Zero-field paramagnetic resonance of trivalent gadolinium in several lanthanide tris(ethylsulfate) nonahydrates and nonadeuterates: Determination of crystal-field and hyperfine-structure parameters*†

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(Received 5 June 1974)

Employing isotopically pure trivalent $^{153}$Gd, $^{157}$Gd, and $^{159}$Gd as impurities in dilute mixed crystals of lanthanum tris(ethylsulfate) nonahydrate $[\text{La(C}_{2}\text{H}_{4}\text{SO}_{4})_{2}\cdot 9\text{H}_{2}\text{O}]$, it has been possible to determine both accurate zero-field splitting parameters of crystal-field origin and hyperfine-structure constants due to magnetic dipole and electric quadrupole interaction terms. The zero-field paramagnetic-resonance spectroscopic method employed herein, has allowed the absolute signs of the hyperfine parameters to be determined. Thus the magnitude and sign of all diagonal parameters of the zero-field effective spin Hamiltonian are accurately known. It is reasoned that the quadrupole hyperfine constant is due, almost entirely, to lattice contributions; moreover, it is shown to be of the expected order of magnitude and sign. The hyperfine anomaly is discussed and is shown to be slightly less than the experimental error limits (three standard deviations). A set of crystal-field parameters are reported for the first time for $^{157}$Gd in crystals of $[\text{La(C}_{2}\text{H}_{4}\text{SO}_{4})_{2}\cdot 9\text{D}_{2}\text{O}]$, which differ significantly from those of the hydrate.

Variations of crystal-field parameters with various lanthanide host ions are also observed. Finally, a few comments concerning zero-field linewidths and their measurement are made.

I. INTRODUCTION

Zero-field paramagnetic resonance (ZFPMR) is an important technique for observing magnetic-dipole transitions in the microwave region of the spectrum. These transitions are usually between energy levels split by crystal-field and hyperfine interactions in transition metal, lanthanide, and actinide ions. To date, such systems have been mainly studied by conventional electron-paramagnetic-resonance (EPR) spectroscopy in the presence of a strong Zeeman perturbation which forces differences in energy levels to match the energy of a source oscillator. In ZFPMR the microwave source is tuned to match the energy spacings of the sample. The data presented in this work emphasize the comparison between high- and zero-field paramagnetic resonance, particularly for improved hyperfine-structure resolution and crystal-field-splitting determinations.†

One of the long-standing problems of EPR spectroscopy of ions in crystals is that of the unexpected and unexplained crystal field and hyperfine structure of S-state ions (i.e., ions with half-filled shells such as $4f^7$), particularly in the lanthanide series. The usual lanthanide S-state ions are Eu$^{3+}$, Gd$^{3+}$, and Tb$^{3+}$; of these three $4f$ S-state ions Gd$^{3+}$ is the most widely and readily obtainable chemically. Its lattice substitutional properties in a variety of trivalent-metal-ion crystal-host materials$^{14}$ makes Gd$^{3+}$ an ideal choice for such a study. In the treatment of gadolinium paramagnetic resonance, a fictitious spin $S = \frac{7}{2}$ is used in conjunction with a spin Hamiltonian to describe the spectrum in an empirical manner. This convenient description is possible because the spectrum always displays the site symmetry of the ion. An exhaustive tabulation has been provided by Buckmaster and Shing.$^{15}$

Many mechanisms have been proposed for the splitting, but no single one can account for the magnitude and sign of the observed splitting. This area of intense research has been many times reviewed and discussed in detail.$^{16,17}$ In order to evaluate a given theoretical mechanism for crystal-field splitting of the $^8S_{7/2}$ ground state of Gd$^{3+}$, the calculated over-all splitting should be compared with the measured value (this work)$^{18}$ 7258, 4±9.6 MHz for $\text{La(C}_{2}\text{H}_{4}\text{SO}_{4})_{2}\cdot 9\text{H}_{2}\text{O}$. Gd$^{3+}$.

All mechanisms proposed to account for crystal-field splitting of Gd$^{3+}$ could, until very recently, be classified into three groups: much too large, much too small, and about the right magnitude but of the wrong sign.$^{18-20}$ However, using an effective operator technique, Newman and Urban$^{21}$ were able to calculate "intrinsic spin-Hamiltonian parameters." It is their contention that, due to geometric factors and a linear superposition of contributions from neighboring ligands, measured crystal-field parameters, especially quadrupolar ones, may or may not have the same sign as calculated (intrinsic) ones. It is clear that these concepts give both sign and rough order-of-magnitude agreement between crystal-field spin-Hamiltonian parameters obtained from theory and experiment.$^{21,22}$ The conclusion is that, at least for YVO$_4$, YPO$_4$, LuPO$_4$, LuAsO$_4$, and CaF$_2$, correlation and relativistic crystal-field effects contribute most significantly to the observed splittings. Indeed, these authors indicate that the previous calculation by Wybourne
when interpreted by their method, gives reasonable agreement between theory and experiment. Since the only quantitative treatment of these concepts has been for Gd$^{3+}$-H$^+$ and Gd$^{3+}$-F$^-$ in CaF$_2$ tetragonal sites, it still remains to be seen if general agreement for a larger number of situations will obtain.

It is possible that results presented herein can make a positive contribution to this area as the measured crystal-field parameters are different for $Ln(C_2H_3SO_4)\cdot 9H_2O$: Gd$^{3+}$ and $Ln(C_2H_3SO_4)\cdot 9D_2O$: Gd$^{3+}$ where $Ln$ is a lanthanide. Several mechanisms are suggested (end of Sec. III) by which this effect could occur.

Curiously enough, hyperfine structure was never conclusively observed for Gd$^{3+}$ in these hosts in spite of its observation in many others. Since the spectral splittings produced are greater in zero than in high field when compared to the observed linewidths, it was possible to measure hyperfine constants for the first time. A detailed analysis of the spectrum and the constants themselves is presented in Secs. III and IV. The constants are discussed from the point of view of the theory of hyperfine interactions.

As a conclusion to this introduction, it is perhaps appropriate to mention at least briefly previous work on Gd$^{3+}$ in the ethylsulfate-host-crystal system. The field has indeed had a long history commencing with studies in the late 1940’s and early 1950’s. The now famous work of Bleaney, Elliott, Stevens, and co-workers genuinely initiates EPR spectroscopy as a general physical technique for the study of solids. Studies over the years have produced many interesting results for these salts. The works that directly bear on the present investigation are an early work of Bleaney and co-workers which suggested that the zero-field splitting (ZFS) could not be characterized by spin-Hamiltonian parameters obtained at high field; a ZFPMR study of Gd$^{3+}$ in lanthanum tris (ethylsulfate) nonahydrate (LES), of large experimental uncertainty ($\pm 25$ MHz in transition frequency), which indicated such a discrepancy did not exist but which observed no hyperfine structure; a high-field experiment which demonstrated gadolinium-gadolinium and gadolinium-water proton coupling; and finally, recent low-field measurements reinvestigate the adequacy of fit of the high-field spin-Hamiltonian parameters to low-field spectra. One interesting aspect of these latter results is the suggestion that field-frequency plots for one of the transitions may be nonlinear in fields below 10 G. Other work has been reviewed in Ref. 39.

II. EXPERIMENTAL METHODS AND CRYSTAL STRUCTURE

The ZFPMR spectra used in this work were taken at room temperature, 77, 4.2, and 2 K in the ZFPMR spectrometer previously described. In each specific case below the microwave structure used to record the spectrum will be given.

Ethylsulfate crystals used in this research were generally prepared according to the method of Erickson. In the case of the deuterated ethylsulfates 99.7-at. %, Merck, Sharp, and Dohme Canada, Ltd, deuterium oxide was used not only for the reaction but also for washing the diethylsulfate. These samples were kept out of the atmosphere as much as possible to minimize isotopic exchange.

The bismuth magnesium double nitrate (BMDN) sample was prepared according to the method of Gerkin and Thorsell. The magnesium was Research Organic/Inorganic Chemical Corp. 99.99% magnesium metal crystals. The bismuth used was Alfa Inorganics 99.9995% bismuth-metal shot. The nitric acid used was duPont reagent grade.

In the case of the isotopically enriched crystals for either of the above hosts, Union Carbide Oak Ridge National Laboratory stable isotope oxides were used. The $^{159}$Gd$_2$O$_3$ used was 99.7% in $^{159}$Gd, 0.08% in $^{160}$Gd, and the balance $I=0$ isotopes. The $^{155}$Gd used was 99.82% in $^{155}$Gd, 0.07% in $^{157}$Gd, and the balance $I=0$ isotopes. The $^{160}$Gd used was 99.99% in $^{160}$Gd.

