Microwave-based quantum control and coherence protection of tin-vacancy spin qubits in a strain-tuned diamond membrane heterostructure

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Robust spin-photon interfaces in solids are essential components in quantum networking and sensing technologies. Ideally, these interfaces combine a long-lived spin memory, coherent optical transitions, fast and high-fidelity spin manipulation, and straightforward device integration and scaling. The tin-vacancy center (SnV) in diamond is a promising spin-photon interface with desirable optical and spin properties at 1.7 K. However, the SnV spin lacks efficient microwave control and its spin coherence degrades with higher temperature. In this work, we introduce a new platform that overcomes these challenges – SnV centers in uniformly strained thin diamond membranes. The controlled generation of crystal strain introduces orbital mixing that allows microwave control of the spin state with 99.36(9)% gate fidelity and spin coherence protection beyond a millisecond. Moreover, the presence of crystal strain suppresses temperature dependent dephasing processes, leading to a considerable improvement of the coherence time up to 223(10) μs at 4 K, a widely accessible temperature in common cryogenic systems. Critically, the coherence of optical transitions is unaffected by the elevated temperature, exhibiting nearly lifetime-limited optical linewidths. Combined with the compatibility of diamond membranes with device integration, the demonstrated platform is an ideal spin-photon interface for future quantum technologies.

INTRODUCTION

Color centers in diamond are a leading platform in quantum technologies, key achievements such as the demonstration of a quantum register [1–3], distant entanglement generation between three nodes [4], quantum teleportation [5], along with myriad landmarks in quantum sensing [6, 7]. In recent years, group IV centers have gained much attention due to their excellent optical properties [8–15]. Their $D_{3d}$ symmetry renders optical transitions insensitive to first-order charge noise [16–18]. Additionally, a favorable Debye Waller factor leads to the majority of photons being emitted into the zero-phonon line, critical for spin-photon entanglement [19]. However, the electronic structure of group IV centers – a spin 1/2 system with two ground state orbital branches – renders the electron spin susceptible to phonon-driven transitions between the two branches [20]. This temperature-dependent spin dephasing can be mitigated by operating at millikelvin temperatures [21, 22] or by engineering the local phonon density of states through nanostructuring [23, 24]. Alternatively, dephasing can be mitigated by qubit engineering such as working with group IV centers with high spin-orbit coupling and thus large orbital splitting [25], or by leveraging spin-strain interaction in randomly- or controllably strained group IV centers[3, 24]. With a spin-orbit coupling significantly higher than those of the silicon vacancy (SiV) and the germanium vacancy (GeV) centers, the SnV center has the highest reported spin coherence time at 1.7 K [26]. However, efficient microwave (MW) control of group IV spins requires the magnitude of spin-strain interaction to be comparable with the spin-orbit interaction, which for SnV necessitates strain approaching 0.1%. This degree of strain is challenging to achieve in microelectrical mechanical structures (MEMS) such as diamond cantilevers, with reported values on the order of 0.015% [23]. Therefore, a controlled process to generate ≈0.1% strain in diamond is desired to improve SnV qubit performance by both increasing the operational temperature and enabling efficient MW driving.

In this work, we utilize heterogeneous integration of diamond membranes to generate strain-tuned SnVs. By bonding SnV incorporated pristine diamond membranes to a glass substrate, we leverage the heterogeneous thermal expansion coefficients of the two materials to generate a uniform, in-plane strain in the diamond to the order of 0.1%. This strain greatly increases the energy splitting between the two orbital levels of the SnV and induces orbital mixing in the spin ground state. We demonstrate MW manipulation of the spin with 99.36(9)% Rabi fidelity at 4.50(2) MHz for 24 dBm MW input power. At 1.7 K, the implementation of dynamical decoupling allows the SnV to reach millisecond
coherence time, which is largely preserved even at 4 K, owing to the strain-induced increased ground state orbital splitting. In combination with near lifetime-limited optical linewidths up to 7 K, our spin-photon interface is compatible with broadly utilized low-infrastructure and cost-effective portable cryogenic systems. Additionally, the demonstrated strained-membrane heterostructure maintains robustness and flexibility for additional photonic, electronic, and micro-electromechanical systems (MEMS) integration. Our SnV-based diamond membrane platform greatly reduces the technological barrier for establishing quantum nodes for networking.

SnVs in strained diamond

This work relies on strain engineering to improve SnV qubit performance. First, we demonstrate that heterogeneous thermal expansion disparities between diamond and glass in a diamond-membrane heterostructure are sufficient to generate uniform strain of the magnitude necessary to beneficially impact SnV. The diamond membranes used in this work were generated via the “smart-cut” method combined with isotopically purified ($^{12}$C) overgrowth. The membrane thickness is nominally 150 nm, with pristine crystal quality and atomically smooth surfaces [27]. To introduce a positive tensile strain inside the diamond membrane, we bond them onto 500 μm-thick fused silica substrates—a material with a low thermal expansion coefficient ($< 1 \times 10^{-6}\text{K}^{-1}$) — using a layer of hydrogen silsesquioxane (HSQ). The schematic of this strain generation method is shown in Figure 1 (a). The device is then annealed at 600°C, beyond the temperature at which the HSQ solidifies to glass, bonding the heterostructure in a ”zero-strain” condition [28]. Due to the mismatch in thermal contraction between diamond and fused silica and the negligible thickness of the diamond membrane compared to that of the fused silica substrate, cooling down the device to cryogenic temperature regime generates a positive (tensile), static strain profile in the diamond membrane with an estimated magnitude of 0.05% to 0.1% (see section 1.3 and 1.4 in SI for details). This passive, uniform, and membrane-compatible strain generation is complimentary to recent demonstrations of electromechanically-induced strain on suspended diamond beams [24, 29].

Figure 1 (b) is the microscope image showing the layout of our diamond-membrane heterostructure device. Prior to the membrane bonding, we patterned and etched a 5 μm deep trench on the fused silica to suspend part of the membrane and mitigate background fluorescence from the HSQ resist. To study MW control of the SnV centers, we patterned and deposited gold coplanar waveguides following membrane bonding.

The strain monotonically increases the orbital splitting of the SnV centers in the membranes, which can be
directly verified in the photoluminescence (PL) spectra at 1.7 K. The energy level diagram of the strained SnV is shown in Figure 1 (c), highlighting the ground state orbital splitting ($\Delta g_s$) and the respective contributions of spin-orbit coupling, strain, and magnetic Zeeman interaction in purple, blue, and green boxes. Figure 1 (d) compares the spectra of a strained (unstrained) SnV center in a diamond membrane (bulk diamond) with $\Delta g_s \approx 1300(850)$ GHz. This particular strained center is used in further optical, microwave, and spin characterizations in this work. Remarkably, we note that all color centers in the membrane are comparably strained. As shown in Figure 1 (e), we observed a distribution of the orbital branches splitting centered around 1500 GHz across different devices with a minimum (maximum) value of 1200(1800) GHz. We carried out density functional theory (DFT) calculations to compute strain-susceptibilities and characterize the SnV spin-strain interaction (see SI); our results show that the increase of the splitting between orbital branches from 850 GHz to $\approx 1500$ GHz due to strain, corresponds to a diamond membrane strain magnitude of 0.075 % (see section 1.2 in the SI for details). The consistent strain generation, in combination with our ability to perform additional integration and nanofabrication following membrane bonding [30, 31], highlights the robustness and versatility of our platform.

**Optical properties of SnV under strain**

To investigate the potential of strained SnV as a spin-photon interface, we first verify that the symmetry of the defect is preserved even under considerable strain by characterizing the optical transitions as a function of the magnetic ($B$) field orientation. Using the (111) crystallographic axis—the high symmetry axis of the SnV as the reference, we rotate the $B$ field in both polar ($\theta$) and azimuthal ($\phi$) angles at the same magnitude (0.2 T). The absolute energy splitting between the two spin-conserving transitions (A1-B2) with respect to $\theta$ and $\phi$ is shown in Figure 2 (a), indicating that large splittings at moderate values of magnetic field are achievable which is ideal for later SnV spin initialization and control. Similarly to the unstrained case, we observe a $\phi$ rotational symmetry of the splitting with respect to (111), which corresponds to the intrinsic spin quantization axis. We further verify that the polarization of the SnV transitions (i.e. dipole operator matrix elements) remain along the (111) direction (see section 3.1 of the SI), as in the unstrained case [18].

