High-resolution optical spectroscopy and magnetic properties of Yb$^{3+}$ in Y$_2$SiO$_5$

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Rare-earth doped crystals are promising systems for quantum information processing. In particular, paramagnetic rare earths could be used to build coherent interfaces with optical and microwave photons. In addition, isotopes with nonzero nuclear spins could provide long-lived states for quantum state storage and processing. Yb$^{3+}$ is particularly interesting in this respect since it is the only paramagnetic rare earth with a spin-$\frac{1}{2}$ isotope, which corresponds to the simplest possible level structure. In this paper, we report on the optical and magnetic properties of Yb$^{3+}$ in the two sites of Y$_2$SiO$_5$, a commonly used crystal for quantum applications. We measured optical inhomogeneous linewidths, peak absorption coefficients, oscillator strengths, and excited-state lifetimes. The Zeeman tensors were also determined in the ground and excited states, as well as the ground-state hyperfine tensor for the $^{171}$Yb$^{3+}$ ($I = \frac{1}{2}$) isotope. These results suggest that Yb$^{3+}$:Y$_2$SiO$_5$ is a promising material for applications like solid-state optical and microwave quantum memories.

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I. INTRODUCTION

Rare-earth (RE) doped crystals are promising solid-state candidates for quantum information processing [1–4]. In particular, they can show extremely narrow optical homogeneous linewidths at low temperature, in the range of a few kHz to less than 100 Hz [5,6]. Combined with inhomogeneous linewidths in the GHz range, this allows optical addressing of ions within ensemble for quantum processors, or quantum memories with large time-bandwidth products. Moreover, several RE ions have nonzero nuclear spins, which can act as long-lived, optically addressable qubits [7,8]. As an example, hyperfine transitions of Eu$^{3+}$:Y$_2$SiO$_5$ can show coherence lifetimes ($T_2$) up to 6 h at 2 K [9] using various dephasing control techniques, and still reach a few ms at 20 K [10]. Recent results include entanglement storage [11,12] and light-matter teleportation at the telecom wavelength [13], single-photon level memories with storage in nuclear spin states [14,15], as well as memories with high efficiency [16,17], and storage time exceeding one minute [18]. Large and switchable interactions between RE ions have also been observed [19], as well as single RE detection [20,21], which opens the way to quantum processing in these systems.

Paramagnetic RE ions, such as Nd$^{3+}$ or Er$^{3+}$, have one more degree of freedom due to their electron spins. This can be used to interface microwave photons to a RE doped crystal through a superconducting resonator [22,23] and obtain quantum memories for superconducting qubits [24–26]. In this case too, nuclear hyperfine transitions can provide long storage time. We recently showed that $^{144}$Nd$^{3+}$:Y$_2$SiO$_5$ hyperfine transitions have ground-state coherence lifetimes up to 9 ms at 5 K, whereas the electron spin $T_2$ is about 100 µs [27]. High-fidelity coherent transfer of microwave excitations to nuclear spins was also demonstrated by quantum state tomography. Ultimately, it should be possible to build coherent interfaces between optical and/or microwave photonics qubits, and long-lived nuclear spin quantum states.

Yb$^{3+}$ ions have attractive properties in this respect. Their 4$f^{13}$ configuration comprises only two multiplets: $^2F_7/2$ (ground) and $^2F_5/2$ (excited) separated by $\approx 10000$ cm$^{-1}$. This energy corresponds to the near infrared, where laser diodes are easily found. The excited state decays usually radiatively, with a lifetime of about 1 ms, which sets a limit for the optical coherence lifetime of 2 ms. There are also two naturally abundant isotopes with nonzero nuclear spin: $^{171}$Yb$^{3+}$ ($I = \frac{1}{2}$) and $^{173}$Yb$^{3+}$ ($I = \frac{3}{2}$), which hyperfine transitions could be used as long-lived qubits. The lower spin angular momenta of Yb$^{3+}$ compared to Er$^{3+}$ and Nd$^{3+}$ ($I = \frac{3}{2}$ for all nonzero spin isotopes) result in simpler energy level structures. This is important since optical initialization and coherent manipulation of spins require selecting specific transitions within the optical inhomogeneous linewidth by potentially complex optical pumping sequences. The lower number of spin states in $^{171}$Yb$^{3+}$ and $^{173}$Yb$^{3+}$ could simplify them considerably.