The lanthanide tris (ethylsulfate) nonahydrates are an isomorphic series of compounds, and as such have a special importance in the EPR of transition ions, especially the rare earths. The trivalent lanthanide ion, $Ln^{3+}$, may be any member of the rare-earth series and sits at a site of $C_{3v}$ symmetry. The nine water molecules form a triangular prism, with the oxgens in the mirror plane at a greater distance from the threefold axis than the oxgens forming the end triangles. Good diagrams may be found in the crystal-structure papers of Ketelaar and Fitzwater and Rundle. The water protons are probably hydrogen bonded to the oxgens of the sulfate groups, but have not been located experimentally. There are two formula units per unit cell, with approximately 18 nearest-neighbor sulfate oxgens about the lanthanide ion. The ethylsulfate anions are arranged in such a manner to make the space group $P6_3/m(C_{3v})$. The waters of hydration may be replaced with D$_2$O when the crystal is synthesized, without deuteration of the ethylsulfate chains.

III. CRYSTAL FIELD

A. $^{160}$Gd and natural gadolinium in La(C$_2$H$_5$SO$_4$)$_3$·9H$_2$O

1. Results

ZFPMR spectra of gadolinium–doped LES where the gadolinium ions have either the naturally occurring isotopic composition or are 99.99% $^{160}$Gd$^{3+}$ ($I=0$) are presented first. These studies were made for three reasons. First, it is necessary to ascertain if hyperfine structure from the two odd
isotopes can be observed superimposed on the 1 = 0 lines when they are present in their natural composition of about 15% each. Second, accurate determination of ZFS is essential in order to test the adequacy of the spin-Hamiltonian formalism in zero field in a more stringent way than has been done in the past. Third, precisely determined crystal-field parameters must be used in the calculation of hyperfine-structure constants (see below).

The form of the spin Hamiltonian for the crystal field of C_{3v} symmetry has been well worked out,\textsuperscript{13,17,39} Using a fictitious spin $\frac{S}{2} = \frac{1}{2}$ to represent the splitting in the $^3S_{1/2}$ ground state it is written in a form given by Elliott and Stevens,\textsuperscript{24}

$$\mathcal{H}_{\text{CF}} = B_2^0\hat{S}_z^2 - B_4^0\hat{S}_z^4 + B_6^0\hat{S}_z^6 + B_8^0\hat{S}_z^8$$ \hspace{1cm} (3.1)

in which the $B_n^0$'s are numerical constants, and the $\hat{O}_n^0$'s are Stevens's operator equivalents as have been extensively tabulated by Hutchings.\textsuperscript{40}

This spin Hamiltonian leads to four distinct Kramers' doublets in zero field. If the small off-diagonal coefficient $B_2^0$ is neglected, their energies are given by

$$\begin{bmatrix}
    E_1 \\
    E_2 \\
    E_3 \\
    E_4
\end{bmatrix} = \begin{bmatrix}
    7 & 1 & -3 & -5 \\
    1 & -13 & -9 & -5 \\
    -3 & -9 & 5 & 9 \\
    -5 & -9 & 9 & 5
\end{bmatrix} \begin{bmatrix}
    b_0^0 \\
    b_4^0 \\
    b_1^0 \\
    b_1^0
\end{bmatrix} = \begin{bmatrix}
    E(a + \frac{1}{2}) \\
    E(a - \frac{1}{2}) \\
    E(a + \frac{1}{2}) \\
    E(a - \frac{1}{2})
\end{bmatrix}, \hspace{1cm} (3.2)

$$

where $b_0^0 = 3b_2^0$, $b_4^0 = 60b_0^0$, $b_1^0 = 1260b_0^0$, and $b_2^0 = 1260b_0^0$, and $M_z$ is the projection of $\hat{S}$ on the crystal threefold axis for the four states. Using simple $\Delta M_z = \pm 1$ selection rules, the three zero-field transitions are

$$\begin{bmatrix}
    E_1 - E_2 \\
    E_2 - E_3 \\
    E_3 - E_4
\end{bmatrix} = \begin{bmatrix}
    T_1 & T_2 & T_3 \\
    6 & 4 & -2 \\
    20 & -10 & 12
\end{bmatrix} \begin{bmatrix}
    b_0^0 \\
    b_4^0 \\
    b_0^0
\end{bmatrix}, \hspace{1cm} (3.3)

$$

with approximate relative intensities as shown.

These three ZFPMR transitions are shown in Fig. 1. Their transition energies as a function of temperature are given in Table I. In order to determine the error in these transition frequencies, their positions have been measured as a function of modulation-field strength, in both resonant cavities and in traveling wave helices, and for $^{157}$Gd as well as natural Gd. Standard deviations in transition frequencies are given in Table I. In each case transition frequencies for the fairly broad lines were picked as the midpoint of the full width at half-height of the transition. For a given transition this value may be picked with a precision several times greater than the standard deviation for the line position.

2. Discussion

It is appropriate to compare the present measurement of the ZFS with that obtained by other workers. The results of this comparison are given in Table I. We compare our directly measured ZFS with that of Dagg, Kemp, and Symmonds\textsuperscript{25} for two temperatures. The two sets of numbers agree within experimental error, primarily due to the large error limits of the other workers.

Present results may also be compared with zero-field transition frequencies predicted by variable-field EPR. The predicted frequencies are given for the high-field measurements of Dagg et al., and for the low-field results of Gerkin and Thorssell.\textsuperscript{38} In the former case, in order to obtain the error limit of the predicted frequency, the square root of the sum of the squares of the error limits on $b_2^0$, $b_4^0$, and $b_0^0$, as obtained from ordinary EPR, are multiplied by the squares of the matrix coefficients in Eq. (3.3) to convert them to errors in transition energies. The two sets of numbers thus obtained agree within the 0.3% experimental error. The uncertainty in the ZFPMR measurements is only slightly less than the uncertainty in the conventional EPR results. This is due to the linewidth in this salt (~25 MHz) and it is not a general statement about the relative precision of ZFPMR. In another salt with sharp lines, the fact that frequencies may easily be determined to about 1 ppm with ZFPMR, whereas fields may be measured to 100 ppm only with difficulty, would demonstrate the higher precision of ZFPMR.

| TABLE I. Comparison of crystal field ($g = 0$ isotopes) transition energies for gadolinium-doped lanthanum ethyl sulfate at two temperatures. |
| --- | --- | --- | --- |
| | ZFPMR (measured)\textsuperscript{a} | ZFPMR (measured)\textsuperscript{b} | EPR (calc.)\textsuperscript{c} | Low-field EPR (calc.)\textsuperscript{d} |
| | (MHz) | (MHz) | (MHz) | (MHz) |
| 14.2 K | 1383.1 ± 2.1 | 3381.4 ± 9 | 3381.4 ± 2.8 | ... |
| T1 | 2503.5 ± 3.6 | 2500.3 ± 26 | 2502.7 ± 4.2 | ... |
| T2 | 1371.9 ± 3.9 | 1366.3 ± 9 | 1366.2 ± 3.9 | ... |
| T3 | 3446.6 ± 2.1 | 3444.3 ± 12 | 3444.0 ± 2.8 | 3442.5 ± 6 |
| T4 | 2550.4 ± 3.6 | 2555.4 ± 10.2 | 2549.1 ± 4.2 | 2547.9 ± 1.2 |
| T5 | 1396.8 ± 3.9 | 1390.1 ± 14.1 | 1391.0 ± 3.9 | 1390.1 ± 7.4 |

\textsuperscript{a}Error limits are taken as three standard deviations; hence $\sigma(T_1) = 0.7$ MHz, $\sigma(T_2) = 1.2$ MHz, and $\sigma(T_3) = 1.3$ MHz.\n
\textsuperscript{b}Error limits as given in their work (Ref. 35).\n
\textsuperscript{c}Obtained from zero-field measurements.\n
\textsuperscript{d}In this work there is an unexplained 26.1-MHz splitting detected between $\Delta M_z = 1$ and $\Delta M_z = -1$ transitions being a method of extrapolation from fields above 10 G. The frequency indicated here is the average of these two (Ref. 38).
The low-field results of Gerkin and Thorsell are somewhat more problematical. These workers observed a 26.1-MHz splitting between the $\Delta M_b = 1$ and the $\Delta M_b = -1$ transition energies for $T_3$ by extrapolation from measurements taken above 10 G. Results from our measurements do not confirm the existence of these splittings. This additional complication may be the reason that the predicted frequencies from low-field EPR lie just outside the error limits of our measurements. The error limits used by these workers of less than 1 MHz may be somewhat optimistic.

