

Determination of tetragonal crystalline electric field parameters for Yb³⁺ and Ce³⁺ ions from experimental g-factors values and energy levels of Kramers doublets

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The tetragonal crystalline electric field parameters for Yb³⁺ and Ce³⁺ ions are expressed via ground multiplet excited doublets energies and parameters defining doublets' wave functions. The crystalline electric field parameters for Yb³⁺ ion in YbRh₂Si₂, YbIr₂Si₂ and KMgF₃ crystals extracted from excited state doublets energies and g-factors of ground state doublet are compared with parameters determined in other works. { No more than 200 words. Use the MRSej style of paragraph named "_MRSej_Abstract" }

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1. Introduction { Use the MRSej style of paragraph named "_MRSej_Section" }

Our work was initially stimulated by investigation of heavy-fermion Kondo lattice compounds. Very peculiar magnetic, thermal and transport properties of 4f-electron based heavy-fermion systems are determined by the interplay of the strong repulsion of 4f-electrons on the rare-earth ion sites, their hybridization with wide-band conduction electrons and an influence of the crystalline electric field. { For first paragraph use the MRSej style of paragraph named "_MRSej_TextNonIndent" }

In this paper we present the detailed calculation of CEF parameters from energies of ground multiplet excited Kramers doublets and g-factors of ground state Kramers doublet. { Use the basic MRSej style of paragraph named "_MRSe_Text_Main" }

2. Diagram of Yb³⁺ g-factors

A free Yb³⁺ ion has a 4f¹³ configuration with one term ²F. The spin-orbit interaction splits the ²F term into two multiplets: ²F_{7/2} with J = 7/2 and ²F_{5/2} with J = 5/2, where J is value of the total momentum **J** = (J_x, J_y, J_z). Multiplets are separated by about 1 eV [4]. The Hamiltonian of the Yb³⁺ ion interaction with the tetragonal CEF could be written via equivalent operators O_k^q(**J**) [4]: { Below we use the MRSej style of paragraph named "_MRSej_Eq" }

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$$H = \alpha B_2^0 O_2^0 + \beta (B_4^0 O_4^0 + B_4^4 O_4^4) + \gamma (B_6^0 O_6^0 + B_6^4 O_6^4),$$
 { One tab after } (1)

where B_k^q are the CEF parameters, α = 2/63, β = -2/1155, γ = 4/27027 [4]. { After equations we use "_MRSej_TextNonIndent" or "_MRSe_Text_Main" }

As follows from the group theory, the two-valued irreducible representation D^{7/2} of rotation group contains two two-dimensional irreducible representations of the double tetragonal group [4].

The former results correspond to bases |5/2>, |-3/2> and |-5/2>, |3/2>, the latter corresponding to bases |7/2>, |-1/2> and |-7/2>, |1/2>. It is convenient to introduce parameters C, A and D:

$$C = 4B_2^0/21 + 40B_4^0/77 - 560B_6^0/429, \quad A = 4B_2^0/7 + 8B_4^0/77 + 80B_6^0/143. \quad (2)$$

Table 1. Energies, wave functions and g-factors of Yb³⁺ ion in tetragonal crystalline electric field. **{ “_MRSej_table_caption” }**

$E_{1,2} = -D \pm C / \cos \varphi_7$	$E_{3,4} = D \pm A / \cos \varphi_6$ { “_MRSej_tableIn” }
$ ^1\Gamma_7' \uparrow, \downarrow\rangle = \pm c_1 \pm 5/2\rangle \pm c_2 \mp 3/2\rangle$	$ ^3\Gamma_6' \uparrow, \downarrow\rangle = \pm a_1 \mp 7/2\rangle \pm a_2 \pm 1/2\rangle$
$ ^2\Gamma_7' \uparrow, \downarrow\rangle = \mp c_2 \pm 5/2\rangle \pm c_1 \mp 3/2\rangle$	$ ^4\Gamma_6' \uparrow, \downarrow\rangle = \mp a_2 \mp 7/2\rangle \pm a_1 \pm 1/2\rangle$
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Since matrices (2) are diagonal in the bases of their eigenvectors we can find the relations between our angular parameters and CEF parameters: $\tan \varphi_7 = C_3/C$, $\tan \varphi_6 = A_3/A$, it is enough to take $-\pi/2 \leq \varphi_7, \varphi_6 \leq \pi/2$.

Table 2. Experimental g-factors of Yb³⁺ ion in tetragonal crystalline electric field given in figure 1.

