Low-lying energy spectrum of the cerium dimer

A. V. Nikolaev
Franklin Institute of Physical Chemistry and Electrochemistry of RAS, Leninskii pr. 31, 119991, Moscow, Russia and Skobeltsyn Institute of Nuclear Physics, Moscow State University, Vorob’evy Gory 1/2, 119991, Moscow, Russia
(Dated: January 22, 2013)

The electronic structure of Ce₂ is studied in a valence bond model with two 4f electrons localized at two cerium sites. It is shown that the low-lying energy spectrum of the simplest cerium chemical bond is determined by peculiarities of the occupied 4f states. The model allows for an analytical solution which is discussed along with the numerical analysis. The energy spectrum is a result of the interplay between the 4f valence bond exchange, the 4f Coulomb repulsion and the spin-orbit coupling. The calculated ground state is even \( \Omega = \Lambda = \Sigma = 0 \) state and the \( ^3\Sigma^+ \) doublet (\( \Omega = \pm 5 \), \( \Lambda = \pm 6 \), \( \Sigma = \mp 1 \)). The calculated magnetic susceptibility displays different behavior at high and low temperatures. In the absence of the spin-orbit coupling the ground state is the \( ^3\Sigma^- \) triplet. The results are compared with other many-electron calculations and experimental data.

PACS numbers: 31.15.xw, 31.15.vn, 33.15.Kr

I. INTRODUCTION

Understanding the electronic nature of chemical bonds of \( f \)-elements is an issue which has been attracting wide attention of investigators working in diverse areas of research. The main question is the involvement of 4f states in the formation of chemical bonds in molecules and in solids. In general, one can speak of two extreme scenarios. That is, either \( f \)-electrons participate in chemical bonding which is the case for 5f-states in the U₂ molecule \(^1\), or they stay essentially localized at each site forming core-like states like in lanthanides. This phenomenon known as the localization-delocalization paradigm in solid-state physics \(^2\) is the subject of intense theoretical considerations and discussion.

In various compounds cerium is believed to display both types of 4f behavior which makes it the key element in these studies. Indeed, in solids Ce and its compounds exhibit a broad range of unusual electronic, magnetic and structural properties called correlation effects \(^3\). While much effort has been invested in understanding of the complex electronic behavior in sophisticated cerium compounds, very little attention has been given to the simplest cerium chemical bond \(^4\) \(^5\), which is realized in Ce₂. Meanwhile, owing to the progress of the spectroscopy of small metal clusters \(^6\) precise experimental data on the cerium dimer \(^6\) have been reported recently, followed by first reliable many-electron calculations \(^6\) \(^7\).

Cao and Dolg \(^4\) have performed a series of calculations of Ce₂ using the complete active space self-consistent field (CASSCF) method, multireference configuration interaction (MRCI) method and coupled cluster approach [CCSD(T)] and concluded that its ground state “superconfiguration” is \( 4f^14f^1(6s6p)^2(5d5p)^4 \). Six low-lying molecular terms \( ^1\Sigma^+, ^1\Sigma^-, ^3\Sigma^+, ^3\Sigma^-, ^3\Pi^+_g, ^1\Delta_g \) and \(^6\Pi \) “virtually degenerate” so that the exact refinement of the ground state is very difficult. The authors then restricted themselves to the \(^1\Sigma^+ \) and \(^3\Sigma^+ \) ground states for which in the scalar relativistic approximation they obtained the following spectroscopic constants: \( R_e = 2.62 \pm 0.02 \) Å, \( D_e = 1.60 \pm 0.41 \) eV and \( \omega_e = 201 \pm 13 \) cm\(^{-1} \). The low-lying excitations were attributed to the core-like properties of 4f states. In the spin-orbital calculation the authors obtained the \( \Omega = 0 \) ground state followed by two low-lying levels \( ^3\Sigma^- \) and \( ^3\Pi_g \).

The nature of the Ce₂ bonding and the role of the 4f-states were also considered in CASSCF studies by Roos et al. in \(^5\). As in \(^4\) it was found that the triple bond molecular configuration \((\sigma_u)^2(\pi_u)^4\) couples in the \(^1\Sigma^+ \) term. Two remaining nonbonding and localized 4f electrons are in \( \phi \)-state \((m = \pm 3)\). These two 4f electrons then can form the configurations \( (4f\phi_u)^2 \), \( (4f\phi_g)(4f\phi_u) \) and \( (4f\phi_u)^2 \), which result in molecular terms with angular momenta \( L_z = 0 \) and \( 6 \). The six calculated low-lying levels again are almost degenerate having approximately the same spectroscopic parameters (without the spin-orbit interaction): \( R_e = 2.63 - 2.66 \) Å, \( \omega_e = 166 - 189 \) cm\(^{-1} \), \( D_e = 2.61 - 2.68 \) eV (The CASPT2 level of theory and a basis set of VQZP quality). The spin-orbit coupling has only small effect on the computed properties, in particular it reduces \( D_e \) to 2.51 eV in good agreement with the experimental estimate \( D_0 = 2.57 \) eV [from the third law]. Calculations of Roos et al. give a longer \( R_e \) and a softer \( \omega_e \) in comparison with the values of Cao and Dolg \(^4\). Both theoretical estimates of \( \omega_e \) \(^5\) \(^6\) are smaller than the experimental result \(^6\): \( \omega_e = 245.4 \pm 4.2 \) cm\(^{-1} \) [the force constant \( k_e = 2.28(1) \) mdyne/Å].

Thus, the standard well-known many-electron procedures \(^4\) \(^5\) correctly describe the Ce₂ chemical bonding but find six nearly degenerate low-lying molecular terms \( ^1\Sigma^+, ^1\Sigma^-, ^3\Sigma^+, ^3\Pi^+_g, ^3\Pi_g, ^1\Pi_g \) instead of a well defined ground state. The difficulty lies in the smallness of the splitting energies of these excitations requiring a special treatment. In the following we formulate a simplified
valence bond (VB) model accounting for the 4f electrons in the cerium dimer, find the important interactions and obtain the “fine” energy spectrum of the electron excitations. Our approach is based on the picture of 4f-4f interactions as rightly found in [4, 5], but requires a more deep insight in the couplings which experience the 4f electrons in the molecule.

II. MODEL

We start with the description of the triple bond of Ce₂. The ground state electronic configuration of the cerium atom is (4f)\(^1\)(5d)\(^1\)(6s)\(^2\) with the \(^1\)G\(_4\) ground term. This implies that the Ce atom is a rare case of the first Hund rule violation [8]. The nature of such behavior lies in “quartic Fermi hole” which effectively separates the valence electrons and decreases their Coulomb repulsion [8]. Already at this stage of the investigation it is clear that the 4f electron is not independent, through the Coulomb repulsion it is strongly coupled with the valence 5d and 6s electrons on the same cerium site.

