Spectroscopic investigations of negatively charged tin-vacancy centres in diamond

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Abstract

The recently discovered negatively charged tin-vacancy centre in diamond is a promising candidate for applications in quantum information processing (QIP). We here present a detailed spectroscopic study encompassing single photon emission and polarisation properties, the temperature dependence of emission spectra as well as a detailed analysis of the phonon sideband and Debye–Waller factor. Using photoluminescence excitation spectroscopy we probe an energetically higher lying excited state and prove fully lifetime limited linewidths of single emitters at cryogenic temperatures. For these emitters we also investigate the stability of the charge state under resonant excitation. These results provide a detailed insight into the spectroscopic properties of the SnV⁻ centre and lay the foundation for further studies regarding its suitability in QIP.

1. Introduction

As the field of quantum information processing (QIP) is rapidly expanding, significant effort has been spent on the search for quantum bit systems providing favourable properties such as individually addressable spins with long coherence times, optical emission spectra with narrow homogeneous and inhomogeneous broadenings and bright single photon emission. At the same time, it is necessary to maintain scalability of such devices which is an advantage inherent to solid state emitters. Among potential candidates, colour centres in diamond were shown to be highly suitable to encounter many of the tasks set in QIP such as coherent manipulation of single spins with long coherence times [3–6], single photon nonlinearities [7], strong light–matter interactions [8] and entanglement of remote spins [9, 10]. More specifically, two colour centres raised strong interest, i.e. the well studied negatively charged nitrogen–vacancy centre (NV⁻) due to its spin coherence times approaching one second [4] and the negatively charged silicon–vacancy centre (SiV⁻) with its narrow, bright zero phonon line (ZPL) [11] and the optically addressable, doubly spin degenerate four level fine structure [12, 13]. Nevertheless, both of them suffer from severe drawbacks. In particular the NV⁻ centre is subject to strong spectral diffusion [14, 15] and low photon emission into its ZPL, while the SiV⁻ centre reaches long spin coherence times only at millikelvin temperatures, requiring the use of a helium dilution refrigerator [5, 6].

For this reason there is an ongoing search for a colour centre combining the advantages of both the NV⁻ and the SiV⁻ centre. A potential candidate for this is the negatively charged tin-vacancy centre (SnV⁻ centre) [16, 17], exhibiting a four level fine structure, similar to that of the SiV⁻ centre. Simulations predict that it shares the same molecular structure, consisting of an impurity atom in a split vacancy conformation of D₃d symmetry, with all group IV—vacancy (G4V) centres (SiV, GeV, SnV, PbV) [18]. The inherent inversion symmetry of this structure renders the G4V centres insensitive to first order stark shifts, resulting in small spectral diffusion. On the other hand, the ground state splitting of the SnV⁻ centre amounts to 850 GHz, which is sufficient to
overcome phonon mediated dephasing processes [19], that are the main source of decoherence for the SiV\(^{-}\) centre [20, 21], already at liquid helium temperatures [16]. This renders the SnV\(^{-}\) centre an interesting candidate for application in QIP.

In this work, we characterise the optical emission properties of single as well as ensembles of negatively charged SnV centres. By investigating the temperature dependence of emission spectra we furthermore prove the suitability of SnV\(^{-}\) defects as temperature sensors at nanoscale. The polarisation of the defect in emission as well as in absorption is providing evidence for an alignment of the defect in the crystallographic \(\{111\}\) direction of the diamond host matrix. At cryogenic temperatures we prove truly Fourier limited linewidths and investigate the stability of the charge state of single emitters. Additionally the temperature dependence of the Debye–Waller factor and a detailed analysis of the phonon sideband (PSB) at cryogenic temperatures reveals a dominant coupling of the centre to phonons of \(a_{1g}\) symmetry. Finally we probe an energetically higher lying excited state using photoluminescence excitation spectroscopy and thereby extend the understanding of the level structure of the SnV\(^{-}\).

