Spin-Orbit Qubits of Rare-Earth-Metal Ions in Axially Symmetric Crystal Fields

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Contrary to the well-known spin qubits, rare-earth-metal qubits are characterized by a strong influence of crystal field due to large spin-orbit coupling. At low temperature and in the presence of resonance microwaves, it is the magnetic moment of the crystal-field ground state which nutates (for several μ s) and the Rabi frequency Ω_R is anisotropic. Here, we present a study of the variations of $\Omega_R(\vec{H}_0)$ with the magnitude and direction of the static magnetic field \vec{H}_0 for the odd ¹⁶⁷Er isotope in a single crystal CaWO₄:Er³⁺. The hyperfine interactions split the $\Omega_R(\vec{H}_0)$ curve into eight different curves which are fitted numerically and described analytically. These "spin-orbit qubits" should allow detailed studies of decoherence mechanisms which become relevant at high temperature and open new ways for qubit addressing using properly oriented magnetic fields.

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Worldwide studies on possible implementation of a spin-based quantum computer showed the significance of scalable single electron spin qubits in, e.g., quantum dots [1–3]. Also of interest are atomic and nuclear spins 1/2 dispersed in solid state matrices with coherence times persisting at relatively high temperature [4–6] or a new class of potentially scalable systems which recently emerged with the observation of Rabi oscillations [7] in the single molecular magnet V₁₅ [8], followed by electronic-spin coherence studies on Fe₈ [9] and Fe₄ [10]. Spin 1/2 qubits are isotropic or quasi-isotropic and their Rabi frequency $\Omega_R = g\mu_B h_{\rm mw}/2h$ depends only on the amplitude of the linearly polarized microwave field $h_{\rm mw}$ (*h* is the Planck constant, μ_B is the Bohr magneton, and $g \sim 2$).

With rare-earth ions in crystals [11], the situation is very different. Contrary to qubits with magnetic moments simply proportional to the spin (single electrons, 3d transition metal ions with zero or quenched orbital moment), the orbital magnetic moment of a rare-earth ion is large and not reduced by the crystal field; this last being much weaker than the spin-orbit coupling. In the general case this leads to a strong magnetic anisotropy and, as it will be shown in this Letter (see preliminary results in [11]), to a strong dependence of the Rabi frequency on the directions of the microwave and static applied fields. The amplitude of the total angular moment $\vec{J} = \vec{L} + \vec{S}$ is generally much larger than 1/2 leading to magnetic moments as large as $10\mu_{B}$ (Ho³⁺) allowing spin manipulations in low driving fields. For odd isotopes, hyperfine interactions of 4f electrons with the rare-earth nuclear spin can be large enough so that nuclear spins are entangled with the total angular moment [12] and modify the Rabi frequency [8].

In this Letter we concentrate on the anisotropy of the Rabi frequency of a single crystal CaWO₄:Er³⁺, a system

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which presents an in-plane magnetic anisotropy favoring the observation of Rabi oscillations and several erbium isotopes (odd and even) allowing the study of coherent dynamics with and without nuclear spin. In the first part we give a theoretical description (numerical and analytical) of what we may call "spin-orbit qubits," and in the second part we present the experimental results confirming quantitatively the theoretical predictions allowing one to understand the effect of strong spin-orbit coupling on Rabi oscillations.

The crystals of CaWO₄:0.05% Er³⁺ used in this study, obtained by the Chokralsky method, are characterized by a body-centered tetragonal scheelite-type structure ($I4_1/a$ space group) with the lattice constants a = b = 0.524 nm and c = 1.137 nm [13]. The Er³⁺ ions substitute for the Ca²⁺ ions inside the eightfold oxygen surroundings, the charge excess being compensated by supplementary substitutions of Na⁺ ions in the crystal. The Hamiltonian operating in the space of states of the ground ${}^{4}I_{15/2}$ multiplet of a single Er³⁺ ion in CaWO₄ (S_4 point symmetry) is given by

$$H = H_{cf} + H_{hf} + H_Z, \tag{1}$$

where H_{cf} is the Hamiltonian of a single rare-earth ion, reduced by the host-matrix crystal field and 4f symmetry:

$$H_{cf} = \alpha_J B_2^0 O_2^0 + \beta_J (B_4^0 O_4^0 + B_4^4 O_4^4 + B_4^{-4} O_4^{-4}) + \gamma_J (B_6^0 O_6^0 + B_6^4 O_6^4 + B_6^{-4} O_6^{-4}).$$
(2)

The O_l^m are the Stevens equivalent operators, α_J , β_J , γ_J the Stevens coefficients [14], and the B_l^m are the crystalfield parameters. For CaWO₄:Er³⁺, $B_2^0 = 231$, $B_4^0 = -90$, $B_4^4 = \pm 852$, $B_4^{-4} = 0$, $B_6^0 = -0.6$, $B_6^4 = \pm 396$, $B_6^{-4} = \pm 75$ cm⁻¹ in the Cartesian system of coordinates with the *z* axis parallel to the *c* axis. The second term in (1),