For the treatment of hyperfine structure presented below, crystal-field parameters appropriate to these zero-field measurements will be necessary. This determination is not possible in closed form since, in $C_{3v}$ symmetry, it is necessary to determine four crystal-field parameters with three zero-field transitions. This situation, however, becomes tractable because the $b_6$ coefficient is so small. If we use a typical value of $-4.25 \times 10^4$ cm$^{-1}$ as determined by Dagg et al.,$^{35}$ the coefficient $B_5^2 = 1.01 \times 10^{-5}$ MHz. The other three crystal-field coefficients may then be varied to fit exactly the three ZFS transitions. Neglect of the off-diagonal matrix elements connecting the $\pm \frac{3}{2}$ levels with the $\pm \frac{1}{2}$ levels via $\hat{D}^6_0$ leaves the best-fit crystal-field parameters invariant to about five significant figures. Indeed, second-order perturbation theory shows that this coefficient can lead to a position shift for the $\pm \frac{3}{2}$ or $\pm \frac{1}{2}$ energy levels of about 0.5 MHz, which is on the order of the standard deviation of transition frequencies. Considering the slow temperature dependence of $B_5^2 (-0.01 \text{kHz/K})$, the 77-K value is employed at both 77 and 4.2 K with little error. The best least-squares-fit crystal-field parameters are given in Table II.

The standard deviations in these parameters are derived from standard deviations in transition frequencies by taking the absolute value of the inverse of the matrix of coefficients in Eq. (3.3). The error is the square root of the sum of the squares of the terms obtained by multiplying this matrix by a column vector of standard deviations of transition frequencies. These are also given in Table II.

B. Natural gadolinium in various $Ln(C_2H_5SO_4)_3 \cdot 9D_2O$

1. Introduction and results

It has recently been pointed out by Gerkin and Thorsell$^{33}$ that crystals of $La(C_2H_5SO_4)_3 \cdot 9D_2O$ show a Gd$^{3+}$ linewidth up to a factor of 2 less than the corresponding hydrate. They therefore used such crystals in place of hydrated crystals in their low-field EPR studies of Gd$^{3+}$ in LES. It is not obvious from ordinary EPR or even low-field EPR, but it is immediately apparent from ZFPMR that the crystal-field parameters for the deuterate are different from those of the hydrate. This difference is immediately obvious using ZFPMR, since one measures the ZFS directly without recourse to computations based on variable-field data.

Inasmuch as the origin of crystal-field splitting of Gd$^{3+}$ is still an open question, it becomes important to pursue the nature of this deuteration effect on the crystal parameters. Accordingly, both
temperature and host (La, Sm, Er) ionic-radius dependence of crystal-field parameters for the deuterate were investigated.

The temperature dependence of ZFS in ethylsulfate hydrates has been discussed by several workers. Dagg, Kemp, and Symmons noted that ZFS is a maximum at about 130 K in LES. Between 77 and 4 K the ZFS is, within experimental error, linearly dependent on absolute temperature. The dependence on temperature from 290 to 77 K for several lanthanide ethylsulfates has also been studied by Gerkin and Thorsell.

To compare deuterate with hydrate in this regard we have studied the lanthanum salt at 77 and 4 K. The results are given in Tables III and IV in which deuterate data may be compared with the corresponding hydrate data. It is immediately obvious from Table IV that the temperature dependence of each is the same. This dependence probably indicates that the gross features of hydrogen bonding in each are quite similar.

ZFS and crystal-field parameters for several deuterated lanthanide ethylsulfates at 77 K have also been studied. The results may be found in Table V. The dependence of $b_2^0$ on the ionic radius of the host lanthanide ion at both 77 K and room temperature is linear. Figure 2 presents the dependence of $b_2^0$ on the radii for hydrate and deuterate. Both are seen to be linear, with best weighted-least-squares slopes of 288.9×10$^{-4}$ cm$^{-1}$ Å$^{-1}$ for the hydrate and 295.6×10$^{-4}$ cm$^{-1}$ Å$^{-1}$ for the deuterate, at 77 K. Considering the errors in $b_2^0$ values and the fact that only three points were used to determine the slope, we judge these two slopes to agree within experimental error.

2. Discussion

It has been shown that within experimental error the deuterate and hydrate crystal fields depend in the same way on temperature and host lanthanide ion radius. We wish to inquire, then, what causes the deuterate to have a 4% stronger quadrupolar crystal-field parameter $b_2^0$. The deuteron differs from the proton in three fundamental ways. First,

### Table II. Best least-squares-fit crystal-field spin-Hamiltonian parameters for $^{16}$Gd$^{3+}$-doped LES at two temperatures, using $b_{2}^{0} = -4.25 \times 10^{-4}$ cm$^{-1}$.

<table>
<thead>
<tr>
<th>Coefficient</th>
<th>Temp. (K)</th>
<th>Value (MHz)</th>
<th>Std. dev. (MHz)$^a$</th>
<th>Value (10$^4$ cm$^{-1}$)</th>
<th>Std. dev. (10$^4$ cm$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$b_2^0$</td>
<td>77</td>
<td>204.418</td>
<td>0.083</td>
<td>6.187</td>
<td>0.027</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>200.665</td>
<td>0.083</td>
<td>66.936</td>
<td>0.027</td>
</tr>
<tr>
<td>$b_4^0$</td>
<td>77</td>
<td>-0.2028</td>
<td>0.0013</td>
<td>-6.765×10$^{-2}$</td>
<td>0.043×10$^{-2}$</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>-0.1996</td>
<td>0.0013</td>
<td>-6.656×10$^{-2}$</td>
<td>0.043×10$^{-2}$</td>
</tr>
<tr>
<td>$b_6^0$</td>
<td>77</td>
<td>1.376×10$^{-3}$</td>
<td>0.072×10$^{-3}$</td>
<td>4.59×10$^{-4}$</td>
<td>0.24×10$^{-4}$</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.373×10$^{-3}$</td>
<td>0.072×10$^{-3}$</td>
<td>4.58×10$^{-4}$</td>
<td>0.24×10$^{-4}$</td>
</tr>
<tr>
<td>$b_2^0$</td>
<td>77</td>
<td>613.25</td>
<td>0.25</td>
<td>204.560</td>
<td>0.083</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>602.01</td>
<td>0.25</td>
<td>200.807</td>
<td>0.083</td>
</tr>
<tr>
<td>$b_4^0$</td>
<td>77</td>
<td>-12.168</td>
<td>0.077</td>
<td>-4.059</td>
<td>0.026</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>-11.973</td>
<td>0.077</td>
<td>-3.994</td>
<td>0.026</td>
</tr>
<tr>
<td>$b_6^0$</td>
<td>77</td>
<td>1.734</td>
<td>0.091</td>
<td>0.578</td>
<td>0.030</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.730</td>
<td>0.091</td>
<td>0.577</td>
<td>0.030</td>
</tr>
<tr>
<td>rms error of least-square fit</td>
<td>77</td>
<td>6.0×10$^{-4}$</td>
<td>(\ldots)</td>
<td>(\ldots)</td>
<td>(\ldots)</td>
</tr>
</tbody>
</table>

$^a$Standard deviations are taken to be the same at both temperatures.

### Table III. Crystal-field parameters and ZFS for LES hydrate and LES deuterate at two temperatures. In each case $b_2^0$ was taken as 0. The error limits are three standard deviations.

