Millisecond Coherence Time in a Tunable Molecular Electronic Spin Qubit

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Supporting Information

ABSTRACT: Quantum information processing (QIP) could revolutionize areas ranging from chemical modeling to cryptography. One key figure of merit for the smallest unit for QIP, the qubit, is the coherence time ($T_2$), which establishes the lifetime for the qubit. Transition metal complexes offer tremendous potential as tunable qubits, yet their development is hampered by the absence of synthetic design principles to achieve a long $T_2$. We harnessed molecular design to create a series of qubits, ($\text{Ph}_4\text{P})_2[V(C_8S_8)_{3}]$ (1), ($\text{Ph}_4\text{P})_2[V(\beta-C_3S_3)_{3}]$ (2), ($\text{Ph}_4\text{P})_2[V(\alpha-C_3S_3)_{3}]$ (3), and ($\text{Ph}_4\text{P})_2[V(C_3S_5)O]_{3}$ (4), with $T_2s$ of $1-4 \mu s$ at 80 K in protiated and deuterated environments. Crucially, through chemical tuning of nuclear spin content in the vanadium(IV) environment we realized a $T_2$ of $\sim 1$ ms for the species ($d_{50-50}\text{Ph}_4\text{P})_2[V(C_8S_8)_{3}]$ (1) in CS$_2$, a value that surpasses the coordination complex record by an order of magnitude. This value even eclipses some prominent solid-state qubits. Electrochemical and continuous wave electron paramagnetic resonance (EPR) data reveal variation in the electronic influence of the ligands on the metal ion across 1–4. However, pulsed measurements indicate that the most important influence on decoherence is nuclear spins in the protiated and deuterated solvents utilized herein. Our results illuminate a path forward in synthetic design principles, which should unite CS$_2$ solubility with nuclear spin free ligand fields to develop a new generation of molecular qubits.

By exploiting the inherent properties of quantum species such as spins, quantum information processing (QIP) offers the potential to fundamentally transform our approach to modeling chemical systems and cryptography. This tantalizing concept relies upon quantum manipulation of the smallest units of a QIP system, qubits. Electronic spins, one of the most intuitive candidate qubit classes, demonstrate promise in a variety of materials and complexes. Even beyond the well-developed ability to control such qubits and their facile synthetic tunability, electronic spin qubits offer the potential to host multiple qubits within a single manifold of spin levels. Yet, despite decades of research, molecular electronic spin qubits continue to suffer from rapid collapse of their superpositions, a process known as decoherence. The key variable that describes the lifetime of the superposition, the coherence time, or spin–spin relaxation time $T_2$, is still at least an order of magnitude lower in the best transition metal complex qubits than required for quantum error correction schemes and realistic applications. The rational design of qubits with long coherence times necessitates careful consideration of the chemical variables that contribute to decoherence. One primary factor frequently implicated in decoherence is the interaction of the electronic spin with environmental nuclear spins, within both the complex and external matrix. To engender longer coherence times in transition metal complexes, we are designing nuclear spin-free ligand environments to eliminate the former source of nuclear spins. This approach restricts our ligand selection and design to ligands composed of carbon, oxygen, and sulfur exclusively, which exhibit 98.9, 99.8, and 99.2% natural abundances of nuclear spin-free isotopes, respectively.

We recently communicated a pulsed EPR investigation of the species [V(C$_8$S$_8$)$_3$]$_2$ which revealed a $\mu$s-length $T_2$ that persisted to above the temperature of liquid N$_2$. Motivated by these promising results, herein we report a two-part, systematic investigation of synthetically accessible parameters on the lifetime of the qubit formed in the vanadium(IV) trisdithiolate platform: first, a comprehensive study of ($\text{Ph}_4\text{P})_2[V(C_8S_8)_{3}]$ (1) and, second, a subsequent evaluation of a series of molecules, where the impact of specific molecular factors on the superposition lifetime could be tested. Incredibly, we observe a 2 order of magnitude range in $T_2$ within this study, including observation of an unrivaled coherence time $T_2 = 0.7$ ms in 1.

