. After that, the fluorescence light is divided up at a 50:50 beam splitter and fiber-coupled to two single-photon counting modules (SPCMs), forming a Hanbury Brown and Twiss interferometer [39]. To reduce the total count rate to avoid measurements close to these detectors’ saturation point, a neutral density (ND) filter (OD = 2.0) was added in front of the beam splitter. A time-to-digital converter (TDC) tracks the arrival of single photons and calculates the average count rate as well as coincidences in each measurement. More information on the experimental apparatus is given in appendix A. Appendix B gives more details on the coincidence measurement and extraction of the pulsed second-order photon correlations.

To observe lifetime variations and photon bunching associated with collective effects, we investigated 104 agglomerates of nanodiamonds with an expansion of A < 1 μm² and 12 agglomerates exceeding this size. For agglomerates within A < 1 μm² being in the order of the laser focus, we raster-scan the sample in x- and y-direction with a step-size of s = 0.3 μm, choose the position of maximum count rate, and perform a single measurement. We group these structures into three different size categories according to the size evaluated from SEM images which were taken subsequently to all lifetime and correlation measurements with the SEM Hitachi SU8000. In this process, the surface of the sample is covered with an iridium layer of 2 nm to achieve a conducting surface. This layer leads to a reflective surface and changes therefore the
optical properties. We ensure that one and the same structure is found again by triangulation of the measurement positions using multiple marker structures on the glass-substrate. Those 104 small agglomerates consisted of 37 structures with a size of $A < 0.05 \mu m^2$, 44 with $0.05 \mu m^2 < A < 0.20 \mu m^2$ and 23 with $0.20 \mu m^2 < A < 1.00 \mu m^2$. For the 12 agglomerates exceeding a size of $A > 1 \mu m^2$, we measure at every position of a raster scan with a step size of $s = 0.4 \mu m$. We further increase the distance between sampling points to $s = 1 \mu m$ for agglomerates exceeding $A > 10 \mu m^2$ to reduce the total measurement time. This procedure allows quantifying variations of the lifetime and collective emission throughout agglomerates. Each measurement sequence consists of $1 \times 10^9$ repetitions of the laser pulse. During the first $1 \times 10^4$ repetitions, photon detection is shut to reach a steady-state and spin-polarization into the $m_t = 0$ state.

3. Transition to bulk material

We trace the transition from optical nano- to bulk-like properties of NV centers by scanning agglomerates with different sizes and performing a fluorescence lifetime measurement (FLIM), as described in section 2. To quantify changes between samples with different size as well as variations within agglomerates, the measurement data is approximated by a model considering the time-dependent excitation of the laser pulse and a subsequent bi-exponential decay according to

$$I(t) = a_1 \exp(-t/\tau_1) + a_2 \exp(-t/\tau_2),$$

where $I(t)$ is the time-resolved intensity detected, $a_1$ and $a_2$ are the amplitudes at time $t = 0$ of a fast decay $\tau_1$ and slow decay $\tau_2$. For individual nanodiamonds we find that the latter value is in the order of the reported fluorescence lifetime $\tau_2 \approx 25$ ns [27, 31, 36], which differs to the bulk lifetime of 12.9 ns [34, 35]. Therefore, we associate this lifetime with a change of optical emission due to a change of the optical DOS given by the effective refractive index.

In figure 3(a), we show the position-dependent slow decay component $\tau_2$ in four agglomerates with varying sizes. All measurement data of 104 small nanodiamond agglomerates within the three size categories and 12 raster scans of larger agglomerates are shown as a function of the agglomerate size-scale $l = \sqrt{\text{area}}$ in figure 3(b). The area on the substrate’s surface of each structure was extracted from SEM images. For the full raster scans, we consider the area of the respective agglomerate and evaluate those positions that show connected fluorescence above a $1/e$-threshold of 36.8% of the maximum count rate measured in the smallest agglomerate. This threshold yields good agreement when evaluating the area...
Figure 3. Lifetime distribution as a function of agglomerate size. (a) Position-dependent fit value of slow decay $\tau_2$ in four different agglomerates with a common color bar. (b) Fit coefficients $\tau_2$ over the size-scale $l = \sqrt{\text{area}}$ of the agglomerate they were measured in. The individual measurements are shown in the color bar used in (a). The region of single measurements without raster scan is shaded in blue, and the four agglomerates shown in (a) are highlighted and numbered according to figure 1. The data for each measurement is shown in appendix D. The count rate weighted average (standard deviation) is shown in thicker blue dots (bars) and was fitted by a phenomenological exponential decay.

(i.e. the size of connected fluorescence above the threshold) of agglomerates with the optical setup compared to the more exact measurements with the SEM.