From the $B$-field scan of the strained SnV, we note that besides the normal A1-B2 splitting maximum along the quantization axis, an additional local maximum at $\theta = 90^\circ$ – the equator plane perpendicular to the quantization axis – is observed, with the relative A1-B2 position being inverted, as verified by coherent population trapping measurements (see SI). This differs from the unstrained case. The novel feature arises from the moderate crystal strain (comparable in magnitude to the spin-orbit coupling) which increases the difference in effective Zeeman shift between ground and excited states, mostly visible for a magnetic field orthogonal to the spin-orbit-dictated quantization axis. As is the case for moderately strained SiV centers [22] for MW-based control, we roughly align the $B$-field towards the quantization axis to achieve highly cycling optical transitions with cyclicity reaching $\eta \approx 2500$ (see section 4.2 of SI). We note that $\eta$ can be as low as 6 when the $B$ field is perpendicular to the quantization axis, which is ideal for Raman-based all-optical control of strained SnV (see section 4.3 of SI). Moreover, by comparing the dependence on $\theta$ of the A1-B2 splitting with calculated results, we are able to determine the Stevens reduction factor $g_L$ for ground and excited states mentioned in [32]. This model is then used to explain the optically detected magnetic resonance (ODMR) frequency of the strained SnV discussed below.

Additionally, our measurements reveal near-transform limited optical linewidths, thereby showing that the application of strain does not alter the excellent coherence properties of the optical transitions, as previously demonstrated with unstrained centers [11, 25]. As shown in Figure 2 (b), the 20 s average scan returns a mean linewidth of 47.4(16) MHz, only 40 % more than the lifetime-limited value of 32.26(19) MHz (4.933(190) ns) optical lifetime, see section 3.2 of SI). The long term frequency stability of the {A1, B2} transitions returns a center frequency stan-
standard deviation of $\sigma_c = 23.8(1)$ MHz and a A1-B2 splitting standard deviation of $\sigma_s = 13.28(6)$ MHz (see section 3.4 of SI). This linewidth and peak stability is comparable to that of other measurements of group IV color centers in nanostructures [3, 13, 33] and thus confirms the excellent potential of these defects for quantum photonic applications.

The resolvable splitting and narrow optical transitions are crucial for the spin initialization and readout of the SnV qubit. The spin initialization curve with subtracted background is shown in Figure 2 (c), indicating a fitted exponential decay constant of 24.2(3) $\mu$s. The initialization pulse duration was set to 200 $\mu$s allowing us to reach a fidelity of 98.8%. We note that with a cyclicity of over 2500, this platform is a prime candidate for single shot readout if the signal counts can be improved via on-chip structures (nanophotonics, fiber couplers or grating couplers, solid immersion lenses) [33–38] or external methods (microcavities) [39–41].

Efficient MW control of the SnV spin

A critical component of a spin-photon interface is high-fidelity spin control, commonly achieved through MW driving of the electron spin. In the case of group IV centers, a MW field can only drive the spin transition in the presence of strain [23, 42]. This arises due to the orthogonality of orbital states associated with the electron spin qubit of group IV centers [18]. Strain that is comparable in strength to spin-orbit coupling relaxes this orthogonality, enabling microwave control. SnV, with larger spin-orbit coupling (850 GHz) and smaller strain susceptibility than SiV and GeV, requires large crystal strain to meet this criteria. This strain requirement goes beyond the achievable magnitude demonstrated via active strain tuning [23] or implantation-induced strain [3].

To demonstrate efficient MW control, we utilize the nominal 0.1% crystal strain in the diamond membrane. We estimate an effective Landé factor $g$ of 1.62 for the transverse microwave field with the external magnetic field roughly aligned to the SnV quantization axis (see section 2.1 in SI). This value is relatively high compared with spin-orbit-dominated regime for unstrained centers ($\leq 0.3$) and is close to the free electron value ($g = 2$). In addition, we tapered the MW waveguide around the measurement area by shrinking its width to 6 $\mu$m to enhance the microwave amplitude, as shown in Figure 1 (b). The distance between the target SnV and the waveguide is $\approx 4$ $\mu$m, ensuring an efficient exposure to the MW driving field (see section 2.1 - 2.3 in SI for details).

We begin the MW control characterization by initializing the spin via optical pumping and scan the frequency of a MW field across the expected spin resonance while monitoring the fluorescence intensity of the spin readout at 1.7 K. In Figure 3 (a) we observe clear signature of optically detected magnetic resonance (ODMR) for the target SnV center. The 81.5 $mT$ external magnetic field is aligned to the quantization axis by polarisation measurements and 3D field scan. The ODMR shows a profile

![Figure 3](image_url)
with two overlapping peaks separated by 628(182) kHz, indicating an interaction between the electronic spin of the SnV with another system in the vicinity, likely a $^{13}$C nuclear spin or the electron spin of a P1 center. Further investigation is needed to understand the nature of this interaction. By driving both power-broadened ODMR transitions, we are able to resonantly manipulate the spin state of the SnV with a Rabi frequency of 4.50(2) MHz. The Rabi oscillation curve and the chevrons (Rabi oscillations with varied driving frequency) are shown in Figure 3 (b) and (c). We observe a long-time averaged Rabi $\pi$-gate fidelity of 99.36(9) %, improving significantly from previously demonstrated optical Raman-based spin control value [26]. We note that the MW power delivered to the device is approximately 24 dBm (250 mW) which is comparable to previous demonstrations on strained SiV [3]. We also characterized the power dependence of the Rabi rate. Starting from a linear dependence, the Rabi rate deviates to sub-linear when the power surpasses 24 dBm due to excessive heating (see section 2.4 in SI), which could be optimized by replacing gold with superconducting metals (such as niobium or NbTiN) to deliver the MW signal.

We further characterize the single qubit gate fidelity of MW control via randomized benchmarking. For this, we use the following set of Clifford gates: \{I, $\pi_x$, $\pi_y$, $\pi_x/2$, $-\pi_x/2$, $\pi_y/2$, $-\pi_y/2$\} (see section 5.1 in SI). To prevent excessive heating effect during benchmarking which would lead to undesired spin decoherence, we apply a slightly slower Rabi rate (2.8 MHz, 18 dBm) which requires no time buffer between gates. The benchmarking result is shown in Figure 3 (d). We extract an average Clifford gate fidelity of 97.7(1) %, indicating power efficient MW control with high fidelity under stringent randomized benchmarking.

**SnV spin coherence properties**

We next utilize microwave control to characterize the SnV coherence at 1.7 K. We perform a Ramsey measurement as shown in Figure 4 (a). The Gaussian envelope of the Ramsey oscillations corresponds to a spin dephasing time $T_2^*$ of 2.5(1) µs. Similar to ODMR, we observe interaction with a proximal spin in the Ramsey measurement, and we verify that this does not originate from the detuning of the MW signal via phase dependent readout (see section 5.2 in SI). Possible decoherence sources could be nearby vacancies and defects in the diamond membrane, as well as surface spins from both sides of the membrane [43].

Advanced pulse sequences, such as dynamical decoupling via CPMG (Carr-Purcell-Meiboom-Gill) and XY pulse sequences [44, 45], allow us to extend the spin coherence to millisecond timescales. The CPMG results are shown in Figure 4 (b). The $T_{2,\text{echo}}$ returns a value of 100(1) µs, which is already longer than 35.5(30) µs measured using all-optical spin echo process (see section 4.3 and 4.4 in SI), in the absence of optically induced dephasing mechanisms. The $T_{2,C\text{PMG}_{128}}$ refocusing microwave pulses, prolongs the SnV spin coherence to 1.57(8) ms. We note that with no signal normalization being applied, the CPMG figure indicates a high signal fidelity of $\approx 86 \%$ for up to 128 pulses. Future developments on the MW driving fidelity including superconducting metals and faster Rabi pulses can further improve the signal fidelity to higher numbers of pulses. We plot the relationship between the $T_2$ and the number of CPMG or XY pulses $N$ in Figure 4 (c) and fit it with $T_2 \sim N^\beta$. The fitting curve returns a sub-linear dependence with a $\beta$ factor of 0.593(8). We observed minimal $T_2$ differences between CPMG and XY sequences. XY sequences are more resilient to control pulse errors compared to CPMG [45], verifying that the observed coherence is not limited by our control (see section 5.4 in SI).