However, the (4f)\(^1\)(5d)\(^1\)(6s)\(^2\) configuration with the closed 6s-shell is not favorable for chemical bonding. Therefore the electronic atomic state of Ce is first transformed to the (4f)\(^1\)(5d)\(^2\)(6s)\(^1\) configuration with the \(^5\)H\(_3\) term situated at 0.299 eV above the \(^1\)G\(_4\) ground term [5]. In the (4f)\(^1\)(5d)\(^2\)(6s)\(^1\) configuration both 5d and 6s shells become open and form the triple bond with six valence electrons: (6sσ\(^2\))(5dσ\(^4\)). The \(^1\)Σ\(_g^+\) bonding state of 5d and 6s electrons (without the 4f states) is well understood [4, 5], in particular it occurs in La₂ where the molecular characteristics (\(D_0 = 2.5\) eV, \(k_c = 2.28\) (1 mdyne/Å [8], the (5d)\(^1\)(6s)\(^2\) initial and (5d)\(^2\)(6s)\(^1\) final [at 0.33 eV] configuration) are very close to those in Ce₂. Although the triple bond is strong, the total energy gain is weakened by 0.6 eV because of the necessity to promote one 6s-electron to the 5d-shell for both cerium atoms.

We consider the first cerium atom at (0,0,0) and the second at (0,0,\(R_c\)) (so that the z-axis is the molecular axis) and find the valence density distribution of the four 5d and two 6s valence electrons. The \(^1\)Σ\(_g^+\) 5d–6s ground state implies an axially symmetric electron density. The conclusion can also be proved by considering the density distributions of two s and four d-electrons in the (6sσ\(^2\))(5dσ\(^4\)) configuration.

Now to the 5d and 6s valence states we add two 4f-electrons. The main assumption supported by [4, 5] is that to a first approximation the 4f electrons do not contribute to the chemical bonding and remain localized. Then it follows that each 4f electron should be either in the \(Y_{1=3}^{\pm 3}\) (\(\Theta, \varphi\) space) or in the \(Y_{3}^{\pm 3}\) (\(\Theta, \varphi\) space) angular orbital state where \(C = -\sqrt{10/\pi}/8\). (Here we deal with the real spherical harmonics \(Y_{l}^{m,t}\) (\(t = c\) for cosine- and \(t = s\) for sine-like azimuthal (\(\varphi\)-dependence) defined in Ref. [10]). That is because in both these states the Coulomb intrasite repulsion between the 4f electron and the bonding 5d and 6s states at the cerium site is minimal (see Appendix A). Indeed, in these \(m = \pm 3\) f—states the angular density dependence is \(\sin^{3}\Theta\) with the maximum at \(\Theta = \pi/2\), while in the binding 5d states maximum lies at \(\Theta = \pi/4\) and in 6s states at \(\Theta = 0\) (considering their overlap). Thus, the occupation of the \(m = \pm 3\) f—states ensures a certain space separation of the populated 4f states from the binding 5d and 6s electrons, which effectively minimizes the intrasite 4f–6s and 4f–6s Coulomb repulsions. The situation is discussed in detail in Appendix A, notice that the conclusion about the occupancy of the 4f\(\phi\) states is also explicitly supported by the calculations of Roos et al. [5] and implicitly by Cao and Dolg [4].

Thus, at each cerium site the 4f electron space is limited to two \(m = \pm 3\) orbitals, see Fig. 1.

\[
\begin{align*}
\psi_{f,c}(\Theta, \varphi, r) &= R_f(r)Y_{1=3}^{3,c}(\Theta, \varphi), \quad (1a) \\
\psi_{f,s}(\Theta, \varphi, r) &= R_f(r)Y_{1=3}^{3,s}(\Theta, \varphi). \quad (1b)
\end{align*}
\]

Here \(R_f(r)\) is the radial part of the 4f states.

Since there is very little overlap between 4f functions belonging to different sites, the valence bond (VB or Heitler-London) [11, 12] approach should give the best description to the 4f electronic problem. Its basis set consists of the two electron 4f states:

\[
\Psi_{\alpha,\alpha}(\vec{r}_1, s_{1z}; \vec{r}_2, s_{2z}) = \psi_{f,\alpha}(\vec{r}_1)u_{1s}\psi_{f,\alpha}(\vec{r}_2)u_{2s}, \quad (2)
\]

where the indices \(\alpha\) and \(\alpha\) refer to two orbital sites at sites \(a\) and \(b\), i.e. \(\alpha, \alpha = c\) or \(s\) (orbital functions defined in Eq. (1a,b) and visualized in Fig. 1), \(s_z\) is the spin projection number (±1/2), \(u_s\) is the corresponding spin function (\(\alpha\) or \(\beta\)).

The two electron wave function \(\Psi_{\alpha,\alpha}(\vec{r}_1, s_{1z}; \vec{r}_2, s_{2z})=\Psi_{\Omega}(\vec{r}_1, \vec{r}_2)\Psi_{S}(s_{1z}; s_{2z})\) is a product of the orbital part \(\Psi_{\Omega,i}\) (\(i = 1 – 4\)), which can be labeled as

\[
|c, s\rangle, \ |s, c\rangle, \ |c, c\rangle, \ |s, s\rangle, \quad (3a)
\]

and the spin function \(\Psi_{S,j}\) (\(j = 1 – 4\)),

\[
(\alpha, \alpha), \ (\alpha, \beta), \ (\beta, \alpha), \ (\beta, \beta). \quad (3b)
\]

(Here we omit cerium site indices \(a\) and \(b\), that is \(|c, s\rangle = |c_a, s_b\rangle\) etc.) Therefore, in total there are 16 two electron
basis states \( t \) (\( t = 1-16 \)), which are then antisymmetrized in respect to the electron permutation.

In atomic units the corresponding valence bond Hamiltonian \([11, 12]\) is given by

\[
H^{VB} = -\frac{1}{2} \nabla_1^2 - \frac{1}{2} \nabla_2^2 + U(\mathbf{r}_1) + U(\mathbf{r}_2) + \frac{1}{\mathbf{r}_{12}} + H^{so},
\]

where \( H^{so} \) is the spin-orbit coupling considered in detail below (Sec. 11C), while

\[
U(\mathbf{r}) = U_a(\mathbf{r}) + U_b(\mathbf{r}).
\]

Here \( U_{a,b}(\mathbf{r}) < 0 \) is an effective 4f attraction to the nucleus \( a \) (or \( b \)) screened by all core electrons.

It is well known that the evaluation of the matrix elements of \( H^{VB} \) in the space of the functions \( \{|\mathbf{r},s\rangle\} \) leads to the following equation:

\[
\langle t_1 | H^{VB} | t_2 \rangle = E_0 \delta_{t_1,t_2} + J_{t_1,t_2} + K_{t_1,t_2}.
\]

where \( E_0 = E[(6s\sigma)^2(5d\pi)^4] \) is the background molecular energy, \( \delta \) is the Kronecker delta, while \( J \) and \( K \) are the direct and exchange matrix elements which we consider in Sec. 11A and Sec. 11B respectively.