<table>
<thead>
<tr>
<th>Temp. (K)</th>
<th>Hydrate</th>
<th>Deuterate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_1$ (MHz)</td>
<td>77</td>
<td>3446.6±2.1</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>3383.0±2.1</td>
</tr>
<tr>
<td>$T_2$ (MHz)</td>
<td>77</td>
<td>2550.4±3.6</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>2503.5±3.6</td>
</tr>
<tr>
<td>$T_3$ (MHz)</td>
<td>77</td>
<td>1396.8±3.9</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1371.9±3.9</td>
</tr>
<tr>
<td>$b_2^0$ (10$^{-4}$ cm$^{-1}$)</td>
<td>77</td>
<td>204.56±0.15</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>200.81±0.15</td>
</tr>
<tr>
<td>$b_4^0$ (10$^{-4}$ cm$^{-1}$)</td>
<td>77</td>
<td>-4.06±0.05</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>-3.99±0.05</td>
</tr>
<tr>
<td>$b_6^0$ (10$^{-4}$ cm$^{-1}$)</td>
<td>77</td>
<td>0.58±0.06</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.58±0.06</td>
</tr>
</tbody>
</table>
the magnetic moment of the deuteron is a factor of 3.25 times smaller than the proton. Second, the deuteron is a spin-one particle and thus has a small quadrupole moment. Third, the deuteron mass is a factor of 2 greater than that of the proton. Considering the smallness of the quadrupole moment and the electrostatic nature of crystal-field theory, a mechanism based on the first two differences would be unlikely. The third difference, however, leads not only to a change in vibrational frequency, but also to a change in the potential in which the hydrogen-bonded proton or deuteron vibrates (see, for example, Hamilton and Ibers42). The hydrogen bond length or angle could change.

Protons or deuterons on the waters of hydration are almost surely hydrogen bonded to oxygens on the sulfate groups. While evidence is not conclusive, it is usually suggested that electric-dipole moments of the water molecules are aligned with negative end toward the lanthanide ion. Hence the nearest neighbors of the central Ln³⁺ ion are oxygens, with hydrogens pointing toward the negative sulfate groups. A lengthening of the hydrogen bond on deuteration might cause not only a change in the lattice constant but also a change in the metal-oxygen distance. Indeed, this system is particularly interesting from the point of view of crystal-field theory since the ligands are neutral. Such a rearrangement in the hydrogen bonding on deuteration would produce a change in the static crystal field. If this small change in structure could be observed experimentally it might then be possible to correlate it with the observed change in crystal-field parameters upon deuteration, if the Gd-O distance does in fact change, this

| TABLE IV. Temperature dependence of crystal-field parameters and ZFS for LEShydrate and LESdeuterate in the region from 77 to 4 K. All dependences have been assumed linear. The error limits given are about three standard deviations. |
|---|---|---|---|---|---|
| | Hydrate | Deuterate |
| \( \frac{dT_z}{dT} \) (MHz K⁻¹) | 0.87 ± 0.06 | 0.92 ± 0.11 |
| \( \frac{dT_x}{dT} \) (MHz K⁻¹) | 0.64 ± 0.10 | 0.65 ± 0.11 |
| \( \frac{dT_y}{dT} \) (MHz K⁻¹) | 0.34 ± 0.11 | 0.38 ± 0.27 |
| \( \frac{db_2}{dT} \) (10⁻³ cm⁻¹ K⁻¹) | 0.051 ± 0.004 | 0.053 ± 0.007 |
| \( \frac{db_4}{dT} \) (10⁻³ cm⁻¹ K⁻¹) | -9.59 ± 1.4 × 10⁻⁵ | ± 2.8 × 10⁻⁵ |
| \( \frac{db_6}{dT} \) (10⁻³ cm⁻¹ K⁻¹) | ± 1.6 ± 10⁻⁵ | ± 3.6 ± 10⁻⁵ |

FIG. 2. Quadrupolar-crystal-field coefficient \( b_2^z \) for three Gd³⁺-doped lanthanide ethyl sulfates vs 4f-electron radii. The open circles are for \( \text{La(C}_2\text{H}_5\text{SO}_4)_3 \cdot 9\text{D}_2\text{O} \). The average 4f radii are those estimated by Templeton and Dauben (Ref. 41), and correspond to Er³⁺, Sm³⁺, and La³⁺.

The crystal-field parameters are from Table V. The experimental error is smaller than the circle size except in the case of the erbium deuterate.

| TABLE V. Crystal parameters and ZFS for three lanthanide ethylsulfate hydrates and deuterates at 77 K with their standard deviations. The calculations assume \( b_2^z = 0 \). The hydrate data are from Ref. 37. |
|---|---|---|---|---|---|---|---|
| | Lanthanum ethylsulfate | Samarium ethylsulfate | Erbium ethylsulfate |
| | Hydrate | Deuterate | Hydrate | Deuterate | Hydrate | Deuterate |
| \( T_1 \) (MHz) | 3440.7 ± 1.9 | 3598.0 ± 3 | 2525.7 ± 2.6 | 3080.5 ± 2 | 2515.1 ± 6.1 | 2685 ± 25 |
| \( T_2 \) (MHz) | 2547.5 ± 1.4 | 2845 ± 3 | 2202.4 ± 1.7 | 2301 ± 2 | 1927.9 ± 4.4 | 2055 ± 25 |
| \( T_3 \) (MHz) | 1390.2 ± 1.2 | 1440.5 ± 10 | 1218.7 ± 1.5 | 1231.7 ± 5 | 1084.4 ± 3.8 | 1075 ± 50 |
| \( b_2^z \) (10⁻³ cm⁻¹) | 204.17 | 212.64 | 175.49 | 185.18 | 152.69 | 161.26 |
| ± 0.08 | ± 0.24 | ± 0.10 | ± 0.15 | ± 0.24 | ± 1.45 |
| \( b_4^z \) (10⁻³ cm⁻¹) | -4.02 | -3.95 | -4.01 | -3.60 | -4.03 | -3.48 |
| ± 0.02 | ± 0.10 | ± 0.03 | ± 0.05 | ± 0.07 | ± 0.55 |
| \( b_6^z \) (10⁻³ cm⁻¹) | 0.51 | 0.56 | 0.53 | 0.088 | 0.57 | -0.40 |
| ± 0.02 | ± 0.13 | ± 0.02 | ± 0.07 | ± 0.06 | ± 0.65 |
would have important consequences for diffraction crystallography. It is customary in x-ray or neutron-diffraction studies to hold the metal-oxygen distance constant when simultaneously refining the structure for protonated and deuterated species.\textsuperscript{43,44} By this simple mechanism, the effect of deuteration might be the same as a change in lattice size with temperature or hydrostatic pressure.\textsuperscript{45,46}

Even if the Gd-O distance does not change, it is possible that the positions of the deuterons do. It is known that the water hydrogens are close enough to the gadolinium ion to allow dynamic nuclear-polarization experiments to be performed.\textsuperscript{36} In \textit{ab initio} calculation of crystal-field parameters by lattice summations, results are quite sensitive to how the water dipole moment is treated.\textsuperscript{47} Generally, it is accommodated by assuming that the water molecule is composed of fractional charges placed so the entire molecule is neutral. Clearly, a change in the water bond angle would cause $b_0^2$ to change. Less-accurate calculations include the nearest-neighbor oxygens only, predicting no effect for a change in hydrogen-bond orientation.

If equilibrium positions of neither oxygens nor hydrogens change when the structure for the hydrate and deuterate are refined independently, one might have to resort to a dynamic crystal-field theory to account for the difference by considering a change in the oxygen-hydrogen vibration frequency.

Measurement of the dependence of the nuclear-quadrupole-coupling parameter on deuteration, host lattice, and hydrostatic pressure would yield information on the contribution of the lattice or crystal electric-field gradient $g_e$ to the total field gradient. A change due to these effects would be difficult to see for the deuterate in LES because its magnitude would be about the same size as the presently available standard deviation of $P$, due to line width.

### IV. HYPERFINE SPECTRA OF 155\textsuperscript{Gd} AND 157\textsuperscript{Gd}

#### A. Results and analysis

Hyperfine structure was not resolved in the naturally occurring gadolinium spectra. In order to determine whether it was really absent, or measure its magnitude if present, isotopically enriched crystals of 0.1-mole\% concentration were prepared. Since $g$ is isotropic to five decimal places,\textsuperscript{316} the hyperfine-structure Hamiltonian in axial symmetry may be represented as\textsuperscript{13}

$$\mathcal{H}_{M_z} = -A\hat{S}_z\hat{I}_z + \frac{1}{2} A(\hat{S}\cdot\hat{I} + \hat{S}\cdot\hat{I}) + \frac{1}{2}(I^2 - \frac{1}{2}I(I+1)), \quad (4.1)$$

in which $A$ is the isotropic hyperfine constant, and $P$ is the nuclear-electric-quadrupole constant, and $\hat{I}$ is the nuclear-spin operator. For these isotopes $I = \frac{3}{2}$ and as before $\hat{S} = \frac{1}{2}$. The Hamiltonian, therefore, is a $32 \times 32$ matrix when represented in the complete set of high-field states, $|M_z M_J\rangle = |M_J\rangle |M_z\rangle$ quantized along the three-fold crystal symmetry axis.