Here, we report on the low-temperature and high-resolution optical spectroscopy of Yb$^{3+}$:Y$_2$SiO$_5$. This crystalline host has been chosen since it shows outstanding properties in terms of narrow optical linewidths and long coherence spin lifetimes when doped with Pr$^{3+}$, Nd$^{3+}$, Eu$^{3+}$, and Er$^{3+}$ [5,27,28]. It is currently the most used host in quantum storage experiments. We measured inhomogeneous linewidths, absorption spectra, and excited-state lifetimes for the two Yb$^{3+}$ sites and found values among the best for RE doped crystals. Magnetic fields are often used to slow down electron spin relaxation [6], which influences optical and spin coherence lifetimes and spectral hole burning efficiency. As a first step towards these dynamical experiments, we determined the ground- and excited-state g tensors from electron paramagnetic resonance and optical spectroscopy. Ground-state hyperfine tensors were also found for $^{171}$Yb$^{3+}$ and $^{173}$Yb$^{3+}$, giving hyperfine structures extrapolated at low magnetic field of a few GHz, suitable for coupling to superconducting resonators. Overall, these results suggest that Yb$^{3+}$:Y$_2$SiO$_5$ is suitable to build coherent interfaces with optical and microwave photons, while offering nuclear spin transitions for long-lived quantum states.
II. EXPERIMENT

We used Y₂SiO₅ (YSO) samples doped at 0.005 at. % (50 ppm) with Yb³⁺ and cut from a boule grown by the Czochralski method. YSO has a monoclinic structure belonging to the Cᵢ (C₂/c) space group. Yb³⁺ can substitute Y³⁺ in two different crystallographic sites, both with a C₁ point symmetry [29,30]. In addition, nonequivalent subsites appear under magnetic fields neither parallel nor perpendicular to b. They are related by the crystal C₂ (b) symmetry axis. Samples were cut along the three principal dielectric axes: b (the C₂ crystallographic axis), a, and c [31]. Ytterbium has five stable even isotopes¹⁶⁸Yb³⁺, ¹⁷⁰Yb³⁺, ¹⁷²Yb³⁺, ¹⁷⁴Yb³⁺, and ¹⁷⁶Yb³⁺ with nuclear spin I = 0 and a total abundance of 69.59%. There are also two odd isotopes,¹⁷¹Yb³⁺ with an abundance of 14.28% and¹⁷³Yb³⁺ with an abundance of 16.13%.

Absorption spectra with 0.1-nm resolution were obtained with a Varian Cary 6000i spectrophotometer. Fluorescence measurements were performed using a Coherent 829 Titanium Sapphire laser pumped by a Coherent Verdi G10 laser, a SpectraPro 750 monochromator (1-nm resolution), and an InGaAs photodiode. Fluorescence lifetimes were measured with a tunable optical parametric oscillator pumped by a Nd:YAG laser (Ekspla NT342B-SH, 6-ns pulse length) as the excitation source, a Jobin-Yvon HR250 monochromator, and an InGaAs photodiode. High-resolution transmission spectra were recorded by scanning a single mode Toptica DL 100 diode laser (1-MHz linewidth) around 980 nm. Continuous frequency scans of about 15 GHz could be performed. The laser beam was collimated with a power of about 1 mW in front of the cryostat. The transmitted signal was detected by a Thorlabs PDA36A photodiode and a reference beam by a Thorlabs PM10A photodiode. A small part of signal was sent to a Toptica Fabry-Perot Interferometer (1-GHz Free Spectral Range at 980 nm). This allowed us to precisely calibrate the spectra frequency scale. The sample was maintained at 10 K in a CTI-Cryogenics closed-cycle cryostat.

Electron paramagnetic resonance spectra were recorded at 9 K with a Bruker ELEXYSYS E500 and an ELEXSYS Super High Sensitivity Probe Head in X band. For the optical determination of Yb³⁺ excited-state g tensors, the crystal was placed between two permanent NdFeB magnets. The field average value was 217 mT with an inhomogeneity of about 10% along the laser propagation axis. To record angular variations, the crystal sat on a pedestal attached to an Attocube ANRx51 stage and was rotated by steps of 10°. The whole assembly was put in a Janis LHe cryostat at 10 K. The transmission spectra were recorded with the setup described above.