Taking into account the orbital degeneracy of 4f states the matrix elements of \( H \) (16×16) where \( H = J, K \) or \( H^{so} \) are conveniently written in the spin block form,

\[
H = \begin{pmatrix}
H_{11} & H_{12} & H_{13} & H_{14} \\
H_{21} & H_{22} & H_{23} & H_{24} \\
H_{31} & H_{32} & H_{33} & H_{34} \\
H_{41} & H_{42} & H_{43} & H_{44}
\end{pmatrix},
\]

where the indices \( s_1, s_2 \) refer to the spin part, Eq. (31). As usual, \( H_{12} = H_{12}^{\dagger} \) etc. Each of the blocks in Eq. (6) can be further represented as a 4×4 orbital matrix of the following structure:

\[
H_{s_1,s_2} = \begin{pmatrix}
a_1 & c_1 & d_1 & e_1 \\
c_1 & a_1 & d_2 & e_2 \\
d_1 & d_2 & a_2 & c_2 \\
e_1 & e_2 & c_2 & a_2
\end{pmatrix},
\]

Here the matrix elements are evaluated between the two 4f–orbital sets \( \{i_1,i_2\} \), Eq. (33) with the fixed spin indices \( \{s_1, s_2\} \), Eq. (25). In our basis with real spherical harmonics, \( a_i, b_i, c_i \) are real, while \( d_i \) and \( e_i \) are imaginary values describing the interaction with an external magnetic field or the spin-orbit coupling. The specific form of the matrix elements is considered below.

A. 4f–4f direct Coulomb repulsion

Here we consider the matrix of the direct Coulomb repulsion:

\[
J_{t_1,t_2} = \int \psi_{t_1}\psi_{t_2}(\mathbf{r}_1)\psi_{t_1}\psi_{t_2}(\mathbf{r}_2) V(\mathbf{r}_1, \mathbf{r}_2) d\mathbf{r}_1 d\mathbf{r}_2,
\]

where \( V(\mathbf{r}_1, \mathbf{r}_2) = 1/|\mathbf{r}_1 - \mathbf{r}_2| \) and the orbital indices \((a_1, a_2)\) belong to the composite index \( t_1 \), while \((o_3, o_4)\) to \( t_2 \), Eq. (2).

From symmetry arguments it can be shown (Appendix B) that there are three types of the matrix elements,

\[
A = \langle c, s | V | c, s \rangle = (c_a, c_a | V | s_b, s_b) = (s_a, s_a | V | c_b, c_b),
B = \langle c, c | V | c, c \rangle = (c_a, c_a | V | c_b, c_b) = (s_a, s_a | V | s_b, s_b),
C = \langle c, s | V | s, c \rangle = (c_a, s_a | V | c_b, s_b) = (c_a, s_a | V | s_b, c_b),
\]

while

\[
\langle c, c | V | s, c \rangle = (c_a, c_a | V | c_b, s_b) = (s_a, s_a | V | c_b, s_b) = 0.
\]

Here we use both standard and so called chemist’s notations [10]. However, it turns out that not all parameters in [8] are independent. Analyzing the azimuthal dependence of the integrands and “the azimuthal selection rule” (Appendix B), one finds

\[
B - A = 2C.
\]

In the \( H = J \) case in Eq. (10) only the diagonal blocks \( J_{s\bar{s}} \) are relevant \((J_{s_1,s_2} \neq 0 \) for \( s_1 = s_2) \) because the matrix elements \( \langle 4 \rangle \) are independent of the spin indices.

Furthermore, the \( J_{a\bar{a}} \) blocks are identical, i.e. \( J_{a\bar{a}} = J_{s\bar{s}} \), Eq. (7). The \( J_{a\bar{a}} \) matrix is given by Eq. (7) where \( a_1 = A, a_2 = B, c_1 = c_2 = C \) and \( d_1 = d_2 = e_1 = e_2 = 0 \).

B. 4f–4f exchange interaction

For the matrix element of the 4f–4f exchange interaction we obtain

\[
K_{t_1,t_2} = -\langle S_{a_1,a_2} U_{a_1,a_2} S_{b_1,b_2} + V_{exx}^{a_1,a_2;b_1,b_2} \rangle d_{t_1}, d_{t_2}, d_{s_1}, d_{s_2}, d_{s_3}, \]

where the composite indices \( t_1 \) and \( t_2 \) refer to \((o_1,a_1,s_1);(o_2,b_2,s_2)\) and \((o_3,a_3,s_3);(o_4,b_4,s_4)\) as described earlier, Eq. (2), and

\[
S_{a_1,a_2} = \int d\mathbf{r} \psi_{o_1,a_1}(\mathbf{r}) \psi_{o_2,a_2}(\mathbf{r}),
\]

\[
U_{a_1,a_2} = \int d\mathbf{r} \psi_{o_1,a_1}(\mathbf{r}) U(\mathbf{r}) \psi_{o_2,a_2}(\mathbf{r}).
\]

Finally, the Coulomb exchange integral in [11] is given by [compare with \( J_{t_1,t_2} \) in Eq. (8)]

\[
V_{exx}^{a_1,a_2;b_1,b_2} = \int \psi_{o_1,a_1}(\mathbf{r}_1) \psi_{o_2,b_2}(\mathbf{r}_2) V(\mathbf{r}_1, \mathbf{r}_2) \psi_{a_2,b_2}(\mathbf{r}_2) \psi_{o_1,a_1}(\mathbf{r}_1) d\mathbf{r}_1 d\mathbf{r}_2.
\]

If two first terms on the right hand side of [11] differ from zero then they are leading, while the third (Coulomb) term is just a minor correction.
In our case there are two 4f functions at each site, Eq. (1a)-(1b), but from symmetry it follows that in all overlap integrals $S_{s_1,s_2}^{a,b}$ there is only one nonzero value,

$$S = \langle c_{a} | c_{b} \rangle = \langle s_{a} | s_{b} \rangle > 0,$$

while $\langle c_{a} | s_{b} \rangle = 0$. Analogously, $U_{\text{exc}} = \langle c_{a} | U_{\text{exc}} | c_{b} \rangle = \langle s_{a} | U_{\text{exc}} | s_{b} \rangle$ and $\langle c_{a} | U_{\text{exc}} | s_{b} \rangle = 0$. It is convenient to introduce the notation

$$\delta E = -2S U_{\text{exc}} > 0$$

for the leading term.