 $H_{hf} = A_I \vec{J} \cdot \vec{I}$, is the magnetic hyperfine interaction for the ¹⁶⁷Er isotope (I = 7/2, 22.9% abundance, hyperfine constant $A_J = -125$ MHz [15]), quadrupolar couplings are negligible. The third term in (1), $H_Z = g_J \mu_B \vec{J} \cdot \vec{H}_0 (g_J =$ 6/5 is the Landé factor of Er^{3+}), is the static Zeeman interaction. We neglect here dipole-dipole interactions between the highly diluted Er^{3+} ions and superhyperfine interactions with nuclear magnetic moments of the ¹⁸³W isotope. These interactions induce negligible changes of Rabi frequencies, but may be principal sources of decoherence [16-19]. The crystal field of CaWO₄ leads to strong in-plane anisotropy of Er^{3+} (*a-b* plane) [15] and effective g factors $g_{\parallel} = g^c = 1.2$, $g_{\perp} = g^a = g^b = 8.4$ [20]. The Rabi frequency can be calculated either in the laboratory frame (LF) by computing the amplitude of probability of transition induced by the microwave field $h_{\rm mw}$ perpendicular to the static field H_0 or in the rotating frame (RF) by computing the splitting at avoided level crossing.

In the LF, we start with the Hamiltonian (1) of a single ¹⁶⁷Er³⁺ ion. By diagonalization of the 128-dimension space electronuclear Hilbert Hamiltonian (J = $15/2 \otimes I = 7/2$), we calculate the Rabi frequencies as a function of the direction of the static field \vec{H}_0 using the expression $\Omega_R = |g_J \mu_B \vec{J}_{pk} \cdot \vec{h}_{mw}| / 2h$ where $\vec{J}_{pk} =$ $\langle \phi_n(\vec{H}_0) | \vec{J} | \phi_k(\vec{H}_0) \rangle$ is a vector varying with the static field \vec{H}_0 and $\phi_{n,k}(\vec{H}_0)$ are the wave functions resulting from the diagonalization of (1). In general, the anisotropy of Ω_R depends on both the directions of \vec{H}_0 and \vec{h}_{mw} . These expressions for spin-orbit qubits in a crystal field generalize the one for simple spin qubits, $\Omega_R = g^{\alpha} \mu_B h_{\rm mw}^{\alpha}/2h$ [21] $(g^{\alpha}$ is the effective g factor in the direction α of the microwave field $h_{\rm mw}^{\alpha}$).

In the RF, we use the time-dependent effective Hamiltonian of the ground states doublet:

$$H(t) = \mu_B \vec{H}_0[g]\vec{S} + \vec{I}[A]\vec{S} + \mu_B \vec{h}_{\rm mw}[g]\vec{S}\cos(2\pi ft),$$
(3)

where f is the microwave frequency, [g] the g tensor, and $[A] = [g]A_J/g_J$ the hyperfine tensor. The NMR paradigm applied to EPR [22,23] allows one to transform (3) into a time independent effective Hamiltonian. The Rabi frequencies are then given by avoided levels splitting. The Rabi frequencies of the I = 7/2 ¹⁶⁷Er³⁺ isotope are derived from the RF energy spectrum obtained numerically for $H_0 || a = b$; Fig. 1. The effective electronuclear spin states $|1/2, m_I\rangle$ and $|-1/2, m_I\rangle$ coupled by the electromagnetic wave ("dressed states") form several avoided level crossings of which splitting is $h\Omega_R$ (decreasing rapidly when the microwave field rotates from the easy a-b plane to the hard c axis). In the case of the I = 0 isotope, when H_0 makes an angle ϕ with the c axis in the x-z plane and $h_{\rm mw}$ is in the same plane, the result can be analytical and given by



FIG. 1 (color online). Energy spectrum of ${}^{167}\text{Er}{}^{3+}$ with I = 7/2 calculated numerically in the rotating frame. The detuning field is, by definition, the difference between the applied field H_0 and the resonant field of the I = 0 isotope. The Rabi frequencies are equal to the splittings at avoided level crossings (dressed states). Allowed EPR transitions ($\Delta m_I = 0$) are gapped and their anisotropy for different directions of the microwave polarization is shown in the inset.

$$\Omega_R(\phi) = \mu_B h_{\rm mw} \frac{g_\perp g_\parallel}{2hg(\phi)},\tag{4}$$

where $g(\phi) = \sqrt{g_{\parallel}^2 \cos^2 \phi + g_{\perp}^2 \sin^2 \phi}$. Much more extended analytical results will be given in a forthcoming publication [24] showing that both LF and RF methods give, in all cases, quite the same results. In the following we will use the LF method which is more convenient for precise calculations.