III. RESULTS AND DISCUSSION

A. Optical spectroscopy

1. Absorption and emission spectra

We first recorded absorption and emission spectra to determine Yb³⁺ crystal-field (CF) level energies. The absorption spectrum (Fig. 1) shows well-resolved lines corresponding to transitions from the lowest CF level ²F⁵/₂(0) of the ground multiplet to the three CF levels ²F⁷/₂(0,1,2) of the excited one. These lines are homogeneously broadened except for the two lowest-energy ones which are likely to be inhomogeneously broadened [32]. A narrow and isolated peak, with a full width at half maximum (FWHM) of 2.8 cm⁻¹, is recorded at 10 505 cm⁻¹ and corresponds to the ²F⁷/₂(0) → ²F⁵/₂(1) transition for site 2. Other transitions above 10 300 cm⁻¹ are much broader with FWHM between 15 and 25 cm⁻¹. The structures in the range 10 450–10 650 cm⁻¹ observed for Yb³⁺ concentrations of a few percent [33] are not seen in this 50-ppm doped sample and are therefore attributed to distorted Yb³⁺ sites.

Low-doping concentration also prevents energy transfer between sites and allows separate recording of each site’s emission spectrum (Fig. 2). These lines correspond to the ²F⁷/₂(0) → ²F⁷/₂(0,1,2,3) transitions and, except for the higher-energy lines, whose widths are instrument limited, are much broader than those observed in absorption. In addition, many partially resolved lines, that we attribute to vibronic transitions, appear in the range 9760–10 060 cm⁻¹ for site

![FIG. 1. Unpolarized absorption spectrum of Yb³⁺:Y₂SiO₅ at 12 K with light propagating along the b axis. Transitions between CF levels are indicated for sites 1 and 2.](image-url)

![FIG. 2. Emission spectra of sites 1 (purple line) and 2 (orange line) of Yb³⁺:Y₂SiO₅ at 10 K excited on the 0 → 2 transitions of Fig. 1 at 919 and 902 nm. Transitions between CF levels are indicated.](image-url)
TABLE I. Optical transitions in Yb$^{3+}$:Y$_2$SiO$_5$. CF level energies; 0-0 transition wavelength (vac), inhomogeneous linewidths, peak absorption coefficient for different light polarizations, oscillator strength; experimental fluorescence decay time ($T_{1s}$) spontaneous emission decay time for the 0-0 transition ($T_1$), $T_{1s}/T_1$ ratio.

<table>
<thead>
<tr>
<th>Energy (cm$^{-1}$)</th>
<th>$^{2}F_{7/2}$</th>
<th>$^{2}F_{5/2}$</th>
<th>$\lambda_{vac}$ (nm)</th>
<th>$\Gamma_{inh}$ (GHz)</th>
<th>$\alpha_0$ (cm$^{-1}$)</th>
<th>$P \times 10^7$</th>
<th>$T_1$ (ms)</th>
<th>$T_{1s}$ (ms)</th>
<th>$T_{1s}/T_1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site 1</td>
<td>0</td>
<td>111</td>
<td>10189</td>
<td>981.463</td>
<td>2.2</td>
<td>3.3 ($E \parallel D_1$)</td>
<td>5.0</td>
<td>0.87</td>
<td>8.5</td>
</tr>
<tr>
<td></td>
<td>499</td>
<td>10391</td>
<td>10874</td>
<td>6.5 ($E \parallel b$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site 2</td>
<td>0</td>
<td>234</td>
<td>10216</td>
<td>978.854</td>
<td>1.7</td>
<td>3.7 ($E \parallel D_1$)</td>
<td>6.4</td>
<td>1.3</td>
<td>6.5</td>
</tr>
<tr>
<td></td>
<td>612</td>
<td>10505</td>
<td>11076</td>
<td>10.3 ($E \parallel D_2$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>970</td>
<td></td>
<td></td>
<td>9.0 ($E \parallel b$)</td>
<td></td>
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</table>

2. This induces an uncertainty in the $^{2}F_{7/2}(2)$ level position, which we determined using the strongest peak at 9982 cm$^{-1}$.

The crystal-field levels for both sites are summarized in Table I and are in good agreement with previous studies [33,34]. CF splittings for the ground and excited multiplets are significantly smaller for site 1, which suggests that it corresponds to the crystallographic site with a coordination number (CN) of 7, in which Y-O distances are larger. Indeed, scalar CF strengths have been found smaller for this site in Ce$^{3+}$:Y$_2$SiO$_5$ in a theoretical study [29]. In the same way, Pr$^{3+}$ ions with the smaller CF splittings have also a higher absorption coefficient, suggesting a higher relative concentration [35]. Since Pr$^{3+}$ ions have a larger ionic radius than Y$^{3+}$ ones, they should occupy preferentially the CN = 7 site, which has a larger volume. Yb$^{3+}$ site attributions should be confirmed experimentally, for example, by electron spin echo envelope modulation that can probe the distances and relative orientations between Yb$^{3+}$ ions and Y$^{3+}$ nuclei [36].

