Enhancing Spin Coherence in Optically Addressable Molecular Qubits through Host-Matrix Control

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Optically addressable spins are a promising platform for quantum information science due to their combination of a long-lived qubit with a spin-optical interface for external qubit control and readout. The ability to chemically synthesize such systems—to generate optically addressable molecular spins—offers a modular qubit architecture which can be transported across different environments and atomistically tailored for targeted applications through bottom-up design and synthesis. Here, we demonstrate how the spin coherence in such optically addressable molecular qubits can be controlled through engineering their host environment. By inserting chromium (IV)-based molecular qubits into a nonisostructural host matrix, we generate noise-insensitive clock transitions, through a transverse zero-field splitting, that are not present when using an isostructural host. This host-matrix engineering leads to spin-coherence times of more than 10 μs for optically addressable molecular spin qubits in a nuclear and electron-spin-rich environment. We model the dependence of spin coherence on transverse zero-field splitting from first principles and experimentally verify the theoretical predictions with four distinct molecular systems. Finally, we explore how to further enhance optical-spin interfaces in molecular qubits by investigating the key parameters of optical linewidth and spin-lattice relaxation time. Our results demonstrate the ability to test qubit structure-function relationships through a tunable molecular platform and highlight opportunities for using molecular qubits for nanoscale quantum sensing in noisy environments.

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I. INTRODUCTION

Quantum information science offers the potential to revolutionize computational, sensing, and networking capabilities [1–5]. Innovations in these areas depend, in part, on optimization of the quantum bit, or qubit, the fundamental unit of quantum information processing, and its environment. Solid-state color centers (such as the nitrogen-vacancy center in diamond) offer a well-developed qubit platform with a robust spin-optical interface for single-qubit initialization and readout [3,6–8] and microwave-frequency spin transitions for coherent control. Introducing these properties into tunable and portable molecular systems combines the key properties of solid-state color centers with opportunities for optimization through bottom-up engineering of both the qubit and its environment [9,10]. These optically addressable molecular spin systems, or “molecular color centers,” provide Angstrom-scale precision and tunability of the local qubit environment [11–15], offering targeted design for applications such as nanoscale quantum sensing [16–18]. Additionally, the ground-state spin hosted in a molecule comprises a portable qubit of <1 nm³ size, such that these systems can be integrated into various host matrices and hybrid material architectures [19–21].

Altering the symmetry or strain environment of solid-state color centers has been shown to significantly enhance qubit properties [22–25]. The portability of molecular
qubits opens up a versatile platform to achieve such enhancements by modifying a qubit’s electronic structure through host-matrix tuning, without altering the chemical composition of the qubit itself. In particular, the sensitivity of the ground-state spin to its local environment suggests that the crystallographic symmetry of the host matrix can be used to induce “clock transitions” which are first-order insensitive to magnetic-field noise and, hence, enhance spin coherence, even in noisy—e.g., nuclear spin-rich—environments. As demonstrated in molecular qubits without a spin-optical interface [26–28] and solid-state spin systems [29–31], such transitions can enhance spin coherence, reducing the need for isotopic control of the nuclear spin environment (e.g., deuteration [32]) or high qubit dilution to prolong coherence.

Here, we create clock transitions in optically addressable molecular spin qubits through host-matrix-induced symmetry control. We use the chromium-based molecular color center—Cr(IV)(o-tolyl)$_4$, 1-Cr [Fig. 1(a)]—to demonstrate the impact of the host environment on the ground-state spin structure. The crystallographic symmetry of 1-Cr and its isostructural, diamagnetic host Sn(IV)(o-tolyl)$_4$, 1-Sn, yields ground-state spin transitions that are first-order sensitive to magnetic-field fluctuations [13]. In contrast, and as we outline below, inserting 1-Cr into a nonisostructural, lower-symmetry host matrix—Sn(IV) (4-fluoro-2-methylphenyl)$_4$, 2-Sn [Fig. 1(b)]—induces clock transitions as a result of the creation of a significant transverse zero-field splitting. These host-induced clock transitions significantly enhance the spin coherence of 1-Cr in 2-Sn (which we refer to as 2) compared to 1-Cr in the isostructural 1-Sn host (which we refer to as 1). We model this behavior from first principles using generalized-cluster-correlation expansion methods and further experimentally demonstrate enhanced optical contrast and spin-lattice relaxation times for these host-matrix engineered molecular color centers. Remarkably, the host modification to achieve this coherence enhancement comprises interchange of just one hydrogen atom on the host ligands with a fluorine atom. In fact, the nuclear spin densities of 1-Sn and 2-Sn are 51.7 and 46.9 N/nm$^3$, where $N$ is the number of nuclear spin-bearing atoms (see Supplemental Material for further details [33]). Thus, the coherence enhancement arises primarily from symmetry control by the host—without requiring control of the nuclear spin bath—offering a pathway for coherence-protected quantum sensing (e.g., of electric fields and strain) in intrinsically noisy environments (e.g., biological systems), all within a versatile molecular platform.