**Spin-photon interface at 4 K**

Finally, we demonstrate that our strained SnV platform shows state-of-the-art spin coherence for Group IV color centers at 4 K. For Group IVs, the dominant decoherence source of the electronic spin is the electron-phonon interaction (phonon-mediated decay) between orbital branches [20, 42]. The electron-phonon interac-
tion rate depends on the temperature-dependent phonon population and the energy splitting $\Delta_{gs}$ between orbital branches. Therefore, enhanced coherence of the group IV centers can be achieved via either cooling down to millikelvin temperature [21, 22], increased energy splitting by using heavier group IV elements [25], engineering of the phonon density of states [46], or strain engineering [24]. Here we utilize both a heavy element (Sn as compared to Si and Ge) and crystal strain in diamond to improve electron spin coherence at elevated temperatures.

The Rabi oscillation of the SnV at 4 K is shown in Figure 5 (a). The fidelity is characterized to be 97.7(5)\%, only slightly lower than the value at 1.7 K due to background heating limitations. We characterize the average gate fidelity via randomized benchmarking at 4 K using the same 2.8 MHz Rabi rate, returning a gate fidelity of 95.7(3)\%, confirming the maintained high performance spin manipulation of the strained SnV at 4 K.

Equipped with high fidelity Rabi control, we investigate the spin coherence of the SnV centers at elevated temperatures. Due to the much larger splitting $\Delta_{gs}$ of the strained SnV ($\approx 1300$ GHz) compared with bulk SnV ($\approx 850$ GHz), electron-phonon dephasing onsets at higher temperatures. Figure 5 (c) shows the $T_{1}^{\text{spin}}$, $T_{2}$, $T_{2,\text{echo}}$ and $T_{2,\text{XY}}$ versus temperature. Fitting the same $\beta$ factor in $T_{2} \sim N^\beta$ using Hahn-echo and XY4 coherence times returns a value of 0.391(8) at 4 K and 0.014 at 4.5 K, indicating that the dominant decoherence mechanism becomes phonon-induced orbital transitions.

From Figure 5 (c) we notice a much lower dephasing time compared with the decay time $T_{1}^{\text{spin}}$ [47]. This feature originates from the fact that only spin-flipping transitions between the lower and upper orbital branch drive $T_{1}^{\text{spin}}$, whereas $T_{2}$ is sensitive to dephasing by the spin-conserving transitions due to different precession frequencies in the orbital branches [23]. In our case, the phonon transitions are highly cycling due to the aligned magnetic field. Nevertheless, $T_{2}$ at 4 K remains at 2.7(1) $\mu$s – comparable to the 1.7 K value, and $T_{2,\text{echo}}$ only decreases slightly to 74(2) $\mu$s, with $T_{2,\text{XY}}$ reaching the depolarization-limited $T_{2} = 223(10)$ $\mu$s. It is worth emphasizing that all of these are record high values for all group IV spin qubits at 4 K to date.

To demonstrate the potential of the strained SnV center as a promising spin-photon interface at elevated temperature, we investigate the temperature dependence of the SnV optical coherence. As shown in Figure 5 (d), we observe that the ZPL linewidth remains unchanged for both A1 and B2 transitions up to 7 K with the maximum linewidth remaining below 52.0(8) MHz—only 60\% higher than lifetime-limited values. In the future, modest Purcell enhancement of SnV emission rates with on-chip nanophotonics or microcavities can generate fully lifetime-limited photons suitable for efficient entanglement generation.

![Figure 5](image-url) FIG. 5. Performance of the strained SnV center at 4 K. (a) Rabi oscillation of the SnV center, showing a gate fidelity of 97.7(5)\% (b) Randomized benchmarking at 4 K, showing an average gate fidelity of 95.7(3)\%. (c) Temperature dependence of the spin decay time $T_{1}^{\text{spin}}$, dephasing times $T_{2}$, $T_{2,\text{echo}}$, and $T_{2,\text{XY}}$. (d) ZPL linewidths of the two spin conserving transitions (A1, B2) with respect to the temperature, showing negligible broadening with the maximum linewidth below 52.0(8) MHz. The transform-limited linewidth is shown with a dashed line.

CONCLUSIONS

In this work, we demonstrate that SnV in strained diamond membranes is a promising platform for quantum technologies. We create simple heterostructures that leverage differences in thermal expansion to passively generate significant strain of 0.05\% to 0.1\% in diamond, enabling efficient, high fidelity microwave control of the SnV spin. The presence of the strain also suppresses the phonon-mediated decay and improves the spin coherence of the SnV at 4 K, which greatly reduces the technological barrier for quantum networking applications. We reach
Rabi $\pi$ gate fidelity of 99.36(9) % (97.7(5) %) with a randomized single qubit gate fidelity of 97.7(1) % (95.7(3) %) at 1.7 K (4K). Dynamical decoupling sequences allow the SnV spin coherence to reach 1.57(8) ms at 1.7 K and 223(10) µs at 4 K. In the future this value can be further enhanced by generating higher strain through heterostructure optimization and/or additional active tuning. Our platform, derived from scalable diamond membrane generation, is compatible with further on-chip integration, such as microwave coplanar waveguides, integrated photonics [31], and MEMS. Finally, 4K cryostats are relatively affordable and less infrastructure-intensive in comparison to cryogen-free 1.7 K and mK dilution-fridge systems. Therefore, the demonstrated spin-photon interface at 4 K can reduce barriers to widespread utilization and deployment of solid-state quantum technologies.

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COMPETING INTEREST


Supplementary information: Microwave-based quantum control and coherence protection of tin-vacancy spin qubits in a strain-tuned diamond membrane heterostructure

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1 Tin vacancy center (SnV\textsuperscript{–}) in strained diamond membranes

1.1 Hamiltonian of the strained SnV\textsuperscript{–}

The SnV\textsuperscript{–} center is a spin-1/2 system. In a mean-field orbital picture, the system has three electrons in four spin orbitals (\{\{|e_x \uparrow\rangle, |e_x \downarrow\rangle, |e_y \uparrow\rangle, |e_y \downarrow\rangle\}\}). Both its electronic ground and excited states are doubly-degenerate; the degeneracy may be lifted by applying strain and/or by spin-orbit interaction. We write the spin Hamiltonian of the system in the minimum model...
of 4 electrons and 3 orbitals, for the ground \((g)\) and excited \((u)\) state \(H_{g,u}\), as the sum of four terms: spin-orbit (SO) interaction \((\hat{H}_{\text{SO}})\); electron-phonon interaction due to the Jahn-Teller effect; strain field, and interaction with an external, static magnetic field \(B\) (Zeeman effect, \(\hat{H}_Z\)). Following Ref\(^1\), we write the term arising from Jahn-Teller distortions in the same form as that describing the strain interaction. Below we merge the two terms into one, that for simplicity we call \(\hat{H}_{\text{strain}}\). Hence the Hamiltonian is written as:

\[
\hat{H}_{\text{sys}} = \hat{H}_{\text{SO}} + \hat{H}_{\text{strain}} + \hat{H}_Z. \tag{1}
\]

In the following three subsections, we discuss each term of the Hamiltonian.

### 1.1.1 Spin-orbit coupling

The component of the orbital angular momentum operator \(\hat{L}_x, \hat{L}_y\) vanish for the Hamiltonian expressed in the \(\{|e_x\rangle, |e_y\rangle\}\) basis\(^1\) and only the following term is non-zero:

\[
\hat{L}_z = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}, \tag{2}
\]

where we have set \(\hbar\) to 1. Therefore, using the \(\{|e_x \uparrow\rangle, |e_x \downarrow\rangle, |e_y \uparrow\rangle, |e_y \downarrow\rangle\}\) basis, the SO Hamiltonian can be represented as:

\[
\hat{H}_{\text{SO}} = \lambda \hat{L}_z \hat{S}_z = \frac{\lambda}{2} \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix} \otimes \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix} = \begin{bmatrix} 0 & 0 & -i\lambda/2 & 0 \\ 0 & 0 & 0 & i\lambda/2 \\ i\lambda/2 & 0 & 0 & 0 \\ 0 & -i\lambda/2 & 0 & 0 \end{bmatrix}. \tag{3}
\]

### 1.1.2 Strain field

The term of the Hamiltonian representing the presence of a strain field can be written as:

\[
\hat{H}_{\text{strain}} = \begin{bmatrix} \varepsilon_{A_1} - \varepsilon_{E_x} & \varepsilon_{E_y} \\ \varepsilon_{E_y} & \varepsilon_{A_1} + \varepsilon_{E_x} \end{bmatrix} \otimes \mathbb{I}_2. \tag{4}
\]

The elements \(\{\varepsilon_{A_1}, \varepsilon_{E_x}, \varepsilon_{E_y}\}\) represent the energy response induced by strain belonging to the different irreducible representations \(A_1, E_x, E_y\) of the \(D_{3d}\) point group of the defect, and are expressed in the SnV\(^-\) center’s local frame, where the \(z\)-axis corresponds to the high symmetry axis of the SnV which is the quantization axis. For example, \(\varepsilon_{A_1} = \langle \Psi | (H - H_0) | \Psi \rangle\), where \(H_0\) is the electronic Hamiltonian in the absence of strain and \(H\) is the electronic Hamiltonian, which includes the strain field applied to the supercell by changing the lattice parameters. Here \(|\Psi\rangle\) represents a Slater determinant expressed in the \(\{|e_x \uparrow\rangle, |e_x \downarrow\rangle, |e_y \uparrow\rangle, |e_y \downarrow\rangle\}\) basis.

