Spectroscopic analysis of hthium terbium tetrafluoride

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The absorption spectra of Tb^{3+} in LiTbF₄ have been recorded in the spectral interval from 4000 to 25000 $cm⁻¹$ and for temperatures between 2.3 and 150 K. This covers the transitions from the ground multiplet F_6 to the multiplets 7F_3 , 7F_2 , 7F_1 , 7F_0 , and 5D_4 . The transitions were predominantly of electric-dipole nature but small contributions of magnetic-dipole nature were seen. The crystal-field splitting was temperature dependent—the reason for this is not completely understood. No experimental evidence for a crystallographic phase transition was found. The energy levels of the ground \overline{F} term were calculated by diagonalizing an effective spin-orbit and crystal-field Hamiltonian in an LS basis. $H = \sum_i \lambda_i (\vec{L} \cdot \vec{S})^i + \sum_i \alpha_i \sum_i B_{im} O_{im}$, where the parameters λ_i are functions of the spin-orbit parameter ζ and the Slater parameter F_2 . The O_{im} and α are Racah operators and reduced matrix elements, respectively. The rare-earth site in LiTbF₄ possesses S_4 symmetry, which allows six crystal-field parameters. ζ and the six B_{im} were varied to obtain the best agreement with the experimentally observed levels. Keeping $F_2 = 434$ cm⁻¹ fixed, a fit with a standar deviation of 12 cm⁻¹ was obtained at 10 K with the following parameters: $\zeta = 1698$ cm⁻¹, $B_{20} = 445$ cm $B_{40} = -761$ cm⁻¹, $B_{44} = 1120$ cm⁻¹, $B_{60} = 4$ cm⁻¹, and $B_{64} = 761 + i609$ cm⁻¹. Although the ground LS term of Tb^{3+} is rather isolated, the term mixing is significant, which is also the case for the multiplet mixing. Even for the ground multiplet the J mixing cannot be ignored.

I. INTRODUCTION

The characteristics of the tripositive rare-earth ions (R^{3*}) doped in LiYF₄ have been studied in numerous works. The main reason for this is the interesting optical properties of these systems, .

For the rare earth heavier than Sm it is possible to grow $\text{LiY}_{1-x}R_xF_4$ crystals for any value of x. Until now, however, most of the attention has been focused on the crystals dilute in R content. Only recently, crystals dense in R have been investigated, and it has mainly been the magnetic properties which have been examined.¹⁻⁶ Especially the features of the systems in the temperature region where the ordered phase appears^{$7-10$} have been investigated.

Approximate crystal-field parameters have been determined from susceptibility and magnetization measurements, $2,5$ but only a few direct spectrory;
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2,5] scopic observations of the crystal-field splitting of the energy levels of R^{3*} in LiRF₄ have been made. $8,11$

The reason why works on crystals dilutely doped with R have been predominant is that for optical applications one is often limited to such systems due to destructive $R-R$ interaction, i.e., concentration quenching of the fluorescence. There are, however, exceptions, one of which is the $LiY_{1}R_{r}F_{4}$ system. Long-lived (5 msec) green fluorescence has been reported¹² from Tb³⁺ in $LiY_{0.5}Tb_{0.25}Gd_{0.25}F_4$ and measurements¹³ on Tb³⁺ in $LiTbF₄$ show that the lifetime of the green light is still longer than 1 msec.

Therefore, it is of interest to know in detail the

energy-level scheme of R^{3*} in the dense crystal, and here I report on the results for the whole ${}^{7}F$ term and the lowest ${}^5\!D_4$ multiplet for the case of Tb^{3*} .