II. RESULTS AND DISCUSSION

A. Host-guest control of molecular color centers

We recently demonstrated optical addressability of molecular spin qubits comprising a chromium (Cr$^{4+}$) ion coordinated by organic ligands in a pseudotetrahedral geometry such as 1-Cr in Fig. 1(a) [13]. The symmetry and tetravalent oxidation state of 1-Cr leads to a spin-triplet ($S = 1$) ground state, while the strong field organic ligands generate a suitable energy level structure for optical-spin initialization and readout. As a result, the ground state can be optically initialized and read out using spin-selective excitation to the spin-singlet ($S = 0$) excited state, combined with photoluminescence (PL) detection, analogous to solid-state color centers [see Fig. 1(c)] [34,35]. Specifically, resonantly exciting these molecular spin qubits with a narrow-linewidth laser initializes the ground state through optical pumping: Selective excitation of a spin sublevel [e.g., $|0\rangle$ in Fig. 1(c)], combined with excited-state decay, transfers population to the other spin sublevels [e.g., $|\pm\rangle$ in Fig. 1(c)]. Similarly, the same selective excitation enables optical-spin readout through the resulting PL: Spin sublevels resonant with the laser are excited and give rise to PL, while the detuned spin sublevels are only weakly excited and, therefore, give rise to weak (ideally vanishing) PL. Combined with microwave control of the ground-state spin, these properties therefore provide a qubit which can

FIG. 1. Host-matrix engineering of optically addressable molecular qubits. (a) Molecular structure of 1-Cr (determined from single-crystal x-ray diffraction) with laser excitation and emission outlined. Hydrogen atoms are omitted for clarity. (b) Single-crystal packing diagram of 1-Cr in its isostructural host, 1-Sn (red, left), and nonisostructural host, 2-Sn (blue, right), showing only positions of metal centers. The cell volumes for the representations of 1-Sn and 2-Sn are 9 and 10 nm$^3$, respectively. Below each cell, we show the molecular structures of the tin host, with hydrogen atoms omitted for clarity. The resulting ground-state spin structures (bottom) show the clock transition ($E > 0$) induced in 2. (c) Energy level diagram of chromium molecular color centers, highlighting resonant excitation to, and photoluminescence (PL) from, the $S = 0$ excited state, and zero-field splitting of the ground-state spin sublevels. (d) PL and photoluminescence excitation (PLE) spectra of 2 at 4 K.
be initialized, coherently controlled, and read out in a fashion which is compatible with single-spin detection.