For the exchange Coulomb interaction $V_{\text{exc}}$, Eq. (12), one finds three types of integrals with nonzero values (compare with Eqs. (7)):

$$A' = \langle c, s | V | c, s \rangle_{\text{exc}} = \langle c_{a} c_{b} V | s_{a} s_{b} \rangle,$$

$$B' = \langle c, c | V | c, c \rangle_{\text{exc}} = \langle c_{a} c_{b} V | c_{a} c_{b} \rangle,$$

$$C' = \langle c, s | V | s, c \rangle_{\text{exc}} = \langle c_{a} s_{b} V | c_{a} s_{b} \rangle.$$

Furthermore, there is the same relation as for the direct matrix elements $A$, $B$ and $C$, Eq. (10), see Appendix B:

$$B' - A' = 2C'.$$

Then, in the full 16x16 matrix $H = K$, Eq. (9), the exchange coupling is effective in the blocks $K^{11\gamma}_{\text{exc}}$ (between states $\alpha, \alpha$), $K^{23\varkappa}_{\text{exc}}$ and $K^{32\varkappa}_{\text{exc}}$ (between the states with $\alpha, \beta$ and $\beta, \alpha$) and $K^{44\varkappa}_{\text{exc}}$ (between states $\beta, \beta$). Each of the blocks has the following (orbital) form:

$$K^{\text{exc}}_{s_1, s_2} = \begin{pmatrix} -C' & \delta E - A' & 0 & 0 \\ \delta E - A' & -C' & 0 & 0 \\ 0 & 0 & \delta E - B' & -C' \\ 0 & 0 & -C' & \delta E - B' \end{pmatrix}.$$

(Here as in Ref. 11 we have taken into account the effect of the half canceling of the $\delta E$-terms in the final expression due to the nonorthogonality of the $f$-functions.)

C. Spin-orbit coupling

Now in the framework of our model we consider the 4f spin-orbit interaction:

$$H_{\text{so}} = \xi \bar{L}_{1} \bar{s}_{1} + \xi \bar{L}_{2} \bar{s}_{2},$$

where $\xi = \xi_{f} \approx 862$ K. This value corresponds to the $\Delta_{\text{so}} = (7/2)\xi = 0.26$ eV splitting between $j = 5/2$ and $7/2$ one electron atomic 4f states. (The energies of the states were obtained by solving the spherically symmetric Dirac equation for atomic cerium states in the framework of the self-consistent field DFT-LDA atomic model.) In our model the 4f quantum space is reduced to two orbital states with $m_{z} = \pm 3$ at each site, and $H_{\text{so}}$ transforms to

$$H_{s}^{\text{so}} = \xi \bar{L}_{1,z} \bar{s}_{1,z} + \xi \bar{L}_{2,z} \bar{s}_{2,z}.$$

TABLE I: Calculated 4f direct (A, B, C) and exchange (A', B', C') Coulomb integrals, the overlap integral $S$, the VB attraction $U_{\text{exc}}$ and the 4f exchange parameter $\delta E$ for various Ce-Ce distances $R$ (in Å). All energy parameters are in Kelvin (K).

| $R$ (Å) | $S$ | $U_{\text{exc}}$ | $\delta E$ |
|--------|-----|----------------|----------|
| 2.62   | 0.07| -14.6          | -14.01   |
| 2.63   | 0.07| -14.6          | -14.01   |
| 2.66   | 0.07| -14.6          | -14.01   |
| 2.70   | 0.07| -14.6          | -14.01   |
| 3.43   | 0.07| -14.6          | -14.01   |
| 3.65   | 0.07| -14.6          | -14.01   |

Therefore, the nonzero spin blocks of the full matrix are $H_{s,s}^{\text{so}} (s = 1 - 4)$, Eq. (18), with the following general structure of each block (see Appendix C):

$$H_{s,s}^{\text{so}} = \frac{3}{2s} \begin{pmatrix} 0 & 0 & n_{11} & n_{21} \\ 0 & 0 & m_{11} & m_{21} \\ -n_{11} & -m_{11} & 0 & 0 \\ -n_{21} & -m_{21} & 0 & 0 \end{pmatrix}.$$

In the $H_{11}^{\text{so}}$ block $n_{11} = m_{11} = 1, n_{21} = m_{21} = -1$, in the $H_{22}^{\text{so}}$ block $n_{12} = m_{12} = -1, n_{22} = m_{22} = 1$, in the $H_{33}^{\text{so}}$ block $n_{13} = m_{13} = 1, m_{13} = m_{23} = -1$ and in $H_{44}^{\text{so}}$ block $n_{14} = m_{14} = 1, n_{24} = m_{24} = -1$.

The spin-orbit coupling, Eq. (19), splits the four single electron states at each cerium site in two doublets: the lowest level with $j_{z} = \pm 5/2$ and the energy $E(\pm 5/2) = -3\xi / 2$ has the eigenfunctions $\psi_{5/2, +} \sim \exp(3mi\phi)\beta$, $\psi_{5/2, -} \sim \exp(-3mi\phi)\alpha$. The other level with $j_{z} = \pm 7/2$ and the energy $E(\pm 7/2) = +3\xi / 2$ has the functions $\psi_{7/2, +} \sim \exp(3mi\phi)\alpha$, $\psi_{7/2, -} \sim \exp(-3mi\phi)\beta$. Correspondingly, two 4f electrons have the following energy spectrum: $E_{s,\alpha,1} = -3\xi / 2$ (4 degenerate states), $E_{s,\beta,2} = 0$ (8 states), $E_{s,\alpha,3} = 3\xi / 2$ (4 states). The inclusion of other interactions modifies the picture.

III. RESULTS AND DISCUSSION

All interaction parameters of the model introduced in Sec. II were evaluated numerically with our valence bond program [13], Table II. There we used the overlapping cerium relativistic atomic 4f functions (the large $j = 5/2$ component for $R_{y}(r)$) calculated within the density functional theory (DFT) in the local density approximation (LDA) [14].

Below we discuss the results of the valence bond model.
with \(H^{so} \neq 0\) and without \(H^{so} = 0\) the spin-orbit coupling.

### A. VB without the spin-orbit coupling

From Table I we see that the largest parameters are \(A\) and \(B\) which correspond to the direct Coulomb repulsion, Sec. II A. Therefore, as a first approximation we consider only the matrix of the direct Coulomb repulsion \(J\). In that case from the structure of the spin matrix it follows that the triplet and singlet states coincide. Having diagonalized the corresponding \(4 \times 4\) orbital matrix analytically, we find its eigenvalues,

\[
E_1 - A = -C, \quad E_2,3 - A = C, \quad E_4 - A = 3C.
\]  

(The eigenvectors are quoted in Eqs. (20a)-(20c), Appendix C.) We conclude that the energy splittings are determined by the parameter \(C\), Eq. (9), which is very small, while \(A\) should be incorporated into the background molecular energy \(E_0 = E[(6s\sigma)^2(5d\pi)^4]\).

Notice that the eigenvalues \(E_2\) and \(E_3\) are degenerate which implies a nonzero orbital moment. In Appendix C we show that indeed the two components are attributed to \(M_z = \pm \hbar\). Taking into account other properties of the eigenvectors one can show that the molecular terms are identified as \(\Sigma^+\), \(6g\) and \(\Sigma^0\), respectively. However, the presented picture is not full because the integral \(C\) is very small, Table I. If \(C\) were of the order of \(A\), the direct Coulomb splitting would have been dominant, but since \(C \approx 0.2\) \(K\) we should consider other relevant interactions.