In order to extract the average lifetime of an agglomerate, we consider the count rate weighted average $\tau_{2,av} = \frac{\sum_i \tau_{2,i} c_i}{\sum_i c_i}$, where $i$ is the number of a measurement taken at a single position, $\tau_{2,i}$ is the lifetime from the fit using equation (1) and $c_i$ is the count rate measured during the pulse sequence detailed in section 2. This average weights the contribution from each NV center excited to the total fluorescence measured in a scan equally. Therefore, the systematic error between a measurement, where the whole agglomerate is excited at once, and multiple measurements where only a part of the agglomerate is excited is reduced. For nanodiamonds of the smallest size category, we find a lifetime ranging from 21 ns to 33 ns with a count rate weighted average of 26 ns, which is in good agreement to the previously reported optical lifetime of 23 ns [36] and 25 ns [27]. With increasing agglomerate size, the variation within each data-set shows, in general, a decrease. The position-resolved scans show that data points with a significantly higher lifetime than average are typically located at the edges of the agglomerates (cf agglomerate 3). The average of fitted $\tau_2$ coefficients shows a reduction from 26 ns towards a value of 19.5 ns. The count rate of individual measurement points varies within and between the agglomerates, which could lead to a pile-up effect at high count rates impacting on the measurement and fit parameters. In appendix C, we have further investigated such a dependence on the count rate and observed no effect in the range of the count rates measured and analyzed for the agglomerate scans. Therefore, we interpret the reduction of $\tau_2$ with increasing agglomerate size as a transition to bulk-like optical properties due to the change of the local optical DOS via the effective refractive index. We determine the length scale $l_{db}$ on which this transition occurs by an exponential fit $A \cdot \exp(-l/l_{db}) + c$ to the weighted average, with a decay constant $l_{db} = 1.8(3) \mu m = 2.8(3) \cdot \lambda_{ZPL}$, where the error is given by the standard deviation of the fit coefficient and $\lambda_{ZPL} = 637$ nm is the emission wavelength of the NV center’s ZPL.

Even though the lifetime does not reach the NV center’s bulk lifetime in the order of 12 ns to 13 ns which is widely reported in the literature [34, 35], we assume that a transition to bulk-like optical properties is observed. This assumption stems from the fact that the graphitic surface [40] of nanodiamonds and small voids between neighboring nanodiamonds lead to a reduction of the effective refractive index in contrast to bulk diamond. Further, the SEM images taken after the measurements suggest that the expansion of agglomerates along x- and y-directions, i.e., the extension along the glass substrate’s surface, is much bigger than the extension along the z-direction, i.e., the laser beam axis. The agglomerates shown in the data above indicate to consist of typically one or two nanodiamond layers. For the largest agglomerate, however, we observe additional crystals on top of these layers (see figure 1(a), right image), which was also evident by an increase in the count rate (see figure 1(b), right image). In this agglomerate, we observe at the position of highest count rate a further significant reduction of $\tau_2$ to 15.2 ns. At the same time, the section without such an extension into z-direction shows lower count rates and $\tau_2 = 20$ ns both in the same order of the previously discussed large planar agglomerates.
In addition to this phenomenological length scale of a steady state, we model the impact of an effective refractive index \( n_{\text{eff}} \) inside an agglomerate on the fluorescence lifetime, as shown in the inset of figure 4. We average the refractive index in two steps. First, a local average is formed within the volume of a sphere with a radius \( r_i \). This local average is in a second step integrated over the volume of the agglomerate, which we approximate by a cylinder. The final result thus differs for different choices of \( r_i \). The details of this calculation is presented in appendix E. Our model neglects local fields \([41]\), which was previously used to explain the fluorescence lifetime discrepancy between bulk- and individual nanodiamonds \([27]\). The lifetime then depends on the refractive index according to

\[
\tau_{\text{rad}}(n_{\text{eff}}) = \tau_{\text{vac}} / n_{\text{eff}}.
\]

Applying this model to the calculated effective refractive index, we find good agreement to our measurement data for length scales \( r_i \in \{130 \text{ nm, 300nm}\} \), as depicted in figure 4. This result reproduces the observation that the macroscopic definition of the refractive index \( n \) can be applied on a length scale of approximately \( \lambda / 4 \) as reported for crystallized Eu\(^{3+}\):Gd\(_2\)O\(_3\) \([42]\). For larger agglomerates (grey shaded region), we observe more than a single layer of nanodiamonds in SEM images, which gives rise to a further reduction of the fluorescence lifetime.

### 4. Collective effects

Collective emission can be observed in different observables. First, an ensemble of collectively emitting quantum sources, which can be described by a macroscopic dipole moment or, equivalently, spin, features a larger emission rate and hence a reduced lifetime \([6, 8]\). Second, for collectively emitting NV centers, the statistics of photons detected is expected to change, which can be observed in the second-order correlation function as enhanced correlation. In reference \([15]\) such superradiant emission of NV centers in individual, highly-doped nanodiamonds at room temperature with dephasing rates of the collective spin in the order of \( \gamma_d/2\pi = 20–40 \text{ MHz} \) for the \( m_s = 0 \)-state has been reported. Even in the presence of large phononic sidebands due to local vibrational modes at room temperature, superradiant emission can be observed. The local vibrational modes decay into global long-wavelength modes \([15, 43]\), erasing any information about the local environment and enabling subsequent superradiant photon emission. Increased optical emission has been reported over different spectral emission regions. However, to ensure the detection of photon bunching due to superradiant emission within an individual spectral domain, a narrow emission band around the ZPL is analyzed in the correlation measurements in reference \([15]\). We follow this procedure and show the emission spectrum measured in an agglomerate in appendix A.