Because of phonon relaxation between CF levels within the multiplets, only the 0-0 [$^{2}F_{7/2}(0)$--$^{2}F_{5/2}(0)$] transition can have a narrow homogeneous linewidth at low temperatures. It is therefore the transition of interest for quantum information processing (QIP) applications [3], and we focus on it in the following. High-resolution absorption spectra for the 0-0 transitions were obtained at zero external magnetic field by scanning a single-mode diode laser. Spectra were obtained for the light electric field $E$ polarized along the $D_2$ and $b$ axes and the wave vector $k$ parallel to $D_1$; for $E$ parallel to $D_1$, $k$ was parallel to $D_2$ (Fig. 3). The lines peak at 10 188.87 cm$^{-1}$ (981.463 nm in vac) and 10 216.06 cm$^{-1}$ (978.854 nm in vac) for sites 1 and 2.

Both lines show a narrow central part, as expected for the $I = 0$ Yb$^{3+}$ isotopes, on top of a broader and weaker structure, in which peaks can be clearly seen in some cases (e.g., site 2, $E \parallel D_2$). We attribute this additional feature to the zero-field hyperfine structures of the $^{171}$Yb$^{3+}$ (abundance 14.3%) and $^{173}$Yb$^{3+}$ (abundance 16.1%) isotopes, which span about 3–4 GHz in the ground state (see Sec. III B). All lines could be well fitted by a combination of Lorentzian lines (see inset in Fig. 3). This has been observed in several RE doped crystals [37,38] and, according to Stoneham [39], corresponds to perturbations by a low concentration of point defects. The FWHM of the central parts of the lines are 2.2 and 1.7 GHz for sites 1 and 2. These values are comparable to those found for Y$_2$SiO$_5$ doped at low levels of RE ions [5] and are related to the difference in ionic radius between Y$^{3+}$(0.892 Å) and Yb$^{3+}$ (0.858 Å) [40]. It is important to note that growth conditions can also have a significant influence on $\Gamma_{inh}$ [41,42].

Large and anisotropic peak absorption coefficients were measured, reaching maximum values of 6.5 cm$^{-1}$ ($E \parallel b$) and 10.3 cm$^{-1}$ ($E \parallel D_2$) for sites 1 and 2, respectively. In both cases, the lower absorption occurs for a light electric field polarized along $D_1$. The average oscillator strengths $P$ were calculated without refractive index corrections and assuming equal Yb$^{3+}$ occupancy for the two sites, because of the close similarity in the absorption spectrum for $E \parallel b$ and $E \parallel D_2$.

FIG. 3. High-resolution absorption spectra of sites 1 (upper graph) and 2 (lower graph) in Yb$^{3+}$:Y$_2$SiO$_5$ at 10 K for light electric field polarized along $b$, $D_1$, and $D_2$ axes. Inset: deconvolution of site 2 spectrum for $E \parallel D_2$ by three Lorentzian curves.
2. Comparison with other rare-earth ions

In the context of quantum processors and quantum memories, RE with strong optical transitions are needed since they can be coherently driven at a faster rate and result in larger opacity [3]. Another important parameter for RE doped crystals coupled to optical cavities is the strength of the transition of interest and its relation to all other possible transitions [45].

As the cavity enhances only resonant transitions, achievable emission rates with Purcell enhancement are proportional to the emission rate through the transition that would be coupled to the cavity, 0-0 in our case. The corresponding radiative emission rates with Purcell enhancement are proportional to the oscillator strength (see previous section). The ratio \( T_{1s}/T_1 \) should be as small as possible, with \( T_{1s} \) as a lower bound. On the other hand, decays trough uncoupled transitions or nonradiative paths represent a loss for the cavity-emitter system and should be minimized. The ratio \( T_{1s}/T_1 \) has therefore to be as small as possible. As shown in Table I, site 2 presents the best values for \( P \), \( T_{1s} \), and \( T_{1s}/T_1 \).

Compared to other RE doped crystals investigated for QIP, its oscillator strength is one of the highest reported in the YSO host [3,45]. It is more than twice the value found in Pr\(^{3+} \), three times that of Er\(^{3+} \), and five times that of Eu\(^{3+} \). This is mainly due to the 0-0 transition originating from the first excited multiplet and the low J value of the ground multiplet of Yb\(^{3+} \). Combined with a relatively short and purely radiative \( T_1 \), the reduced number of possible transitions favors a strong 0-0 transition. Still, some RE-host combinations allow larger oscillator strengths, like Nd\(^{3+} \):YVO\(_4\) (8 \( \times \) 10\(^{-6} \)) or Pr\(^{3+} \):Y\(_2\)Al\(_2\)O\(_3\) (1.5 \( \times \) 10\(^{-6} \)) [45].