2. Investigated samples

The predominantly investigated sample NI58 in this study is an (001) electronic grade bulk diamond homogeneously implanted with tin ions employing an implantation energy of 700 keV and a dose of \(2 \times 10^{13}\) ions \(\text{cm}^{-2}\). The lateral homogeneity of implantation was achieved by scanning the ion beam (diameter of 1 cm\(^{2}\)) repeatedly over the sample. Afterwards, an annealing at 2100 °C with a pressure of 7.7 GPa is performed analogous to the procedure reported in [16]. In figure 1 a fluorescence scan of the diamond sample with 532 nm excitation is shown, revealing a crescent-shaped fluorescence pattern of SnV\(^{-}\) centres. Inhomogeneous removal of the implanted diamond surface during the annealing process provides a smooth transition from very dense ensembles of SnV\(^{-}\) centres (upper right corner) to regions where single emitters can be found (Region A in figure 1). By this means, we are able to study the spectral properties of ensembles as well as single SnV\(^{-}\) centres implanted and annealed under the same conditions in one sample. An exemplary room temperature spectrum of such an emitter is also shown in figure 1 with its characteristic zero phonon line (ZPL) at 620 nm. For comparison, a second sample SC500_01 is investigated, for which tin ions were implanted in an (001) electronic grade bulk diamond with an energy of 80 keV and varying doses resulting in an mean implantation depth of 26 nm (obtained by Monte-Carlo simulations, Stopping and Range of Ions in Matter, SRIM). After implantation, the sample is annealed at 1200 °C for 4 h in high vacuum (10\(^{-6}\) mbar). Both samples were cleaned in boiling tri-acid (1:1:1 mixture of sulfuric, perchloric and nitric acid) for 2 h at 500 °C. Sample SC500_01 was additionally oxidized at 450 °C in an air atmosphere.

All samples are investigated in a home built confocal microscope with various excitation sources. Experiments at cryogenic temperatures make use of a liquid helium flow cryostat (Janis Research, ST-500LN) or a closed cycle system (attodry2100, attocube).
3. Single photon emission and fluorescence lifetime

In order to prove that our investigations are performed on single SnV− centres, we measure the intensity autocorrelation function $g^2(\tau)$ of each centre in sample N158. As it can be seen from the $g^2$ measurement in figure 2(a), we are able to clearly separate single SnV− centres. It has to be emphasized that the plotted data is raw data without any background correction applied. Saturation measurements (figure 2(b)) reveal a vanishing background emission. The value of $g^2(0) = 0.05$ can be fully explained by the dark counts of the avalanche photo diodes (APD; SPCM-AQRH-14, Excelitas) being used in the experiment and thereby proving perfect single photon emission. The saturation count rates (intensities) of the single emitters investigated in sample N158 vary between 80 and 150 kcts s−1 for saturation intensities of 200–600 μW. No discernible linear background contribution is visible. The measurements are carried out using off-resonant 532 nm excitation.

**Figure 2.** (a) Raw data of the autocorrelation measurement on a single SnV− emitter revealing a $g^2(0)$ value of 0.05. The deviation from 0 can be fully explained by the dark counts of the APDs. (b) Typical saturation measurement of a single SnV− emitter. Saturation count rates vary between 80 and 150 kcts s−1 for saturation intensities of 200–600 μW. No discernible linear background contribution is visible. The measurements are carried out using off-resonant 532 nm excitation.

From the photon correlation measurement in figure 2(a) we infer a lifetime of the upper fluorescent level of approximately 13 ns. To further explore the timescale of the fluorescence decay, we measure the radiative decay lifetime for ensembles of different densities by time correlated single photon counting in sample N158. The value of $g^2(0)$ = 0.05 can be fully explained by the dark counts of the avalanche photo diodes (APD; SPCM-AQRH-14, Excelitas) being used in the experiment and thereby proving perfect single photon emission. The count rates are one to two orders of magnitude larger than for SiV− centres in unstructured electronic grade bulk diamond [5, 6]. Furthermore, we compare the saturation count rates to the largest values for NV− centres in bulk diamond known to us [22]. Taking into account the different efficiencies in collecting photons and the Debye–Waller factors of 4% (60%) of the NV− (SnV−) centres, respectively, we find that for SnV− centres the single photon count rates into the ZPL exceed the corresponding value for NV− centres by roughly one order of magnitude, which potentially enables high repetition rates in QIP schemes.