In our experiment, we find fast-decaying fluorescence lifetimes of our nanodiamond agglomerates using the fitting procedure according to equation (1). The data is presented and discussed in more detail in appendix F. In brief, we observe average fast components of the lifetime in the order of \( \tau_1 = 4 \text{ ns} \) in all agglomerates whereas the amplitude of this fast component \( a_1 \) increases with the agglomerate size. However, a lifetime reduction and non-exponential decay from NV centers in diamond can be due to several effects...
ranging from collective emission of NV centers [15], but also non-radiative decay channels [31, 44, 45], population of different $m_I$-states [35–37], emission from the neutral charge state [46], charge state instabilities [47], or blinking [48]. We discuss each of those contributions in appendix F and conclude that the lifetime measurements do not allow differentiating between the influence of dark decay and collective emission. However, if the observed non-exponential decay was only due to dark decay channels we would expect to observe the absence of photon bunching in correlation measurements. By contrast for collective emission, we expect to see photon bunching in the pulsed correlation measurement and the absence of photon bunching in a cw measurement. Such a discrepancy between pulsed and cw excitation is consistent with the interpretation of collective, superradiant effects. The formation of a collective spin is known to feature a finite coherence time, which was reported to be $1/\gamma_d \approx 4–8$ ns for highly-doped nanodiamonds at room temperature [15]. The pulsed measurement with the sequence described in section 2 allows recording the correlation at much shorter times in the order of 500 ps, being one order of magnitude smaller than the expected coherence time of the collective spin. By contrast, the cw excitation averages over long times, where a collective spin is expected to dephase and cease. We, therefore, investigate the second-order correlation function in detail.

4.1. Second-order correlation function

The second-order correlation at zero time delay $g^{(2)}(0)$ can be written via the photon distribution as [49]

$$g^{(2)}(0) = \frac{\langle \hat{a}^+ \hat{a}^+ \hat{a} \hat{a} \rangle}{\langle \hat{a}^+ \hat{a} \rangle \langle \hat{a}^+ \hat{a} \rangle} = 1 + \frac{\Delta n^2 - \langle n \rangle}{\langle n \rangle^2},$$

(3)

where $\langle n \rangle = \langle \hat{a}^+ \hat{a} \rangle$ is the expectation value and $\Delta n^2 = \langle n^2 \rangle - \langle n \rangle^2 = \langle (\hat{a}^+)^2 \hat{a}^2 \rangle + \langle \hat{a}^+ \hat{a} \rangle - \langle \hat{a}^+ \hat{a} \rangle$ is the variance of the photon number expressed with creation and annihilation operators $\hat{a}^+$ and $\hat{a}$. Equation (3) shows that the second-order correlation depends on the first and second moment of the photon distribution. The measurement of the second-order photon correlation with a Hanbury Brown–Twiss interferometer in the time domain is a standard tool to observe collective effects. The experimental setup used has moderate detection efficiencies and harnesses SPCMs as click-detectors for photon counting. Therefore, the second-order correlation function can be extracted from a coincidence measurement precisely, whereas the measurement of the full photon number distribution needs high detection efficiencies and number resolving detectors [50]. We measure coincidences after pulsed excitation and approximate the value by integration over a time window of $t' = 0.5$ ns as explained in detail in appendix B. We refer to this value as $g^{(2)}(0)$. As explained above, we expect to see photon bunching decaying on the order of a few nanoseconds [15] in this pulsed measurement as a signature of collective emission of NV centers due to super-Poissonian photon statistics ($\Delta n^2 > \langle n \rangle$) [49]. The measurement data of such a single measurement showing this expected behaviour is presented in figure 1(b) and in appendix B in figure 11(b).

The position-dependent measurements of $g^{(2)}(0)$ are shown in figure 5(a). We measure photon bunching ($g^{(2)}(0) > 1$) as well as photon anti-bunching (1) in all agglomerates. From these measurements, we see that the amount of photon bunching reduces and shows fewer fluctuations with increasing size. All data points are presented as a function of the count rate measured at the same spatial position and time in figure 5(b). We differentiate between the measurements of small nanodiamond agglomerates measured at a single position, planar agglomerates, and the largest agglomerate, which shows the most substantial lifetime reduction. We observe higher photon bunching for lower count rates and a higher variance in the data. In large structures, we find the strongest photon bunching located at the edges of the agglomerate where the count rate is low and almost constant values of $g^{(2)}(0) \approx 1.06$ in the central planar sections where the count rate is high. Such a photon bunching can be explained by forming either thermal emission or collective emission of quantum emitters at the ZPL. We note that photon bunching due to thermal emission cannot be detected within the time resolution of the detectors due to the relatively large range of transmitted wavelengths by the ZPL filter (637(3) nm). Additionally, no indication of sharp emission resonances except for the ZPL of NV centers was observed in cw and pulsed fluorescence spectra.

To differentiate between these different physical effects, we perform a second measurement of the second-order correlation in cw-mode for the largest agglomerate sample. A comparison of pulsed and cw second-order correlation is shown in appendix G. In the cw-measurement, we observe no bunching or slight anti-bunching, expected from an ensemble of quantum emitters acting as individual emitters. As explained initially in this section, this discrepancy between pulsed and cw excitation is a signature of coherent collective emission. Further, we have ruled out systematic errors by verifying the pulsed measurement scheme and evaluation on a nanodiamond with a low concentration of NV centers (see appendix H). We find similar results for pulsed and cw measurement of the second-order correlation.
function in this control sample. Therefore, we conclude that on short time scales, a collective emission of NV centers is taking place in nanodiamonds with high NV concentrations. We mention here that the observation of super-Poissonian photon statistics does not rule out the presence of an additional dark decay channel.

To obtain further insight, we adopt a model of collective emission in single-nanodiamond crystals [15] and apply it to the agglomerates of nanodiamonds investigated here. We assume the nanocrystals to consist of many spectral domains, where each domain acts collectively, but different domains emit independently from each other. A sketch of the model is presented in figure 6(a).