We can write each term of Eq. 4 as a linear combination of the components of the strain tensor \(\varepsilon\):

\[
\begin{align*}
\varepsilon_{A_1} &= t_\perp (\varepsilon_{xx} + \varepsilon_{yy}) + t_\parallel \varepsilon_{zz}, \\
\varepsilon_{E_x} &= d(\varepsilon_{xx} - \varepsilon_{yy}) + f \varepsilon_{xz}, \\
\varepsilon_{E_y} &= -2d \varepsilon_{xy} + f \varepsilon_{yz}, \tag{5}
\end{align*}
\]
where $\epsilon_{xx}, \epsilon_{yy}, \epsilon_{zz}$ represent the diagonal components of the strain tensor in the $x, y, z$ directions and $\epsilon_{xy}, \epsilon_{yz}, \epsilon_{zx}$ represent the shear strain components; $t_\perp, t_\parallel, d,$ and $f$ are partial derivatives written as $\frac{\partial \epsilon_{A_1}}{\partial (\epsilon_{xx}+\epsilon_{yy})}$, $\frac{\partial \epsilon_{A_1}}{\partial \epsilon_{zz}}$, $\frac{\partial \epsilon_{E_x}}{\partial (\epsilon_{xx}-\epsilon_{yy})}$, $\frac{\partial \epsilon_{E_z}}{\partial \epsilon_{zz}}$, respectively. These four strain-susceptibility parameters completely describe the strain-response of the ground and excited electronic states. In the following, we ignore the diagonal term $\epsilon_{A_1}$, which amounts to a global emission wavelength shift. Hence, the strain Hamiltonian has the following form:

$$\hat{H}_{\text{strain}} = \begin{bmatrix} -\epsilon_{E_x} & \epsilon_{E_y} \\ \epsilon_{E_y} & \epsilon_{E_x} \end{bmatrix} \otimes I_2 = \begin{bmatrix} -\epsilon_{E_x} & 0 & \epsilon_{E_y} & 0 \\ 0 & -\epsilon_{E_x} & 0 & \epsilon_{E_y} \\ \epsilon_{E_y} & 0 & \epsilon_{E_x} & 0 \\ 0 & \epsilon_{E_y} & 0 & \epsilon_{E_x} \end{bmatrix}. \quad (6)$$

### 1.1.3 Zeeman effect

Due to the $D_{3d}$ symmetry of the defect, the orbital component $H_{Z,L}$ of the Hamiltonian $H_Z$ only includes a term $\hat{L}_z B_z$, with a pre-factor $q^2$, called in the literature effective reduction factor, accounting for: (i) electron-phonon interaction (so-called Ham term), and (ii) the symmetry of the defect being lower than $O(3)$ (so-called Steven’s factor). Note that both terms have different values for the ground and excited states and hence the $q$ parameter is different in the ground and excited states. The $H_Z$ Hamiltonian is written as the sum of an orbital $H_{Z,L}$ and spin component $H_{Z,S}$

$$\hat{H}_Z = \hat{H}_{Z,L} + \hat{H}_{Z,S} = q \mu_B \gamma_L \hat{L}_z B_z + \mu_B \hat{S} \cdot \mathbf{B} - 2 \mu_B \delta f \hat{S}_z B_z, \quad (7)$$

where $\mu_B$ is the Bohr magneton and $B_x, B_y, B_z$ are the components of the external, static magnetic field along the crystal frame $x, y, z$ directions, respectively. The last term on the right hand side of Eq. 7 originates from correcting with a factor $\delta$ the electronic Landé $g$ factor to account for spin-phonon interaction mediated by spin-orbit coupling. For all the experimental interpretations, we’ll only use the Ham factor and set the Steven’s factor as 1 except section 4.1 where we gave estimates on the possible values of Steven’s factor.

### 1.2 Strain susceptibility

In the presence of a strain field, the degeneracy of the ground ($gs$) and excited ($es$) states is lifted and we call $\Delta_{gs(es)}$ the energy difference between the two states split by the degeneracy. By diagonalizing the strain Hamiltonian defined in Eq. 6, we obtain:

$$\Delta_{gs(es)} = 2 \sqrt{[d_{gs(es)}(\epsilon_{xx}-\epsilon_{yy}) + f_{gs(es)}\epsilon_{zx}]^2 + [-2d_{gs(es)}\epsilon_{xy} + f_{gs(es)}\epsilon_{yz}]^2}, \quad (8)$$

where the strain-susceptibilities are computed from density functional theory (DFT) calculations. We performed DFT calculations employing both the PBE$^3$ and SCAN$^4$ functionals, and a 511-atom supercell with a [0.5, 0.5] occupation number for the $|e_x \downarrow\rangle, |e_y \downarrow\rangle$ orbitals. We approximated the splittings by the energy difference of the corresponding Kohn-Sham (KS) orbitals. The strain susceptibilities $d_{gs}, d_{es}, f_{gs}, f_{es}$ can be obtained from Eq. 8 by varying the
lattice parameters of the supercell to generate \((\epsilon_{xx} - \epsilon_{yy})\) and \(\epsilon_{zx}\) strain, respectively. Our results are summarized in Table. S1. Note the similarity of results obtained with the two different functionals.

<table>
<thead>
<tr>
<th>Functional</th>
<th>(d_{gs})</th>
<th>(d_{es})</th>
<th>(f_{gs})</th>
<th>(f_{es})</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBE</td>
<td>0.787</td>
<td>0.956</td>
<td>-0.562</td>
<td>-2.555</td>
</tr>
<tr>
<td>SCAN</td>
<td>0.834</td>
<td>0.921</td>
<td>-0.563</td>
<td>-2.592</td>
</tr>
</tbody>
</table>

Table S1: Computed strain susceptibilities (see text) of the SnV\(^{−}\) defect in diamond, in units of PHz/strain, obtained with the PBE and SCAN functionals.

1.3 Strain magnitude simulation

We use COMSOL to simulate the strain profile of the suspended area measured in experiments. Since the strain expression \(\epsilon\) in section 1.1 is defined in local SnV frame while the simulation result \(\tilde{\epsilon}\) returns to the lab frame, a combination of rotation matrices are applied. From \(\langle 100 \rangle\) to \(\langle 110 \rangle\) to \(\langle 111 \rangle\), the rotation operators are \(\hat{R}_y(\theta)\) and \(\hat{R}_z(\theta)\), respectively:

\[
\epsilon = \hat{R}_y(54.7^\circ) \hat{R}_z(45^\circ) \tilde{\epsilon} \hat{R}_z(45^\circ) \hat{R}_y(54.7^\circ)
\]

Here \(\hat{R}_y(\theta)\) and \(\hat{R}_z(\theta)\) refer to:

\[
\hat{R}_y(\theta) = \begin{bmatrix}
\cos(\theta) & 0 & \sin(\theta) \\
0 & 1 & 0 \\
-\sin(\theta) & 0 & \cos(\theta)
\end{bmatrix}, \quad \hat{R}_z(\theta) = \begin{bmatrix}
\cos(\theta) & -\sin(\theta) & 0 \\
\sin(\theta) & \cos(\theta) & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