II. THEORY

A. Free Tb^{3+} ior

The ground configuration of the free Tb^{3+} ion has eight 4f electrons outside closed orbitals. The next configuration, which is a $4f^75d$, is approximately 60 000 cm⁻¹ above the $4f^8$ configuration according to Dieke.¹⁴ Thus, for the lowest less according to Dieke. $^\mathrm{14}$ Thus, for the lowest level: of the ground configuration, the configuration mixing is small, and for these levels it is a good approximation to do all the energy calculations within the $4f^8$ configuration

In this configuration the ground LS term is the rather isolated ${}^{7}F$. The width of the ${}^{7}F$ term is 6000 cm⁻¹ and the next term, which is a ^{5}D , is situated about $20\,000\,$ cm⁻¹ above. In $^5\!D$ the lowest multiplet $^5\!D_{\phi}$ is rather isolated and acts as the upper fluorescence level in Th^{3+} . While for the 7F term the term mixing is small, this is not true for the other terms. The configuration extends to about 150000 cm^{-1} , and above 25 000 cm⁻¹ the classifi cation of the levels is somewhat uncertain.

Although the term mixing for ${}^{7}F$ is moderate, the Lande interval rule for the spin-orbit splitting is not well fulfilled, so it is necessary to introduce even this slight mixing in the calculations. 'The L and S quantum numbers are in this way no longer really good quantum numbers contrary to J . The spin-orbit coupling splits the ground term in seven

TABLE I. Crystal-field splittings of J multiplets in S_4 symmetry.

multiplets with $J = 0, 1, \ldots, 6$, where ${}^{7}F_{6}$ is the ground multiplet.

B. Tb³⁺ ion in a crystal with S_4 symmetry

 $LiTbF₄$ crystallizes in the tetragonal scheelite structure. The Tb³⁺ ion is on a site having S_4 symmetry with respect to the crystal as a whole, but almost D_{2d} symmetry with respect to the eight nearest-neighbor F^* ions. In S_4 symmetry the J multiplets split up as shown in Table I, where Γ_1 , Γ_2 , Γ_3 , and Γ_4 refer to the notation of Koster et $\frac{\Gamma_2}{al}$.¹⁵

The Γ_3 and Γ_4 representations are related by time-reversal symmetry, and the eigenvalues for states transforming according to these representations will therefore be degenerate. In the following they will be named by $\Gamma_{3,4}$

The crystal field mixes the multiplets slightly (less than $10\%)$, causing *J* to be not exactly a good quantum number. Therefore, it is necessary to take J mixing into account in the calculations of the energies of the levels. This, however, does not change the transformation properties of the associated states.

C. Selection rules for S_4 symmetry

The two most significant contributions to the transitions between the levels are the electric and magnetic dipole interaction with the applied electromagnetic radiation. The electric dipole transitions are parity forbidden within a single configuration, so they occur only due to configuration mixing. Still, for the rare earths they are, in. general, stronger than the magnetic dipole transitions.

Thus, calculation of transition strengths requires knowledge of configuration mixing, which is difficult to determine. Qn the contrary, separation of allowed and. forbidden transitions can be' done entirely on group-theoretical grounds. The selection rules for electric and magnetic dipole transitions in $S₄$ symmetry are given in Table II.

TABLE II. Selection rules in S_4 symmetry: (a) electric dipole transitions, (bj magnetic dipole transitions. The π spectrum has the electric polarization parallel to

D. Model Hamiltonian

As stated in Secs. IA-IC the configuration mixing is neglected in the calculations, whereas the mixing of LS terms and of J multiplets is taken into account by using a generalized spin-orbit Hamiltonian and by diagonalizing the entire LS matrix of the ground term.

Following Karayianis¹⁶ the effective spin-orbit operator to order p is given by

$$
H_{\rm so} = \sum_{i=1}^{p} \lambda_i (\vec{L} \cdot \vec{S})^i , \qquad (1)
$$

where the parameters λ_i are given by

$$
\lambda_i = \zeta \sum_{m=i}^p \left(\frac{\zeta}{F_2} \right)^{m-1} \wedge_{i \, m}(N) \;, \eqno(2)
$$

where ζ is the spin-orbit parameter and F_2 is a Slater parameter. $\Lambda_{i,m}(N)$, where N is the number of f electrons, are reduced matrix elements. They depend on the radial part of the wave functions only through the ratios F_4/F_2 and F_6/F_2 , which are wave function insensitive. The parameters Λ
are given by Karayianis.¹⁶ are given by Karayianis.¹⁶