For 1, the axial zero-field splitting is $D = 3.63$ GHz, which splits the $m = 0$ sublevel, $|0\rangle$, from the $m = \pm 1$ sublevels, $|\pm 1\rangle$ [Fig. 1(b)]. In the tetragonally symmetric crystal environment of 1, the Cr$^{4+}$ site contains a fourfold improper rotational axis ($S_4$) that enforces the transverse zero-field splitting $E$ to be $\approx 0$, resulting in spin transitions that are first-order sensitive to magnetic fields [Fig. 1(b)]. However, if $|E| > 0$, the degeneracy of the zero-field states is broken: The $|\pm 1\rangle$ states hybridize to form the non-degenerate levels $|\pm\rangle = (1/\sqrt{2})(|+1\rangle \pm |-1\rangle)$, which have no first-order magnetic moment. Therefore, systems with $|E| > 0$ exhibit transitions which are first-order insensitive to magnetic fields around $B \approx 0$ [Fig. 1(b)], where $B$ is an external magnetic field. To achieve $|E| > 0$ with 1-Cr, we introduce 1-Cr into a nonisocritical host matrix, 2-Sn [Fig. 1(b)], that contains no threefold (or higher) rotational axes such that $E$ is no longer symmetry constrained to zero. In fact, the metal centers in 2-Sn do not contain any symmetry elements, such that the point group contains any symmetry elements, such that the point group is the transverse zero-field splitting, despite the minor modification of the host matrix in 2 generates a significant transverse zero-field splitting, despite the minor modification—i.e., single-site fluorination—to the host-matrix ligands. In fact, 2 displays the largest possible transverse zero-field splitting for its axial zero-field parameter, i.e., $|E| = |D|/3$, highlighting the significant symmetry breaking afforded by the host matrix. We note that, in this system where $|E| = |D|/3$, two of the spin transitions are degenerate at zero field, so we observe two resonances—at $D - E = 2E$ and $D + E$. This degeneracy is lifted under an applied magnetic field. We further note that, since the signs of $D$ and $E$ are not determined in our experiments and do not influence our results, we take $D, E > 0$ for concreteness. The field-frequency ODMR map [Fig. 2(a)] highlights the insensitivity of the spin transitions to magnetic field: To first order, the energies of the $|0\rangle, |\pm\rangle$ sublevels do not shift with increasing field, in contrast to the linear Zeeman shift exhibited for $E = 0$ in 1.

**B. Host-matrix-induced clock transitions in a molecular color center**

We then determine the ground-state zero-field splitting parameters of 2 through continuous-wave optically detected magnetic resonance (cw ODMR). Under continuous optical excitation, applying a microwave frequency on resonance with a transition between spin sublevels of 2 increases the PL due to the mixing of the “bright” and “dark” spin sublevels, i.e., the sublevels which are resonant with and detuned from the laser, respectively. Figure 2(a) shows the cw ODMR spectrum as a function of both the magnetic field and microwave frequency, from which we extract $D = 5.55$ GHz and $E = 1.85$ GHz. Importantly, compared to 1—in which $E = 0$—the use of a lower-symmetry host matrix in 2 generates a significant transverse zero-field splitting, despite the minor modification—i.e., single-site fluorination—to the host-matrix ligands. In fact, 2 displays the largest possible transverse zero-field splitting for its axial zero-field parameter, i.e., $|E| = |D|/3$, highlighting the significant symmetry breaking afforded by the host matrix. We note that, in this system where $|E| = |D|/3$, two of the spin transitions are degenerate at zero field, so we observe two resonances—at $D - E = 2E$ and $D + E$. This degeneracy is lifted under an applied magnetic field. We further note that, since the signs of $D$ and $E$ are not determined in our experiments and do not influence our results, we take $D, E > 0$ for concreteness. The field-frequency ODMR map [Fig. 2(a)] highlights the insensitivity of the spin transitions to magnetic field: To first order, the energies of the $|0\rangle, |\pm\rangle$ sublevels do not shift with increasing field, in contrast to the linear Zeeman shift exhibited for $E = 0$ in 1.

**FIG. 2.** Host-matrix engineered clock transitions in 2. (a) Continuous-wave optically detected magnetic resonance performed on a single crystal of 2 as a function of the magnetic field and microwave frequency overlaid with simulated spin transition frequencies (dashed lines). The ODMR spectrum yields $D = 5.55$ GHz and $E = 1.85$ GHz. (b) Calculated spin-sublevel energies as a function of the magnetic field highlighting their magnetic-field insensitivity. (c) Pulsed ODMR at zero magnetic field demonstrating an optical contrast of approximately 40%. Inset: pulsed ODMR sequence comprising optical initialization (init), microwave, and optical readout (read) pulses.
C. Host-matrix enhanced spin coherence