	YbRh ₂ Si ₂ [1]	YbIr ₂ Si ₂ [2]
$ g_{\parallel} $	0.17(7)	0.85(1)
$ g_{\perp} $	3.561(6)	3.357(5)

EPR spectra of Yb³⁺ ions { “_MRSej_SubSection” }

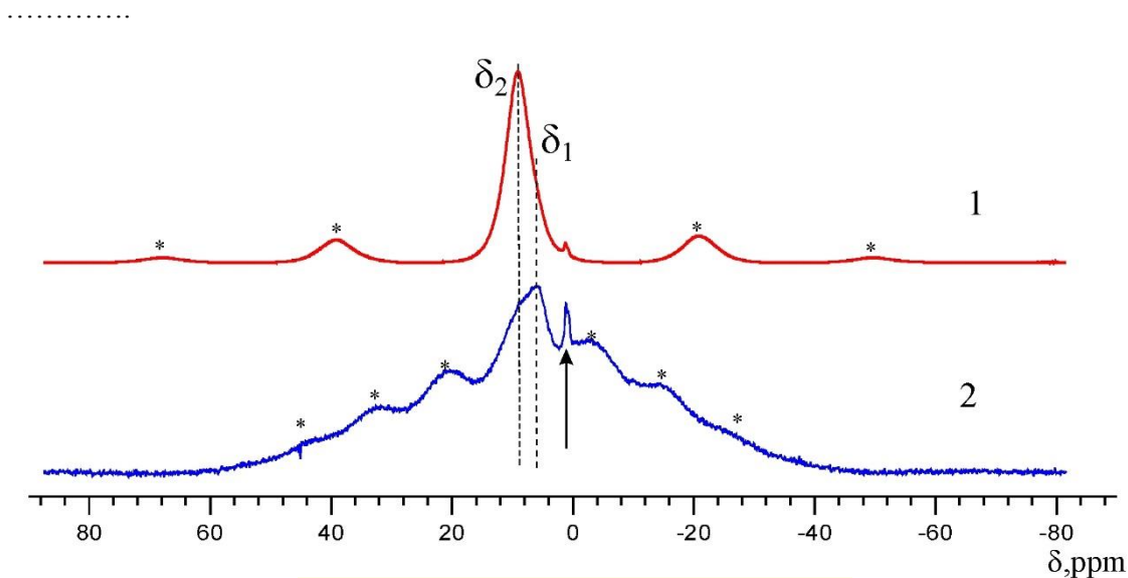
The Zeeman energy $g_J \mu_B \mathbf{HJ}$ in the basis $|\uparrow\rangle, |\downarrow\rangle$ of each doublet could be represented by matrix

$$H_{Zeeman} = g_{\parallel} \mu_B H_z S_z + g_{\perp} \mu_B (H_x S_x + H_y S_y), \quad (3)$$

where \mathbf{H} is the magnetic field, \mathbf{S} is the effective spin operator with $S = 1/2$, μ_B is the Bohr magneton, g_{\parallel} and g_{\perp} are g-factors when the field is applied parallel and perpendicular to the tetragonal z-axis, respectively (tab. 1) The field is applied parallel and perpendicular to the tetragonal z-axis, respectively.

EPR spectra in cubic symmetry case

In the case of cubic symmetry $\tan \varphi_7 = -\sqrt{3}$, $\tan \varphi_6 = -\sqrt{35}$, $c_1 = \sqrt{3}/2$, $c_2 = -1/2$, $a_1 = \sqrt{7/12}$, $a_2 = -\sqrt{5/12}$. In accordance with expansion $\Gamma_8 = \Gamma_7' + \Gamma_6'$ [4] the doublets $^2\Gamma_7'$ and $^3\Gamma_6'$



{ Figures and tables are inserted into tables }

Figure 1. The diagram of g-factors of Yb³⁺ ion in tetragonal crystalline electric field and experimental g-points taken from literature (tab. 2). **{ “_MRSej_figure_caption” }**

Using the least squares method the experimental values of g -factors (tab. 2) and experimental energy of whole 2F term levels have been taken into account. Obtained CEF parameters satisfy the experimental energy scheme of 2F term very well, but are reproduced by our expressions (9) only approximately (tab. 5).