The crystal-field operator is given by

$$
H_c = \sum_i \alpha_i(L) \sum_m B_{im} O_{im}(L) , \qquad (3)
$$

where $O_{i,m}(L)$ are Racah operator equivalents and $\alpha_i(L)$ are reduced matrix elements. In S_4 symmetry the only nonvanishing crystal-field parameters are B_{20} , B_{40} , B_{44} , B_{60} , and B_{64} , where B_{44} and B_{64} are complex. By proper choice of the coordinate system one can make one of them-for example, B_{44} -real, leaving six crystal-field parameters to be determined.

III. EXPERIMENTAL PROCEDURE

A. Crystal preparation

Single crystals of pure LiTbF, and of $LiYF_4$ doped with 10% Tb were grown by spontaneous crystallization from the melt as described by
Laursen and Holmes.¹⁷ The crystals were or Laursen and Holmes. 17 The crystals were orien tated by x -ray technique. Slabs about 1 mm thick were cut with faces perpendicular to an a axis for the polarized spectra and with faces perpendicular to the c axis for the axial spectrum. The slabs were mounted so that the angle between the crystal axis and the polarization axis was less than 2° (in fact, the spectra were rather insensitive to small deviations between the directions of the polarization and the crystal axes).

B. Apparatus

The absorption spectra were recorded using a modified Zeiss MM12 double monochromator. In the relevant spectral region it was possible to achieve a spectral resolution of 6 cm^{-1} . A Glan prism was used as polarizer.

For low-temperature measurements two cryostats were available. The first one, where the crystal was placed directly in the pumped liquid helium, was used for temperatures below 4.² K. The other one, where the crystal was placed in a stream of helium gas, was used for temperatures above 4.² K, In the first cryostat the temperature was determined from the He pressure. In the second the temperature was measured with an Au-Fe vs Cromel thermocouple, which had a resolution of 1 K. The temperature stability of the flow cryostat was ± 1 K.

C. Absorption spectra

The absorption spectra for polarized light for Tb^{3*} in LiTbF₄ were recorded in the region from 4000 to $25\,000\,{\rm cm^{-1}}$ $(2.5\text{--}0.4\,\mu\rm{m})$, which covers the transitions from the ${}^{7}F_{6}$ ground multiplet to the four highest multiplets of the ground term $(J =$ $0, 1, 2, 3$ and to the lowest 5D_4 multiplet.

The spectra were recorded for different temperatures between 2.3 and 150 K. However, for temperatures higher than 100 K, the positions of most of the lines were difficult to determine because the spectra were smeared out due to overlapping with many new lines.

L Transitions within the ground term

At 10 K, which is well above the Curie point T_c $= 2.87 \text{ K}$, which is well above the curle point T_c
= 2.87 K,⁷ only the ground state, which is a quasidoublet made up of two levels transformingaccording to Γ_2 of S_4 (Laursen and Holmes¹⁷), is populated. As shown in Fig. 1 one finds at this tempera-

FIG. 1. Absorption spectra at 10 K for transition from the ground state to ${}^{7}F_3$, ${}^{7}F_2$, ${}^{7}F_1$, and ${}^{7}F_0$ of Tb³⁺ in LiTbF₄. The π -polarized, the σ -polarized, the axial, and the reference spectrum are shown, The arrows indicate the magnetic dipole transitions.

ture for each of the two polarizations four strong absorption lines in the spectral region, where transitions to the ${}^{7}F_{0}$ - ${}^{7}F_{3}$ multiplets are expected.