In this model, an important figure of merit is the collective-domain size $N$, i.e., the number of NV centers acting collectively in a single domain, following previous work [6, 8, 15]. In a simple approach, one could interpret an increased photon bunching indicated by an increased second-order correlation as a sign of increased super-Poissonian emission due to larger collective domains. In this case, the observation of the highest photon bunching in individual nanodiamonds and at the borders of agglomerates would imply that collective domains are smaller in central positions of agglomerates. However, this contradicts the observation of increased fast optical decay in larger agglomerates.

In our model, we assume multiple domains, where $D$ is the total number of domains contributing to the emission. We model a single collective domain of $N$ emitters with Dicke-states $|J, M\rangle$ as a spin-ladder of spin $1/2$ systems, where $J$ describes the maximum total spin of a domain and $M \in \{-J, -J+1, \ldots, J\}$ are the eigenvalues along the quantization axis, denoting the number of excitations in the system as depicted in figure 6(a). We calculate the second-order correlation and the expectation value $\langle n_N \rangle$ and variance $\Delta n^2_N$ of the photon number. The calculation and more detail on the model are presented in appendix I. We furthermore assume that single nanodiamonds and agglomerates comprise many domains, where each domain acts independently from all other domains, i.e., their photon distributions are statistically independent. Each domain $i = 1, \ldots, D$ is described by the respective expectation value $\langle n_{Ni} \rangle$ and variance $\Delta n^2_{Ni}$ of the individual domain’s photon number. Thus, we calculate the total photon distribution’s first and second moment of multiple domains as the sum of individual domains according to

$$\langle n \rangle = \sum_{i=1}^{D} \langle n_{Ni} \rangle$$

and

$$\Delta n^2 = \sum_{i=1}^{D} \Delta n^2_{Ni}.$$
Figure 6. (a) Model for collective emission in nanodiamond agglomerates. NV centers are assumed to form \( i \in \{1, \ldots, D\} \) spectral domains (shown in different colors) of \( N \) two-level emitters. Each domain is described individually by Dicke-ladder-states. (b) Calculated auto-correlation function \( g^{(2)}(0) \) and (c) Fano factor \( F \) as a function of the total number of domains \( D \) for different domain sizes \( N \) under the assumption of a maximally mixed state. (d) Square root of Fano factor \( \sqrt{F} \) as a function of the domain size \( N \).

The second-order correlation at zero time delay of \( D \) domains with different domain sizes \( N_i \) can be calculated using equation (3). Assuming an initial maximally mixed state \( P_{J\rightarrow M}(0) = 1/(N + 1) \) [15], we have calculated the correlation function of different domain sizes \( N \) over the total number of such domains \( D \), as depicted in figure 6(b).

The values calculated for \( D \) domains of independent quantum emitters with a domain size \( N = 1 \) yield the well-known relation of \( g^{(2)}(0) = 1 - 1/D \). However, \( g^{(2)}(0) \) depends on the domain size \( N \) as well as the number of such domains \( D \). Therefore, the second-order correlations approach \( g^{(2)}(0) \approx 1 \) for a large number of independent domains \( D \) for all domain sizes \( N \). We emphasize that the theory data shown in figure 6(b) cannot serve as a look-up table for the experimental data in figure 5(b), because in experimental realizations the distribution of domains and domain sizes is unknown. The theoretical data show the general reciprocal behaviour of the second order correlation to the system size. I.e. the difference \( d_c \) to an uncorrelated system \( d_c = g^{(2)}(0) - 1 \) reduces by a factor of 2 when the system size increases by the same factor. This behaviour is independent of the initial state and individual domain sizes, which are both fixed in the theory plot. By contrast, in the experiment with various agglomerate sizes, the number of domains excited by the laser beam has high variations. We have no measurement tool available to infer these quantities exactly, and, as explained above, a simple comparison of measured correlation functions between such different samples cannot provide complete information on the average collective domain sizes. In order to still trace the emergence of collective effects in experimental data, in the following we establish an alternative method which is independent on the number of domains and based on the number fluctuations of the photon distribution.

4.2. Fluctuations

To have a figure of merit that is (i) independent of the number of contributing domains \( D \) and has (ii) a linear scaling for collective domain sizes \( N \), we introduce the square root of the Fano factor

\[
\sqrt{F} = \sqrt{\frac{\Delta n^2}{\langle n \rangle}}
\]

for quantification. The Fano factor \( F \) measures deviations from a Poissonian distribution, where a value larger (smaller) than unity indicates super-(sub-) Poissonian fluctuations. Based on the discussion above, we calculate and plot the Fano factor assuming a maximally mixed state \( P_{J\rightarrow M}(0) = 1/(N + 1) \) in different domain sizes \( N \) over the total number of such domains \( D \) in figure 6(c). We find that the Fano factor is independent of the number of contributing domains \( D \) and shows almost quadratic scaling with domain...
We emphasize that the scaling is strongly dependent on the initial assumption on the Dicke-ladder state. For the occupation of the highest ladder state, linear scaling of the Fano factor with the domain size \( N \) is calculated. Considering the square root of the Fano factor \( \sqrt{F} \), we find a quantity that shows linear scaling for the collective domain size \( N \) while being independent of the number of contributing domains \( D \) as depicted in figure 6(d).