In order to obtain an approximate analysis of the zero-field spectra, it would be very useful to have approximate analytical expressions for the zero-field energy levels. It was shown above that the off-diagonal term in the crystal field is quite small, so it will be temporarily neglected. Zero-field states can then be treated as pure high-field states $|M_z M_J\rangle$. For convenience, elements of diagonal in the hyperfine operator if $M_J \neq \pm \frac{3}{2}$ will also be neglected. This approximation is worse than the neglect of off-diagonal crystal-field terms, but it will be adequate for an approximate set of hyperfine constants from which to initiate a least-squares analysis of the exact diagonalization of the full $32 \times 32$ matrix. These off-diagonal elements cause shifts on the order of 3.5 MHz. Under these two approximations the matrix breaks up into submatrices of size no greater than $2 \times 2$. The energy-level formulas are given in Table VI. Using $\Delta M_J = \pm 1, \Delta M_z = 0$ selection rules one may then calculate the spectrum. A 15-line spectrum is predicted, with crystal-field transition $T_1$ splitting.
TABLE VII. Hyperfine transitions for \( S = \frac{3}{2}, I = \frac{3}{2} \) according to a \( \Delta M_g = \pm 1, \Delta M_I = 0 \) selection rule. The energy level formulas are from Table VI. The first subscript of a \( T_n \) indicates from which crystal-field transition it is derived.

<table>
<thead>
<tr>
<th>( T_n )</th>
<th>( T_{11} = E_{11} - E_{12} - T_1 + \frac{1}{2} A )</th>
<th>( T_{12} = E_{12} - E_{11} + T_1 + \frac{1}{2} A )</th>
<th>( T_{13} = E_{13} - E_{11} - T_1 - \frac{1}{2} A )</th>
<th>( T_{14} = E_{14} - E_{11} + T_1 - \frac{1}{2} A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( T_2 )</td>
<td>( T_{31} = E_{11} - E_{31} - T_3 + \frac{1}{2} A )</td>
<td>( T_{32} = E_{12} - E_{32} + T_3 + \frac{1}{2} A )</td>
<td>( T_{33} = E_{23} - E_{33} - T_3 - \frac{1}{2} A )</td>
<td>( T_{34} = E_{34} - E_{33} + T_3 - \frac{1}{2} A )</td>
</tr>
<tr>
<td>( T_3 )</td>
<td>( T_{35} = E_{33} - E_{32} = T_3 + \frac{1}{2} A )</td>
<td>( T_{36} = E_{33} - E_{32} = T_3 + \frac{1}{2} A )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( T_4 )</td>
<td>( T_{37} = E_{33} - E_{32} = T_3 + \frac{1}{2} A )</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

into four lines numbered \( T_{11} \) to \( T_{14} \), transition \( T_2 \) splitting into four lines numbered \( T_{21} \) to \( T_{24} \), and transition \( T_3 \) splitting into seven lines numbered \( T_{31} \) to \( T_{37} \). The transition energies are given in Table VII.

Of course, the ordering of these transitions in energy will depend on the signs and magnitudes of the parameters. One may easily see that \( T_1 \) and \( T_2 \) are each split by the same amount and consist of four equally spaced lines, each with a total spread of \( 3A \). Their positions are independent of \( P \) to this order of approximation and the spectrum is invariant to the sign of \( A \). It is therefore necessary to look to the \( T_3 \) transition splitting to determine \( P \) and the sign of \( A \) and \( P \). Note that the splitting are asymmetrical about \( T_3 \) and hence the absolute and relative signs of \( A \) and \( P \) may be determined (see Table VII). As previously mentioned, Bleaney et al. \(^{25} \) were able to determine the absolute signs of the crystal-field parameters; absolute signs are thereby obtained for all diagonal parameters in the spin-Hamiltonian \( (A, P, B^2) \).

Figure 3 presents spectra for \( ^{157}\text{Gd} \)-doped LES at 4 K. It is obvious that the hyperfine splitting is not resolved for \( T_1 \) and \( T_2 \). There are, however, five lines present for \( T_3 \). In order to assign the lines, the approximate expressions of Table VII are needed. A series of plots of these expressions for \( T_{3n} \) can be computed for a wide range of values of \( A \) and \( P \). Each such plot is for \( P \) ranging from \(-100 \) to \( 100 \) MHz for constant \( A \), such as is illustrated in Fig. 4 for \( A = 18 \) MHz. On examining such plots for \( A \) ranging from \(-25 \) to \( 25 \) MHz, only one set of approximate values fits the observed spectra for each isotope and different modulation strengths. The assignment, then, of the five-line spectrum is as follows. The broad center line is assumed to consist of three unresolved transitions. The remaining four lines are assigned as \( T_{35}, T_{37}, T_{31}, \) and \( T_{33} \) in order of increasing frequency. The transition frequencies for the four lines which are invariant to modulation-field intensity, are given in Table VIII.

Least-squares best-fit parameters are calculated by numerical diagonalization of the \( 32 \times 32 \) Hamiltonian matrix with all off-diagonal elements included.
The best-fit crystal-field parameters from $^{160}$Gd, as determined in the previous Sec. III, are employed. Any possible isotopic dependence of the crystal-field parameters (for Gd isotope) is thereby neglected. Such a dependence has been proposed by Marshall for Gd$^{3+}$ in charge-compensated ThO$_2$, but the difference between the $I = 0$ isotope and the odd isotopes is just larger than his experimental error. Since our lines are much broader than those in ThO$_2$, any such effects are unlikely to be observed in this system.

The rms error in the fit is defined as $\delta$ and is also given in Table VIII. After the least-squares routine has converged on a minimum $\delta$, there are various sets of local minima which give approximately the same minimum value, due to roundoff error in the computation. A final value for $A$ and $P$ may then be taken as the average of those belonging to the local minima. The standard deviation of these local minima average of $\delta$, $A$, and $P$ is defined as $\sigma[\delta]$, $\sigma[A]$, and $\sigma[P]$ respectively. Values of $A$ and $P$ so calculated are given in Table IX. Predicted transition frequencies from them and the values of $\delta$ and the $\sigma[\delta]$s are given in Table VIII. Note that in three of the four cases, the rms error in calculated frequency for each transition is greater than the standard deviation of frequency measurements (2 MHz at 77 K, 1.2 MHz at 4 K). Note also that the observed frequencies at 77 K are always higher than the calculated ones. One could reduce $\delta$ for these cases by lowering the value of $T_2$; however, there is no justification for changing experimental values in this manner, so the small systematic error in calculated frequencies must simply be accepted.

In order to calculate the standard deviations in parameters $A$ and $P$, local linear approximations to the derivatives of the transition frequencies with respect to $A$ and $P$ are used. These standard deviations in parameters $\sigma[A]$ and $\sigma[P]$ are propagated directly from the standard deviations in observed transition frequencies. The standard deviations thus calculated appear in Table IX. For some calculations involving both $A$ and $P$ it may be necessary to calculate the error in a resultant function. This computation requires the covariance of $A$ and $P$ as well as their standard deviations, and this covariance is estimated to be $1.56 \times 10^{-1}$ MHz at 77 K and $1.26 \times 10^{-1}$ MHz at 4 K. A description of the statistical procedures may be found in a recent monograph, $^{49}$ Of course $\sigma[A]$ and $\sigma[P]$ are much smaller than $\sigma[A]$ and $\sigma[P]$, as would be expected. It should be emphasized that the standard deviations quoted here are indicative of the linewidth of Gd$^{3+}$ in this salt, rather than indicative of the accuracy of ZFPMR.

Having determined least-squares best-fit hyperfine constants, it is of interest to determine, in a qualitative way, the fit of the relative intensities. This is a complicated problem in ZFPMR because of the influence of the modulation magnetic field.

Since the modulation magnetic field may vary over the sample volume, the line-shape parameters for the low-field spectrum may be different than for the zero-field spectrum. While it has been demonstrated that such factors play little or no role in the determination of zero-field transition frequencies, $^{15}$ the over-all shape and appearance of the

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**TABLE VIII.** Transition frequencies and other parameters for gadolinium-doped lanthanum ethylsulfate at two temperatures for best least-squares-fit hyperfine parameters.