This is in agreement with the transitions being of electric dipole nature. π transitions from the Γ , ground levels to $\Gamma_1({}^7F_0)$, $\Gamma_1({}^7F_1)$, $\Gamma_1({}^7F_2)$, and Γ , (7F_2) lead to four lines. σ transitions from Γ , to $\Gamma_{3,4}(^{7}F_{1}), \ \Gamma_{3,4}(^{7}F_{2}), \text{ and two } \Gamma_{3,4}(^{7}F_{3}) \text{ give four}$ lines, while the Γ_2 to Γ_2 transitions are not electric dipole allowed in S_4 symmetr:

In addition to these strong lines there are also some weak lines in the spectra. The transitions from Γ ₂ to Γ _{3,4} can be seen in the π spectrum whereas the Γ_2 to Γ_1 transitions cannot be seen in the σ spectrum. This indicates that there are also magnetic dipole transitions present —and that the lines are not due to misorientation of the crystal (compare with Table II). The magnetic dipole transitions from Γ_2 to Γ_2 are allowed in the σ spectrum. Experimentally one also finds two weak lines in the σ spectrum corresponding to two of the four possible Γ_2 to Γ_2 transitions. The two other lines should be found in regions, where they are difficult to recover due to the strong lines and noise. A few other very weak lines in the spectra must be due to impurities-in the crystal.

As a verification of the nature of the transitions, the axial spectrum was recorded. In the axial spectrum one should find the electric dipole transitions of the σ spectrum and the magnetic dipole transitions of the π spectrum. Thus, in the axial spectrum only Γ_2 to $\Gamma_{3,4}$ transitions are to be seen. This is what is found experimentally (see

 $\boldsymbol{0}$

 $\pmb{0}$

TABLE III. Experimental positions of the \overline{F} levels, which it has been possible to determine, and calculated positions of all the levels. All values are in cm⁻¹ $=$

Fig. 1).

At higher temperatures more levels of the ground multiplet become populated. Already at about 25 K the transitions from the first excited level $(\Gamma_{3,4})$ can be seen in the spectra, even if the relative population of this level is only around 3 \times 10⁻³. As the temperature is further increased, more and more lines appear in the spectra. By analyzing the spectra at temperatures up to 100 K, it has been possible to determine the energies of the seven lowest levels of the ground multiplet and

 3, 4'

 $\mathbf{0}$

 $\mathbf{0}$

of the two remaining Γ_2 levels of 7F_3 and 7F_2 . The positions of the experimentally determined lines of the ground LS term are listed in Table III.

 $\overline{2}$

The experimental linewidths of the electric dipole transitions from the ground quasidoublet are $10-20$ cm⁻¹, with an instrumental resolution of about 6 cm⁻¹. The lines are not much broader at higher temperatures, but the lines are getting weaker due to depopulation of the ground state, and some of them become disturbed by new lines. Some of the lines, which according to the theory

FIG. 2. π - and σ -polarized absorption spectra for transition from ${}^{7}F_6$ to ${}^{7}F_2$ of Tb³⁺ in LiTbF₄ at four different temperatures. $T_1 = 10 \text{ K}$, $T_2 = 30 \text{ K}$, $T_3 = 50 \text{ K}$, and T_4 = 70 K. The seven lowest levels of ${}^{7}F_6$ at 50 K are inserted in the figure with the ground $2\times \Gamma_2$ coinciding with the low-temperature lines to explain the origin of the warm-up lines. 3, $4 \rightarrow 2$ means the position where a transition from a $\Gamma_{3,4}$ level of $^7F_{6}$ to a Γ_{2} level of ${}^{7\!}F_2$ should be found, etc. The indicated transitions are those allowed by electric dipole interaction in S_4 symmetry. Those marked with a dot are forbidden in D_{2d} symmetry.

in principle should be present, are experimentally absent. This. however, may be due to low transition probability. It is more important that no strong inexplicable lines are experimentally found. As an example, the spectra of the transitions from the excited states to ${}^{7}F_{2}$ are shown in Flg. 2,

As mentioned earlier, the rare-earth ion is on a site of nearly D_{2d} symmetry, in which some S_4 symmetry-allowed transitions are forbidden. The two levels of the ground quasidoublet transform in D_{ad} like Γ_3 and Γ_4 , respectively (notation of Koster et $al.^{15}$. Because both levels are well populated, there will be no changes in the selection rules corresponding to transitions from the ground state.