However, the Fano factor is no direct measurement observable of the photon distribution. To access the Fano factor, we calculate it under the assumption that the expectation value of the photon number \( \langle n \rangle \) scales linearly with the total count rate detected in a pulsed measurement. We mention here that this approximation is only exact, assuming (i) a vanishing excitation pulse-width in time and (ii) \( i = 1, \ldots, N \) independent quantum emitters each emitting with \( \langle n_i \rangle = 1 \). In the experimental sequence, the laser pulse has a width of 500 ps, which is significantly smaller than the typical lifetime of the NV center. This justifies the proportionality of the measured count rate to the number of NV centers excited. However, for a collective domain, the mean photon number \( \langle n \rangle \) will exceed the number of its constituents and scale quadratically during emission, as described by Dicke [6]. Since small domain sizes were reported in [15], we expect the calculated square root of the Fano factor \( \sqrt{F} \) to show a relatively small deviation. In this case, the calculated quantity of \( \sqrt{F} \) can be understood as a quantitative measure for the formation of collective effects.

To estimate the mean photon number \( \langle n \rangle \), we use the measured count rate and normalize it to the count rate measured with the same sequence for a nanodiamond of the control sample housing 10 NV centers shown in appendix H. This allows us to find a bias and scaling for the Fano factor and makes our results comparable with further studies in different experimental setups. Moreover, we calculate the variance as \( \Delta n^2 = (g^{(2)}(0) - 1)\langle n \rangle^2 + \langle n \rangle \) from the measured quantities of the count rate and second-order correlation function.

Figure 7 shows the calculated square root of the Fano factor \( \sqrt{F} \) as a function of the position for four agglomerates. Our analysis leads to non-physical Fano factors \( F \leq 0 \). However, the error given by evaluating the second-order correlation measurement allows \( F > 0 \) for all those measurements. In contrast to the measured second-order correlation function shown in figure 5(a), we find an increase of the calculated square root of the Fano factor \( \sqrt{F} \) with the agglomerate size. We observe small values of \( \sqrt{F} \) in all agglomerates located on the edges and an increasing modulus towards central positions. From these measurements, we deduce that the average collective domain size, which scales at least linearly with \( \sqrt{F} \),
increase in nanodiamond agglomerates. For the largest agglomerate, we find maximum $\sqrt{F} = 6.75$ at the position of maximum count rate, minimum $\tau_2$ lifetime coefficient, and extension into the $z$-direction. This value exceeds the maximum of nanodiamonds of the smallest size category by a factor of 2.5. The discussion above suggests that the average collective domain size increases at least by the same amount. Further, large values of $\sqrt{F}$ indicating larger collective domains are observed in regions of high nanodiamond density. They might suggest a formation of such domains over multiple nanocrystals nearby.

In addition, the square root of the Fano factor $\sqrt{F}$ allows drawing wide-reaching conclusions on the emission properties compared with other observables. Therefore, we evaluate $\sqrt{F}$ for each spatial measurement position as a function of other measurement quantities, i.e., the fit coefficient $\tau_1$ of the optical decay and the second-order correlation function. Here, we differentiate between three classes: individual nanodiamonds, planar agglomerates, and the largest agglomerate with larger extension into the $z$-direction, as shown in figure 8.

As described above, the Fano factor is calculated from the count rate and time-integrated second-order correlation function $g^{(2)}(0)$. The highest values of the second-order correlation functions are measured in single nanodiamonds and small agglomerates. However, they do not coincide with the locations of the highest calculated Fano factors. This depicts the discrepancy between the measured photon bunching and the average collective domain size of the discussion above. The maximum square root of the Fano factor $\sqrt{F}$ = 6.75 is found at $g^{(2)}(0) = 1.07$ in the largest agglomerate. In each class, we observe a larger photon bunching in the presence of a smaller count rate leading to smaller values for $\sqrt{F}$ indicating smaller collective domains.

Further, prominent collective effects, i.e., high Fano factors, are observed in agglomerates where the fast decay component is in the order of $\tau_1 = 4$ ns in all three classes. These measurement results show that the measured fast decay component in the order of $\tau_1 = 4$ ns can be attributed to collective emission of NV centers. The increasing amplitude of this component in larger agglomerates shown initially in figure 1 and discussed in detail in appendix F is consistent with observing a larger average collective domain size. Decoherence on a time scale of a few nanoseconds and the number of emitters localized within a wavelength-cube and hence contributing collectively both limit the decay time to shorter values. The decoherence time is in the order of 4 ns to 8 ns [15] being equal to the decay time. At the same time, the laser spot extension is in the order of the wavelength. This suggests that in our system the number of collectively excited emitters is close to the optimum for the doping of our nanodiamonds at hand.
5. Conclusion

In conclusion, we have traced the quantum-optical emission properties of NV ensembles in doped nanodiamonds of different agglomeration states. We observe the transition to continuous, bulk-like fluorescence-emission properties with increasing agglomerate size to occur on a phenomenological length scale of $1.8 \, \mu m = 2.8 \cdot \lambda_{ZPL}$. A theoretical model for the fluorescence lifetime of single-layer agglomerates shows good agreement to the measurement data when a length scale of $r_i \in \{130 \, nm, 300 \, nm\}$ is used for the definition of the macroscopic refractive index $n$.

Furthermore, we observe collective, superradiant emission in pulsed-measurement sequences. While the second-order correlation function does not yield a clear signal of collective effects, introducing the Fano factor as a novel quantity based on the number fluctuations of the photon statistics, we reveal superradiant emission even for an ensemble of collective domains with varying size. We observe high Fano factors in the emission of NV centers in agglomerates, where a high amount of nanodiamonds is illuminated within the region of the laser focus, which might stem from the formation of collective domains over multiple nanocrystals. Further, the collective emission was attributed to an additional fast optical decay in the order of $\tau_1 \approx 4 \, ns$.