In COMSOL simulation, we use the actual three dimensional (3D) geometry for the diamond membrane and the trench. The temperature-dependent thermal expansion ratio for diamond and fused silica are obtained from these references:\(^5-8\). The initial strain-free temperature is set to 450 °C which is the HSQ healing temperature\(^9\), while the final temperature is set to 4 K. We note that thermal expansion ratios for both fused silica and diamond become negligible below 30 K, thus the simulated strain profile is nearly constant within the temperature range of interest (1.7 K to 7 K) in this study. The simulated structure and the strain distributions of \(E_{xx}\) and \(E_{yy}\) are shown in Figure S1. Since the off-diagonal shear strain is 2-3 orders of magnitude smaller than the diagonal tensile strain, we use the following matrix to represent the simulated strain value:

\[
\tilde{\epsilon} = \begin{bmatrix}
1.3e^{-3} & 0 & 0 \\
0 & 6.8e^{-4} & 0 \\
0 & 0 & -2.5e^{-4}
\end{bmatrix}, \quad \epsilon = \begin{bmatrix}
1.6e^{-4} & -1.8e^{-4} & 5.8e^{-4} \\
-1.8e^{-4} & 9.9e^{-4} & -2.5e^{-4} \\
5.8e^{-4} & -2.5e^{-4} & 5.8e^{-4}
\end{bmatrix}
\]

We note that although \(\tilde{\epsilon}\) only includes diagonal elements, the transformed strain tensor \(\epsilon\) in SnV local frame contains non-negligible off-diagonal elements which could affect the properties of the SnV center through both \(d\) and \(f\) parameters. By comparing the simulated branch splitting
value using equation 8 and PBE results with the actual experimental values, we observed the actual strain to be 0.55 times the simulated value, as shown in Figure S1 (d). This magnitude mismatch could come from either the approximation of energy splittings from KS orbitals being inaccurate, the mismatch of the thermal expansion ratios between COMSOL simulation and reality, or an even lower softening temperature of HSQ rather than the healing temperature. More comprehensive studies of the energy response to strain would require a higher level of method, e.g., embedding theory, which we left for future investigations. In the following calculations, we add this 0.55 pre-factor to the simulated strain tensor to best capture the system properties.

Figure S1: Strain profile of the diamond membrane. (a) The 3D structure of the simulated device. The total length of the carrier wafer is limited to 350 um. (b-c) The $E_{xx}$ ($E_{yy}$) strain profile on the membrane. The position of the SnV center characterized in the main text and SI is labelled as a red star. (d) The energy splitting between two orbital branches in ground and excited states. The simulated strain and the experimentally observed strain magnitudes are labelled in dashed red and green lines, respectively.

1.4 Strain magnitude discussion

We qualitatively categorize the strain magnitude to different regimes via the ground state energy splitting $\Delta_{gs}$. In the spin-orbit regime, this energy splitting is nearly constant, while the splitting
is linear with the external strain when in high-strain regime. Guided by that, we use $\Delta_{gs} = 1200 \text{ GHz}$ as the boundary between the spin-orbit regime and the intermediate regime, and $\Delta_{gs} = 2600 \text{ GHz}$ to identify intermediate and high strain regime. For our work, those values corresponding to strain magnitudes of 0.055% and 0.143%. Here to plot the optical transitions, the magnetic field is set to 80 mT along the quantization axis ($\langle 111 \rangle$ direction) and the strain profile is set to be the same as COMSOL simulated profile but with an additional scaling factor. The calculated relative energy difference of the four $\{A1, A2, B1, B2\}$ transitions are plotted in Figure S2. Compared with the defined low strain “spin-orbit” regime and the high strain regime, our experimentally observed result sits in between, indicating a non-trivial intermediate region where neither spin-orbit coupling or strain shall we treated as perturbation terms. We note that unlike SiV centers,$^{11}$ SnV obtains different quenching $q$ factors at ground (0.471) and excited states (0.125)$^2$, leading to a non-zero splitting between spin-conserving transitions. We also extrapolate a qubit frequency $\omega_s$ of 2.1 GHz which is lower than the ODMR frequency reported in the main text (2.755 GHz). This mismatch could originate from the slight difference between the displayed and the real magnetic field due to hysteresis and the deviation of the effective reduction factors under strain from that in Ref$^2$.

Figure S2: Spin-conserving transition frequencies with respect to different strain magnitudes. The relative strain profile stays the same with only the scaling factor sweeping. The magnitude of the strain is defined by the norm of the tensor. The $B$ field is set to be 80 mT along the quantization axis ($\langle 111 \rangle$) which is in line with the experimental configuration.

2 MW control of the SnV

2.1 MW magnetic response

At zero strain, the SnV spin qubit cannot be manipulated by microwave due to different orbits associated with the spin level. As stated in$^{11}$, the presence of the crystal strain introduces
orbit superposition to SnV’s spin qubit eigenstates, allowing for the coherent control of SnV via external microwave field with qubit frequency \( \omega_s \). Here we use the electronic \( g \) factor to characterize the ability of the MW field (AC B filed) to the spin state of the SnV, including both spin and orbit response of the external magnetic field:

\[
g = \frac{2}{\mu_B} \langle e1 \downarrow | (\hat{H}^{ac}_{Z,L} + \hat{H}^{ac}_{Z,S}) | e2 \uparrow \rangle
\]

(12)

\[
\hat{H}^{ac}_{Z,L} + \hat{H}^{ac}_{Z,S} = \begin{bmatrix}
B^{ac}_z & B^{ac}_x - iB^{ac}_y & -i\eta B^{ac}_z & 0 \\
B^{ac}_x + iB^{ac}_y & -B^{ac}_z & 0 & -i\eta B^{ac}_z \\
i\eta B^{ac}_z & 0 & B^{ac}_z + iB^{ac}_y & -B^{ac}_z \\
i\eta B^{ac}_z & 0 & -i\eta B^{ac}_z & B^{ac}_x - iB^{ac}_y
\end{bmatrix}
\]

(13)

Here the \( B^{ac} \) is a vector with unitary length indicating the direction of the oscillating B field of the microwave. The \( \langle e1 \downarrow | \) and \( | e2 \uparrow \rangle \) are the two spin states of the SnV under external, static B field. The Ham reduction factor \( q \) of the ground state is set to 0.471 according to\(^2\). First we investigate the effect of strain magnitude to the transverse and longitudinal \( g \) factor. The result is shown in Figure S3 (a), indicating a \( g \) factor of 1.64. Here the static B field set to be 80 mT along the quantization axis, which is in line with our experimental setup. We then investigate the angular dependence of the transverse \( g \) factor with different static field orientations using our experimentally observed strain profile. We note that the \( g \) factor has a weak angular dependence, indicating a consistently efficient MW driving efficiency regardless of the static B field orientation, highlighting the robustness of the strained SnV centers.

Figure S3: MW \( g \) factor calculation. (a) The transverse and longitudinal \( g \) factor with respect to the strain magnitude. (b) The angular dependence of the \( g \) factor. The static B field is set to be 80 mT along the quantization axis \( \langle 111 \rangle \).

2.2 Device info

In this work we utilized on-chip coplanar waveguide (CPW) to deliver microwave signals to target SnV centers. Compared with wire-bonded metal striplines\(^12\), lithography-defined CPW...
offers deterministic and reproducible microwave power and magnetic field strength at target location. We designed our waveguide to match the impedance (50Ω) of other electronics in the setup. Ignoring the local dielectric variation near the diamond membrane region, we designed the layout of the CPW based on the permittivity of the fused silica (3.7). The width of the center and the gap is set to 60 µm and 6 µm, respectively. To enhance the local field strength near the SnV region on the membrane, the center of the CPW is reduced to 6 µm. The ground lines of the waveguide is designed to across the membrane to compensate for the trench design, offering a balanced microwave delivering mode. We used a two-port microwave transmission design, demonstrating the potential of driving centers in multiple on-chip devices in the future. The two-port design also allows the microwave signal to be transmitted and dissipated outside of the chamber, relaxing the requirements for reflected signal management (such as circulator). The microscope image of the whole CPW design is shown in Figure S4 (a), with the transmission data of an identical device shown in Figure S4 (b). We show that the transmission loss is low from dc to 15 GHz, with the thermal loss \((P_{\text{in}} - P_{\text{out}} - P_{\text{reflected}})\) around the operation point (2.75 GHz) to be 1.5 dB.

Figure S4: Microwave CPW. (a) Microscope image of the CPW. The bonding pad for wire bonding is not shown. (b) The transmission of the CPW measured via a probe station using a vector network analyzer (VNA). The S11 is the reflection spectrum while the S21 is the transmission. The operation frequency (2.75 GHz) is labelled as a black, dashed line.