We now illustrate how this behavior significantly enhances the spin coherence \( T_2 \) in 2 compared to 1. As a prerequisite for optically detected spin-coherence measurements, we perform pulsed ODMR by applying the pulse sequence outlined in Fig. 2(c). A laser pulse initializes the subensemble of spins, which are then rotated by a microwave \( \pi \) pulse before being read out through the PL with a second laser pulse. (\( \pi \)-pulse lengths are determined from Rabi oscillation measurements—see Supplemental Material [33].) Figure 2(c) shows the pulsed ODMR spectrum as a function of the microwave frequency at \( B = 0 \) for the (doubly degenerate) low-frequency transitions (at \( D = E = 2E = 3.7 \) GHz). We find an ODMR contrast of 40\% (defined such that the maximum possible contrast is 100\%; see Supplemental Material [33]), which is approximately an order of magnitude improvement from our previous demonstration of pulsed ODMR with 1. To measure the spin coherence of 2, we replace the microwave \( \pi \) pulse in the pulsed ODMR sequence with a Hahn echo—i.e., \( \pi/2 - \tau - \pi - \tau \) sequence, where \( 2\tau \) is the free-evolution time—followed by an additional \( \pi/2 \) pulse to project coherences onto populations for optical readout [Fig. 3(b)]. At zero magnetic field, we measure a ground-state spin-coherence time \( T_2 = 10.6 \pm 0.2 \) \( \mu \)s, despite the nuclear spin-rich environment and the relatively high Cr concentration (approximately 1\%, \( \approx 1-2 \times 10^{19} \text{Cr}^{3+}/\text{cm}^3 \)). (For further details on the role of the transition degeneracy in these experiments, see Supplemental Material [33].) In contrast, we measure a significantly shorter \( T_2 = 2.0 \pm 0.1 \) \( \mu \)s for 1 at zero magnetic field, indicating the effectiveness of the clock transition in 2 for enhancing spin coherence.

To further understand the dependence of zero-field spin coherence on the transverse zero-field splitting, we investigate two other molecular systems: Cr(IV)(2,3-dimethylphenyl)_4 diluted in Sn(2,3-dimethylphenyl)_4 (3) and Cr(IV)(2,4-dimethylphenyl)_4 diluted in Sn(2,4-dimethylphenyl)_4 (4). The additional methyl group on the ligands of these compounds induces lower-symmetry crystal packing than 1, and, consequently, \( E \approx 0.5 \) GHz in both cases [13], providing additional test beds of the role of the transverse zero-field splitting in enhancing spin coherence, here from tuning the qubit rather than the host matrix. Figure 3(c) plots the zero-field coherence time for 1, 2, and 3, showing that \( T_2 \) increases with increasing \( E \) (see Supplemental Material [33] for a discussion of the role of \( D \) on \( T_2 \)). 4 shows a similar \( T_2 \) as 3 due to its very similar \( E \) value (see Supplemental Material [33]). Generally, these four systems highlight paths to engineer even longer coherence times through independently optimizing both the host matrix and the chemical composition of the qubit.

To further understand the spin coherence of these molecular color centers interacting with their nuclear spin bath, we use first-principles generalized cluster correlation expansion (gCCE) calculations with Monte Carlo bath state sampling using the PyCCE package [38]. Starting from the crystal structure for these compounds, we calculate the electron-nuclear hyperfine couplings of the Cr-containing molecule using density functional theory (DFT) with the SCAN functional. Using DFT-computed spin densities, we calculate the interactions between the Cr center and nuclear spins in the host matrix and use point dipole-dipole interactions between nuclear spins [33]. The calculated zero-field \( T_2 \) as a function of transverse zero-field splitting \( E \) shows good agreement with the experimental values [Fig. 3(c)]. Since the calculations consider only the nuclear spin bath, they highlight that Cr electronic spins or electric-field- or strain-induced noise are not a major limitation on the coherence in our experiments [33]. Interestingly, the calculations also allow us to determine the distance at which nuclear spins play a significant role in determining the coherence. By varying the number of nearest-neighbor molecules included in the calculations, we find that \( T_2 \)
converges when 3–4 nearest neighbors are included, corresponding to a radius of approximately 1.5 nm around the Cr center (see Supplemental Material for further details [33]).