At higher temperatures, however, some of the transitions would be forbidden in D_{2d} symmetry. These are marked with a dot in Fig. 2. It is seen that none of these lines is strong, but some of them are definitely present above 50 K. Below 50 K D_{2d} symmetry cannot be excluded on basis of the present data.

The spectra recorded as a function of temperature show that the crystal-field-split multiplets are contracting as the temperature is increased, and do so by an amount much larger than expect-

FIG. 3. Position of the'transition from the ground state to ${}^{7}F_0$ of Tb³⁺ in LiTbF₄ as a function of the temperature. Data points with error bars are experimental values. \bullet are calculated values. The error bars indicate the relative uncertainty. The absolute uncertainty of the lines is 5 cm^{-1} .

ed from the thermal expansion of the lattice. In addition to this smooth temperature contraction of the multiplets, it seems that the whole term contracts slightly $(2-3 \text{ cm}^{-1})$ in a small temperature region around 50 K. As an example, the temperature dependence of the ${}^{7}F_{0}$ line is shown in Fig. 3.

The temperature dependence of the multiplets is small below 50 K, but from 50 to 100 K it is as large as $0.1 \text{ cm}^{-1}/\text{K}$ for some of the lines. This is equivalent to a contraction of the crystal-field splitting of about 0.03% /K. This strong temperature dependence is in agreement with the shift of the two lines of the ground multiplet seen by Holmes et al.² in neutron scattering.

For those few lines, which it has been possible to follow from 100 to 150 K, it seems that the temperature dependence is still present with about the same strength as below 100 K. It has been checked that the observed temperature effect is not due to strain induced by the mounting of the crystal.

The contraction could be due to a second-order phase transitions, but experimentally there is no evidence of the presence of a phase with lower symmetry—which in this case should be C_2 , where the $\Gamma_{3,4}$ degeneracy is removed and the Γ_1 to Γ_1 and Γ , to Γ , (in S_4 notation) electric-dipole transitions are allowed in the σ spectrum.

As a supplement, the line shifts for 10% Tb in $LiYF₄$ were examined. It was found that the multiplet contraction was much weaker $\langle 0.02 \text{ cm}^{-1}/\text{K} \rangle$ than for the dense crystal, and the term contraction at 50 K was not detectable $\langle 1 \text{ cm}^{-1} \rangle$.

'To study the effect of the magnetic ordering on the absorption spectra, the spectra were recorded down to 2.3 K. Qualitatively, there was no change in the spectra, but for some of the lines there were line shifts in the temperature interval from

FIG. 4. Experimental position of the transition from the ground state to ${}^{7}F_0$ of Tb³⁺ in LiTbF₄ as a function of the temperature around T_c , where LiTbF₄ order magnet ically.

2.7 to 3.0 K. This is in agreement with the fact that the crystal orders ferromagnetically below 2.⁸⁷ K.' As shown in Fig. ⁴ the shift of the line corresponding to absorption to ${}^{7}F_0$ is $3.5 \pm 1 \text{ cm}^{-1}$ at 2.3 K. The ordering causes the ground quasidoublet to split proportional to the magnetization into an $M_J = +6$ and an $M_J = -6$ level, of which only the lowest is well populated.

For an Ising system the shift at 0 K should be equal to $k_B \theta$, where θ is the asymptotic paramagnetic Curie temperature, assuming that in the ordered phase the domains are long thin needles along the c axis. Holmes $et \ al.^1$ give $\theta = 3.60$ ± 0.1 K (θ_{dipolar} = 3.97 K) and thus a shift of 2.5 ± 0.1 cm^{-1} is expected. The observed shift seems somewhat higher than this, similar to what has been

FIG. 5. π - and σ -polarized absorption spectra at 10 K for the transitions from the ground state to 5D_4 of Tb³⁺ in LiTbF₄.