### 2.3 MW field simulation

We use COMSOL to simulate the magnetic field acting on nearby color centers. In the simulation, we set the microwave drive power to 24 dBm, the microwave drive frequency to 2.75 GHz, and set the characteristic impedance of the coplanar waveguide to 50 Ohm. The simulated structure and the magnetic field distribution are presented in Figure S5. Based on the simulation results, we expect the effective B field applied to the color centers ranges from 0.2 T to 0.4 T, corresponding to a transverse B field of 0.12 T to 0.23 T.

### 2.4 Heating effect of the system

To investigate the power dependence of the SnV Rabi oscillation, we sweep the MW drive power and extract the Rabi frequency. We observe the expected \(\sqrt{p}\)-behaviour for low drive powers
Figure S5: Simulation of the microwave field in diamond membrane. (a) The 3D structure of the simulated device. The length (width) of the suspended diamond membrane is set to 50 µm (25 µm). (b) The magnetic field in diamond membrane, as a function of distance to the coplanar waveguide. The simulated depth is 40 nm from the top surface of diamond, and the simulated position is labeled as a blue arrow in (a).

\[ p < 24 \text{ dBm}, \] but a clear deviation for larger drive powers. All the power and voltages are referred to the estimated value on the device, extracted by a separate calibration measurements in transmission geometry. We note that no increase in cryostat temperature is observed during the pulsed Rabi measurement.

\[
\begin{align*}
\text{Power (dBm)} & \quad 10 & 15 & 20 & 25 & 30 \\
\text{\(\Omega/2\pi\) (MHz)} & \quad 2 & 4 & 6 & 8 & 10
\end{align*}
\]

Figure S6: Fitted Rabi-frequency \(\Omega/2\pi\) over MW amplitude and power, showing the expected \(\sqrt{p}\)-behaviour for low drive powers \(p < 24 \text{ dBm}\).

The effect of heating on the emitter can be modeled as depicted in figure S7 (a) where we follow the approach taken in Ref\textsuperscript{13}. The emitter is treated as a point-like object at a fixed distance \(X\) from the microwave line. Compared to the SnV center the extent of the gold strip is well approximated as infinite such that we can model this as a 1D problem. Assuming the gold heats and cools instantaneously at the beginning and end of a microwave pulse, a solution to
the 1D heat equation yields a temperature increase at the SnV center $\Delta T_{SnV} \propto \text{erf} \left( \frac{d}{\sqrt{4\alpha t}} \right)$ where $\alpha$ is the thermal diffusivity in diamond. Figure S7 (b) shows that the temperature at the emitter at asymptotically approaches the temperature of the microwave line. When higher Rabi frequencies are used another interesting effect is observed for more complex pulse sequences. Figure S7 (c) shows the effect of a sequence of pulses with a constant interpulse spacing $\tau_0$. If $\tau_0$ is significantly smaller than the time per pulse, the heat cannot flow away fast enough such that a net heating effect is observed per applied pulse. This means at high Rabi frequencies the coherence time of the spin can depend on the time between pulses.

Figure S7: (a) The SnV center as a point like object at a fixed distance $x$ from the microwave gold transmission line. This is the model underlying the 1D heat equation solution. (b) Exemplary temperature increase vs time at the emitter position due to continuous microwave drive. The value is expressed as a fraction of the initial temperature difference approaching equilibrium. (c) Exemplary pulse sequence applying 10 successive microwave pulses each with intermediate spacing of $\tau_0$ and $30\tau_0$, respectively. The maximum temperature differs significantly.
3 Additional optical properties of the strained SnV

3.1 Polarization of the SnV

We probe the polarization of the strained SnVs by inserting a motor-mounted half-wave-plate and a linear polarizer in the detection path. The total intensity of the C-peak and the D-peak over polarization angle are shown in Fig. S8a) and Fig. S8b), respectively. No magnetic field was applied. $0^\circ$ in the graph indicates the magnet $x$-axis. The solid line is a fit of the expected polarization, linear for the C-peak and circular for the D-peak, projected into the lab-frame according to the model in Ref. 1. Both figures indicate a polarization behaviour commensurate with bulk group IV color centers, showing that the polarization is not changed when introducing strain.

![C-Peak emission polarisation](image)

(a)

![D-Peak emission polarisation](image)

(b)

Figure S8: Intensity of the (a) C-peak and (b) D-peak PL emission over linear polarisation in the lab-frame.

3.2 Optical lifetime

We extract the optical lifetime of the SnV by driving the C-transition at zero magnetic field with a single EOM-sideband and turning it off abruptly. The fall time is limited to 200 ps by the EOM. The decay time of the single-exponential is $4.933(190)$ ns which is similar to the bulk value, as shown in Fig. S914.
3.3 Power saturation

We extract the initialization rate, optical cyclicity and saturation power by prior knowledge of the optical lifetime and by sweeping the laser power\textsuperscript{15}. The initialisation rates are fitted by $\frac{1}{\eta^2} \frac{p}{p_{\text{sat}}} \frac{1}{1 + p/p_{\text{sat}}}$ and we extract a saturation power of $7.96 \text{ nW}$ and an optical cyclicity of $\eta \approx 2018$. For the microwave spin control measurement we operate at a saturation parameter of $s = p/p_{\text{sat}} \approx 10$ for the initialization and readout pulses.

3.4 Long term stability of PLE

We acquire PLE for more than 11 hours to test the long-time stability of the SnV (see Fig. S11). We observe a very good frequency stability and only modest spectral wandering.
Figure S11: Time evolution of the PLE line shape. Each vertical cut is the average of fast PLE scans over 1 s, with each shot of the measurement containing both green and red excitation. The solid white line is the fitted emitter resonance frequency.
We fit each acquired PLE trace and extract the common mode shift of the spin-conserving transition\(^\text{16}\) (Fig. S12a). The Gaussian distribution of the shot-to-shot center frequencies has a standard deviation of $\sigma =23.8(1)$ MHz. Similarly, the distribution of extracted spin-conserving splittings (Fig. S12b) has a standard deviation of only $\sigma =13.28(6)$ MHz.

![Figure S12: (a) Histogram of the mean C-peak frequency from shot-to-shot. (b) Histogram of the spin-conserving splitting from shot-to-shot. The grey line is a fit with a Gaussian distribution.](image)

4 Optical control of the SnV spin

4.1 Optical splitting with external B field

We scan the magnetic field over the whole sphere at fixed magnitude. The path between the approximately equidistant points is numerically minimised. The hysteresis of the $B$-field is on the order of 10\% as estimated from linear sweeps along a single magnet axis.

The splitting of the $A_1, B_2$ optical transitions with varying $B$ fields can be computed by diagonalizing the system Hamiltonian $H_{sys}$ of Eq. 1, and the results are shown in Fig S13 (b) and (c). When constructing the Hamiltonian, we considered the Steven’s term $g_L$ in the reduction factor $q$ as a free parameter. The Steven’s term, as discussed in subsection 1.1.3, originates from the defect symmetry being lower than $O(3)$. Here we determined the range of $g_L$ by matching the experiments. We plot the difference of the splitting when the $B$ field is aligned with the defect quantization axis ($\theta_B = 0$), and aligned along the equator ($\theta_B = \pi/2$) with varying $g_L \in [0, 1]$ in Fig S13 (c). The white region in the plot (values close to zero) corresponds to the two splittings being close in energy, matching the experimental observations. Therefore our calculations enabled the narrowing down of the the possible values of Steven’s factor to $g_{L,gs} \in [0.5, 1.0]$ and $g_{L,es} \sim 2g_{L,gs} - 1$. 

\(^{14}\)
Figure S13: Experimental measured and simulations of the spin-conserving optical transitions $A_1, B_2$ with varying external magnetic field $B$. (a) Scanning of the splittings of $A_1, B_2$ transitions with varying $B$ field directions. The magnitude of $B$ is set to 0.2 T. The $x, y$ axis represents the azimuthal ($\phi$) and polar angle ($\theta$) of the $B$ field in the Lab frame. The two poles on the plot represent directions along the quantization axis of the SnV$^-$ and the belt represents the equator. (b) Simulated splittings of the $A_1, B_2$ transitions by diagonalizing the system Hamiltonian along a chosen path of varying $B$ fields, where the path is depicted as a red arrow in (a). The $x$ axis represents the polar angle of the $B$ field in the defect frame. Simulation agrees qualitatively with experiments with the magnitude of splitting underestimated by 0.4 GHz. (c) The differences between the $A_1, B_2$ splittings at $\theta_B = 0$ and $\theta_B = \pi/2$ with varying Steven's reduction factor. The white region corresponds to pairs of Steven's reduction factor for ground and excited states, when taken into the diagonalized Hamiltonian, that match the experimental observations.
4.2 Optical cyclicity of the SnV

We coarsely align the $B$-field by matching it to the polarisation of the optical dipoles (see SI section 3.1) and obtain an optical cyclicity of $\eta \approx 2018$. The cyclicity has a single local maximum close to the pole of the emitter axis, such that we can increase it by sweeping two of the three magnet axes independently. We extract the cyclicity by measuring the decay rate of one of the spin-conserving transitions. The frequency of the sidebands driving the transitions is fixed, noting that the change in $B$-field magnitude corresponds to a change in spin-conserving splitting within one optical linewidth. One can see only a modest increase in cyclicity in Fig. S14a and S14b, such that we conclude that strain limits the maximum achievable cyclicity. Nevertheless, the error introduced by the finite cyclicity will be negligible in spin-photon entanglement protocols due to the relatively high value and enable single-shot readout with nanostructures or microcavities.