The observation of collective emission in a poly-crystalline solid-state system at room temperature paves the way towards application of superradiance in a robust and versatile system. Further steps will include studies with varying excitation pulse widths and amplitudes to shed light on initial Dicke-state formation and the dephasing of the macroscopic collective spin. Spatial information about the size of collective domains can be accessed by a change of the excitation region either via super-resolution techniques such as stimulated emission depletion spectroscopy [51, 52] or via changes in the optical setup towards larger/smaller beam waists. The integration of NV centers into a photonic environment featuring the coupling to a single optical mode such as optical cavities [53] and waveguides [54–58] could allow observation and control of collective interaction on larger length-scales. Combining such devices with other established methods such as spin-to-charge conversion of NV centers [59] could enable further control of the collective emission direction and coupling in a collective system on the meso-scale. Beyond the fundamental understanding, the highly entangled Dicke states involved render collective emission attractive for applications in quantum metrology [17–20].

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Data availability statement

All data that support the findings of this study are included within the article (and any supplementary files).

Appendix A. Measurement setup

The microscope objective (Nikon, CFI Plan Apochromat Lambda 60XC) used in the measurements has a numerical aperture of $NA = 0.95$ and a magnification of 63 together with the tube lens in front of the camera. The correction ring was adjusted for the cover-glass thickness of 0.17 mm. The objective focuses the laser on a diffraction-limited spot. The spatial fluorescence response from a large agglomerate detected with the EMCCD camera is shown in figure 9. We observe an almost Gaussian shape of the fluorescence intensity which is broadened due to scattering of the nanocrystals. The FWHM of the measured intensity distribution is $w_{FWHM} = 1.1 \, \mu m$. In the agglomerate scans with SPCMs, a single measurement point is therefore integrating over this region due to scattering. The two SPCMs (lasercomponents, COUNT-T-100) used in the HBT setup have a dead-time of 45 ns and showed a maximum count rate of 0.124 MHz during the measurements corresponding on average to less than 0.006 counts in the time window of the dead-time. The count rate when only the substrate without a diamond nearby is illuminated by the laser was measured as 600 Hz, the dark count rate (laser turned off) was 250 Hz. The TDC is a TimeTagger20 from Swabian Instruments, which was programmed to simultaneously record data for FLIM (histogram mode), the second-order correlation function (correlation mode), and the count rate. A bandpass filter is used to
transmit the light around the ZPL of the fluorescence spectrum. Such a fluorescence spectrum measured in an agglomerate and the filtered spectrum are presented in figure 10.

**Appendix B. Correlation function measurements**

In order to approximate the second order correlation at zero time delay \( g^{(2)}(0) \) in a pulsed measurement, we follow the method detailed in [15]. In brief, we measure the time difference of coincidences \( c(t) \) in an interval of \([-1.5625 \, \mu s, 1.5625 \, \mu s]\) showing in total 15 repetitions of the pulse sequence with a bin-size of 25 ps. We fit all peaks separately with the function 
\[
    f(t) = A \exp\left(-\frac{|t - t_i|}{\tau}\right) + c
\]
in order to find the center \( t_i \) of the \( i \)th peak. Afterwards, a time window \( t' \) centered on this respective local maximum is used to compare the coincidences per time window of the 0th peak with the other 14 peaks labeled as \([-7, \ldots, -1, 1, \ldots, 7]\). The second-order correlation function is then the integration over an infinitesimal time
\[
    g^{(2)}(0) = \lim_{t' \to 0} \int_{-t'/2}^{t'/2} \frac{c_0(t)}{\sum_{i \neq 0} c_i(t)} \, dt,
\]
We approximate \( g^{(2)}(0) \) using a finite time window \( t' \) as
\[
    g^{(2)}(0)|_{t'} = \int_{-t'/2}^{t'/2} \frac{c_0(t)}{\sum_{i \neq 0} c_i(t)} \, dt,
\]
and calculate the error via error propagation of the standard deviation of \( c_i(t) \). Measurement data and the extracted correlation function over different widths of the time window \( t \) are depicted in figure 11.

Ever decreasing time windows lead to a more precise approximation of the limit value but vice versa to an increased uncertainty, stemming from the higher variance of coincidences for smaller time windows of

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**Figure 9.** (a) Spatial fluorescence response of an agglomerate upon laser excitation on an EMCCD camera. The \( x \) and \( y \)-axis are displayed in the scale of the sample. (b) Data along the \( x \)-axis at \( y = 19.4 \, \mu m \) (blue) and Gaussian fit to the data points (orange).

**Figure 10.** Fluorescence spectrum of NV centers measured in a diamond agglomerate (blue). The orange curve gives the transmitted spectrum through the filter used in all measurements.
peaks $[−7, \ldots, −1, 1, \ldots, 7]$. The time window chosen to approximate the value of $g^{(2)}(0)$ is $t′ = 0.5$ ns in accordance to reference [15]. We refer to this approximated value as $g^{(2)}(0)_{0.5 ns}$ in the main text.