From the photon correlation measurement in figure 2(a) we infer a lifetime of the upper fluorescent level of approximately 13 ns. To further explore the timescale of the fluorescence decay, we measure the radiative decay lifetime for ensembles of different densities by time correlated single photon counting in sample N158. The density in this sample is an indicator for the depth of the ensemble within the diamond lattice since diamond was removed inhomogeneously during sample preparation, see section 2. We define the depth of an investigated ensemble by the distance from the surface at which the implantation probability (obtained by Monte-Carlo simulations, SRIM) has dropped to approximately one percent of the distribution maximum (which lies at 168 nm depth for implantation without removal of diamond). This is illustrated in figures 3(a) and (b) for the case of 200 and 140 nm ensemble depth. The ensemble depth in the experiment is determined by first measuring the count rates in a spot in the high density region where no diamond was removed during sample preparation. This value is used to normalise the area of the Gaussian implantation distribution. For every investigated ensemble we measure the count rates and compare this to our normalised distribution, i.e. until which depth the distribution has to be integrated to obtain the measured count rates. It has to be noted that the count rates are measured at excitation powers far below saturation and therefore the change in lifetime is not affecting the value significantly. In figure 3(c) the lifetime of a large set of different ensembles is plotted against the ensemble depth below the diamond surface. Ensembles being deeply situated within the crystal lattice (>100 nm) exhibit lifetimes on the order of 5 ns as it was theoretically predicted and experimentally observed before in several studies [16–18, 23, 24]. With decreasing ensemble depth the lifetime increases due to a reduction of the photonic local density of states for large fractions of the ensemble being located in proximity to the surface as it was seen for NV centres before [25]. At an ensemble depth of about 40 nm the lifetime distribution reaches a maximum and decreases steeply when approaching zero distance to the surface, which can be explained by non radiative decay channels opening up for emitters in very close vicinity to the surface. This is in agreement with [26] having observed similar variations in lifetime of NV− centres with the dimension of the nanodiamonds they are incorporated in. For the investigated single emitters, lifetimes vary between 7 and 25 ns which is significantly
into the ground states $K_1$. With rising temperature the thermal population in the surface, which is most likely the source for the large variations in lifetime.

4.1. Temperature dependent spectroscopic investigation of $\text{SnV}^-$ centres

In order to characterise the electron–phonon interaction of $\text{SnV}^-$ centres, we first measure temperature dependent spectra of single defects in a temperature range from 10 K up to 250 K in sample NI58. As it can be seen from figure 4, at 10 K only two optical transitions C and D occur. These correspond to the decay from the lower excited state $3$ into the ground states $2$ and $1$. With rising temperature the thermal population in the upper excited state $4$ increases. The energy separation in the excited state doublet averaged over several measurements yields $3030(100)$ GHz. Thus transitions A and B only become visible at elevated temperatures as was also seen in [16]. The $\text{SnV}^-$ thus exhibits the same fine structure as the $\text{SiV}^-$ for which a $D_{3d}$ symmetric defect orientation in the diamond lattice was proven [12]. The increasing linewidth of the transitions follows a $T^3$ power law (figure 5(a)), which was theoretically suggested and experimentally confirmed for $\text{SiV}^-$ centres as well, indicating a low strain crystal environment [19, 27]. The lineshift is very well described by a $T^2 + T^4$ power law with a vanishing $T^5$ contribution. The $T^3$ power law for a ZPL shift according to linear e-symmetric phonon coupling which was derived by Jahnke et al [19] is not reproducing the data well (see figure 5(b)). However, the accurate description of the linewidth broadening with temperature makes the $\text{SnV}^-$ centre a suitable candidate for low temperature sensing on the nanoscale. Its large ground state splitting of $830(30)$ GHz (lowest value for ensemble as well as for single emitter measurements) is advantageous in comparison to the $\text{SiV}^-$ centre due to the fact that the four transitions merge at comparatively higher temperatures making the fits more reliable at temperatures below 75 K and thereby covering a large temperature range with high resolution.

The same measurements are repeated for ensembles of $\text{SnV}^-$ centres in sample NI58. To this end, a region on the sample is chosen where the linewidth at a temperature of 5 K is below the resolution limit of the spectrometer of 10 GHz. Even though the resolution limited linewidth could possibly hint at the investigated number of $\text{SnV}^-$ centres being small, we are not able to see any saturation effects of the ensemble with 10 mW of green laser light where count rates exceed 20 Mcts s$^{-1}$. For comparison, typical saturation powers for the single emitters investigated in the previous section are on the order of 200 µW, exhibiting average saturation count rates of 120 kcts s$^{-1}$. The $\text{SnV}^-$ ensemble spectrum at cryogenic temperatures consists of the same characteristic fine structure as the single $\text{SnV}^-$ centres. As it is shown in figure 6, the broadening of the linewidths of peaks C and D is very well reproduced by a pure $T^3$ law as in the single emitter case. Since adding a $T^6$ or $T^7$ term leads to longer than what was observed before [16, 17, 23, 24]. In sample NI58 all single emitters are in close proximity to the surface due to the sample fabrication. Removal of diamond during annealing procedure introduces damage on the surface of the diamond and therefore leaves single emitters in a damaged environment close to the surface, which is most likely the source for the large variations in lifetime.

4. Zero phonon line emission

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no improvement of the fit, we conclude that the ensemble is situated in a very little strained crystal environment [19]. Furthermore, also the shift of the optical transitions C and D for the ensemble is shown in figure 6. Both of them obey a $T^2 + T^4$ law, again agreeing with the data for the single SnV$^-$ centre. As for the single centre, the very accurately described dependence of linewidth and line position on temperature renders ensembles of SnV$^-$ centers excellent candidates for thermometry on nanoscale. This was proposed by Alkahtani et al but not investigated below 120 K [28]. Furthermore, our investigations hint at very low strain in still dense ensembles of SnV$^-$ centres which makes them very promising for application in QIP where a high optical density is required while maintaining narrow emission linewidths.