<table>
<thead>
<tr>
<th>Parameter (MHz)</th>
<th>$^{152}$Gd</th>
<th>$^{153}$Gd</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{31}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>obs.</td>
<td>1225.0, 1222.5</td>
<td>1206.8, 1208.0, 1208.0</td>
</tr>
<tr>
<td>calc.</td>
<td>1230.2, 1315.0</td>
<td>1334.6, 1332.0</td>
</tr>
<tr>
<td>$T_{37}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>obs.</td>
<td>1465.9, 1460.6</td>
<td>1451.2, 1448.5</td>
</tr>
<tr>
<td>calc.</td>
<td>1457.9, 1552.0</td>
<td>1532.5, 1534.0</td>
</tr>
<tr>
<td>$T_{33}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>obs.</td>
<td>4.3, 1465.1</td>
<td>4.9, 1465.1</td>
</tr>
<tr>
<td>calc.</td>
<td>4.3, 1465.1</td>
<td>4.9, 1465.1</td>
</tr>
<tr>
<td>$\sigma[\delta]$</td>
<td>$5.2 \times 10^{-3}$</td>
<td>$9.9 \times 10^{-3}$</td>
</tr>
<tr>
<td>$\sigma[A]$</td>
<td>$5.1 \times 10^{-2}$</td>
<td>$7.7 \times 10^{-2}$</td>
</tr>
<tr>
<td>$\sigma[P]$</td>
<td>$1.4 \times 10^{-1}$</td>
<td>$4.1 \times 10^{-1}$</td>
</tr>
</tbody>
</table>

*The standard deviation of transition frequencies is taken to be 2.0 MHz at 77 K, and 1.2 MHz at 4 K.
*The standard deviation of the rms error of the fit.
*The standard deviation of the rms error of the fit.
*The standard deviation of the A values described in footnote c. The best fit A is the mean of these A values.
*The standard deviation of the various P values described in footnote c. The best fit P is the mean of these P values.
TABLE IX. Best-fit hyperfine constants at two temperatures. For definition of symbols, see text.

| Isotope | Temp. (K) | A (MHz) | Δ(A) | P (MHz) | Δ(P) | Units
|---------|-----------|---------|------|---------|------|--------
| 154Gd  | 77        | 18.26   | 0.37 | -52.86  | 1.26 | MHz
| 154Gd  | 4         | 18.10   | 0.22 | -53.09  | 0.76 | MHz
| 154Gd  | 77        | 14.59   | 0.37 | -48.01  | 1.26 | MHz
| 154Gd  | 4         | 14.33   | 0.22 | -51.16  | 0.76 | MHz
| 157Gd  | 77        | 6.09    | 0.12 | -17.63  | 0.42 | 10^4 cm^{-1}
| 157Gd  | 4         | 6.034   | 0.07 | -17.71  | 0.25 | 10^4 cm^{-1}
| 157Gd  | 77        | 4.86    | 0.12 | -16.01  | 0.42 | 10^4 cm^{-1}
| 157Gd  | 4         | 4.78    | 0.07 | -17.06  | 0.25 | 10^4 cm^{-1}
| 157Gd  | 77        | 6.55    | 0.13 | -18.96  | 0.45 | G
| 157Gd  | 4         | 6.49    | 0.08 | -18.04  | 0.27 | G
| 155Gd  | 77        | 5.23    | 0.13 | -17.22  | 0.45 | G
| 155Gd  | 4         | 5.14    | 0.08 | -18.35  | 0.27 | G

*Based on an averaged standard deviation of 2 MHz for transition frequencies at 77 K, and 1.2 MHz at 4 K.

Conversion of MHz to G was accomplished using a g factor of 1.9919 (Ref. 39(a)).

spectra vary greatly with modulation-field strength. The most stringent test of the quality of calculated parameters is thus a complete synthesis of observed total spectrometer output. Therefore, many calculated spectra have been plotted assuming various effective modulation fields and various Gaussian linewidths for the zero- and low-field spectra. A typical result is shown in Fig. 5(a). While the fit is qualitatively correct in many respects, it is not an exact fit (compare with Fig. 3). Of course, spectra for low fields may depend on the angles between modulation field and crystal axes, and, as such, on whether single crystals were used. To illustrate the angular effect, Fig. 5(b) contains a spectrum with the modulation field perpendicular to the crystal axis. It is clear that for an exact computation of the spectrum would have to integrate over the entire sample volume to account for the field magnitude and angular variations.

Approximate expressions for transition energies indicated that T₁ and T₂ should each be split into four equally spaced lines with total spread 3A. The theoretical line-shape calculations show that observed spectra do not contradict this conclusion. Indeed, starting with the best fit A, one may calculate the spectrum shown in Fig. 5(c), after suitable choice of line shapes. The "doublet" spectrum arises from the large linewidth in this salt.

It is particularly interesting to examine a salt with sharper lines but still characterized by a similar Hamiltonian. The expected zero-field spectrum for Bi₂Mg₂(NO₃)₁₂·24H₂O, bismuth magnesium double-nitrate (BMDN), is very similar, even though the symmetry is approximately C₃v (exactly C₃v) rather than C₃v. The corresponding hyperfine split T₂ transition is shown in Fig. 6. Total spread in the spectrum is consistent with recent ENDOR measurement of the hyperfine constants of this sample by Butti et al. The T₂ transition, from which a measurement of the quadrupole parameter P could be made, lies at about 750 MHz. The recent availability of crystal-field parameters for a variety of gadolinium-doped lanthanide double nitrates (LMND) makes it particularly interesting to determine the series varia-

FIG. 5. Computer-synthesized ZFPMR spectra of Gd³⁺-doped La(C₂H₆SO₄)₃·9H₂O at 4 K, using the best least-squares fit spin-Hamiltonian parameters. Net absorption in arbitrary units is shown with net zero-field absorption directed up and net low-field absorption directed down. Effective conditions were chosen to give qualitative agreement with line intensity and shape. (a) T₁ₘ transitions at an effective modulation field of 5 G directed parallel to the crystal symmetry axis. The effective zero-field linewidth is 25 G and the effective low-field linewidth is 60 G. Compare with Fig. 3(c). (b) Tₙₜ transitions at an effective modulation field of 10 G directed perpendicular to the crystal symmetry axis. The effective zero-field linewidth is 25 G and the effective low-field linewidth is 50 G. Compare with Fig. 3(c). (c) T₉ₕ transitions at an effective modulation field of 5 G directed parallel to the crystal symmetry axis. The effective zero-field linewidth is 20 G and the effective low-field linewidth is 40 G. Compare with Fig. 3(b).
bution, denoted $A_{\text{BDSLc}}$ in the notation of Sandars and Beck,\textsuperscript{56e} is due to mixing of higher electronic states into the ground state by spin-orbit and spin-spin interactions. This has the effect of destroying spherical symmetry for the half-filled and closed shells and thus some $f$ or other $I \neq 0$ electrons can appear at the origin. For the $^5S_{1/2}$ state of europium these authors find the spin-orbit operator dominant, $A_{\text{BDSLc}}$ is found to be opposite in sign to the experimental $A$ and "the Casimir effect" must be used to bring the calculated value of $A$ closer to the experimental one. In order to estimate the value of $A_{\text{Cas}}$, one must numerically solve the radial Dirac equation. Even by using the best relativistic wave functions for europium available at the time, they can reduce the discrepancy between $A_{\text{BDSLc}}$ and experiment by only one-half.

Fortunately, there is an experimental way to tell whether core polarization or relativistic effects give rise to $A$ for half-filled shell ions. One may define the so-called hyperfine anomaly $\Delta$ which is given by the equation

$$155A/157A = \left[ g_e(155)/g_e(157) \right] (1 + \Delta),$$

(4.2)

in which the $g_e$'s are the nuclear $g$ values for the indicated isotopes. $\Delta$ is essentially a measure of how the distribution of nuclear magnetization inside the nucleus differs for each isotope. This isotope effect on the nuclear-magnetization distribution is probed through interaction with $s$ electrons inside the nucleus. Our interest is not necessarily in nuclear physics, but rather in the fact that core polarization will lead to a hyperfine anomaly and relativistic effects will not.\textsuperscript{58}

Table X presents a survey of measurements of $A$ and calculated $\Delta$'s for Gd$^{3+}$ in a variety of hosts. Included is our measurement of $A$ in LES, which is the largest value for Gd$^{3+}$ in a diamagnetic host. The hyperfine-structure anomalies are calculated using a ratio of $g_e$ values of $0.7633 \pm 0.0035$ from electron-nucleus double resonance (ENDOR)\textsuperscript{59} in CeO$_2$. It is seen that $\Delta$ is zero within experimental error for diamagnetic hosts even for precise ENDOR determinations. Large errors in $\Delta$ are primarily due to error in the $g_e$ ratio; hence, ENDOR determinations show little improvement in accuracy over EPR determinations. Improvement in the accuracy of $\Delta$ must await a better measurement of the ratio of the $g_e$'s.