**Appendix C. Count rate dependence of fit parameters**

We have investigated the dependence of the fit parameters $\tau_1$, $\tau_2$, and $a_1$ in equation (1) on the count rate of the respective measurement, repeating a FLIM measurement in an agglomerate on a single position and constant laser power. Further, we have applied different grey filters in front of the detectors to vary the count rate. We have normalized and subsequently fitted the data according to equation (1) and plotted the extracted fit coefficients $\tau_1$, $\tau_2$, and $a_1$ over the count rate in figure 12.

We observe constant values within the statistical error of the fit for low count rates up to $≈ 0.25$ MHz. The maximum count rate measured in the agglomerate scans is 0.124 MHz. We, therefore, rule out an impact of the count rate in all presented measurements used for the analysis. Even higher count rates lead to a significant change in the lifetime measurements and fit parameters. Especially, the fit parameter $\tau_2$ increases for such high count rates, which is in clear contrast to our observations of a lifetime reduction in large agglomerates, where the highest count rates are measured.

**Appendix D. Small nanodiamond agglomerates**

The individual measurements of nanodiamonds where a single measurement at maximum count rate was taken instead of a full confocal scan are presented in figure 13. Small lifetimes in the order of 21 ns are observed for any size category, whereas the highest observed lifetimes in each category reduces.
Appendix E. Model for the effective refractive index

We model the influence of the surrounding environment by means of the effective refractive index $n_{\text{eff}}$ as depicted in the inset of figure 4. We assume a cylinder-shaped agglomerate placed on a glass substrate with fixed height of $h = 100 \text{ nm}$ and a variable radius $R$. Further, we model the agglomerate to consist of multiple nanodiamonds with air gaps in between with a filling factor of $f = 0.74$, which is the value for close packing of equal spheres. Therefore, the local refractive index of the agglomerate is calculated as $n_{\text{aggl}} = fn_{\text{diamond}} + (1-f)n_{\text{air}} = 2.05$, where $n_{\text{diamond}} = 2.42$ and $n_{\text{air}} = 1.00$. We calculate the effective refractive index $n_{\text{eff}}(R', h')$ at position $(R', h')$ inside the agglomerate ($R' \in \{0, R\}, h' \in \{0, h\}$) by integrating over the local refractive indices inside a sphere centered at $(R', h')$ with a variable radius $r_i$ as

$$n_{\text{eff}}(R', h', r_i) = \int_0^{2\pi} \int_0^{r_i} n_{\text{loc}}(R', h', r, \phi, \theta) r^2 d\phi d\theta.$$  \hspace{1cm} (E1)

Where $n_{\text{loc}}(R', h', r, \phi, \theta)$ is the local refractive index of the material, i.e. $n_{\text{aggl}} = 2.05$, $n_{\text{air}} = 1.00$ or $n_{\text{glass}} = 1.52$, and $r_i$ is the length scale for the influence of the environment on the effective refractive index. In the simulation, we approximate this integral as a discrete sum over sampling points with a resolution of 10 nm. The effective refractive index of an agglomerate with radius $R$ is then calculated as the sum of all effective refractive indices $n_{\text{eff}}(R', h', r_i)$ in the region of the agglomerate.

Appendix F. Fast optical decay

In the literature the additional fast optical decay is attributed to several processes. These include collective effects [15], dark decay channels created by impurities and surface charge traps [31, 44, 45], as well as different ISC rates from the optical excited state’s spin projections $m_s = 0$ or $m_s = \pm 1$ [35–37].

Further effects, such as blinking and charge state instabilities, have been reported. Blinking is observed on significantly larger time scales in the millisecond regime for NV centers in nanodiamonds smaller than the investigated ones [48]. Charge state dynamics of NV-ensembles in nanodiamonds of similar size have been investigated in [47], and it has been shown that the negatively charged NV-centers dominate the emission and a conversion to the neutral NV0 state appears in the millisecond regime due to optical excitation. Therefore, both effects cannot explain the fast optical decay within the nanosecond regime.
Figure 14. (a) Fast decay lifetime $\tau_1$ of the fit model in dependence of the spatial position for the four agglomerates discussed in the main text and (b) coefficients $\tau_1$ of all agglomerates in dependence on the agglomerate size measured in. (c) Fast decay fit amplitude $a_1$ in dependence of the spatial position for the four agglomerates discussed in the main text and (d) in dependence of the agglomerate size.

The extracted fit-coefficients of the fast decay lifetime $\tau_1$ as well as the amplitude $a_1$ of the fit model (equation (1) in the main text) for the four agglomerates discussed in the main text is shown in dependence of the position in figures 14(a) and (c). All data are shown in dependence of the agglomerate size in figures 14(b) and (d).

We observe on average $\tau_1 \approx 4$ ns in all agglomeration-states. The variation in the data is decreasing for larger agglomerates. In the largest agglomerate, we observe higher values of $\tau_1$ at positions of extension into the $z$-direction. Such small lifetimes are far below the reported decay of $m_s = \pm 1$-states in nanodiamonds (12.8 ns [36]) as well as bulk diamond (7.3–7.8 ns [35, 37]). Furthermore, also the neutral NV0 charge state shows larger fluorescence lifetimes [46]. Therefore, we rule out that the fast decay $\tau_1$ stems from increased ISC of $m_s = \pm 1$-states or neutral NV0 centers.

Further, an increase of $a_1$ on small length-scales within the small size categories of single scans is observed. This effect is also evident in figure 1(c) of the main text, where the zoom-in section compares the contribution of the fast decay and the smallest nanodiamond-agglomerates show less pronounced fast decay. Again, the variation in the data decreases with increasing agglomerate size. The largest agglomerate is an exception and shows higher amplitudes $a_1$ at those positions where we see an extension into the $z$ direction.