To the best of our knowledge, the behavior of ground-state electronic spin coherence in the low magnetic field regime (from 0 to approximately 100 mT) is largely unexplored in molecular systems, but, as demonstrated in solid-state color centers, is an important domain for applications in quantum information science [39,40]. To explore this regime in molecular color centers, we measure $T_2$ as a function of the magnetic field for 1, 2, and 3. In each case, $T_2$ decreases with increasing magnetic field between 0 and 30 mT [Fig. 3(d)]. The good agreement between the experimental data and the gCCE calculations indicates that this behavior arises from the magnetic-field-dependent dynamics of the nuclear spin bath, combined with moving away from the clock transitions of 2 and 3. A similar drop in $T_2$ with field due to nuclear spin bath dynamics—albeit at a lower field scale—has been studied in the nitrogen-vacancy center in diamond (which has $E = 0$) [41]. In this case, $T_2$ drops with magnetic field to a minimum at approximately 0.1 mT due to nuclear spin bath dynamics, before recovering at approximately 10 mT, when the nuclear Zeeman splitting dominates over the electron-nuclear-nuclear spin interactions [41]. Our observations in molecular color centers can be assigned to a similar mechanism—albeit with a larger characteristic field scale due to the stronger spin interactions [42]—combined with the increased noise sensitivity for 2 and 3 as they shift from their zero-field clock transitions. Because of the higher field scale involved for these molecular color centers, we see only a reduction in $T_2$ in the measured field range. We note, however, that electron spin resonance measurements performed on 1 at higher magnetic fields ($\approx$400 mT) yield $T_2 \approx 2.5 \mu s$ [15], consistent with the theoretical prediction of a high-field recovery in $T_2$ [41] (see Supplemental Material for further details [33]).

D. Key parameters to optimize molecular spin-optical interfaces

Having shown how host-matrix engineering can enhance the coherence of molecular color centers, we now explore additional key properties of 2. The optical linewidth is a crucial parameter in these molecular systems: It determines the readout and initialization fidelity by setting the spin selectivity of the excitation. Quantifying this linewidth is, therefore, an important step to further optimize molecular color centers. To measure the homogeneous optical linewidth (i.e., that of the subensemble of spins probed under resonant excitation), we perform a two-laser-tone experiment. We apply a fixed laser tone, at frequency $f_L$, along with a second laser tone, detuned by $\Delta f_L$, which we sweep. When the difference in laser frequencies, $\Delta f_L$, matches the spin transition frequencies, the second laser tone excites population that is shelved in other (dark) spin sublevels, thus increasing the PL. Similarly, when $\Delta f_L = 0$, the PL is lower than for a finite detuning, since the population is already shelved in the dark sublevels by the first laser tone. The linewidths of these spectral peaks and holes enable us to determine the homogeneous optical linewidth of the subensemble—which determines the spin-optical contrast—from the inhomogeneously broadened ensemble (Fig. 4). To mitigate the slope in the PL traces [Fig. 4(c)] caused by the inhomogeneous broadening, we use a differential ODMR measurement: We apply a fixed microwave drive $f_{MW}$ at the frequency of one of the spin transitions and measure the ODMR signal as we sweep $\Delta f_L$. From a fit to these measurements, we extract an optical linewidth of $\approx 3$ GHz for 2 [Fig. 4(d)]—see Supplemental Material for further details [33]]. Importantly, since this is comparable to the zero-field splitting parameters, this indicates promise for significantly improving molecular spin-optical interfaces by lowering linewidths. Future work will focus on understanding the electron-phonon and electron-electron dephasing mechanisms contributing to this linewidth (i.e., optical coherence) [43]. We note that similar spin-flip, intraconfigurational optical transitions in coordination compounds exhibit homogeneous linewidths on the order of 10–100 MHz [44,45], suggesting avenues for future improvements through, e.g., lower temperatures, selective deuteration [46], or reduced Cr$^{4+}$ concentration.