\( T_{1s} \) is also short, similar to what is found in Pr\(^{3+} \):YSO (5.7 ms) and much shorter than for Er\(^{3+} \) (54.6 ms) or Eu\(^{3+} \) (120 ms) doped YSO. Compared to systems with very high oscillator strengths, such as Nd\(^{3+} \):YVO\(_4\), \( T_{1s} \) is, however, about 13 times longer. Finally, \( T_{1s}/T_1 \) is very close to the lowest value observed for YSO, 4.8 in Er\(^{3+} \):YSO and about one order of magnitude better than for Pr\(^{3+} \) (34.5) and Eu\(^{3+} \) (63.2). Only Tm\(^{3+} \):LiNbO\(_3\) shows a significantly lower ratio (2.25). In summary, the 0-0 transition of Yb\(^{3+} \) ions in site 2 of YSO shows among the best properties for RE doped crystals in terms of oscillator strength, spontaneous decay \( T_{1s} \), and \( T_{1s}/T_1 \) ratio.

![Image](image-url)

**FIG. 4.** Site 1 EPR spectrum obtained for \( B \) at 160° from \( D_1 \) in \( D_1D_2 \) plane in Yb\(^{3+} \):Y\(_2\)SiO\(_5\). Lines corresponding to the \( I = 0, \frac{1}{2}, \) and \( \frac{3}{2} \) isotopes are observed and calculated from a spin Hamiltonian model (see text).

**B. Magnetic properties**

We first determined Yb\(^{3+} \) ground-state Zeeman and hyperfine tensors by EPR, complementing previous studies where only particular orientations of the magnetic field were investigated [30,34]. Due to the two crystallographic sites, which generally divide in two subsites under a magnetic field, and the two Yb\(^{3+} \) isotopes with \( I \neq 0 \), many lines were observed for magnetic fields \( B \) in the range 50–1000 mT. Figure 4 shows site 1 lines for \( B \) at 160° from \( D_1 \) in \( D_1D_2 \) plane, a configuration where the two subsites, related by a \( C_2 \) symmetry along \( b \), are equivalent. Apart from the Zeeman line at 103.5 mT, the two transitions corresponding to \( ^{171}\text{Yb}^{3+} \) (\( I = \frac{3}{2} \)) are observed at 76.1 and 131.2 mT. The other lines located between 64.5 and 144.6 mT are attributed to the hyperfine structure of \( ^{171}\text{Yb}^{3+} \) (\( I = \frac{5}{2} \)). In this case, some transitions with \( \Delta M_I \neq 0 \) are clearly seen and could be partly due to quadrupole interactions (Fig. 4 and Fig. S1 in the Supplemental Material [46]). For comparison, the positions of lines corresponding to \( \Delta M_I = 0 \) and deduced from \( ^{171}\text{Yb}^{3+} \) hyperfine tensor (see below) are indicated. The lines were very narrow at low magnetic field and FWHMs as low as 12 MHz could be recorded, which is favorable to coupling with high-quality-factor microwave resonators [23].

The \( g \) tensors of the \( I = 0 \) isotopes were determined for both sites from angular variations of the Zeeman lines in the three perpendicular planes \( bD_1, bD_2, \) and \( D_1D_2 \) (Figs. 5 and S2). All line positions were simultaneously fitted to the Hamiltonian \( H = \mu_B \mathbf{B} \cdot \mathbf{g} \cdot \mathbf{S} \), where \( \mu_B \) is the Bohr magneton and \( \mathbf{S} \) is an effective \( \frac{1}{2} \) spin operator. The \( C_2 \) symmetry linking the subsites was also taken into account. The effective spin approach is possible because of the large ground-state CF splittings (100–200 cm\(^{-1} \)) compared to the Zeeman one (0.3 cm\(^{-1} \)). Moreover, at low temperature, only the lowest-energy CF doublet is populated. For each plane, misalignment between the crystal and the laboratory frame was also allowed for by introducing three variable angles in the fit. Excellent agreement with experimental data was obtained, as shown in Figs. 5 and S2. The \( g \) tensors are given in the Supplemental Material [46] and corresponding principal values and Euler angles for the principal axes are found in Table II.
TABLE II. Magnetic properties of \( \text{Yb}^{3+}: \text{Y}_{2}\text{SiO}_5 \). Principal values of the \( \mathbf{g} \) and \( \mathbf{A} \) (\(^{171}\text{Yb}^{3+}\), in MHz) tensors and Euler angles (in degrees) defining the principal axis orientations (\( zxz \) convention).