As conclusion of the evaluation of the temperature dependent ZPL emission we find a dominant $T^4$ contribution for the ZPL line shift of a single SnV$^-$ centre. An additional $T^2$ term has to be added to accurately describe the ensemble measurements at temperatures below 100 K, which might be associated with inhomogeneous broadening effects. According to Hizhnyakov et al [29–31] a $T^4$ law is expected for quadratic electron–phonon coupling of $a_{1g}$ phonons and was also experimentally confirmed for SiV$^+$ centres [27]. In section 5 we present further evidence for the electron–phonon coupling of the optical transition being dominated by $a_{1g}$ phonons.

The broadening of the ZPL line is caused by single phonon absorption and Raman scattering between sublevels of ground and excited states, respectively, for which a $T + T^3$ law has been derived [19]. The sublevels split due to spin–orbit interaction and the corresponding doubly degenerate orbitals are also subject to Jahn–Teller distortion in which the $e_g$ phonons will only couple these degenerate orbitals both in the electronic ground and excited state [18]. Thus, $a_{1g}$ phonons do not contribute to the ZPL broadening and there only remains the

Figure 4. Spectra of a single emitter under off-resonant 532 nm excitation at temperatures in between 10 K and 250 K. At 10 K only the C- and D-Transitions are visible, at temperatures around 75 K also A and B occur due to increasing thermal population in the upper excited state. With rising temperature, a line shift as well as line broadening is observable.

Figure 5. Lineshift and linewidth for increasing temperature are depicted for the C-transition of a single emitter. Dots are experimentally obtained values, the solid lines are fits regarding to the power laws mentioned in the text. The measurements are carried out using off-resonant 532 nm excitation.
effect of phonons. This is in accordance with our observations where the linear contribution $\mu_T$ would only be visible for high resolution measurements at very low temperatures.

4.2. Comparison of photoluminescence spectra for different annealing temperatures

The effect of the annealing temperature after ion implantation on the room temperature linewidth is addressed in [16] showing a monotonous decrease of room temperature ZPL linewidth with annealing temperature. We here find that the annealing temperature furthermore very strongly influences the inhomogeneous distribution of ZPL lines. In figure 7, a comparison between the two investigated samples is shown for ensembles of comparable density of SnV$^-$ centres. As obvious from figure 7, the annealing procedure at $\approx 2100$ °C greatly reduces the inhomogeneous distribution induced by the lattice damage that is caused during implantation of the heavy tin atoms. The linewidth of the ZPL transitions of the whole ensemble in sample NI58 (figure 7(a)) is 15 GHz, which is similar for the individual emission lines (10–50 GHz) occurring in sample SC500_01. The line positions of the C and D transitions in sample NI58 are within a 150 GHz error margin the same for single emitters and ensemble, while they are spread over a large range (>10 nm) in sample SC500_01. We want to emphasize that contrary to the SiV$^-$ centre, where annealing temperatures of $\approx 1200$ °C are sufficient to produce low strain ensembles [27], for the SnV$^-$ centre higher annealing temperatures are required during sample preparation in order to reduce inhomogeneous broadening. The influence of annealing temperature on strain is also visible for room temperature emission. In figure 8 typical room temperature spectra of single centres are compared for both samples. The ZPL features considerably narrower linewidth (2–3 nm) for the low-pressure-low-temperature (LPLT) annealed sample (figure 8(a)) than for the HPHT annealed NI58 (5–6.5 nm, figure 8(b)). Furthermore, for the HPHT annealed sample we find a very homogeneous distribution of ZPL centre wavelengths around 619.6 nm whereas the LPLT annealed sample shows a broad distribution of ZPL centre wavelengths (610–630 nm). In the latter sample we also find reduced emission into the phonon sideband. A reason for this could be a different phonon coupling for different types and strengths of strain in the two samples, resulting in differing phonon broadening. Similar room temperature distributions of ZPL centre
wavelengths and linewidths were reported for SiV\textsuperscript{−} centres in nanodiamonds, were it is common that large strain is exerted on the colour centres [11, 32].