The EPR measurements have been performed using a conventional Bruker EPR spectrometer E580. This spectrometer works at both continuous wave (cw) or time resolved (TR) mode in the X band. All experiments were carried out at the frequency of f = 9.7 GHz. The detection of the signal was performed using a cavity working with the TE_{011} mode with Q factor of about 4000 in the cw mode (perfect coupled cavity) and 200 in the TR mode (overcoupled cavity). The sample-cavity ensemble is surrounded by a ⁴He flux cryostat and the temperature was controlled by an Oxford instrument ITC from 2.5 to 300 K. Two pulse sequences have been used: (i) the usual "spinecho sequence" which strongly reduces the decoherence associated with inhomogeneous cw linewidth and (ii) the Rabi oscillations technique consisting of a spin-echo sequence preceded by an excitation pulse. This technique allows one to obtain the time evolution of the averaged magnetization proportional to $\langle S_z(t) \rangle$, i.e., the Rabi oscillations if the time scale is such that coherence wins over relaxation (here the z axis $\|\vec{H}_0$).

Figures 2(a) and 2(b) give an example of measured Rabi oscillation and its Fourier transform whereas Fig. 2(c) shows the eight electronuclear transitions of the I = 7/2



FIG. 2 (color online). (a) Rabi oscillation recorded on the nuclear-spin free Er^{3+} isotope in CaWO₄ ($H_0 = 325.1 \text{ mT}$, $h_{\text{mw}} = 0.26 \text{ mT}$) and (b) its Fourier transform. (c) Contour plot of the Rabi frequency distribution measured in the I = 7/2 isotope while H_0 tilted by 12° from the *c* axis is swept from 220 to 440 mT (colored isomagnetization scale increases from blue to red). The eight Rabi frequencies with smaller intensity correspond to the transitions $(1/2, m_I) \rightarrow (-1/2, m_I)$ of the I = 7/2 isotope ($\Delta m_I = 0$) and the single one with larger intensity to the I = 0 isotope. Calculated Ω_R are represented by symbols going from black square ($m_I = -7/2$) to green circle ($m_I = 7/2$).

isotope, plus the single one of the nuclear-spin free isotope. In this experiment \vec{H}_0 was tilted by $\phi = 12^\circ$ from the *c* axis in order to resolve the eight electronuclear transitions. The symbols correspond to the $\Omega_R = 6.61, 7.30, 7.95, 8.57, 9.14, 9.65, 10.1, 10.5$ MHz calculated from diagonalization of the electronuclear Hamiltonian (1) (LF method) using the published crystal-field parameters and hyperfine constant only [15]. The RF expression (4) generalized to the case $I \neq 0$ and linear in A_J gives the values $\Omega_R = 6.63, 7.22, 7.82, 8.42, 9.02, 9.6, 10.21, 10.80$ MHz [24]. The small differences between the two methods are caused by the neglect of higher terms in A_J . Both sets of values agree very well with experimental data.

Figure 3 shows the evolution of the Rabi frequency of each electronuclear transition when the crystal rotates from $\theta = 0^{\circ} (\vec{H}_0 || c')$ to $\theta = 90^{\circ} (\vec{H}_0 || b)$ (see inset). Above $\theta \approx$ 30° (not shown) all the Rabi frequencies $\Omega_R(m_I, \theta)$ tend asymptotically to the value $\Omega_R \sim 15$ MHz predicted by the usual equation $\Omega_R = g^{\alpha} \mu_B h_{mw}^{\alpha}/2h$ [21] independently of m_I . At $\theta < 30^{\circ}$ these quasidegenerated frequencies $\Omega_R(m_I, \theta)$ split off in a way strongly dependent on the algebraic value of m_I . It is the small tilt angle ϕ which allows one to resolve the transitions with different m_I (i.e., for $\phi = 0$, Ω_R does not depend on m_I). Nonmagnetic isotopes show a similar variation lying between the blue (down-pointing) triangles and green (up-pointing) triangles with $\Lambda_R = \Omega_R(H_0 || c') / \Omega_R(H_0 \perp c') \approx 0.77$. In these experiments a rotation of the resonance field \vec{H}_0



FIG. 3 (color online). Evolution of Ω_R when the static field H_0 rotates by an angle θ around the microwave field h_{mw} . Colored continuous lines are computed numerically. Inset: Geometry of the experiment where the sample defined by its axes a, b, c is tilted by a small angle $\phi \sim 7^\circ$ around the b = b' axis. The dotted line represents the conventional equation $\Omega_R = g^{\alpha} \mu_B h_{mw}^{\alpha}/2h$ [21].