In the initial part of our study, we investigated the stability of the spin superposition of the qubit 1 at the central EPR resonance (Figures 1, S1) by measuring $T_2$ as a function of solvent and temperature (Figure 2). Specifically, we measured 0.5 mM solutions of 1 in four different solvent systems: 1:1 d$_7$-
The lifetime of the superpositions of a dissolved electronic spin qubit. The solvents except the nuclear spin-free CS₂, of \((\text{Ph}_4\text{P})_2[\text{V}(\text{C}_8\text{S}_8)_3] \) in DMF/butyronitrile (PrCN), PrCN, and a 0.01 mM solution in dimethylformamide (DMF)/d₈-toluene (Tol), DMF/Tol, DMF/PrCN, and a 0.01 mM solution of \((\text{Ph}_4\text{P})_2[\text{V}(\text{C}_8\text{S}_8)_3] \) (1') in CS₂ (see Figures 2, S2). In all solvents except the nuclear spin-free CS₂, \(T_2\) remains relatively constant from 10 to 60 K, after which the decay rate displays strong temperature dependence. The temperature independence of \(T_2\) at low temperature suggests that nuclear spin-mediated decoherence is significant in this regime, since phonon mediated processes are strongly temperature dependent. At higher temperatures the \(T_2\) values coalesce toward \(\leq 1\) μs, indicating an increasing role of thermal vibrations and spin–lattice relaxation in decoherence, mediated by the spin–orbit coupling of the vanadium(IV) ion. The concentration dependence of \(T_2\) for 1 was evaluated in PrCN, showing no change in \(T_2\) at 10 K from concentrations of 1 to 0.05 mM. On this basis we further conclude that intermolecular interactions are suppressed at the concentrations utilized in this report.

Elucidation of the phonon contributions to decoherence was accomplished through analysis of the spin–lattice relaxation time, \(T_1\), in each of the solvent systems from 10 to 120 K (Figures S3, S4). For \([\text{V}(\text{C}_8\text{S}_8)_3]^2^-\), the magnitudes of \(T_1\) across all solvents illustrate substantial temperature dependence, decreasing by several orders of magnitude from milliseconds at 10 K (6.5(1)–36.2(2) ms) to under 5 μs at 120 K. The values of \(T_1\) are nearly indistinguishable across solvent systems, indicating that anolyte–solvent intermolecular interactions are not dominant in dictating spin–lattice relaxation, in accordance with other transition metal complex systems. Note, noticeable differentiation occurs at 10 K likely due to differing thermal properties of the frozen glasses. In systems where phonon contributions, librational motion, or efficient molecular tumbling limit \(T_2\), the maximum value of \(T_2\) is \(T_1\). In our system, however, we observe \(T_1 \gg T_2\) within the entire low temperature range, which precludes phonon limitation of \(T_2\) for \([\text{V}(\text{C}_8\text{S}_8)_3]^2^-\) in all solvents except CS₂. Additionally, the value of \(T_2\) at low temperature does not trend with \(T_1\), as evidenced by \(T_1(\text{PrCN}) > T_1(\text{CS}_2)\) yet \(T_2(\text{CS}_2) \approx T_2(\text{PrCN})\). Further comparison of the \(T_1\) and \(T_2\) data for the 1:1 DMF/Tol, 1:1 d₈-DMF/d₈-Tol, and CS₂ solvent systems also reveals nearly identical \(T_1\) values despite vastly different \(T_2\).

Acquisition of data across a range of solvents and temperatures provides us with sufficient evidence to conclude that nuclear spins in the solvent system are the dominant mediator of low temperature decoherence here. The strong influence of nuclear spin on coherence explains the dramatic magnitude of \(T_2\) for 1' in CS₂. In this system, \(T_2\) demonstrates strong temperature dependence over the entire temperature range, which indicates that, following elimination of nuclear spins, the only remaining sources of decoherence are \(T_1\)-related effects. Remarkably, at 10 K, the value of \(T_2\) is 2 orders of magnitude greater than for the other solvent systems, with \(T_2 = 675(7)\) μs (≈0.7 ms) in CS₂. It is vital to note that previous studies of protiated transition metal qubits in CS₂ did not demonstrate this profound enhancement by exclusion of environmental nuclear spins. We attribute the magnitudes here to the crucial combination of the nuclear spin free ligand field and environment which together enable the 2 order of magnitude increase.