<table>
<thead>
<tr>
<th>Site</th>
<th>( g_x )</th>
<th>( g_y )</th>
<th>( g_z )</th>
<th>( \alpha )</th>
<th>( \beta )</th>
<th>( \gamma )</th>
<th>( A_x )</th>
<th>( A_y )</th>
<th>( A_z )</th>
<th>( \alpha_A )</th>
<th>( \beta_A )</th>
<th>( \gamma_A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site 1 Ground state</td>
<td>-0.31</td>
<td>-1.60</td>
<td>6.53</td>
<td>252.8</td>
<td>113.8</td>
<td>0</td>
<td>-2140</td>
<td>-5302</td>
<td>247</td>
<td>67</td>
<td>122</td>
<td></td>
</tr>
<tr>
<td>Excited state</td>
<td>-0.8</td>
<td>1.0</td>
<td>3.4</td>
<td>77</td>
<td>84</td>
<td>173</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site 2 Ground state</td>
<td>-0.13</td>
<td>-1.50</td>
<td>6.06</td>
<td>59.1</td>
<td>11.8</td>
<td>347.4</td>
<td>2</td>
<td>1490</td>
<td>-4760</td>
<td>51</td>
<td>11</td>
<td>12</td>
</tr>
<tr>
<td>Excited state</td>
<td>-1.0</td>
<td>1.4</td>
<td>-3.3</td>
<td>234</td>
<td>157</td>
<td>190</td>
<td></td>
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</table>

Figure 6 shows the calculated effective \( g \) factors for magnetic fields oriented in the \( bD_1, bD_2, \) and \( D_1D_2 \) planes. For site 1, the largest principal value is 6.53, with the principal axis oriented close to the \( D_1 \) axis. For site 2, the largest principal value is similar, 6.06, but corresponds to an orientation close to the \( b \) axis. This explains the main features of the angular variations of Fig. 5. For both sites, the two other principal values are much smaller. This results in low-\( g \) factors in the \( bD_2 \) and \( D_1D_2 \) planes for sites 1 and 2, respectively.

As mentioned above, EPR lines were fitted allowing for crystal misalignment in each plane. For site 1, misalignment angles were below 2°. Surprisingly, these angles were much larger in the case of site 2, reaching for example 7° for the \( D_1D_2 \) variation. Moreover, fitting site 1 transitions with site 2 angles resulted in poor agreement with experimental data. This rules out a higher sensitivity of site 2 transitions to crystal misalignment. This discrepancy is particularly clear in the \( D_1D_2 \) angular variation (Fig. S2). For magnetic fields around 412 mT (but for different crystal orientations), subsite lines are separated by only 1.2 mT for site 1, whereas site 2 lines are separated by 21.3 mT. This result suggests that the distortion introduced by \( \text{Yb}^{3+} \) substituting \( \text{Y}^{3+} \) in site 2 results in an effective \( C_2 \) axis direction significantly different from the host crystal. This observation should be further investigated, as it is of importance for predicting particular energy level structures, such as clock transitions [47–49].

The \( \mathbf{g} \) tensor of the \( ^2\text{F}_{5/2}(0) \) excited state for the \( I = 0 \) isotopes was determined using optical measurements, by recording transmission spectra under a magnetic field of about 200 mT. The low 0-0 inhomogeneous linewidth allowed observing four partially resolved transitions in most orientations for both sites. Figure 7(a) shows a transmission spectrum for site 1 with \( B || D_2 \). In this case, the ground state Zeeman splitting is larger than the excited one and the energy difference corresponding to the effective \( g \) factors of the ground and excited states is indicated in Fig. 7(a). Effective \( g \) factors were determined in the three \( D_1D_2, bD_1, \) and \( bD_2 \) planes by rotating the crystal with respect to the magnetic field (Figs. S3 and S4). The values for \( B || b \) are in agreement with those reported in [34]. In a first step, the ground-state \( \mathbf{g} \) tensor measured by EPR was compared to the experimental data to check for sample misalignment. The corresponding angles were then used to fit the excited-state Zeeman splittings to the spin Hamiltonian in the same way as for the ground state. The calculated effective excited-state \( g \) factors are shown in Figs. 7(b) and 7(c).