One can ask whether the upper bound on $\Delta$, set by its experimental error, is small enough to decide that core polarization is not playing a major role in the appearance of hyperfine structure. The hfs anomaly for Eu$^{3+}$ was determined by Baker and Williams\textsuperscript{60} to be $(-0.78 \pm 0.11)\%$, in a situation for which it has been suggested that core polarization accounts for about 80% of the hyperfine constant. One must therefore conclude that present error
limits on $\Delta$ are not as yet small enough for a meaningful
determination of the extent of core polarization in Gd$^{3+}$ to be made. Such a determination must
await more precise experiments to measure the various contributions to $A_i$, and must await accurate
unrestricted Hartree-Fock and relativistic wave functions to calculate these contributions from first
principles.

It is unfortunate that the ZFPMR experimental error is greater than the difference between the
measured $A$ values for 77 and 4 K. Theories have been developed for the temperature dependence of the
hyperfine constant for $S$-state ions, which yield a $2\%$ decrease in $A$ for Eu$^{3+}$ in CaF$_2$ on going
from 4 to 77 K, due to temperature-dependent configuration mixing by the orbit-lattice interaction.
ZFENDOR should allow precise enough measurements to determine temperature dependence for
lower than cubic symmetry by reducing the problem of broad lines in LES. Of course, this tempera-
ture dependence may be especially well studied in ThO$_2$ and BMDN for which magnetic resonance
lines are considerably sharper than in LES.

2. Quadrupole Interaction

The electric-quadrupole term in the Hamiltonian represents the energy of interaction of the electric-
field gradient (EFG) at the nucleus with the nuclear-electric-quadrupole moment and is written as

$$e_q = e_q (1 - \gamma_e) + e_q (1 - R_q),$$

in which $e_q$ is the EFG due to the external crystal field about the ion, $e_q$ is the EFG due to the val-
ence-shell electrons, and $e$ is the charge on the electron. The two parameters $\gamma_e$ and $R_q$ are
Sternheimer antisheilding factors which account for the effect of distortion of the closed shells of
electrons by the gradients. They may have either sign and hence may reverse the direction of the field
gradient.

Since $e_q$ is proportional to the quadrupolar crystal-field parameter $A_q^0$, it must vanish in cubic
symmetry. (A discussion of the relation between $A^I$'s and $B^I$'s can be found in Ref. 40.) Many of the host in Table X are cubic, and hence observation of a small quadrupole term in their EPR would be due to the second part of Eq. (4.3). However, in the nonrelativistic limit of LS coupling, the second term in Eq. (4.3) would also vanish due to the spherical symmetry of the half-filled shell. We know of only one case for which a quadrupolar-interaction term has been measured for Gd$^{3+}$ in cubic symmetry; namely, the ENDOR investigation in ThO$_2$ by Hurrell, 64 He was able to obtain spectra for the $^{157}$Gd$^{3+}$ isotope only and measured the quadrupole constant for the appropriate cubic spin Hamiltonian $B = -0.687 \pm 0.18$ MHz. The other two rare-earth S-state ions have somewhat larger constants. 65

Evans, Sandars, and Woodgate 66 have calculated the various contributions to the quadrupole constant for the S-state atoms, $^{55}$Mn, and both odd europium isotopes. They ascribe nonzero contributions to the quadrupole interaction to a breakdown of LS coupling, and to a breakdown of the nonrelativistic approximation. Using relativistic wave functions it is possible to calculate a contribution called the Casimir effect indicated by $B_{\text{Cas,}}$, which in both cases was found to dominate the breakdown of LS coupling.

Since the quadrupole interaction is so small in cubic symmetry for which the first part in Eq. (4.3) vanishes, it may be expected that the first term will dominate the second in symmetry lower than cubic. There are only three cases for which this interaction has been measured. Using a standard quadrupole term of the form $P[J_x^2 \frac{1}{2}(J_z^2) - J_x^2]$, Danner, Ranon, and Stamires 67 and Rannon and Stamires 68 were able to measure $|^{157}P| = 53.7 \pm 0.3$ MHz and $|^{155}P| = 50.4 \pm 0.3$ MHz in YPO$_4$ by ordinary EPR. It is interesting to note that the ZFPMR studies of Kahle et al., 66 did not reveal the existence of hyperfine structure in the spectrum of this salt. Using high-field ENDOR, Butti et al., 54 measured $^{157}P = 26.527 \pm 0.003$ MHz and $^{155}P = 24.905 \pm 0.002$ MHz in BMDN. For convenience, the value obtained in this work is restated: $^{157}P = -53.09 \pm 0.76$ MHz and $^{155}P = -51.16 \pm 0.76$ MHz in LES. The relatively large reported error (3o) is mostly determined by the linewidth of Gd$^{3+}$ transitions in LES.

$P$ may be written as $3Qe^2q/4l(2l-1)$. Hence, a ratio of $P$ values for two isotopes should equal the ratio of quadrupole moments for each, if they have the same spin and are positioned in the same EFG. Thus one can compare the ratio of $P$'s with the ratio of nuclear-electric-quadrupole moments, $R(Q) = \frac{|^{157}Q|}{|^{155}Q|}$. There are four measurements of $R(Q)$. They are the optical determination by Kaliteevskii 69 of $1.28 \pm 0.02$, the optical determination by Speck 70 of $0.80 \pm 0.02$, the Mössbauer determination by Prange 71 of $0.78 \pm 0.06$, and the accurate atomic beam measurement by Unsworth 72 of $1.06534 \pm 0.00003$.

The ratio of $P$ values, $R(P) = \frac{|^{157}P|}{|^{155}P|}$, from above are $1.065 \pm 0.012$ for YPO$_4$, $1.06513 \pm 0.00021$ for BMDN, and $1.038 \pm 0.030$ for LES. The optical determinations are mutually inconsistent. Even though one of them is consistent with the Mössbauer value, all three are inconsistent with the accurate beam value. Moreover, all three $R(P)$'s agree with the beam measurement within experimental error.

It is interesting to note that except possibly for sign, the $P$ values for YPO$_4$ and LES agree with each other within experimental error. If $eQ_z$ were the only gradient contributing to the quadrupole term, then the $A_0^2$ values for each salt would have to be the same if $\gamma_0$ were the same for both. In fact, estimates lead one to conclude that this is roughly correct with $90 \text{ cm}^{-1} \approx |A_0^2| |Q_z| \approx 100 \text{ cm}^{-1}$ for both YPO$_4$ and LES. 73 While $b_0^2$ for these systems are quite different [$b_0^2(YPO_4) \approx 3, 5 b_0^2(\text{LES})$] it must be remembered that $A_0^2$ is the important crystal-field parameter for the quadrupole-interaction constant. It can be demonstrated that the relationship between these two parameters is neither direct nor simple. Indeed, higher-order and indirect terms may invalidate lattice summations and even simple decomposition of mechanisms; it is perhaps possible that in low-symmetry (noncubic) crystals an apparent $q_z$ is enhanced. This latter point would most simply explain the fact that $|P(YPO_4)| = |P(\text{LES})|$ for Gd$^{3+}$. It would be particularly useful if ZFPMR could be employed to obtain a sign for $P$ in YPO$_4$.

Finally, it is appropriate to inquire whether the quadrupole coefficients for LES are of about the right magnitude in an absolute sense. The EFG at the La$^{3+}$ nucleus of diamagnetic LES has been measured by NMR and, based on this determination, a procedure has been developed for estimating the ionic contribution to $P$ for any lanthanide ion in LES. 74 Also, the lattice contribution to $P$, $P_{\text{latt.}}$, for $^{155}$Gd$^{3+}$ in neodymium ethylsulfate (NdES) has been measured by nuclear-alignment techniques. 75 Block and Shirley 76 find $P_{\text{latt.}}$ for $^{155}$Gd$^{3+}$ to be $-43.2 \pm 1.5$ MHz for $^{155}$Gd in NdES using a quadrupole moment of $1.56$ b. Since this isotope has the same spin as those of interest here, one may predict $^{155}P$ equal to $-44.0 \pm 7.3$ MHz using $^{155}Q = 1.59 \pm 0.16$ b. 76 Considering the approximations in the above calculation and the fact that the host lanthanide ion is different, observed and predicted magnitude and absolute sign of $P$ are judged to be in agreement.