Figure S14: Optical cyclicity at (a) fixed $B_y$ and $B_z$, with swept $B_x$ and (b) fixed $B_x$ and $B_z$, with swept $B_y$.

The optical cyclicity, as a branching ratio between spin-conserving and spin-flipping transitions, depends on both system properties and the external optical field. Theoretically, we can only investigate the system property side. Here we investigate an alternative problem—spontaneous emission rate ratio between spin-conserving and spin-flipping transitions—optical cyclicity with absence of the external optical excitation.

According to Ref$^1$, we use optical dipole matrices to calculate the emission rate of the two transitions. The rate (probability) can be expressed using Fermi’s Golden Rule:

$$P = 2\pi \rho \left| \langle \psi_f | e \cdot \hat{r} | \psi_i \rangle \right|^2 = 2\pi \rho \left| \langle \psi_f | \hat{p} | \psi_i \rangle \right|^2$$  \hspace{1cm} (14)

Where $\rho$ is the density of states where we set to 1, the $|\psi_f\rangle$ and $|\psi_i\rangle$ are the final and initial state of the SnV which we assign to the excited state minimum $|e_{A \downarrow}\rangle$ and ground states $|e_1 \downarrow\rangle$ ($|e_2 \uparrow\rangle$) for spin conserving (flipping) transitions. The transition probability (rate) $P$ is related to the optical dipole $\hat{p}$ which is defined as:
Using the above definition, we can calculate the spin flip ratio which is the inverse of spontaneous cyclicity
\[
\frac{1}{\eta} = \frac{P_{\text{flipping}}}{P_{\text{conserving}}}
\]
with respect to the strain magnitude and the $B$ field polar angle $\theta$, as shown in Figure S15. The operation point of the MW-based (all-optical) control of the SnV spin qubit is highlighted in white (black) stars, showing a cyclicity of $\geq 2000$ if $\theta < 4^\circ$ and a cyclicity of $\approx 20$ if $\theta > 85^\circ$, in a rough agreement with the experimentally observed values. We note that the presence of the moderate-level strain will make the overall cyclicity lower than the strain-free case, but the achievable value is still compatible with single shot readout requirements if the signal count can be improved by device design or setup optimization.

Figure S15: The calculated cyclicity of the SnV with respect to the $B$ field polar angle $\theta$ and the strain magnitude. The operation point of the MW-based (all-optical) control of the SnV spin qubit is highlighted in white (black) stars.

4.3 All-optical spin control and operation at perpendicular $B$-fields

We implemented the all-optical control technique shown in Ref.\textsuperscript{15} on Device 1 on a strained SnV with a ground state splitting of $\Delta_{GS} = 1384$ GHz. We extracted an optical lifetime of $7.04(10)$ ns which is compatible for an SnV in proximity of a surface\textsuperscript{17}. We set the magnetic field to $|B| = 100$ mT perpendicular to the emitter axis. From the saturation power measurement in Fig. S16 we extract a saturation power of $p_{\text{sat}} = 4.82(81)$ nW and a cyclicity of $\eta = 5.78(36)$.

The low saturation power and low cyclicity indicate that efficient all-optical control should be possible in principle.
We implement an optical lambda system on the spin-conserving transition A1 and spin-flipping transition A2 and measure coherent population trapping (CPT) by driving both transitions simultaneously (Fig. S17). Fitting the data with the model in Ref.\textsuperscript{15,18}, we get an excited state decay rate of $\Gamma/2\pi = 26.52(91)$ MHz, only a factor of 1.17 larger than the transform-limited linewidth $\Gamma_0/(2\pi) = 22.60(5)$ MHz.

For all-optical Raman control, we choose to operate at a single-photon detuning of $\Delta = 1.5$ GHz. At the lowest laser sideband powers of $p = 455$ nW, we get an intrinsic ODMR linewidth of $\delta f = 1/T_{2*} = 899(54)$ kHz (Fig. S18). The qubit frequency of $f_{\text{qubit}} = 2.321$ GHz, yields a $g$-factor of $g = 0.83$, meaning that perpendicular fields can couple to the SnV efficiently due to strain.

We sweep the Raman drive time $T$ at laser sideband powers of $p = 1012$ nW and extract a Rabi frequency of $\Omega/2\pi = 450(47)$ kHz (Fig. S19) and a $\pi$-gate fidelity of $F_\pi = 83(2)$ %, similar to Ref.\textsuperscript{15}.

Ramsey measurements (Fig. S20) yield an inhomogeneous dephasing time of $T_{2*} = 1.13(7)$ $\mu$s and a Hahn-Echo measurements (Fig. S21) a dephasing time of $T_2 = 35.5(30)$ $\mu$s.
Figure S18: All-optical pulsed ODMR at perpendicular $B$-field directions.

Figure S19: All-optical spin control at perpendicular $B$-field directions and sideband power of $p = 1012$ nW.

We additionally measured the spin decay time $T_1$ at the perpendicular field orientation and found much shorter times on the order of 100 µs (Fig. S22).
Figure S20: All-optical Ramsey measurement showing an inhomogeneous dephasing time of $T_{2s} = 1.13 \pm 0.07 \, \mu$s

Figure S21: All-optical Hahn-Echo measurement showing a dephasing time of $T_2 = 35.45 \pm 2.96 \, \mu$s

Figure S22: Spin decay $T_1$ over inverse temperature $1/T$. 

20
4.4 Spin $T_1$ analysis

Phonon-induced depolarization of group IV centers’ spin is the dominant source of decoherence. Therefore, improving the spin decay time $T_1$ is the central task to enhance the coherence of the system. As discussed in Ref. 11, the spin decay time $T_1$ in the group IV has two meanings, the orbital relaxation time $T_{1\text{orbit}}$ between the energy levels in different orbital branches but with the same spin projection, and the $T_{1\text{spin}}$ between the qubit levels with frequency $\omega_s$. Ultimately, the $T_{1\text{spin}}$ is the factor that directly relates to the coherence of the SnV, so we limit our $T_1$ discussion to $T_{1\text{spin}}$ only.

There are three phonon-induced $T_{1\text{spin}}$ decay paths, including direct single phonon, resonant two phonon (Orbach process) 19, and off-resonant two phonon (Raman process). Similar to SiV, the SnV shows much slower single phonon and Raman spin decay, so we focus on the Orbach process and study its dependence with the $B$ field orientation ($\theta$) and the strain magnitude. Adapted from Ref. 19, we write the decay rate $\gamma_{1\text{spin}}^2$ as follows:

$$\gamma_{1\text{spin}}^2 \propto \frac{\Delta_{gs}^3 \exp(h\Delta_{gs}/k_BT)}{1 - \sum |\langle e_i | H_{AC}^\epsilon | e_j \rangle|^2}$$

(16)

where $i$ represent the states of the lower orbital branch ($|e_1\downarrow, e_2\uparrow\rangle$) and $j$ represent the states of the upper orbital branch ($|e_3\downarrow, e_4\uparrow\rangle$). The $H_{AC}^\epsilon$ denotes an AC strain field which correlates to the phonon interaction in the crystal. We used balanced magnitude for the $H_{\epsilon E_x}^{AC}$ and $H_{\epsilon E_y}^{AC}$ by setting the $H_{\epsilon}^{ac}$ as follows:

$$H_{\epsilon}^{ac} = e \begin{bmatrix} -1 & 0 & 1 & 0 \\ 0 & -1 & 0 & 1 \\ 1 & 0 & 1 & 0 \\ 0 & 1 & 0 & 1 \end{bmatrix}.$$  

(17)

The relative decay rate at temperature 4 K with the maximum normalized to 1 is shown in Figure S23, with MW-based (small $\theta$) and all-optical (large $\theta$) operation points for spin control of the SnV highlighted in black (white) stars. We observe a ratio of 500 to 1200 between the two $T_{1\text{spin}}$, which is roughly inline with our experimental values measured at 6 K (MW-based control $T_{1\text{spin}}=2.5$ ms, all-optical control $T_{1\text{spin}}=1.65$ ms). We note that as a pre-factor, the temperature would not change the decay rate ratio between the two configurations. This ratio reiterates the fact that the Orbach process is the dominant factor for the $T_{1\text{spin}}$ decay.