Now in addition to the direct Coulomb repulsion \(J\) we consider the \(4f-4f\) exchange interaction \(K\), Sec. II B. A small overlap \(S\) between two relevant \(4f\) functions splits the singlet and triplet states. If there were only one orbital for each site, the spin singlet state would have been lower in energy than the spin triplet state. Indeed, in the single orbital valence bond problem \(11,12\) we have

\[
E_s = E_0 - \delta E, \quad E_T = E_0 + \delta E,
\]  

where \(\delta E\) is given by Eq. (14). In our case of two \(4f\) orbitals at each site the reverse is true. This can be clearly seen if for the time being we ignore the exchange repulsion \((A' = B' = C' = 0)\), and treat the electron site exchange as a perturbation. As an example we consider the state \(21a\) with the eigenstate \((C7a)\). Its orbital part \(\Sigma^+\) is not a perturbation. The energy levels \(E_0\) for all states are summarized in Table II, which holds even if \(\delta E\) is not a perturbation. The energy levels are also schematically shown in Fig. 4.

Thus, the situation in \(Ce_2\) could be an interesting example of the triplet ground state. In practice however the relatively strong (in comparison with \(\delta E\)) spin-orbit coupling changes the electron spectrum.

### B. VB with the spin-orbit coupling

In this section we discuss the most general case, when the full valence bond Hamiltonian \(H^{VB}\) is given by Eq. (13).

Surprisingly, an analytical solution can be figured out even for this case except for the small Coulomb repulsion which accompanies the \(4f\) exchange transitions \([i.e. \ A' = 0, B' = 0, C' = 0, Eq. (15a)-(15c)]\). The analytical expressions for the energy levels are quoted in Table III. Notice that the three lowest states are those found in Ref. 4 \([\Omega = (\Omega_1, 0, 5\Omega_1)]\) and Ref. 8 (Table 6).

For completeness we can include the \(4f\) electron exchange repulsion \((A', B', C')\) and compute the energy levels numerically. The calculated energy values for characteristic internuclear distances \(R\) and the parameters of Table I are quoted in Table IV. Since \(A', B'\) and \(C'\) are very small, they introduce only a minor correction to the analytical expressions.

| \(\Sigma^+\) | \(\Sigma^0\) | \(\Sigma^-\) |
|----------|----------|----------|
| \(\Sigma^+\) | \(-C + \delta E\) | \(3C - \delta E\) |
| \(\Sigma^0\) | \(C - \delta E\) | \(C + \delta E\) |
| \(\Sigma^-\) | \(3C - \delta E\) | \(3C + \delta E\) |

### Table II: The analytical expressions for the energy levels of \(Ce_2\) in the VB model with the spin-orbit coupling \((A' = 0, B' = 0, C' = 0)\). \(P\) is the state parity.

| \(E\) | \(\delta E\) | \(J_u\) | \(L_z\) | \(S_z\) | \(P\) |
|-------|----------|----------|----------|----------|----------|
| 1     | \(-\sqrt{9\Omega^2 + 4C^2}\) | \(C - \delta E\) | 1 | 0 | 0 | 0 | \(g\) |
| 2     | \(-\sqrt{9\Omega^2 + 4C^2}\) | \(C + \delta E\) | 1 | 0 | 0 | 0 | \(u\) |
| 3     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(g\) |
| 4     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(u\) |
| 5     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(g\) |
| 6     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(u\) |
| 7     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(g\) |
| 8     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(u\) |
| 9     | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(g\) |
| 10    | \(\Omega\) | \(\Omega\) | 1 | 0 | 0 | 0 | \(u\) |
Usually, for labeling of molecular states one uses the standard many-electron spectroscopic notation \( ^{2S+1}A_\Omega \). However, in the \( \Sigma = S_z = 0 \) states the inclusion of the spin-orbit coupling mixes up singlet \((\Sigma = 0)\) and triplet \((\Sigma = 1)\) states and the partial weights of singlet and triplet contributions in the final \( S_z = 0 \) states are nearly equal. (The effect was found in \( ^4 \Sigma \) \( ^5 \Sigma \) ) In our model for the lowest three states we obtain:

\[
\Psi_1(g_s) = a\Psi(3\Sigma_g^-; S_z = 0) - b\Psi(1\Sigma_g^+; S_z = 0), \quad (23a)
\]

\[
\Psi_2 = -a\Psi(1\Sigma_u^-; S_z = 0) + b\Psi(3\Sigma_u^+; S_z = 0), \quad (23b)
\]

\[
\Psi_3,1 = a\Psi(3\Sigma_u^-; S_z = 1) + b\Psi(3\Sigma_u^+; S_z = 1), \quad (23c)
\]

\[
\Psi_3,2 = b\Psi(3\Sigma_u^-; S_z = -1) + a\Psi(3\Sigma_u^+; S_z = -1). \quad (23d)
\]

Here \( a = 0.7072 \), \( b = 0.7070 \), \( \Psi(3\Sigma_g^-; S_z = 0) \) is the \( S_z = 0 \) component of the \( 3\Sigma_g^- \) triplet etc. Notice that the \( 3\Sigma_u^- \) and \( 3\Sigma_u^+ \) states in \( \Psi_3 \) form the \( 3\Sigma_u \) level \((\Omega = 5)\), see Fig.2 below. In the following we will use the short notation \((1,3)\) for the spin multiplicity \((2S + 1)\) of the strongly mixed states. For example, the ground state is labeled as \( ^{3}\Sigma_{0}(g) \) \((or \ ^{3}\Sigma_{0}(g) \) etc.

The exchange Coulomb interaction \((A', B', C' \neq 0)\) splits the doublets 5 and 6 \((\Omega = \Lambda = 6, \Sigma = 0, g,u)\), Table I

\[
1^{3}6_{0}(g,u) \rightarrow 1^{3}\Sigma_{0}(g,u) + 1^{3}\Sigma_{0}(g,u). \quad (24)
\]

In the split lines \((5a, 5b, 6a, 6b)\) in Table VI \(\Omega = 0, \Lambda = 0 \) and \(\Sigma = 0\). Although the ordering of the three lowest levels is the same as in Ref. 4, the energy splittings are different. At \( R = 2.62 \) \(\text{Å} \) we have \( E_2 - E_1 = E_3 - E_1 = 32.7 \) \( K \), while in the spin-orbit calculation of Cao and Dolg \( E_2 - E_1 \leq 1 \) \( \text{cm}^{-1} \) and \( E_3 - E_1 \leq 2 \) \( \text{cm}^{-1} \).

It is worth noting that the levels \( 2, 3 \) and \( 9, 10 \) are virtually degenerate. According to Table III the energy difference between them is \( 2C^2/\zeta \ll (\zeta, \delta E) \).