It is possible to evaluate $P$ in another way. If $A_0^2$ is a quadrupolar-lattice sum, one can write (using the notation of Edmonds).
\[ P_{\text{int.}} = 3QA_{24}^{0}G_{Ph}/(2l(2l-1)), \] (4.4)

in which \( \gamma_{y} = 1 - \gamma_{x} \) and is assumed to be \( \pm 21.5 \) for LES. The negative sign is usually chosen to agree with antishielding calculations. Experimental crystal-field parameters from optical experiments are defined as \( V_{2}^{0} = \gamma_{x}A_{24}^{0}(\langle r^{2} \rangle) \). \( \langle r^{2} \rangle \) is the expectation value of the radius of the open-shell 4\( f \) electrons. Edmonds’ \( \gamma_{y}A_{24}^{0} \) is equal to Freeman and Watson’s \( A_{24}^{0} \). Powell and Orbach\(^{77}\) obtain \( V_{2}^{0} = 100.6 \text{ cm}^{-1} \) and Freeman and Watson’s best Hartree-Fock estimate of \( \langle r^{2} \rangle \) is \( 0.785a_{0}^{2} \). \( \gamma_{x}A_{24}^{0} \) for Gd then becomes \( 128a_{0}^{2} \text{ cm}^{-1} \) (\( a_{0} \) is the Bohr radius). Finally, Edmonds estimates \( \gamma_{y}(\text{La})/\gamma_{x}(\text{La}) \) equal to \(-550 \), which yields \( A_{24}^{0}y_{y}(\text{Gd}) = -550A_{24}^{0}y_{x}(\text{Gd}) = (-7.04 \times 10^{4})a_{0}^{2} \text{ cm}^{-1} \), with the measurement of \( 155\text{Q} \)
\begin{equation}
\text{linewidth} = 1.59 \times 10^{4} \text{ MHz}.
\end{equation}

This estimate is just over a factor of 2 from our measurements of \( 155\text{Q} \). Allowing for generous error limits at each step in the above procedure, and allowing for the fact that no account of \( q_{5} \) has been made, once again the measured value and sign of \( P \) appear reasonable. For gadolinium in crystals of less than cubic symmetry, the hyperfine spectrum may be expected to be dominated by the quadrupole interaction, in contrast to the more usual situation, for which \( P \) is much smaller than \( A \). This quadrupole interaction is composed almost exclusively from lattice contributions.

V. LINESWIDTHS

Linewidths for Gd-doped LES are essentially the same for ZFPR transitions as EPR transitions (\(-20-30 \text{ MHz}\)). Thus, inhomogeneity in the external field contributes in only a small way to the linewidth. Also, since the linewidth is largely independent of temperature and the strength of the external field, one may rule out broadening by spin-lattice relaxation as the cause of broad lines. Indeed, the observed linewidth of 25 MHz sets a 12.7 nsec lower bound on the relaxation times. The spin-lattice relaxation time has been investigated by several workers\(^{78}\) and found to be four or five orders of magnitude slower than this lower limit.

The linewidth of gadolinium in various hosts is known to depend on concentration, at concentrations above about 0.1 mole\(^{\circ} \). \(^{79,80}\)

The residual linewidth in LES is caused by inhomogeneous broadening of the resonance line; this broadening is much greater than in other hosts. One can presume it to result from two factors; namely, the statistical variation of the crystal-field parameters due to lattice strains and dipolar interaction with the neighboring nuclear moments.

On this basis, then, one might be willing to suggest that the broadening in gadolinium-doped LES must be entirely due to dipolar broadening. This is, however, not the case, as Nd-doped LES shows a linewidth of 8-10 MHz for many of its lines.\(^{4,10}\)

Since transitions are not observed between crystal-field levels in Nd (the first low-lying crystal-field level is 100 \text{ cm}^{-1} above the ground state), inhomogeneities in the crystal potential must play a greatly reduced role in the Nd line broadening. Hence a major part of the linewidth for gadolinium must arise from variation in the crystal field. If relaxation times contributed significantly to the linewidth, one would expect Nd to have wider lines than Gd because the relaxation time of the former is considerably faster.

The effect of the surrounding nuclear magnetic moments associated with water of hydration in LES can be seen in several ways. The linewidth of Gd-doped BMDN is on the order of 10 MHz. This comparative sharpness can be explained by the fact that there are only six neighboring nitrogens in BMDN, but there are 18 neighboring protons in LES. Additionally, the magnetic moment of \( ^{14}\text{N} \) is seven times smaller than that of \( ^{1}\text{H} \). The protons of the water molecules are tightly coupled to the gadolinium ion, since one may observe their dynamic nuclear orientation.\(^{36}\) Sharpening of the spectrum on deuteration is accounted for by the magnetic moment of the deuteron being 3.25 times smaller than that of the proton. The fact that there is only a 20-50\% sharpening is probably associated with the substantial inhomogeneities of the crystal field in LES. It is thus clear that both dipolar and crystal-field mechanism contribute roughly equally to the observed Gd\(^{3+} \) in LES zero-field linewidth.

In the series of ethylsulfates in which the host ion is varied, a great variety of linewidths are observed. In this instance observed linewidths are a complicated function of the magnetic moment and relaxation time of the host lanthanide. The detailed explanation of the trends as a function of host, temperature, and transition involved must await a suitable theory for paramagnetic relaxation of rare-earth S-state ions in either zero or high field.

VI. CONCLUSION

Both crystal-field and hyperfine interactions in Gd\(^{3+} \)-doped LES hydrate have been studied. It was shown that the zero-field spin Hamiltonian agrees with the high-field spin Hamiltonian to a higher degree of accuracy than has previously been achieved. The available signal-to-noise ratio enabled both the magnitude and absolute sign of the dipole- and electric-quadrupole hyperfine interaction constants for this system to be determined for the first time. These constants had previously gone unobserved during 23 years of conventional EPR experiments. These parameters were discussed from the viewpoint of the theory of hfs, and
their magnitudes and signs were argued to be reasonable in comparison with other systems. Indeed, it is shown that for systems of lower than cubic symmetry, the quadrupole interaction should dominate the dipole interaction in the spectrum. While the size of $P$ can be determined in high field by placing $\mathbf{H}$ perpendicular to the crystal symmetry axis, its sign remains indeterminate except at extremely low temperatures ($\sim 0.001$ K).

Finally, a previously unnoticed effect was reported. The crystal field parameters of gadolinium in various lanthanide ethylsulfates change on deuteration of the waters of hydration. Some mechanisms were discussed which cause this change as well as line widths for the isomorphic series of lanthanide ethylsulfates. For the diamagnetic LES the $\text{Gd}^{3+}$ line widths are shown to arise from at least two separate and distinct factors: crystal-field inhomogeneities and magnetic dipole-dipole (local-field) coupling with neighboring nuclei.

*Supported in part by grants from The Research Corporation, Army Research Office-Durham, Office of Naval Research, and the National Science Foundation.
†Presented in part at the Twenty-ninth Symposium on Molecular Structure and Spectroscopy, 10 June 1974 (Paper T9).
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\(14\) See, for example, G. H. Dieke, Spectra and Energy Levels of Rare Earth Ions in Crystals (Interscience, New York, 1968).
\(18\) Through this paper all errors are rms errors, defined as three times the standard deviation $\sigma$.
\(26\) B. Bleaney, R. J. Elliott, H. E. D. Scofield, and E. S. Trenam, Philos. Mag. 42, 1062 (1951).
\(29\) H. A. Buckmaster, Can. J. Phys. 34, 150 (1956); 34, 341 (1956).
E. R. Bernstein and G. M. Dobbs

53A. Zalkin, J. D. Forrester, and D. H. Templeton, J. Chem. Phys. 39, 2881 (1963). The reason for the sharp lines compared to the ethylsulfates is not understood, except to say that since the trivalent gadolinium is surrounded by the nitrate groups, its nearest neighbors are not hydrogen bonded. The Ga-rays substitutes for the trivalent bismuth and the structure may be considered as [Bi(NO₃)₄]₂⁺Mg(H₂O)₆⁻·6H₂O.
79Yu E. Pol'skii, Fiz. Tverd. Tela 6, 842 (1964) [Sov. Phys.–Solid State 6, 651 (1964)].
80W. Low and D. Shaltiel, J. Phys. Chem. Solids 6, 315 (1958). It is a general feature of S-state ion high-field spectra that linewidths increase for higher Ms values, due to the increasing effect of the crystal field.