TABLE IV. Experimental positions in cm⁻¹ of the 5D_4 levels;

	Experimental values
	(10 K)
9	20666
	20654
3, 4	20641
$\overline{2}$	20 589
	20 580
3,	20 571
	20 56 1

found in LiHo \mathbf{F}_4 (Battison *et al.*⁹) in which case the deviation might be explained by the large hyperfine interaction in Ho^{3*} . This is not possible in the case of Tb³⁺ in LiTbF₄, where the total hf splitting is 0.3 cm^{-1} .

2. Transitions to the 5D_4 multiplet

In the region around $20\,600$ cm⁻¹ three lines in the π spectrum and two lines in the σ spectrum were found at low temperatures, as shown in Fig. 5. They are due to transitions from the ground state to the three Γ_1 levels and two $\Gamma_{3,4}$ levels of the 5D_4 multiplet. The two Γ_2 levels of 5D_4 were found by examining the warm-up spectra. The experimental line positions of 5D_4 are listed in Table IV.

IV. CALCULATIONS

A. Pitting procedure

The energy levels of the ground LS term were calculated using the model described in Sec. IID. The six crystal-field parameters and the spinorbit parameter ζ were varied, until the best agreement with the 19 experimentally observed levels was obtained.

It is not relevant to let $F₂$ vary freely, since this parameter primarily must give the correct configuration splitting. However, the fitting was tried with several $F₂$ values between 434 cm⁻¹ given by Wybourne¹⁸ and the "smoothed" value 417
cm⁻¹ given by Karayianis.¹⁶ The result was not cm⁻¹ given by Karayianis.¹⁶ The result was not very sensitive to the $F₂$ value, but the best initial fit was obtained with $F_2 = 434$ cm⁻¹. This value was used in the remaining calculations.

In order to get an LS term splitting in good agreement with the experiments, it was necessary to use an effective spin-orbit Hamiltonian including three terms $[p = 3 \text{ in Eq. (1)}].$ Note, however that the number of fitting parameters is independent of the number of these terms. As experimental data for the ${}^{7}F_{5}$ and ${}^{7}F_{6}$ multiplets have not been

obtained, it is especially important not to use more free parameters in the fitting procedure than dictated by the underlying physics.

Setting $Im B_{64}$ equal to zero and fitting only five crystal-field parameters gave almost as good agreement as when all six crystal-field parameters were allowed to vary freely. This is what might be expected from the fact that the rare-earth ion is on a site of nearly D_{2d} symmetry, where all crystal-field parameters should be real. The final fittings, however, were done using all six crystal-field parameters.

B. Results

The fitting procedure was carried out at five temperatures between 10 and 100 K. (For the lines only known experimentally at one temperature, a temperature dependence similar to the other lines was assumed.) As shown in Fig. 6 the crystalfield parameters were strongly temperature dependent. This was not the case for the ζ parameter. ζ was found to be 1698 cm⁻¹ for all tempera tures lower than 100 K, where it was 1700 cm^{-1} . The uncertainty of ζ was less than 2.5 cm⁻¹. Wybourne¹⁸ gives a free-ion value of 1705 cm^{-1} for Tb". In crystals the spin-orbit coupling strength may differ from the free-ion value due to the coupling to the crystal field.

In the fittings a standard deviation of 12 cm^{-1} was obtained. The standard deviation is given by

$$
S = \left(\sum_{i=1}^{19} \frac{(\Delta E_i)^2}{19 - 6}\right)^{1/2}.
$$
 (4)

The calculated energy levels at 10 K are given in Table III and the crystal-field parameters at 10 K in Table V. 'The calculated position of the lowest $\Gamma_{3,4}$ level of the ground multiplet is 108 cm^{-1} at 10 K and 100 cm⁻¹ at 100 K. Holmes et al.²

FIG. 6. Crystal-field parameters at the Tb³⁺ site in LiTbF_A as a function of temperature. The sign of Im B_{64} is unknown. The lines through the points are only for visual aid.