If the spin-orbit coupling is dominant (which is the case for \( Ce_{2} \)), the twelve energy levels are clearly divided in three groups:

\[
\begin{align*}
(1) & \quad ^{3}1\Sigma_{0}(g), ^{1}3\Sigma_{u}(u), ^{3}6_{0}(u). \quad (\sim E_{so,1}) \quad (25a) \\
(2) & \quad ^{3}1\Sigma_{1}(g,-), ^{1}3\Sigma_{0}(g,+), ^{1}3\Sigma_{0}(g,-), ^{1}3\Sigma_{0}(u,-), ^{3}\Sigma_{0}(u,+), ^{3}\Sigma_{0}(u,-). \quad (\sim E_{so,2}) \quad (25b) \\
(3) & \quad ^{3}1\Sigma_{0}(g), ^{3}6_{7}(u), ^{3}3\Sigma_{0}(u). \quad \quad (\sim E_{so,3}) \quad (25c)
\end{align*}
\]

Here the levels are given in the increasing energy order as in Tables III and IV with some additional characteristics \((the \ parity \ and \ the \ mirror \ reflection, \ when \ possible) \) in parentheses. If the other interactions are omitted \((i.e. \delta E = 0, C = 0, \ and \ A' = B' = C' = 0)\) then the energy spectrum is reduced to only three levels \((E_{so,1} = -3\zeta, E_{so,2} = 0, E_{so,3} = +3\zeta)\) discussed in Sec. II C.

The relation between three important characteristic parameters of \( Ce_{2} \) is the following \( \zeta \gg \delta E \gg C \). However, the analytical and numerical solution can be used to study the model for any parameter set. It is clear that such a scenario is not relevant for \( Ce_{2} \), but it can possibly occur in other electronic systems with localized \( f \) electrons. In particular, it is instructive to consider the case of a small spin-orbit coupling: \((C, \delta E) \gg \zeta \). For that relation \((provided \ that \ A' = B' = C' = 0)\) one can still use the analytical solution, Table III but of course the order and level grouping will be different. The splittings and relations between energy levels in the absence of the spin-orbit coupling \((\zeta = 0)\) are schematically shown in Fig.2.

In Fig. 2, the most realistic subcase \((all \ relevant \ interactions \ included)\) is \((B+C)\). If on top of it a small spin-orbit coupling is introduced, then the spin triplet states there are split according to the following scheme:

\[
\begin{align*}
^3\Sigma_{0} & \rightarrow ^3\Sigma_{0}(g,-) + ^3\Sigma_{1}(g,-), \quad (26a) \\
^3\Sigma_{u} & \rightarrow ^3\Sigma_{0}(u,+), ^3\Sigma_{0}(u,-), \quad (26b) \\
^3\Sigma_{u} & \rightarrow ^3\Sigma_{0}(u,+), ^3\Sigma_{1}(u,-), \quad (26c) \\
^3\Sigma_{u} & \rightarrow ^3\Sigma_{0}(u,+), ^3\Sigma_{1}(u,-). \quad (26d)
\end{align*}
\]
IV. CONCLUSIONS

For the cerium dimer we proposed a model of two interacting 4f electrons which can explain and refine the low-lying energy spectrum of Ce$_2$ observed in many electron computational studies [4,5]. The model is a modified valence bond approach where two relevant 4f orbital states at each cerium site are explicitly taken into account, Eqs. (1a), (1b), and Fig. 1. The other valence 6s and 5d states are grouped in a conventional triple chemical bond with the (6s$\sigma_g^2$(5d$\pi_u^4$) closed molecular shell.

In the framework of the 4f model we can clearly trace the origin of the energy levels and establish intrinsic relations between them. For the most important interactions the problem of finding the 4f energy spectrum is solved analytically. Apart from the spin-orbit coupling ($\zeta = 862$ K) the largest interaction is the direct Coulomb repulsion of the two localized 4f electrons (described by parameters $A, B$) but the effective Coulomb parameter responsible for the energy splitting is very small ($C \sim 0.2$ K), Table 1. The next important interaction is the valence bond exchange of the two localized 4f electrons ($\delta E \sim 14$ K) which is also accompanied by small exchange Coulomb repulsion ($A', B' \sim 1.5$ K, $C' \sim 0.07$ K).

As a result, the even ground state level with $\Omega = \Lambda = \Sigma = 0$ (composed of $3\Sigma_0(g)$ and $1\Sigma_0(g)$ states, Eq. (23a)) is separated from first excited levels with $\Omega = \Lambda = \Sigma = 0$ (composed of $3\Sigma_0(u)$ and $1\Sigma_0(u)$ states, Eq. (23b)) and $4\Sigma_5(u)$ by an energy gap of $\sim 30$ K. The ordering of these levels is in agreement with calculations of Cao and Dolg [4], although the energy splittings are different. The $J_z = \Omega = \pm 5$ excited states support magnetic moments, giving rise to the magnetic susceptibility dependence $\chi$ shown in Fig. 3. The magnetic susceptibility clearly displays two different regimes. At large temperatures ($T > 30$ K) it follows the Curie law indicating of “free” magnetic moments. At small temperatures ($T < 20$ K) $\chi$ deviates from the Curie law and rapidly decreases as $T \to 0$. Such low $T$ behavior and disappearance of the “free” 4f magnetic moments of cerium is caused by the population of the nonmagnetic ground state ($\Omega = \Lambda = \Sigma = 0$) at the expense of the excited magnetic ones. The conclusion can be checked by experimental measurements.

We have also studied the energy spectrum in the absence of the spin-orbit coupling, Fig. 2. Here, the unusual feature is the $3\Sigma_g^-$ triplet ground state.

The peculiarities of the electronic structure of the simplest cerium bond possibly can clarify the picture of phase transitions in solid cerium [18].

I am grateful to K.H. Michel, B.N. Plakhutin, A.V. Tsyvashchenko, E.V. Tkalya, A.V. Avdeenkov and B. Verberck for valuable discussions.

Appendix A: Appendix A

According to the recent many electron calculations (CASSCF) of Ce$_2$ [5] each cerium site has a single 4f electron which is virtually nonbonding and preserves its localized nature. Furthermore, without the spin-orbit coupling it occupies one of the two orbitally degenerate 4f states with the highest azimuthal angular momentum ($m = \pm 3$). Although it is not mentioned explicitly, the same orbital degeneracy follows from the low-lying molecular term spectrum computed by Cao and Dolg [4]. Below we show that the lowering of the $m = \pm 3$ states is caused by the Coulomb repulsion between the 4f states and the bonding electron states.

We consider a simplified model with two 5d and one 6s bonding electrons at each cerium site and use the method of multipole interaction [19]. According to [4,5] one 6s electron and two 5d electrons from each Ce site contribute to the (6s$\sigma_g^2$(5d$\pi_u^4$) triple bond. That implies the occupation of the $Y_{1c}^{1c}$ and $Y_2^{1s}$ orbital d-states ($m = \pm 1$). The resultant electron density $((Y_2^{1s}(\Theta, \phi))^2 + (Y_2^{1c}(\Theta, \phi))^2)$ is axially symmetric. Expanding it in the multipolar series one finds that the first nonspherical component responsible for splittings is given by the $Y_{l=2}^0$ angular dependence of the 5d-state density, $\rho_{d,t=2}(r, \Theta, \phi) = 2C_{d,t=2}^0R_d^2(r)Y_{l=2}^0(\Theta, \phi)$, (A1)

where $R_d(r)$ is the 5d atomic function radial dependence, and $C_{d,t=2}^0 = 0.090112$ is the angular density matrix element [18].