5 MW control of the SnV spin

5.1 Randomized benchmarking

The gates are chosen from the Clifford group and are $\{I, \pi_x, \pi_y, \pi_x/2, -\pi_x/2, \pi_y/2, -\pi_y/2\}$. We randomly choose $(N-1)$-gates and use the last gate to undo the sequence, followed by a z-basis measurement. The last gate is part of the Clifford group. We get the $\pi$-gate from Rabi measurements and adjust the time $t_\pi$ accordingly. The identity is implemented as wait-time for
Figure S23: The calculated Orbach decay rate of the SnV with respect to the $B$ field polar angle $\theta$ and the strain magnitude. The operation point of the MW-based (all-optical) control of the SnV spin qubit is highlighted in black (white) stars.

t_\pi$, whereas $\pi/2$-gates have a duration of $t_{\pi/2}$. No buffer times are used which would make the qubit prone to dephasing errors, but the drive amplitude is reduced such that local heating effects is not a limiting factor. All randomized benchmarking measurements were taken at a Rabi frequency of $\Omega/(2\pi) = 2.8$ MHz. A total of 10 randomized sequences were applied each time to average out over different implementations. The fidelity $F$ is extracted by fitting the readout with $A \ast F^N + B$, from which we get the error per Clifford gate$^{20}$.

5.2 Ramsey $T_{2\ast}$ at different qubit frequency

Figure S24: Ramsey measurement of the inhomogeneous dephasing time $T_{2\ast}$ at a qubit frequency of 3.694 GHz.

We change the applied magnetic field magnitude from 81.5 to 117 mT and measure the Ramsey dephasing time $T_{2\ast}$. We extract $T_{2\ast} = 2.63(14)$ $\mu$s at the qubit frequency of 3.694 GHz (see Fig.
S24), indicating that $g$-factor fluctuations as reported in Ref.21 are not limiting the observed $T_2^*$.  

### 5.3 Ramsey measurements with phase-readout

Figure S25: Ramsey measurement of $T_2^*$. Both the delay-time $\tau$ and the phase of the second $\pi/2$-pulse are varied.

Fig. S25 shows the measurement from which we extract $T_2$. Ramsey and the modulation of the qubit frequency in Fig. 4 (a) in the main text. We fit for every time delay a modulation of the phase by $A \cos \phi + B$, where $A$ is the visibility shown in the main text and $B$ is the mean value for all time delays averaged over all phases. We then extract the inhomogenous dephasing time $T_2$ by fitting $A$ over delay time $\tau$ with an Gaussian envelope $\propto \exp ((-\tau/T_2)^2)$. With this technique, we can distinguish with certainty a real modulation of the qubit frequency (loss in coherence and visibility) versus a detuning error (no total loss of visibility, but no readout at certain delays and phases). The origin of the beating pattern needs further investigation. The MW-electronics were tested for any modulation. Likely candidates could be surface spins or substitutional nitrogen centres (P1-centres) with a large gyromagnetic ratio close to the one of a free electron ($g = 2$) resulting in relatively large couplings even at large distances.
5.4 XY-sequences

Figure S26: State fidelity $F$ over total delay time $\tau$ for XY-pulse sequences.

Fig. S26 shows the coherence over total time delay $\tau$ in the XY-family of dynamical-decoupling sequences. We observe an almost identical scaling of coherence time over the number of $\pi$-pulses (see main text Fig. 4 c). Because of the much smaller sensitivity of XY- pulse sequences with regards to rotation- or offset errors compared to CPMG sequences, we conclude that our control does not limit the coherence\textsuperscript{22}.

6 Experimental details

6.1 Device fabrication

The diamond membrane substrate was generated via He\textsuperscript{+} implantation and annealing. 450 nm-thick diamond overgrowth layer was introduced in a diamond chemical vapor deposition (CVD) chamber, followed by a $^{120}$Sn implantation with $2 \times 10^8$ cm\textsuperscript{-2} dose and 40 nm target depth. Individual membranes were patterned via lithography and electrochemically etched to undercut the graphitized layer. The target membrane was then transferred to a HSQ-coated fused silica substrate with a 5 $\mu$m-deep etched trench to the generate suspended area. The substrate was annealed at 600 $^\circ$C for 8 h in argon atmosphere. Membrane was thinned down to 160 nm via ICP RIE etching using Ar/Cl\textsubscript{2}, O\textsubscript{2}/Cl\textsubscript{2}, and O\textsubscript{2} recipes. The microwave coplanar waveguide was lithographically defined, followed by Ti and Au deposition with thicknesses of 10 nm and 200 nm, respectively. Excess resist was lift-off using 80 $^\circ$C NMP solutions.

6.2 Measurement setup

All the measurement data in this work were taken in Cambridge, UK. The device was studied in a closed-cycle cryostat (attoDRY 2100) with a base temperature of 1.7 K at the device and...
in which the temperature can be tuned with a resistive heater located under the sample mount. Superconducting coils around the sample space allow the application of a vertical magnetic field from 0 to 9 T and a horizontal magnetic field from 0 to 1 T. Unless explicitly stated otherwise, all measurements were conducted at $T = 1.7 \text{ K}$. The optical part of the set-up consists of a confocal microscope mounted on top of the cryostat and a microscope objective with numerical aperture 0.82 inside the cryostat. The device is moved with respect to the objective utilizing piezoelectric stages (ANPx101/LT and ANPz101/LT) on top of which the device is mounted. Resonant excitation around 619 nm is performed by a second harmonic generation stage (ADVR RSH-T0619-P13FSAL0) consisting of a frequency doubler crystal pumped by a 1238 nm diode laser (Sacher Lasertechnik Lynx TEC 150). The frequency is continuously stabilized through feedback from a wavemeter (High Finesse WS/7). The charge environment of the SnV- is reset with microsecond pulses at 532 nm (Roithner LaserTechnik CW532-100). PL measurements were done with a Teledyne Princeton Instruments PyLoN:400BR eXcelon CCD and SpectraPro HRS-750-SS Spectrograph. Optical pulses are generated with an acousto-optic modulator (Gooch and Housego 3080-15 in the 532 nm path and AA Opto Electronics MT350-A0.2-VIS) controlled by a delay generator (Stanford Research Instruments DG645). For resonant excitation measurements, a long-pass filter at 630 nm (Semrock BLP01-633R-25) is used to separate the fluorescence from the phonon-sideband from the laser light. The fluorescence is then sent to a single photon counting module (PerkinElmer SPCM-AQRH-16-FC), which generates TTL pulses sent to a time-to-digital converter (Swabian Timetagger20) triggered by an arbitrary waveform generator (Tektronix AWG70002A). Photon counts during “initialize” and “readout” pulses are histogrammed in the time-tagger to measure the spin-population. Sidebands driving both resonantly transitions as well as off-resonant all-optical control were generated by an amplitude electro-optic modulator (Jenoptik AM635), and the amplitude, phase, and frequency of the sidebands are controlled by a 25 Gs/sec arbitrary waveform generator (Tektronix AWG70002A). The EOM is locked to its interferometric minimum with a lock-in amplifier and PID (Red Pitaya, STEMlab 125-14) and using a freely available Lock-in+PID application with a feedback loop on the signal generated by a photodetector (Thorlabs PDA100A2).

Microwave pulses are generated with the second channel of the arbitrary waveform generator and amplified with a low-noise amplifier (Minicircuits ZX60-83LN12+) and a high-power amplifier (Minicircuits ZVE-3W-83+). Microwave signals inside the cryostat are delivered via the in-built pico-coax cables, self-soldered cables and a customised PCB. The signal is transmitted through a second line and terminated outside of the cryostat with 50 Ohms.
Figure S27: Sketch of the experimental set-up described in detail in the text.

References


