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Determination of tetragonal crystalline electric field parameters for Yb³⁺ and Ce³⁺ ions from experimental *g*-factors values and energy levels of Kramers doublets {*Use style*" *MRSej Title*" }

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The tetragonal crystalline electric field parameters for Yb^{3+} and Ce^{3+} ions are expressed via ground multiplet exited doublets energies and parameters defining doublets' wave functions. The crystalline electric field parameters for Yb^{3+} ion in $YbRh_2Si_2$, $YbIr_2Si_2$ and $KMgF_3$ 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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Keywords: crystalline electric field parameters, *g*-factors, Yb-based intermetallides, heavy-fermion systems {"_*MRSej_Key"*}

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 4*f*-electron based heavy-fermion systems are determined by the interplay of the strong repulsion of 4*f*-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 exited 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 $4f^{13}$ configuration with one term ²*F*. The spin-orbit interaction splits the ²*F* term into two multiplets: ²*F*_{7/2} with J = 7/2 and ²*F*_{5/2} with J = 5/2, where *J* is value of the total momentum $\mathbf{J} = (J_x, J_y, J_z)$. 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 $O_k^q(\mathbf{J})$ [4]: {*Below we use the MRSej style of paragraph named " MRSej Eq.*"}

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

where B_k^q are the CEF parameters, $\alpha = 2/63$, $\beta = -2/1155$, $\gamma = 4/27027$ [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 $|5/2\rangle$, $|-3/2\rangle$ and $|-5/2\rangle$, $|3/2\rangle$, the latter corresponding to bases $|7/2\rangle$, $|-1/2\rangle$ and $|-7/2\rangle$, $|1/2\rangle$. It is convenient to introduce parameters *C*, *A* and *D*:

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

Table 1. Energies, wave functions and g-factors of Yb^{3+} ion in tetragonal crystalline electric field.{"_MRSej_table_caption'' }

$E_{1,2} = -D \pm C / \cos \varphi_7$	$E_{3,4} = D \pm A / \cos \varphi_6 \{ \text{``_MRSej_tableIn''} \}$
$ {}^{1}\Gamma_{7}^{t}\uparrow,\downarrow\rangle = \pm c_{1} \pm 5/2\rangle \pm c_{2} \mp 3/2\rangle$	$ {}^{3}\Gamma_{6}^{t}\uparrow,\downarrow\rangle = \pm a_{1} \mp 7/2\rangle \pm a_{2} \pm 1/2\rangle$
$ ^{2}\Gamma_{7}^{t}\uparrow,\downarrow\rangle = \mp c_{2} \pm 5/2\rangle \pm c_{1} \mp 3/2\rangle$	$ {}^{4}\Gamma_{6}^{t}\uparrow,\downarrow\rangle = \mp a_{2} \mp 7/2\rangle \pm a_{1} \pm 1/2\rangle$

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 $-\pi/2 \le \varphi_7$, $\varphi_6 \le \pi/2$.

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

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

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

EPR spectra of Yb^{3+} ions {" **MRSej SubSection**"}

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

where **H** is the magnetic field, **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

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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 ${}^3\Gamma_6'$



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"*}

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

4. Calculation of CEF parameters for Yb³⁺ ion. Comparison with another papers

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

$$B_{2}^{0} = \frac{1}{8}b + \frac{3}{4}b_{6}\cos\varphi_{6} + \frac{1}{4}b_{7}\cos\varphi_{7},$$

$$B_{4}^{0} = -\frac{1}{4}b + \frac{1}{32}b_{6}\cos\varphi_{6} + \frac{5}{32}b_{7}\cos\varphi_{7}, \quad B_{4}^{4} = -\frac{7\sqrt{35}}{32}b_{6}\sin\varphi_{6} - \frac{35\sqrt{3}}{32}b_{7}\sin\varphi_{7},$$

$$B_{6}^{0} = -\frac{13}{160}b + \frac{39}{320}b_{6}\cos\varphi_{6} - \frac{91}{320}b_{7}\cos\varphi_{7}, \quad B_{6}^{4} = \frac{117\sqrt{35}}{320}b_{6}\sin\varphi_{6} - \frac{273\sqrt{3}}{320}b_{7}\sin\varphi_{7},$$
(9)

In paper [7] CEF parameters of Yb³⁺ ion in KMgF₃ crystal have been found (tab. 5). Using the least squares method the experimental values of *g*-factors (tab. 2) and experimental energy of whole ${}^{2}F$ term levels have been taken into account. Obtained CEF parameters satisfy the experimental energy scheme of ${}^{2}F$ 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 Yb³⁺ 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 Yb³⁺ ion in HfSiO₄ is ${}^{3}\Gamma_{6}^{t}$ and both parallel and perpendicular g-factors have a negative sign. The ground state doublet of Yb³⁺ ion in KMgF₃ is ${}^{4}\Gamma_{6}^{t}$, the sign of g_{\parallel} is positive,

the sign of g_{\perp} is negative. In CaF₂ crystal the tetragonal center of Yb³⁺ is in state ${}^{1}\Gamma_{7}^{t}$ and the sign of g_{\parallel} is positive but the sign of g_{\perp} can be both positive and negative.

4. Summary

For Yb³⁺ and Ce³⁺ 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 C₆₀ EPR spectra.

Determination of tetragonal CEF parameters for Yb³⁺ and Ce³⁺ ions from experiment ... Short title

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