$$C_{d,t=2}^0 = \langle 2, 1s|2, 0|2, 1s \rangle = \langle 2, 1c|2, 0|2, 1c \rangle$$

$$= \int d\Omega Y_{l=2}^{1c}(\Omega)Y_{l=2}^0(\Omega)Y_{l=2}^{1c}(\Omega). \quad (A2)$$

Here $\Omega$ stands for the polar angles ($\Theta, \phi$). Now one can calculate the matrix of the Coulomb interaction of the
5d density with the 4f electron. (In principle it will have the \( \ell = 2 \) and \( \ell = 4 \) contributions \[19\], but the \( \ell = 4 \) term is small and in the following will be ignored.) The matrix is diagonal in the space of the 4f orbitals with the diagonal elements given by

\[
V_{f,\tau} = V_{df} C_{f,\ell=2}^0(\tau). \tag{A3}
\]

Here \( \tau = (m, c) \) or \((m, s) \) (\( m = 0 \) to 3) is the azimuthal dependence of the 4f orbital functions, \( C_{f,\ell=2}^0(\tau) = (3, \tau | 2, 0, 3, \tau) \) [compare with (A2)], while

\[
V_{df} = 2C_{f,\ell=2}^0 v_{2,df}, \tag{A4a}
\]

\[
v_{2,df} = \frac{4\pi}{5} \int dr dr' \frac{R_d^2(r')}{r'^2} \frac{R_f^2(r)}. \tag{A4b}
\]

\( r_c (r_o) \) is the largest (the smallest) value of \( r \) and \( r' \).

\( v_{2,df} \) describes the quadrupolar \((\ell = 2)\) component of the 5d - 4f intrasite Coulomb repulsion \[19\]. With the 5d and 4f atomic radial dependencies (DFT-LDA atomic calculations \[14\]) we obtain \( v_{2,df} = 7.1054 \) eV and \( V_{df} = 1.2806 \) eV. The \( C_{f,\ell=2}^0(\tau) \) coefficients and the corresponding 4f energy splittings are quoted in Table V (Notice that due to the z-axial symmetry \( C_{f,\ell=2}^0(m, c) = C_{f,\ell=2}^0(m, s) \)).

Thus, we conclude that the 5d - 4f Coulomb repulsion favors the lowering of the \( m = \pm 3 \) f-states. However, there are two other interactions which act in the opposite direction: the screened attraction to the other nucleus and the 5d - 4f exchange interaction.

The other nucleus attraction leads to an axially symmetric Coulomb potential which has the form of (A3), but with an interaction constant of the opposite sign. The 5d - 4f exchange interaction is more complex. In terms of the intrasite Coulomb multipole repulsion it corresponds to the odd \((\ell = 1, 3, 5)\) transition density contributions, and the dipolar \((\ell = 1)\) contribution is rather strong \[19\]. However, this exchange interaction leads to the high spin \((S = 2)\) ground state (the \(5^3H_3\) term of the 4f5d6s configuration) of the cerium atom, which in the real molecule is effectively suppressed by the triple bond formation with a zero spin state of the binding 5d and 6s states. Therefore, the bond formation mechanism diminishes the 4f - 5d exchange interaction, while the 4f - 5d Coulomb repulsion \[19\] remains and lowers the \( m = \pm 3 \) f-states in respect to the others \[4, 5\].

### Appendix B: Appendix B

The relevant Coulomb integrals, Eq. (10), can be simplified by considering their azimuthal parts, i.e. the azimuthal dependence of the corresponding \( \psi_{a1,a}(\vec{r}_1) \psi_{a1,a}(\vec{r}_2) \) functions standing on the right and their counterparts \( \psi_{a2,b}(\vec{r}_2) \psi_{a1,b}(\vec{r}_2) \) on the left from the Coulomb interaction \( V(\vec{r}_1, \vec{r}_2) = 1/|\vec{r}_1\vec{r}_2| \). In the following we define \( \vec{r}_a = \vec{r}_1 - \vec{R}_a \) and \( \vec{r}_b = \vec{r}_2 - \vec{R}_b \). The important observation is that the Coulomb integral differs from zero only if the azimuthal density dependencies on the left and on the right side have the same functional form. This conclusion can be derived by various ways, in particular it follows from the multipole expansion of the Coulomb interaction

\[
V(\vec{r}_a, \vec{r}_b) = \sum_{l,\tau} v_l(r_a, r_b) Y_l^\tau(\hat{r}_a) Y_l^{\tau}(\hat{r}_b). \tag{B1}
\]

Here the angular functions \( Y_l^\tau(\hat{r}_a) \) and \( Y_l^{\tau}(\hat{r}_b) \) have the same azimuthal dependence \((\tau)\) which is a trivial conclusion if they refer to the same site: \( \vec{R}_a = \vec{R}_b \). In case of two different sites (aligned along the z-axis) one can expand a spherical harmonic centered at one site in terms of the spherical harmonics centered at the other site \[13\]. It turns out that the procedure conserves the azimuthal dependence of the spherical harmonics translated along the \( z \)-axis \[13\], which proves our initial statement.

In the following in deriving relations between the Coulomb integrals, Eq. (10), we use chemist’s notations for the matrix elements \[10\]. Further, we explicitly single out the azimuthal dependence \( \rho_{a1, a}(\varphi) \) of the density functions \( \rho_{a1, a}(\vec{r}_a) \rho_{a1, a}(\vec{r}_b) \) with \( a_1, a_3 = c, s \), Eq. (1a,b). We then arrive at three different dependencies:

\[
\rho_{c,c}(\varphi) = (\cos 3\varphi)^2 = \frac{1}{2} + \frac{1}{2} \cos 6\varphi, \tag{B2a}
\]

\[
\rho_{s,s}(\varphi) = (\sin 3\varphi)^2 = \frac{1}{2} \frac{1}{2} \cos 6\varphi, \tag{B2b}
\]

\[
\rho_{c,s}(\varphi) = \cos 3\varphi \sin 3\varphi = \frac{1}{2} \sin 6\varphi. \tag{B2c}
\]

From the azimuthal selection rule it follows that \( \langle c_a, c_b|V|c_b, s_b \rangle = D = 0 \), Eq. (9) and (10), and \( \langle c_a, c_b|V|c_b, c_b \rangle = \langle s_a, s_a|V|s_b, s_b \rangle = B \).