We next measure the resulting optical contrast of 2—which provides a lower bound on the spin polarization—by applying an optical pulse (2 ms long) and measuring the emitted photons during this pulse, followed by a wait time much greater than the spin-lattice relaxation time for ground-state equilibration before the next repetition of
the experiment. Over the course of the optical pulse, the PL decreases as spins are optically pumped from the probed bright spin sublevel to the dark spin sublevels. The difference in PL at the beginning and the end of the pulse provides a lower bound on the spin polarization of 65% [Fig. 5(a)]. This is a marked improvement on $I$, where we previously observed a contrast of 14% [13]. We then measure the spin-lattice relaxation time $T_1$ of the ground-state spin by applying an optical initialization pulse, followed by a variable relaxation time $T$, before measuring the relaxation of the spins through a readout pulse during which we collect the PL. By varying the relaxation time, we measure $T_1 = 1.21 \pm 0.02$ ms [Fig. 5(b)], a fivefold improvement compared to our previous measurements of $I$.

The enhanced optical contrast and $T_1$ of $2$ suggests either improved thermalization and/or crystal quality relative to $1$. For example, measuring $T_1$ as a function of the temperature for $2$ shows that $T_1$ drops by more than a factor of 4 between 4 and 7 K (see Supplemental Material [33]), highlighting the dramatic influence of temperature on $T_1$. Thus, a fivefold enhancement of $T_1$ for $2$ relative to $1$ could result from a decrease in effective temperature at the sample of a few Kelvin. Similarly, a reduction in local temperature should reduce the optical linewidths [47], thus improving the contrast. Additionally, the increased $|D|$ of $2$ can improve the spin selectivity of resonant optical excitation. Thus, improving sample thermalization and modifying $|D|$ to larger, yet still measurable, values offer clear routes to optical contrast approaching 100% for molecular color centers. Overall, these measurements show how key molecular qubit properties can be engineered through host-matrix control.

### III. CONCLUSION

This work demonstrates how symmetry engineering through atomistic host-matrix control of a molecular spin qubit can significantly enhance both spin-coherence and spin-optical interfaces. Through host-based symmetry control, we have demonstrated spin-coherence times exceeding 10 $\mu$s for optically addressable molecular spins in a nuclear and electron spin-rich environment. These results combine the advantages of noise-protected coherence with optical qubit initialization, coherent control, and readout in a versatile molecular architecture. By exploring the influence of the host matrix on optical contrast, homogeneous optical linewidths, and spin-lattice relaxation times, our results highlight further directions to improve molecular spin-optical interfaces, e.g., by using lower temperatures to limit thermal vibrations and engineering vibrational modes through isotope control and modification of the chemical makeup of the host matrix or qubit. The ability to transport molecular qubits between different environments, and tune these hosts with atomic-level precision, highlights exciting opportunities for further control over a range of qubit properties, e.g., spin-orbit coupling [48], as well as integration with photonic or phononic devices [49,50]. In particular, the enhanced coherence demonstrated at zero magnetic field could be used to sense electric fields, strain, or temperature at the nanoscale while retaining insensitivity to magnetic-field noise.

Our results demonstrate the promise of rapidly advancing molecular color centers through an iterative feedback loop of bottom-up design, targeted chemical synthesis, qubit measurement, and accurate first-principles calculations. For example, our results point to even longer coherence times using $E$ values of approximately 10 GHz, which should be synthetically achievable [51]. Such $E$ values would enable accessible ground-state spin control while further enhancing optical contrast through increased ground-state spin splittings. Overall, the flexibility of molecular color center design combined with the application of accurate theoretical and computational tools offers promise for optimizing spin-optical interfaces, for applications ranging from nanoscale quantum sensing to long-distance entanglement distribution.

The data underlying this paper are available at [52]. Crystallographic data for $2$-Sn (CCDC code: 2189798) can be obtained free of charge [53].

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S. L. B., D. W. L., D. E. F., and D. D. A. are inventors on patent application No. 63008589 submitted by the University of Chicago that covers chemically tunable optically addressable molecular-spin qubits and associated methods. S. L. B., P. D., D. W. L., M. O., G. G., D. E. F., and D. D. A. are inventors on patent application No. 22-T-121 by the University of Chicago that covers enhancing quantum properties of optically addressable molecular qubits through host-matrix engineering.

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