4. Calculation of CEF parameters for Yb^{3+} ion. Comparison with another papers

Let us calculate the CEF parameters for the given excited state doublets energies $\Delta_1 < \Delta_2 < \Delta_3$. It follows from (3) that we find:

$$B_2^0 = \frac{1}{8}b + \frac{3}{4}b_6 \cos \varphi_6 + \frac{1}{4}b_7 \cos \varphi_7,$$

$$B_4^0 = -\frac{1}{4}b + \frac{1}{32}b_6 \cos \varphi_6 + \frac{5}{32}b_7 \cos \varphi_7, \quad B_4^4 = -\frac{7\sqrt{35}}{32}b_6 \sin \varphi_6 - \frac{35\sqrt{3}}{32}b_7 \sin \varphi_7, \quad (9)$$

$$B_6^0 = -\frac{13}{160}b + \frac{39}{320}b_6 \cos \varphi_6 - \frac{91}{320}b_7 \cos \varphi_7, \quad B_6^4 = \frac{117\sqrt{35}}{320}b_6 \sin \varphi_6 - \frac{273\sqrt{3}}{320}b_7 \sin \varphi_7,$$

In paper [7] CEF parameters of Yb^{3+} ion in KMgF_3 crystal have been found (tab. 5). Using the least squares method the experimental values of g -factors (tab. 2) and experimental energy of whole 2F term levels have been taken into account. Obtained CEF parameters satisfy the experimental energy scheme of 2F term very well, but are reproduced by our expressions (9) only approximately (tab. 5).

In this case g_{\parallel} and g_{\perp} are related by the equation $g_{\parallel} + 2g_{\perp} + 8 = 14p_3^2$, but as the admixture of excited ${}^2F_{5/2}$ multiplet is small ($p_3 = 0.00551$ [7]) we obtain previous relation $g_{\parallel} + 2g_{\perp} + 8 = 0$. On the diagram (fig. 1) we marked experimental values of Yb^{3+} g -factors in several crystals (see also tab. 2). This allows us to estimate the signs of g -factors and to make assumptions about the ground state Kramers doublet on the basis of measured absolute values of g -factors. For example, it is evident that the ground state doublet of Yb^{3+} ion in HfSiO_4 is ${}^3\Gamma_6'$ and both parallel and perpendicular g -factors have a negative sign. The ground state doublet of Yb^{3+} ion in KMgF_3 is ${}^4\Gamma_6'$, the sign of g_{\parallel} is positive, the sign of g_{\perp} is negative. In CaF_2 crystal the tetragonal center of Yb^{3+} is in state ${}^1\Gamma_7'$ and the sign of g_{\parallel} is positive but the sign of g_{\perp} can be both positive and negative.

4. Summary

For Yb^{3+} and Ce^{3+} ions all possible sets of tetragonal crystalline electric field parameters that satisfy the given experimental energy scheme of ground multiplet are defined.

The earlier published CEF parameters for Yb^{3+} ion in YbRh_2Si_2 and YbIr_2Si_2 crystals calculated with the use of least squares method could be obtained from our formulas (see tab. 5).

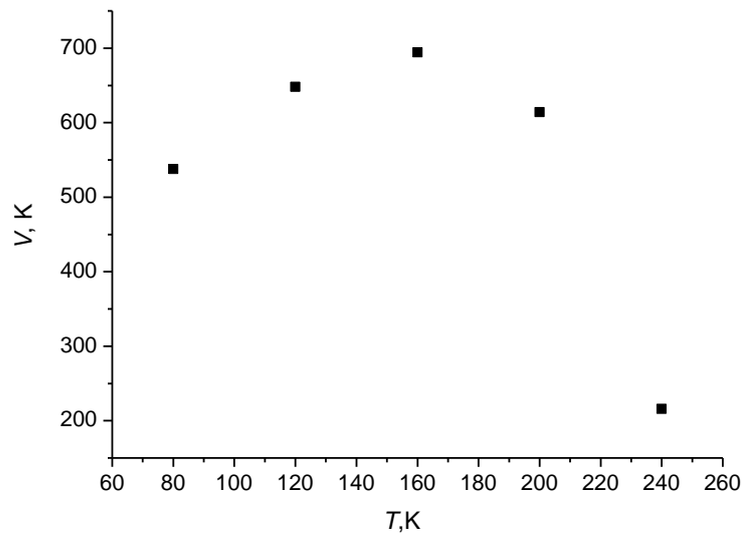


Figure 2. Temperature dependence of the strength of the potential V (see (6)), obtained from the simulation of C_{60} EPR spectra.

Acknowledgments

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