In order to prove the relation (10) we consider the structure of the general Coulomb integral,

\[
(I_{a,b}^c|M_a|M_b) = \sum_M \int d\Omega f(\Theta) I_A(M_a|M_b), \tag{B3}
\]

where \( \Theta \) stands for \( (\Theta_a, \Theta_b, r_a, r_b) \), while \( I_A(M_a|M_b) \) is the azimuthal integral with two identical \( \varphi \)-dependencies \( M_a \) and \( M_b \). For example, \( I_3(6a_1|6b_1) \) with \( M_a = M_b = M = 6 \) stand for the azimuthal integral between \( \cos 6\varphi \) and \( \cos 6\varphi \) or between \( \sin 6\varphi \) and \( \sin 6\varphi \):

\[
I_{A,c}(M_a|M_b) = \int_0^{2\pi} d\varphi_a \int_0^{2\pi} d\varphi_b \cos(M\varphi_a)V(r_a, r_b) \cos(M\varphi_b). \tag{B4}
\]
Each term on the right hand side of (B3) has the following form:

\[
\int_Ω dΩ f(\Theta) I_A(M_a|M_b) = \int \sin Θ_a dΘ_a \int r_a^2 dr_a \int \sin Θ_b dΘ_b \int r_b^2 dr_b C^2 [P_3^c(\cos Θ_a)P_3^c(\cos Θ_b)]^2 
\]

\[
R_I^c(r_a) R_I^c(r_b) I_A(M_a|M_b). 
\]

(B5)

Here \( P_3^c(x) \) is the associated Legendre polynomial of two electron states (Eqs. (3a) and (3b)) can be obtained from these relations. Explicitly, the matrix of \( L_z \) has the nonzero blocks \( L_z(s,s) \) (s = 1 - 4) [see Eq. (B9)] of the form

\[
L_z(s,s) = \mu_B \begin{pmatrix} 0 & 0 & 3i & -3i \\ 0 & 0 & 3i & -3i \\ -3i & -3i & 0 & 0 \\ 3i & 3i & 0 & 0 \end{pmatrix}, 
\]

where \( \mu_B \) is the Bohr magneton.

Further, the 16 \times 16 spin matrix \( S_z \), Eq. (B1), has two nonzero diagonal blocks \( S_z(1,1) \) and \( S_z(4,4) \) of the form

\[
S_z(s,s) = ±\mu_B \hat{I}_z, 
\]

(C3)

where the plus sign refers to the \( \alpha, \alpha \) block \( (s,s = 1,1) \) and the minus sign to the \( \beta, \beta \) block \( (s,s = 4,4) \), while \( \hat{I}_z \) stands for the unit 4 \times 4 matrix.

The molecular interaction with an external magnetic field \( H \) is given by

\[
V_{mag} = -\vec{\mathcal{M}} \vec{H}, 
\]

(C4)

with the molecular magnetic moment

\[
\vec{\mathcal{M}} = 2\vec{S} + \vec{L}. 
\]

(C5)

For the magnetic field \( H \) applied along the \( z \)-axis we obtain

\[
V_{mag}^z = -\hbar(2S_z + L_z), 
\]

(C6)

where \( \hbar = H\mu_B \). The relations (C2), (C3) and (C6) are used to calculate the \( z \)-component of the orbital, spin and full momentum for each two-electron level.

For example, by means of Eq. (C2) we can find the orbital quantum number required for the characterization of the degenerate states, which are considered in Sec. II A. There we have discussed the three energy levels given by Eq. (21a)-(21c). The corresponding eigenvectors are

\[
(-1,1,0,0)/\sqrt{2}, 
\]

(C7a)

\[
(1,1,0,0)/\sqrt{2}, \quad (0,0,-1,1)/\sqrt{2}, 
\]

(C7b)

\[
(0,0,1,1)/\sqrt{2}. 
\]

(C7c)

In the presence of the external magnetic field the Hamiltonian \( V(f - f) \) of the direct Coulomb repulsion (see Sec. II A) is modified:

\[
H = E_{bond} + V(f - f) + H_L, 
\]

(C8)

where the last term is the orbital polarization given by \( H_L = -L_z h \) (the first term \( E_{bond} \) is the binding energy of \( Cc_2 \) which is not important for our discussion). From (C8) we obtain four eigenvalues \( (i = 1 - 4) \)

\[
E_i - E_0 = -C, \quad C - 6h, \quad C + 6h, \quad 3C. 
\]

(C9)

and the corresponding eigenvectors: \((-1,1,0,0)/\sqrt{2}, (i,i,-1,1)/\sqrt{2}, (i,i,1,-1)/\sqrt{2}, (0,0,1,1)/\sqrt{2}\).

Notice, that at zero magnetic field \( (h \to 0) \) \( E_2 = E_3 \) as in (21c). On the other hand, from (C9) it is clear that two components correspond to \( M_z = \pm 6 \) and therefore in this doublet \( L_z = 6 \). In other two non-degenerate states \( L_z = M_z = 0 \) (i.e. they are \( \Sigma \)-states).
[1] L. Gagliardi, B.O. Roos, Nature 433, 848 (2005).
[2] R.C. Albers, Nature 410, 759 (2001).
[3] P. Fulde, *Electron Correlations in Molecules and Solids* (Springer Heidelberg, 1995).
[4] X. Cao and M. Dolg, Mol. Phys. 101, 1967 (2003).
[5] B.O. Roos, R. Lindh, P.-A. Malmqvist, V. Veryazov, and P.-O. Widmark, J. Phys. Chem. A 112, 11431 (2008).
[6] X. Shen, L. Fang, X. Chen, and J.R. Lombardi, J. Chem. Phys. 113, 2233 (2000).
[7] J.R. Lombardi and B. Davis, Chem. Rev. 102, 2431 (2002).
[8] J.D. Morgan III and W. Kutzelnigg, J. Phys. Chem. 97, 2425 (1993).
[9] E. F. Worden, R. W. Solarz, J. A. Paisner, and J. G. Conway, J. Opt. Soc. Am. B 68, 52 (1978); W. C. Martin, R. Zalubas, and L. Hagan, *Atomic Energy Levels - The Rare Earth Elements*, Natl. Stand. Ref. Data Ser., Natl. Bur. Stand. (U.S.) 60 (1978).
[10] C.J. Bradley and A.P. Cracknell, *The Mathematical Theory of Symmetry in Solids*, (Clarendon, Oxford, 1972).
[11] W. Heitler and F. London, Z. Physik 44, 455 (1927).
[12] *Valence Bond Theory*, D.L. Cooper (editor), Theoretical and Computational Chemistry, Vol. 10 (Elsevier, Amsterdam, 2002).
[13] A.V. Nikolaev and K.H. Michel, J. Chem. Phys. 122, 064310 (2005).
[14] R.G. Parr, W. Yang, *Density-Functional Theory of Atoms and Molecules*, (New York, Oxford University Press, 1989).
[15] D.C. Koskenmaki and K.A. Gschneidner, Jr., *Handbook on the Physics and Chemistry of Rare Earths*, ed. K.A. Gschneidner, Jr., and L. Eyring (Amsterdam: North-Holland, 1978), p. 337.
[16] A. Szabo and N.S. Ostlund, *Modern Quantum Chemistry*, (McGraw-Hill, Dover, 1989).
[17] L.D. Landau and E.M. Lifshitz, *Quantum Mechanics - Non-relativistic theory*, v. 3, (Pergamon Press, Oxford, 1965), Chap. 83.
[18] A.V. Tsyashchenko, A.V. Nikolaev, A.I. Velichkov, A.V. Salamatin, L.N. Fomicheva, G.K. Ryasny, A.A. Sorokin, O.I. Kochetov, M. Budzynski, and K.H. Michel, Phys. Rev. B 82, 092102 (2010).
[19] A.V. Nikolaev and K.H. Michel, Phys. Rev. B 66, 054103 (2002).