Based upon this extraordinary result, we pursued nutation experiments designed to determine whether this candidate qubit can be placed into any arbitrary superposition of the states depicted in Figure 1. In these measurements, a variable-length microwave (mw) pulse (a nutation pulse or tipping pulse) was applied to 1' in CS₂ followed by two mw pulses to detect the tipped angle induced by the nutation pulse (Figure 3). Note that the applied dc magnetic field quantizes the alignment of the spin into down \((M_z = −1/2)\) and up \((M_z = 1/2)\).
+1/2) orientations. Thus, a spin alignment tipped away from the quantized orientations constitutes a superposition of the two levels. The composition of the superposition is dictated by the tipping angle, which is dependent on the power ($B_1$) and length of the mw pulse. With increasing pulse length, these data display a continuous oscillation (a Rabi oscillation), with frequency $\Omega_R$ (the Rabi frequency), as the composition of the newly created superposition cycles through all possible combinations of the two levels (see Figures 3, S5). Note that the decay of the Rabi oscillation occurs on a time scale considerably shorter than the obtained $T_2$ under the same conditions. This observation does not indicate significant destabilization of the qubit by direct manipulation. Rather, the decay indicates a weakening of the echo intensity due to homogeneous and inhomogeneous broadening mechanisms, as well as a distribution of effective $B_1$ fields experienced across the investigated sample. 10

The nutation data highlight two important aspects of the quantum control of $[\text{V(C}_8\text{S}_8\text{)}_3]^{2-}$. Crucially, the time length between adjacent minima and maxima denotes the spin-flip time, which corresponds to a single-qubit NOT computational operation. In $I'$ these times fall from 47 to 19 ns with increasing $B_1$ (see Figures 3, S5), with the potential to be even shorter, following instrumental optimization. A useful figure of merit to characterize the qubit, $Q_M$, defined as the ratio of $T_2/\Omega_R$ and represents the number of spin-flip operations that could be performed with a qubit in the time length of $T_2$. In $[\text{V(C}_3\text{S}_4\text{O})_3]^{2-}$ as isolated in CS$_2$, the fastest operation time determined yields $Q_M \approx 36,000$, which is unrivaled by an order of magnitude among candidate transition metal complex qubits. We note that this value is significantly higher than the threshold required for quantum error correction, an important result if an assembly were realized that contained a multitude of these qubits. 17 Thus, such a magnitude of $Q_M$ implies genuine utility for $I'$ given the combination of the multiple qubits afforded by the electronuclear hyperfine coupling 12 to the access to all superpositions of these qubits, and extremely long $T_2$ values.

To evaluate the universality of the nuclear spin-free design principle and study the tunability of the VS$_6$ platform, we synthesized and studied the complexes (Ph$_4$P)$_2[\text{V(}\beta\text{-C}_3\text{S}_5\text{)}_3] (2), (\text{PhP})_2[\text{V(}\alpha\text{-C}_3\text{S}_5\text{)}_3] (3),$ and $(\text{PhP})_2[\text{V(C}_3\text{S}_4\text{O})_3] (4)$, each with different nuclear spin-free ligand fields (Figure 4a, Tables S1–S4). Within this set of complexes, electron donation was varied across the series in both $\sigma$- and $\pi$-orbitals, as evidenced by cyclic voltammetry experiments which suggest a trend of increasing electronic donating behavior from C$_8$S$_8^{2-} < \beta$-C$_3$S$_5^{2-} < \alpha$-C$_3$S$_5^{2-} < $C$_3$S$_4$O$^{2-}$ (Figure S6). The second factor of potential tunability in 1–4 is the variation in spin delocalization with ligand identity, as indicated by variation in

Figure 3. Rabi oscillation for $I'$ in CS$_2$ at 20 K which shows the ability of $[\text{V(C}_8\text{S}_8\text{)}_3]^{2-}$ to assume any arbitrary superposition of the $|\pm 1/2, -1/2\rangle$ levels. The blue line is a guide for the eye, while spin orientations for specific nutation pulse lengths are depicted. Inset: Spin-flip operation time as a function of $B_1$ attenuation.