4.3. Polarisation of single SnV\textsuperscript{−} centres
An interesting figure of merit is the polarisation of single SnV\textsuperscript{−} centres, which gives insight into the orientation of the dipole within the diamond lattice with respect to the crystal axes. The polarisation of the emitters in sample NI58 is measured in excitation as well as in emission. This is done by employing a linear polariser followed by a half-wave, a quarter-wave plate and finally a dichroic mirror in the excitation pathway. The quarter-wave plate remains at a fixed position throughout the measurements and is used to compensate the phase shift induced by the dichroic mirror. Rotating the half-wave plate and at the same time recording the emitted count rates, reveals the excitation dipole pattern of the single SnV\textsuperscript{−} centres. In order to also characterise the polarisation of the emitted photons, a reversed configuration is applied, where the dichroic mirror is followed by a quarter-wave, a half-wave plate, and, finally, a linear polariser. Both absorption and emission dipoles are shown for one exemplary SnV\textsuperscript{−} centre in figure 9 where the count rates are only corrected for the dark counts of the APD. As opposed to the results of [17] it can be seen that the projections of the dipoles into our measurement plane (diamond (001) plane) perfectly overlap and the orientation is at an angle of 45° with respect to the surface edges of the sample cut along the [100] and [010] direction. This points towards an alignment of the high symmetry axis of the SnV\textsuperscript{−} in (111) direction which is in excellent agreement with former findings for the SiV\textsuperscript{−} centre [12, 33]. All measured SnV\textsuperscript{−} centres exhibit the same or a perpendicular emission pattern with overlapping absorption and emission dipoles. Visibilities of up to 85% in emission and 87% in excitation are measured for the investigated set of SnV\textsuperscript{−} centres. The deviation from full contrast is most probably caused by a contribution of weaker x and y dipoles [12]. This is further emphasised by the results of [23] where they find the C-transition being slightly and the D-transition strongly elliptically polarised.
4.4. Photoluminescence excitation spectroscopy and stability of tin-vacancy centres under resonant excitation

The spectral quality of the SnV\(^-\) emission is further determined by the optical coherence of the ZPL at low temperatures. To explore this, we perform photoluminescence excitation (PLE) spectroscopy on the C-transition of single emitters in sample NI58 using excitation with a narrowband dye laser (Matisse DS, SIRAH) and detecting fluorescence from the excited state into the PSB (filtered in a window from 655 nm to 665 nm). As it can be seen in figure 10, we are able to find purely Fourier-limited linewidths for single SnV\(^-\) centres similar to the findings in [24] and significantly narrower than in [23]. For 12 evaluated emission lines we measure a mean linewidth of 31 MHz, slightly above the Fourier limit due to power broadening. For the emitter in figure 10, the fluorescence lifetime is determined using time correlated single photon counting to be 7.61(3) ns yielding a Fourier limited linewidth of 20.9(1) MHz, being in excellent agreement with the measured value of 18(3) MHz. The excitation power for the measurement in figure 10(b) is 200 pW. We find that even slightly larger excitation powers below 1 nW cause termination of the emission, presumably by ionisation from the excited state. Such a termination is exemplarily shown in figure 11(a) where a steep drop of fluorescence is visible when approaching the line centre. The charge can be brought back by applying an off-resonant green laser (532 nm, 40 \(\mu\)W). This is achieved experimentally by coupling the green and resonant lasers into the same fibre to achieve perfect spatial overlap in our confocal microscope and thereby address the same emitter with both colours. We explain this phenomenon with a two-photon ionisation process of SnV\(^-\) to SnV\(^0\) in which the first photon excites the SnV\(^-\) centre resonantly, while the second photon transfers the electron from the excited state to the conduction band. This is in agreement with the ionisation occurring only when approaching the resonance maximum (see figure 11(a)) and the calculated \((-|0\rangle\) ionisation energy of 3.2 eV in [18] which is
exceeded by two 620 nm (2.0 eV) photons. Application of the 532 nm (2.33 eV) laser leads to a conversion of SnV0 back to SnV− in a single photon process matching again the (0 − −) ionisation energy of 2.3 eV derived in [18]. However, the transition line is shifted in this process. Most probably this shift is related to the second order sensitivity of the SnV− centre to an altered charge environment. Constantly adding a green laser while scanning the resonance supports this assumption, as the transition is broadened by spectral jumps to 580(10) MHz as the green laser is creating a dynamic equilibrium of moving charges (see figure 11(b)). Similar effects of spectral diffusion were observed for the SiV− centre in [34].

Since the charge instability could be caused by the emitters being located very close to the surface of the diamond sample, we further investigate the second sample SC500_01. The emitters in this sample are situated more than 20 nm deep in the diamond lattice but still show the same instabilities under resonant excitation and spectral diffusion occurs in the same manner. We therefore conclude that the negative charge state of the SnV centre is not stable under resonant excitation in an electronic grade diamond host matrix and manipulation of the Fermi level, e.g. by co-implantation of electron donors might be required in order to overcome this problem.