Table II gives the principal values and axes directions of the excited-state \( \mathbf{g} \) tensors for both sites (tensors in the crystal reference frame are given in the Supplemental Material [46]). The largest values are close, 3.4 and 3.3 for sites 1 and 2, and lower than those of the ground state by a factor of about 2. This can be qualitatively understood by considering the expression of the \( g_{zz} \) component for a pure \( M_J \) crystal-field level: \( g_{zz} = 2g_JM_J \), where \( g_J \) is the Landé factor. With \( M_J = \frac{1}{2} \) and \( \frac{3}{2} \), we find \( g_{zz} = 8 \) and 4.3, which reproduces approximately

![FIG. 5. Angular variation of EPR transitions in the \( bD_1 \) plane (\( I = 0 \) isotopes) for sites 1 and 2 in \( \text{Yb}^{3+}: \text{Y}_{2}\text{SiO}_5 \). Crosses: experimental data; lines: fit to a spin Hamiltonian model.](image)

![FIG. 6. Calculated ground-state effective \( g \) factors in the \( bD_1, bD_2, \) and \( D_1D_2 \) planes for sites 1 and 2 in \( \text{Yb}^{3+}: \text{Y}_{2}\text{SiO}_5 \).](image)
FIG. 7. (a) Transmission spectrum of site 1 in Yb$^{3+}$:Y$_2$SiO$_5$ under a magnetic field of 217 mT along the $D_2$ axis ($T = 10$ K). Energy separations corresponding to the ground- and excited-state effective $g$ factors are indicated. (b), (c) Calculated excited-state effective $g$ factors in the $bD_1$, $bD_2$, and $D_1D_2$ planes for sites 1 and 2.

the experimental ratio of 2. The principal axes corresponding to the largest principal $g$ values are oriented close to the $D_1$ and $b$ axes for sites 1 and 2. These orientations are close to those obtained for the ground state, which explains that the angular variations and effective $g$ factors in the $D_1D_2$, $bD_1$, and $bD_2$ planes look similar [Figs. 6, 7(b) and 7(c), and S3 and S4].

It was shown in Er$^{3+}$:YSO that a long optical coherence lifetime could be obtained with strong fields applied in direction where large effective $g$ factors for the ground and excited states are observed in both sites [6]. At low temperatures, the upper Zeeman levels are strongly depopulated in this configuration, and Er$^{3+}$ spin flips are suppressed. Dephasing of the optical transitions between the lowest Zeeman levels is thus strongly decreased too. In Er$^{3+}$:YSO, this could be obtained with a field in the $D_1D_2$ plane, for which subsites are equivalent. From Figs. 6 and 7, it is clear that a similar strategy in Yb$^{3+}$:YSO requires a magnetic field in the $bD_1$ plane, which results in magnetically nonequivalent subsites and, for example, a reduced optical density. In the $D_1D_2$ plane or along the $b$ axis, where subsites are magnetically equivalent, one site has always a low effective $g$ factor. Additional experiments are, however, needed to fully explore this question, as spin dephasing mechanisms may have complex dependencies with respect to RE concentration, temperature, and magnetic field magnitude and orientation [50].

We finally determined the ground-state hyperfine interaction tensors $A$ for $^{171}$Yb$^{3+}$ ($I = \frac{1}{2}$) from EPR experiments. The angular variations of the corresponding EPR lines in the $D_1D_2$, $bD_1$, and $bD_2$ planes were fitted to the Hamiltonian

$$H = \mu g B \cdot g \cdot S + I \cdot A \cdot S - \mu_n g_n B \cdot I,$$

(2)

where $\mu_n$ is the nuclear magneton and $g_n$ the nuclear $g$ factor ($g_n = 0.9885$). We used the $g$ tensors and misalignment angles previously determined from the transitions of the $I = 0$ isotopes. Figure 8(a) shows the angular variation of the two transitions between hyperfine levels in the $bD_1$ plane for the two subsites of site 2. Experimental data are very well reproduced by the fit, as in the case of the other angular variations (Fig. S5). The $A$ tensors are given in [46] for sites 1 and 2, and the corresponding principal values and principal axes are gathered in Table II.