Figure 4. Evaluation of a series of VS$_6$ qubits. (a) Molecular structures of the complexes as they appear in the crystal structures of 1–4. Green, yellow, red, and gray spheres represent vanadium, sulfur, oxygen, and carbon atoms, respectively. (b) Nutations for 1–4 that verify quantum control in each member of the series. Data were recorded in 1:1 DMF/Tol at 20 K, and 14 dB attenuation of $B_1$. The spin-flip operation time of 52 ns is highlighted. (c) Temperature dependences of $T_2$ for 1–4 in 1:1 DMF/Tol are nearly indistinguishable.
the hyperfine coupling constant A with ligand identity. Analyses of the cw EPR data (see the Supporting Information, Table S5, and Figure S7) suggests localization of the unpaired electrons in vanadium d orbital, which is further only weakly adjusted by ligand set. Note, the d orbital is energetically distinct from the other 3d orbitals in the pseudo trigonal antiprismatic coordination environments of 1−4.24,25 Thus, small distortions to the coordination geometry of the vanadium(IV) ion, as may occur from molecular vibrations, are hypothesized to be particularly ineffective at modulating the energy of the d orbital. If true, then this aspect of the electronic structure of 1−4 may suppress the ability of the spin−orbit coupling of the vanadium(IV) ion to mediate fast electronic spin decoherence.18,24

The impacts of the foregoing electronic structure factors on the viabilities of 1−4 for QIP were evaluated in a preliminary manner by the determination of T2 and T1 (see Tables S6−S9, Figures S9−S13). T1 and T2 are nearly indistinguishable across 1−4 in 1:1 DMF/Tol, 1:1 DMF/PrCN, and 1:1 d2-DMF/d2-Tol except at the lowest temperatures, where slight differences are seen (Figures S10−S13). However, none of the trends implied by the CV or cw-EPR experiments reproduce the trends in T2 or T1 for 1−4. Importantly the temperature dependences of these parameters and the operation times of the complexes (Figures 4, S14, S15) are remarkably similar across the series of complexes. The observed similarity in the data highlight the viability of each of 1−4 as potential qubits, and demonstrate the generalizability of the nuclear spin-free synthetic design principle to a wide range of nuclear spin-free ligand fields. However, the data highlight the overwhelming impact of the environmental nuclear spins on these qubits. On the basis of these results, we conclude that further evaluations of the specific electronic factors that affect coherence in transition metal complexes must first exclude nuclear spins. The insolubility of 2−4 in CS2 precludes the investigation of the coherent spin dynamics in the absence of nuclear spins, and this fact highlights CS2 solubility as an important design parameter for future transition metal qubits.

On the basis of these experiments, we conclude that the solubility in CS2 and the nuclear spin-free ligands cooperate to create exceedingly long coherence times. This synergistic effect offers a blueprint for the future design of tunable, coordination complex-based qubits. Indeed, this relatively high degree of tunability confers special advantages to the present species over nitrogen-doped C60, despite comparable T2 values.25,26 The remarkable lifetime of the spin qubit 1′ in CS2 lifts the achievable limit of tunable, transition metal complex qubits to the range of solid-state qubit candidates, eclipsing some prominent examples such as the nitrogen vacancy center in non-isotopically enriched diamond, defects in silicon carbide, and Er-doped CaWO4.27−30 Electronic and nuclear spin qubits in isotopically purified environments consistently reach T2 values equal to or greater than ms in magnitude31−34 and we anticipate the realization of comparable lifetimes if enrichment were similarly pursued here.

The foregoing results demonstrate a considerable step forward in the design of transition metal complexes useful for studying spin-based quantum computation. Prior to this report, a clear limitation to the use of transition metal complexes for QIP was their generally low coherence times. Importantly, the magnitude of T2 observed here, which eclipses all other transition metal complex qubits, and some solid state candidates (Figure 5),35 demonstrates the broad viability of coordination compounds as qubits. Thus, these results substantially improve the prospects of developing transition metal complexes with coherence times appropriate for more advanced QIP studies, including creating a hybrid electron−nuclear quantum memory31 with the 51V nuclear spin, or exploiting the full range of transitions within the hyperfine manifold for qubit manipulation.

■ ASSOCIATED CONTENT

Supporting Information

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Experimental details and spectroscopic, crystallographic, and electrochemical data (PDF)

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All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

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