5. Analysis of the phonon sideband and Debye–Waller factor

Being a solid state emitter embedded in a diamond host matrix renders the SnV− centre sensitive to coupling to phononic modes. Since these electron–phonon interactions are of great importance for the spectral purity of the defect emission, a thorough evaluation of the phononic sideband of single SnV− centres is necessary. In figure 12(a) the sideband spectrum of a single SnV− in sample NI58 at a temperature of 30 K is depicted. There are six observable peaks shifted by 46 meV, 76 meV, 109 meV, 122 meV, 148 meV, 181 meV with respect to the ZPL position. The structure of these peaks is in excellent agreement with the predicted phonon sideband by Thiering and Gali [18], when only taking into account eg symmetric phonons coupling to the centre (solid blue line in figure 12(a)). If comparing merely the position of the peaks, the spectrum for the combined coupling of ae and eg phonons (solid green line in figure 12(a)) also fits, but the peak height is less well predicted. The simulations are normalised to the maximum of the dominant sideband peak at 660 nm. Since also the Huang–Rhys factor is calculated for both phonon coupling schemes, the next section is dedicated to a thorough evaluation thereof. It has to be pointed out further that no quasi-local mode caused by the eigenmotion of the tin atom, which should occur at a frequency shift of 29 meV from the ZPL (see [17]), can be observed experimentally. This is a further hint at the high quality of the sample since the local oscillator transition corresponding to a coupling to eg phonons should be symmetry forbidden in an unstrained crystal environment [35] which in the investigated sample was created by the high temperature annealing. The local mode was observed for SiV− and GeV− centres [36, 37] in material with residual strain.

The Debye–Waller factor is an important number when evaluating colour centres with respect to their suitability in QIP, as for most of the protocols only photons emitted via the ZPL transition can be used. Therefore, the success probability of such QIP experiments depends directly on the number of photons emitted without phonon interactions. For this reason, we here evaluate the Debye–Waller factor within a temperature range from 5 K up to 300 K in sample NI58 by fitting the obtained spectra in the range from 610 to 740 nm with Voigt profiles in order to account for the Gaussian spectrometer response function. Then the area of the ZPL fit is divided by the total area of the fit to the whole spectrum yielding the Debye–Waller factor. The result is shown in figure 12(b).
in figure 12(b). The solid line is a fit to the data according to the equation $\text{DW}(T) = \exp(-S(1 + \frac{2\pi^2 T^2}{3 T_{\text{cutoff}}^2}))$ derived in [38] with $S$ being the Huang–Rhys factor and $T_{\text{cutoff}}$ the cutoff temperature for the phonons actually coupling to the SnV$^-$ centre. This cutoff temperature is common to the Debye temperature, as found in [39]. However, it is not taking into account all phonons present in the diamond lattice but only those involved in the centre-phonon interaction. The fit yields a Huang–Rhys factor of 0.57(2) corresponding to a Debye–Waller factor of $\text{DW}(0 \text{ K}) = 0.57(1)$ being in close agreement with the theoretically proposed $\text{DW}_{eg}(0 \text{ K}) = 0.63$ for only $a_{\text{tg}}$ phonons coupling to the centre [18]. The Debye–Waller value for the combined coupling of $a_{\text{tg}}$ and $e_g$ symmetric phonons yields $\text{DW}_{a_{\text{tg}}+e_g}(0 \text{ K}) = 0.41$, again agreeing to a lesser extend with our experiment. The cutoff temperature finally amounts to 680(40) K, which corresponds to an effective phonon frequency of 59(4) meV. The frequency of the acoustic $a_{\text{tg}}$ phonons considered in [18] was only estimated in a range between 60 and 100 meV which includes 59(4) meV, while the $e_g$ phonon frequency was calculated to 75.6 meV, far off from our measured value and therefore again pointing towards a predominant impact of $a_{\text{tg}}$ phonons to the centre-phonon interaction. With this third indicator we can conclude that the lattice coupling between the SnV$^-$ centre and its host matrix is dominated by phonons of $a_{\text{tg}}$ symmetry.