implies a change of its amplitude in order to keep at resonance, and this raises the question to know whether the observed variations of $\Omega_R(m_I, \theta)$ are mainly due to changes in direction or amplitude of the vector field \vec{H}_0 . In order to answer this question, we plot in Fig. 4 the evolution of Ω_R vs $||\vec{H}_0||$. The large dispersion observed on Ω_R at given $||\vec{H}_0||$ definitively confirms that it is, as expected, a directional effect. All the measured Rabi frequencies of Figs. 3 and 4 are very well reproduced by our crystal-field model without any fitting parameter using only published values of parameters, as described above. In particular, the



FIG. 4 (color online). Evolution of Ω_R with Ω_R vs $\|\tilde{H}_0\|$ while the resonant field is rotated. The symbols represent the same data and the lines are calculated numerically and analytically as in Fig. 3. The dotted line represents the conventional equation $\Omega_R = g^{\alpha} \mu_B h_{mw}^{\alpha}/2h$ [21].

ratio Λ_R varies from 0.58 to 0.90, whereas, in the absence of orbital contribution, $\Lambda_R(m_I) = 1$ for any value of m_I .

Interestingly, if the coordinate system is such that the microwave field \vec{h}_{mw} has only one nonzero component $(h_{\rm mw}^{\alpha})$, the scalar-product equation $\Omega_R = |g_J \mu_B \vec{J} \cdot \vec{h}_{\rm mw}|$ simpler form: $\Omega_R(\vec{H}_0, \vec{h}_{mw}) =$ the /2h takes $G(\alpha, \beta)\mu_B h_{\rm mw}^{\alpha}/2h$ where $G(\alpha, \beta) =$ $g_J |\langle \phi_1(H_0^\beta) | J_\alpha | \phi_2(H_0^\beta) \rangle|$ depends on both directions α and β of the static and microwave fields $(\vec{\alpha} \cdot \vec{\beta} = 0)$. This expression traduces the fact that when H_0 rotates around $h_{\rm mw}$ the crystal-field/Zeeman competition varies leading renormalization of wave functions entailing variations of $G(\alpha, \beta)$, i.e., of Ω_R . As mentioned above, this is true only if the crystal is tilted so that \vec{h}_{mw} is not along a crystallographic axis ($\phi \neq 0$, inset of Fig. 3). In the opposite case where \vec{h}_{mw} is along a crystallographic axis (a axis in Fig. 3), Ω_R is isotropic and follows the usual expression $\Omega_R = g^{\alpha} \mu_B h_{\rm mw}^{\alpha} / 2h$ [21] represented by a dashed line in Figs. 3 and 4. This renormalization of wave functions appears clearly in the analytical expressions for the nonmagnetic and magnetic isotopes: Ω_R depends on both g_{\parallel} and g_{\perp} if $\phi \neq 0$ whereas for $\phi = 0$ it depends on g_{\perp} only, and $\Omega_R(m_I)$ associated with the different m_I are clearly different from each others unless $\phi = 0$ or $\pi/2$ [24].

In conclusion, we have shown with the example of Er^{3+} ions diluted in the single-crystalline host matrix CaWO₄ that crystal field deeply modifies the coherent quantum dynamics of what we call "spin-orbit qubits." In particular, it introduces dramatically large variations of the Rabi frequency Ω_R when the static or/and dynamical field (\dot{H}_0) or $h_{\rm mw}$) deviates from the crystallographic axes. This effect, directly connected with modifications of the wave functions of resonant states, is amplified or depressed by the hyperfine interaction of the odd isotope (I = 7/2)which removes the eightfold degeneracy of Ω_R by increasing or decreasing it when $m_I > 0$ or $m_I < 0$. All the experimental data, showing a rich pattern of coherent oscillations, are interpreted quantitatively without requiring any fitting parameter giving a clear picture of this new type of crystal-field-dependent qubits. These qubits are interesting for several reasons. (i) The coherent nutation of the ground-state magnetic moment deriving from crystal-field effects acting on $\vec{J} = \vec{L} + \vec{S}$ (and $\vec{J} + \vec{I}$ with odd isotopes) is associated with, not yet well studied, symmetry and temperature-dependent spin-lattice decoherence mechanism; in particular, the transfer of quantum dynamics from the spin bath to the nutating system is probably partial, implying the existence of residual decoherence [16–19]. (ii) Despite significant spin-phonon coupling, relatively long living coherence is observed (\approx 50 µs at 2.5 K in CaWO₄:Er³⁺) showing that the coupling with crystal-field environment is not redhibitory. (iii) The magnetic moment generally much larger than 1/2 allows spin manipulations in low driving field vectors (amplitude and direction). (iv) Rare-earth qubits inserted in a semiconducting film should be scalable with selective addressing (application of weak local field pulses created by nanoline current adding algebraically to uniform static field) and couplings (controlled carrier injection through a gate voltage).

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