TABLE V. Crystal-field parameters in cm^{-1} . It is not possible to determine the sign of $\text{Im}B_{64}$ with the methods referred to here. The first column gives the values obtained in this work, when the parameters are fitted inside the \overline{F} term. The uncertainties correspond for the given model to an uncertainty pn the experimental data of about 5 cm^{-1} . The second column gives the values obtained by Holmes *et al.* (Ref. 2) by neutron-scattering and by susceptibility measurements. The third column gives the values obtained in this work, when the parameters are fitted inside the \overline{F}_6 multiplet.

find the same levels at 106 and 98 cm⁻¹, respectively, by inelastic neutron scattering. Good agreement is also found for the next $\Gamma_{3,4}$ level where the calculated values are 175 and 165 cm^{-1} , and the experimental values from the inelastic neutron scattering are 169 and 161 cm^{-1} .

V. DISCUSSION

(4) A. Crystal field

As shown in Table V the spectroscopically determined B parameters at 10 K differ rather much from those found by Holmes *et al.*² by fitting to the susceptibility in the temperature range 5-300 K and to a few levels of the ground multiplet. The temperature dependence found (Fig. 6) indicates that a detailed comparison between these two sets of parameters cannot be undertaken.

The spectroscopically determined B parameters give splittings of the ground multiplet —when diagonalizing within the whole LS term-which is in good agreement with the seven lowest levels, which it has been possible to determine experimentally. As seen from Fig. 7 the calculated positions of both the remaining three high-lying levels as well as the seven lower-lying levels are in semiquantitative agreement with the experimental splittings for Tb^{3*} doped in $CawO₄$ given by Wort-
man.¹⁹

On the other hand, when diagonalizing only within the ground multiplet, but with B parameters found by fitting to the whole LS term, some of the levels come out wrong, differing as much as 50 cm⁻¹ from the experimental values. This indicate

FIG. 7. Splittings of the ground multiplet of Tb^{3+} . (I) Experimentally determined for Tb^{3+} doped in CaWO₄ (Ref. 19). (II) Calculated for Tb³⁺ in LiTbF₄ by diagonalizing the entire LS matrix of the ground term. (III) Calculated for Tb³⁺ in LiTbF₄ with the same parameters as in II , but diagonalizing only the J matrix of the ground multiplet. {IV) Calculated for Tb^{3+} in LiTb F_4 with the parameters of Holmes et al. $(Ref. 2)$.

that even for the ground multiplet, J mixing is significant. If the B parameters are fitted inside the ground multiplet only, the B values given in the third column of Table V are obtained. (Because of six parameters being available for fitting of l0 levels, the fit is perfect to within ¹ cm^{-1} .) It is clearly seen that the set of B parame ters are model dependent to a degree which is significant in relation to a quantitative discussion of experimentally determined properties. Still, it is remarkable that the order of all the levels is the same for all four cases depicted in Fig. 7. This must be due to the circumstance that the different interactions are of such relative strengths that the LS description as well as the J description are both good enough for a quantitatively correct discussion of at least the lowest multiplet.

When the splitting of the ground multiplet is calculated using the parameters given by Holmes $e t$ $al.,²$ it is clear from Fig. 7 that the three highes levels have too low energies, whereas the others fit very well.

The last fact is mainly due to the clamping of the lowest $\Gamma_{3,4}$ levels to the positions obtained by

the neutron-scattering results, whereas the low value of the total splitting is connected to the low value of B_{20} . B_{20} is, however, almost tied to the . difference between the reciprocal susceptibilities $\chi_{\scriptscriptstyle\rm II}^{-1}$ and $\chi_{\scriptscriptstyle\rm I}^{-1}$ at "high" temperatures. At 300 K this high-temperature limit is approached, and therefore the value of B_{20} , obtained by Homes et al.,² can be regarded as approximately correct for T $\simeq 300$ K. In this connection it is noteworthy that if the temperature dependence of B_{20} observed in the spectroscopic results persists up to room temperature, the value of B_{20} will be lowered to about 360 cm"' at this temperature. Yet, because of the significant but partly unknown temperature variation of all the B 's, we shall defer at this stage to make a detailed comparison with the susceptibility data in the whole temperature range 4-300 K.