### 6. Excited state spectroscopy

The evidence in literature [16, 17, 23, 24] and in the present work strongly indicate a molecular model of the SnV$^-$ centre as inversion symmetric D$_{3d}$ point group. It thus should feature the same electronic structure as the well investigated SiV$^-$ and GeV$^-$ centres [18]. The existence of $E_g$ and $E_u$ ground and excited states are confirmed by PL experiments (see figure 4) and Zeeman splittings in external B-fields [23, 24]. In order to further explore the SnV$^-$ electronic level structure, we here perform PLE measurements on an ensemble in sample N158. The excitation radiation is provided by a white light laser (SuperK Extreme EXW-12, NKT photonics), which we scan from 480 nm up to 595 nm while keeping its bandwidth fixed to 5 nm and maintaining constant power. At room temperature, a spectrum is taken for every excitation wavelength and the counts constituting the ZPL of the SnV$^-$ are evaluated by fitting the spectra with Voigt profiles. Figure 13(a) shows a broad Lorentzian shaped resonance centered around 2.348(4) eV with a width of 190(12) meV. We attribute this resonance to a higher lying excited state with $A_{2u}$ symmetry resulting from a broad $a_{\text{tg}}$ band. This band is created by an $a_{\text{tg}}$ defect level situated in the valence band and therefore being broadened due to strong mixing with the diamond band. Similar results were found for the SiV$^-$ centre in diamond [40, 41]. The width of the $A_{2u}$ resonance (190 meV) being narrower than the corresponding state of the SiV$^-$ centre (400 meV) indicates a position of the $a_{\text{tg}}$ orbital closer to the valence band edge in agreement to the predictions of [18]. At 2.12 eV another peak is observable which corresponds to the most prominent sideband peak of the SnV$^-$ at 122 meV. This is indicated in figure 13(a) by overlapping the PLE measurement with the mirror image of the SnV$^-$ PL spectrum with respect to the ZPL position. The peak at 2.12 eV is a result of addressing the corresponding higher lying vibrational ladder of the excited state and, furthermore, stating that the ground and excited states vibronic structure are of similar nature.
The measurement is repeated at 2 K, detecting the SnV\(^-\) ZPL by filtering with a 620 ± 10 nm bandpass filter. As shown in figure 13 (b), the absorption peak remains centered at 2.349 ± 3 eV but with a decreased width of 120 ± 8 meV. This narrowing is due to a reduced homogeneous phonon broadening at low temperatures. The second peak is again coinciding with the most prominent sideband peak of the SnV\(^-\).

7. Summary and Conclusion

In this work, we have investigated the spectroscopic properties of the SnV\(^-\) centre. We found that the main dipole of the SnV\(^-\) centre is aligned along the \(\{111\}\) direction of the diamond lattice, thereby indicating the high symmetry axis of the defect. The electron–phonon scattering characteristics of the centre were investigated by measurement of ZPL line shifts and broadenings as function of temperature, by comparing the simulated (see [18]) and the measured PSB as well as by the temperature dependence of the Debye–Waller factor of a single SnV\(^-\) centre. From the results we were able to show that the electron–phonon interaction is dominated by phonons of \(a_{1g}\) symmetry and that Debye–Waller factors can reach up to 60% at 10 K. This value for the Debye–Waller factor is slightly smaller as in the case of the SiV\(^-\) centre, but still more than a magnitude larger than for the NV\(^-\) centre. By performing PLE spectroscopy over a large wavelength range, we revealed a higher lying excited state at 2.35 eV resulting from an \(a_{1g}\) defect level lying in the valence band and strongly coupling to the diamond modes. In the same measurement we found that the vibrational ladder of the ground state strongly resembles the one of the excited state indicating similar electronic wave functions. Finally, single centres with truly Fourier limited linewidths were found with their usefulness being limited by a dominant ionisation process resulting in termination of the fluorescence when approaching the resonance maximum of the fine structure transitions and non negligible spectral diffusion. This can be overcome in future experiments by carefully controlling the Fermi level of the surrounding diamond host, for example by co-implantation of electron donors. Stabilising the negative charge state of the SnV\(^-\) centre would further allow for measuring the ground state spin coherence time and comparing it to the theoretical expectation.