In general, the effect leading to the fast decay in the order of $\tau_1 \approx 4$ ns is more prominent in agglomerates. However, from the lifetime measurements we cannot differentiate between the influence of two possible contributions, i.e. dark decay and collective emission. Therefore, we focus on the analysis of the second order correlation in the main text.
Figure 15. (a) Measured cw second order correlation function at zero time delay $g^{(2)}(0)$ and (b) difference of approximated second order correlation from the pulsed measurement (see figure 5(a) in the main text) and cw second order correlation function $g^{(2)}(0) - g^{(2)}(0)$ in dependence of the position.

**Appendix G. Comparison of cw and pulsed second-order correlation**

For the largest agglomerate discussed in the manuscript the correlation function in dependence of the position has been detected in cw-mode as shown in figure 15(a) in a second measurement. The difference of results in pulsed and cw-mode is shown in figure 15(b).

As mentioned in the main text, we measure in general a cw second order correlation $g^{(2)}(0) < 1$. The difference between cw and pulsed measurement shows a trend to larger values at positions of comparatively small count rate at the edges of agglomerates.

**Appendix H. Control sample**

In order to rule out systematic errors of measurements performed with the pulsed sequence, nanodiamonds with few NV-centers but equal size were investigated with both measurement schemes. For these nanodiamonds we expect no formation of collective domains leading to anti-bunching and the agreement of both methods to infer the normalized second order correlation function $g^{(2)}(0)$. The integration time for both measurements has been increased but the applied pulse sequence has been left unchanged to all other pulsed measurements. The comparison of both measurements is depicted in figure 16. The minimum value of the measured second order correlation function is shifted by approximately 10 ns stemming from the different arm lengths of the HBT interferometer which employs different lengths of optical fibers.

Normalization of the cw measurement presented in figure 16(a) was achieved by the formula

$$g^{(2)}(t) = G^{(2)}(0) \Delta T \frac{b}{N_1 N_2},$$

where $G^{(2)}(t)$ is the measured correlation histogram, $b$ is the bin-width and $N_1$ and $N_2$ are the number of events in each channel. The pulsed measurement was normalized with the method detailed in appendix B. For the investigated nanodiamond we find good agreement within the error bars of both measurement methods and conclude that this nanodiamond houses 10 NV centers. From the measured count rate of those 10 NV centers, we infer the number of NV centers contributing in the agglomerate scans, in order to normalize the calculated Fano factor.

**Appendix I. Photon distribution of single collective domains**

In order to calculate the second order correlation we use the expectation value $\langle n \rangle$ and the variance $\Delta n^2$ of the photon number. Since we are only interested in the second order correlation at zero time delay $g^{(2)}(0)$, we assume a vanishing excitation pulse to excite the emitters into the highest $J = N/2$-subspace and neglect decoherence processes of the collective spin to lower subspaces. In the model, we describe this by an initial population of these Dicke-states $P_{J,M}(0)$. The collective state after excitation is then given by $\rho_N(0) = \sum_M P_{J,M}(0) |J,M\rangle \langle J,M|$. 
Figure 16. Comparison of measurement methods in nanodiamonds housing few NV centers. (a) Measured cw second order correlation function; (b) coincidence measurement of pulsed excitation. (c) Approximated second order correlation at zero time delay for different time windows \( t' \) of the pulsed measurement.

For such a single Dicke-ladder of size \( N \) we calculate the expectation value \( \langle n_N \rangle \) as

\[
\langle n_N \rangle = \langle J^+ J^- \rangle = A
\]

and the variance \( \Delta n_N^2 \) as

\[
\Delta n_N^2 = \langle J^+ J^- J^+ J^- \rangle + \langle J^+ J^- \rangle - \langle J^+ J^- \rangle \langle J^+ J^- \rangle = B + A - A^2,
\]

where we introduced the quantities \( A = \langle J^+ J^- \rangle \) and \( B = \langle J^+ J^- J^+ J^- \rangle \). We calculate \( A \) and \( B \) by taking the trace

\[
A = \text{Tr}(\rho_N(0)A)
\]

\[
= \sum_{M=-J}^{M=J} P_{JM}(0) \langle J^+ J^- J^+ J^- | J, M \rangle
\]

\[
= \sum_{M=-J}^{J} P_{JM}(0) [J(J+1) - M(M-1)],
\]

\[
B = \text{Tr}(\rho_N(0)B)
\]

\[
= \sum_{M=-J}^{M=J} P_{JM}(0) \langle J^+ J^- J^+ J^- | J, M \rangle
\]

\[
= \sum_{M=-J}^{J} P_{JM}(0) [J(J+1) - M(M-1)][J(J+1) - (M-1)(M-2)].
\]

Entering \( A \) and \( B \) leads to the expectation value

\[
\langle n_N \rangle = \sum_{M=-J}^{J} P_{JM}(0) [J(J+1) - M(M-1)]
\]
and the variance

\[
\Delta n^2 = \sum_{M=-j}^{j} P_{JM}(0) [J(J+1) - M(M-1)] [J(J+1) - (M-1)(M-2)]
+ \sum_{M=-j}^{j} P_{JM}(0) [J(J+1) - M(M-1)]
- \left( \sum_{M=-j}^{j} P_{JM}(0) [J(J+1) - M(M-1)] \right)^2.
\]

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