The calculated splitting Δ in the paramagnetic phase of the two Γ ₂ levels spanning the ground quasidoublet is 1.9 cm^{-1} , which is higher than the values given by Holmes et $al.^2$ (1.3 cm⁻¹), by Magvalues given by Holmes *et al.*² (1.3 cm⁻¹), by M
ariño *et al*.¹¹ (\simeq 1 cm⁻¹), and (for Tb³⁺ diluted in LiYF₄) by Laursen and Holmes¹⁷ (0.93 cm⁻¹). Calculation with B parameters fitted inside the ground multiplet gives a value of 1.5 cm⁻¹ for Δ .

The splitting Δ is, for Tb³⁺ in a Scheelite-crystal, strongly sensitive primarily to $Im B_{\alpha\alpha}$. If a value of $\Delta \simeq 1$ cm⁻¹ is considered the most reliable. this will therefore point to a somewhat lower value of $\text{Im} B_{64}$ than that obtained from the term fitting.

8. Temperature dependence of the spectrum

Four features concerning the temperature dependence of the level positions have been found experimentally. (i) A small shift of the lines takes place in a narrow temperature region around 50 K. (ii) The crystal-field splittings show a large continuous decrease in the temperature region $50-$ 100 K. There is indication that this decrease continues to higher temperatures. (iii) The temperature effects are much less pronounced in a diluted system. (iv) At no temperature do the data indicate that additional crystal-field parameters are necessary in order to explain the level positions. This means that a transition to a lower rare-earth site symmetry is very unlikely.

 A ls-Nielsen *et al.*⁷ have determined the structure of $LiTbF_4$ at 100 K and at room temperature by neutron diffraction. They find that the crystal structure is very stable, although one coordinate of the fluorine ions is changed about 0.25% from 100 K to room temperature. The thermal expansion is small and anisotropic. The structure has not been determined below 100 K, where the spectroscopic data indicate that something more interesting happens.

In the scheelite structure (space group $I4_1/a$) there exist three degrees of freedom in the fluorine positions. The first fluorine can—apart from physical constraints —be placed arbitrarily in the unit cell, whereas the remaining fluorines are fixed by the symmetry requirements. Further the fluorines probably give the dominant contribution to the crystal-field energy of the $4f$ electrons of the R^{3*} ions. This coupling, together with the possibility of changing the fluorine positions without breaking the space-group symmetry. should in general (i.e., for all ${\rm Li}R\, \mathrm{F}_4$ systems) lead to temperature-dependent properties. The magnitude of the effect is difficult to predict, but qualitatively a. smooth temperature dependence might be expected.

The temperature dependence of the experimental data for $LipF_4$ indicates, however, the existence of a characteristic transition temperature T_t around 50 K. The nature of the transition is not clear. Speculatively, the continuous character of the transition and the constancy of the B's below T_t

might be caused by the fluorine system being trapped in positions corresponding to local D_{2d} symmetry at the R^{3+} sites. This is possible without breaking the global S_4 symmetry of the R^{3*} site . or the space-group symmetry. However, there would still exist two degrees of freedom for configurational changes in the fluorine system. Finally the cooperative nature of the phenomena —which is indicated by the weakening of the effect in a diluted system —is understandable, because each fluorine ion makes significant contributions to the crystal-field energy on at least two R^{3+} sites.

In contrast to the temperature dependence observed for LiTbF₄, Battison *et al.*⁹ found no temperature dependence for Ho^{3+} in LiHo F_4 for those transitions which they examined.

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