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References

[1] Atatüre M, Englund D, Vamivakas N, Lee S Y and Wrachtrup J 2018 Nat. Rev. Mater. 3 38–51
[2] Awaschalom D D, Hanson R, Wrachtrup J and Zhou B B 2018 Nat. Photon. 12 516–27
[3] Bar-Gill N, Pham I M, Jarmola A, Budker D and Walsworth R L 2013 Nat. Commun. 4 1743
[4] Aboeleish M H, Craemer J, Bakker M A, Kalb-N, Markham M, Twitchen D J and Tamminiai T H 2018 Nat. Commun. 9 2552
[5] Becker J N, Pingault B, Groß D, Günögean M, Kukharchyk N, Markham M, Edmonds A, Atatüre M, Bushev P and Becher C 2018 Phys. Rev. Lett. 120 053603
[6] Sukachev D D, Sipahigil A, Nguyen C T, Bhaskar M K, Evans R E, Jelezko F and Lukin M D 2017 Phys. Rev. Lett. 119 223602
[7] Sukachev A et al 2016 Science 354 847–50
[8] Weinzel C, Görlitz J, Becker J N, Walsmley I A, Poem E, Nunn J and Becher C 2019 Phys. Rev. Lett. 122 063601
[9] Sipahigil A, Goldman M L, Togan E, Chu Y, Markham M, Twitchen D J, Zibrov A S, Kubanek A and Lukin M D 2012 Phys. Rev. Lett. 108 143601
[10] Hensen B et al 2015 Nature 526 68 EP
[11] Neu E, Steinmetz D, Riedrich-Möller J, Gsell S, Fischer M, Schreck M and Becher C 2011 New J. Phys. 13 025012
[12] Hepp C et al 2014 Phys. Rev. Lett. 112 036405
[13] Becker J N, Görlitz J, Arend C, Markham M and Becher C 2016 Nat. Commun. 7 13512
[14] Fu K M C, Santori C, Barclay P E and Beausoleil R G 2010 Appl. Phys. Lett. 96 121907
[15] Wolters J, Sazdak N, Schell A W, Schröder T and Benson O 2013 Phys. Rev. Lett. 110 027401
[16] Iwasaki T, Miyamoto Y, Taniguchi T, Sinyaee P, Metsch M H, Jelezko F and Hatano M 2017 Phys. Rev. Lett. 119 253601
[17] Tchernij S D et al 2017 ACS Photonics 4 2580–6
[18] Thiering G and Gali A 2018 Phys. Rev. X 8 021063
[19] Jahnke K D, Sipahigil A, Binder J M, Doherty M W, Metsch M, Rogers L J, Manson N B, Lukin M D and Jelezko F 2015 New J. Phys. 17 043011
[20] Pingault B, Becker J N, Schulte C H H, Arend C, Hepp C, Godde T, Tartakovskii A I, Markham M, Becher C and Atatüre M 2014 Phys. Rev. Lett. 113 263601
[21] Rogers L J et al 2014 Phys. Rev. Lett. 113 263602
[22] Lesik M, Tetienne J P, Tallaire A, Achard J, Mille V, Gicquel A, Roch J F and Jacques V 2014 Appl. Phys. Lett. 104 113107
[23] Rugar A E, Dory C, Sun S and Vučkovic J 2019 Phys. Rev. B 99 205417
[24] Trusheim M E et al 2018 arXiv:1811.07777
[25] Mohtashami A and Koenderink A F 2013 New J. Phys. 15 043017
[26] Inam F A, Edmonds A M, Steel M J and Castellolet S 2013 Appl. Phys. Lett. 102 253109
[27] Arend C, Becker J N, Sternschulte H, Steinmüller-Nethl D and Becher C 2016 Phys. Rev. B 94 045203
[28] Alkatahmi M, Cojocaru I, Liu X, Herzig T, Meijer J, Küpper J, Lühmann T, Akimov A V and Hemmer P R 2018 Appl. Phys. Lett. 112 241902
[29] Hizhnyakov V, Kaasik H and Sildos I 2002 Phys. Status Solidi b 234 644–53
[30] Hizhnyakov V, Boltrushko V, Kaasik H and Sildos I 2003 J. Chem. Phys. 119 6290–5
[31] Hizhnyakov V, Boltrushko V, Kaasik H and Sildos I 2004 J. Lumin. 107 351–8
[32] Linder S, Sommer A, Muzha A, Krueger A, Gines L, Mandle S, Williams O, Londero E, Gali A and Becher C 2018 New J. Phys. 20 115002
[33] Neu E, Fischer M, Gsell S, Schreck M and Becher C 2011 Phys. Rev. B 84 205211
[34] Nguyen C T et al 2019 arXiv:1907.13200v2
[35] Londero E, Thiering G, Razinkovas L, Gali A and Alkauskas A 2018 Phys. Rev. B 98 035306
[36] Dietrich A, Jahnke K D, Binder J M, Terai R, Isoya J, Rogers L J and Jelezko F 2014 New J. Phys. 16 113019
[37] Ekimov E A, Lyapin S G, Boldyrev K N, Kondrin M V, Khmelnitskiy R, Gavva V A, Kotereva T V and Popova M N 2015 JETP Lett. 102 791–6
[38] Brand J, Weinzierl G and Friedrich J 1981 J. Chem. Phys. Lett. 84 197–200
[39] Plakhnik T, Doherty M W, Cole H J, Chapman R and Manson N B 2014 Nano Lett. 14 4989–96
[40] Gali A and Maze J R 2013 Phys. Rev. B 88 235205
[41] Häußler S et al 2017 New J. Phys. 19 063036