For both sites, the ratio between the tensor elements $A_{ij}/g_{ij}$ is nearly constant for the large $A_{ij}$ and $g_{ij}$ values ($A_{ij}/g_{ij} \approx -0.0264$ for site 1, $A_{ij}/g_{ij} \approx -0.0259$ for site 2), as expected for a pure $J$ multiplet. Indeed, $A_{ij}/g_{ij} = A_J/g_J$ where $A_J$ does not depend on $M_J$. As a result, the $g$ and $A$ tensors of both sites have nearly the same orientation as $J$ mixing by the crystal field should be very small given the large separation between the $^2F_{7/2}$ and $^2F_{5/2}$ multiplets ($10,000$ cm$^{-1}$) compared to the multiplet splittings ($<1000$ cm$^{-1}$). The larger deviation between the $g$ and $A$
principal axes orientations for site 2 could be due to an enhanced mixing effect, consistent with the stronger crystal field of the CN = 6 environment (see Sec. III A). The large hyperfine interaction results in zero-field splittings of 3.7 and 3.1 GHz for sites 1 and 2, which can be tuned by several GHz using small magnetic fields oriented along the large effective g factors (Figs. 9 for site 1 and S6 for site 2). Moreover, clock transitions could appear when the Zeeman and hyperfine interactions start to be comparable. These transitions, also known as ZEFOZ transitions (zero first-order Zeeman shift), are insensitive to magnetic field fluctuations, which can increase their coherence lifetime [47,49]. As an example, a transition insensitive to fluctuations in the direction of the applied magnetic field, i.e., a partial clock transition, is given for comparison (blue lines).

We also deduced the hyperfine tensor for $^{173}\text{Yb}^{3+}$ by scaling $^{171}\text{Yb}^{3+}$ A tensor by the ratio (−0.27) of the nuclear gyromagnetic factors ($g_n = −0.27$ for $^{173}\text{Yb}^{3+}$). This accounted for the observed angular variations of $^{173}\text{Yb}^{3+}$ strongest EPR transitions [Figs. 8(b), S7, and S8], although small energy shifts and extra $\Delta M_I \neq 0$ transitions are observed (Figs. 4 and S1). This could be partly due to the quadrupole interaction, which was not included in the model Hamiltonian. The calculated zero-field splittings are about 3.5 GHz for both sites and show complex behaviors at low fields, including many transitions insensitive to fluctuations in the direction of the applied magnetic field (Fig. S6). However, this should be taken only as an indication since the quadrupole interaction was not included.

IV. CONCLUSION

Optical properties of a 50-ppm doped Yb$^{3+}$:Y$_2$SiO$_5$ crystal have been studied in the context of applications in quantum information processing. In particular, a detailed study of the transition between the lowest crystal-field levels of the $^2\text{F}_{7/2}$ and $^2\text{F}_{5/2}$ multiplets has been carried out at low temperature, allowing the measurements of inhomogeneous broadening $\Gamma_{\text{inh}}$, peak absorption coefficients of polarized light, oscillator strengths $P$, and excited-state lifetimes $T_1$.

For Yb$^{3+}$ ions in site 2, we found of $P = 6.4 \times 10^{-7}$, one of the largest values observed for a rare-earth ion in Y$_2$SiO$_5$, while $\Gamma_{\text{inh}} = 1.7$ GHz. A relatively strong spontaneous emission rate was also found for the 0-0 transition: $T_{1r} = 6.5$ ns and a ratio $T_{1r}/T_1 = 5.0$. These values also compare favorably with those obtained for other RE doped materials.

We also determined the ground- and excited-state Zeeman tensors of the $I = 0$ isotopes from angular variations obtained, respectively, with EPR and optical transmission under a magnetic field of about 200 mT. The largest principal values for both sites are close to $g = 6$ for the ground state and $g = 3$ for the excited state. The corresponding principal axes are close for the ground and excited states and oriented, respectively, along $D_1$ and $b$ for sites 1 and 2. Yb$^{3+}$ is the only paramagnetic rare-earth ion with a $I = \frac{1}{2}$ isotope, which could be advantageous for optical addressing of spin transitions. We determined the ground-state hyperfine tensor for the $^{171}\text{Yb}^{3+}$ ($I = \frac{1}{2}$) isotope from EPR measurements. Principal values as large as 5 GHz are observed, leading to calculated total zero-field splittings of 3.7 and 3.1 GHz for sites 1 and 2. With magnetic fields oriented along directions of large effective g factors, ground-state transitions can be tuned by several GHz with fields of tens of mT. In regions were the Zeeman and hyperfine interactions are comparable in strengths transitions insensitive to magnetic field fluctuations are predicted, which could lead to increased coherence lifetimes. Extrapolated hyperfine tensor for $^{173}\text{Yb}^{3+}$ ($I = \frac{3}{2}$) isotope is in reasonable agreement with EPR angular variations. Calculated zero-field splittings are in the same range as for $^{171}\text{Yb}^{3+}$, with a much complex behavior under magnetic field.

In conclusion, these measurements suggest that Yb$^{3+}$:Y$_2$SiO$_5$ could be used for quantum information processing with optical, electron, and nuclear spin degrees of freedom.

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