Trigonally Distorted Hexacoordinate Co(II) Single-Ion Magnets

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Abstract: By simple reactions involving various cobalt(II) carboxylates (acetate and in situ prepared pivalate and 4-hydroxybenzoate salts) and neocuproine (neo), we were able to prepare three different carboxylate complexes with the general formula [Co(neo)(RCOO)] (R = –CH₂ for 1, (CH₃)₂C– for 2, and 4OH-C₄H₆– for 3). The [Co(neo)(RCOO)] molecules in the crystal structures of 1–3 adopt a rather distorted coordination environment, with the largest trigonal distortion observed for 1, whereas 2 and 3 are similarly distorted from ideal octahedral geometry. The combined theoretical and experimental investigations of magnetic properties revealed that the spin Hamiltonian formalism was not a valid approach and the L-S Hamiltonian had to be used to reveal very large magnetic anisotropies for 1–3. The measurements of AC susceptibility showed that all three compounds exhibited slow-relaxation of magnetization in a weak external static magnetic field, and thus can be classified as field-induced single-ion magnets. It is noteworthy that 1 also exhibits a weak AC signal in a zero-external magnetic field.

Keywords: single-ion magnets; magnetic anisotropy; cobalt(II)

1. Introduction

Hexacoordinate Co(II) complexes often tend to possess very large magnetic anisotropies arising from the direct contribution of spin orbit coupling to the ground state. If the coordination geometry is regular octahedron, the ground state is ⁴T₁₈g. Then, angular momentum is the main contributor to the zero-field splitting (ZFS), because the spin-orbit coupling operator transforms under ⁴T₁₈g, thus it directly mixes with the ground state. Magnetic anisotropy is then very large and of the easy-plane type [1], the classic spin Hamiltonian description loses validity, and low-lying excited states appear [2]. The design of highly anisotropic magnetic molecules exploits changing of the regular octahedral geometry, which can be achieved either by elongation/compression of the metal–ligand bonds or by trigonal distortion. Both types of distortion can lead to double orbitally degenerate ground states (⁴E₉ for compressed octahedron, D₄ᵥ, ⁴E’’’ for trigonal prism, D₃h), manifesting themselves again by a large contribution of spin-orbit coupling to the ground state. However, contrary to the ⁴T₁₈g state [3,4], the anisotropy of E-states is of the axial character [5]. Thus, Co(II) complexes with doubly degenerate ground states are highly interesting for synthesis of single-ion magnets (SIMs), which are a class of singe molecule magnets (SMMs) [6] containing only one paramagnetic center [7]. Among all SIMs based on 3d transition metal ions [8–15], those with trigonal prismatic environment around the Co(II) center have a special position, because they often exhibit slow relaxation of magnetization in the absence of an external magnetic field [16–22]. It must be noted that so-called zero-field SIMs (ZF-SIMs) are still very rare for complexes of 3d transition metals, which is because of the rather specific requirements needed for occurrence of ZF-SIMs; i.e., large and axial magnetic anisotropy with negligible rhombicity [23]. Thus, in line with the
matter discussed above, the trigonal Co(II) complexes with doubly degenerate ground state are ideal candidates for ZF-SIMs. Besides this class of Co(II) compounds, there are only a few examples of 3d metal based ZF-SIMs: linear two-coordinate Fe(I) [24,25], Co(II) [10] complexes, pentacoordinate Fe(III) [26], or tetracoordinate Co(II) complexes [27–32].

In this paper, we focused our attention on trigonal distortion and, as the object of our research, we used very common carboxylate complexes [33] with one bidentate N-donor chelating ligand. Recently, we reported on two [Co(neo)(PhCOO)$_2$] polymorphs (neo stands for neocuproine), which differed in their trigonal distortion and magnetic properties. Both compounds behaved as SIMs in a weak external magnetic field ($B = 0.1$ T), so-called field-induced SIMs [34]. In the present paper, we show that, for [Co(neo)(RCOO)$_2$] complexes, we can achieve a significant change in the trigonality of the coordination polyhedron by variation of carboxylate ligands (RCOO–).

2. Materials and Methods

2.1. Materials

Co(NO$_3$)$_2$·6H$_2$O, Co(ac)$_2$·4H$_2$O, neocuproine, sodium pivalate hydrate, sodium 4-hydroxybenzoate, and solvents (MeOH, diethyl ether (Et$_2$O)) were supplied by VWR International (Stˇ rˇ brná Skalice, Czech Republic), Sigma-Aldrich (Prague, Czech Republic), Lach-Ner (Neratovice, Czech Republic), and Litolab (Chudobín, Czech Republic).

2.2. Synthesis

2.2.1. Complex [Co(neo)(ac)$_2$] (1)

To the solution of Co(ac)$_2$·4H$_2$O (0.48 mmol, 120 mg) in 5 mL of methanol, 100 mg of neocuproine (0.48 mmol) was added. The solution was ultrasonicated for 15 min. The violet solution was filtered through a paper filter and crystallized by slow diffusion of Et$_2$O in a closed flask. Then, 130 mg of 1 was isolated by filtration (yield = 70%) as violet crystals, which were dried in a desiccator under reduced pressure (overnight). IR (ATR, $v$, cm$^{-1}$): 408 w, 437 w, 550 w, 618 w, 675 m, 733 w, 778 w, 813 w, 846 w, 865 m, 937 w, 1007 w, 1037 w, 1162 w, 1226 w, 1299 w, 1383 w, 1421 s, 1440 s, 1501 m, 1556 s, 1593 m, 1617 w, 3001 w, 3055 w.

2.2.2. Complexes [Co(neo)(piv)$_2$] (2) and [Co(neo)(4OH-benz)$_2$]·2CH$_3$OH (3)

Compounds 2 and 3 were both prepared using the following method. To the solution of Co(NO$_3$)$_2$·6H$_2$O (70 mg, 0.24 mmol) in 15 mL of methanol, 50 mg of neocuproine (0.48 mmol) was added. The solution was heated up to 50°C, then 2 molar equiv. of the corresponding sodium salt was added (60 mg of sodium pivalate hydrate, 2; 77 mg of sodium 4-hydroxybenzoate, 3). The violet solution was filtered through a paper filter and crystallized isothermally. Then, 49 mg (yield = 43%) of 2 and 52 mg of 3 (yield = 36%) were obtained as violet crystals a couple of days later. The crystals were dried in a desiccator under reduced pressure (overnight).

2: IR (ATR, $v$, cm$^{-1}$): 355 m, 407 w, 551 w, 609 w, 656 w, 681 w, 733 w, 776 w, 791 w, 810 w, 864 w, 900 w, 940 w, 1002 w, 1032 w, 1157 w, 1225 m, 1299 s, 1351 w, 1377 w, 1421 s, 1447 m, 1486 s, 1593 m, 1593 m, 1593 m, 1593 m, 2865 w, 2926 w, 2967 w.

3: IR (ATR, $v$, cm$^{-1}$): 406 w, 504 w, 549 w, 630 m, 658 w, 701 w, 728 w, 783 m, 855 m, 1030 m, 1099 w, 1142 w, 1166 m, 1227 m, 1284 m, 1373 s, 1396 s, 1503 m, 1534 w, 1568 w, 1593 s, 2818 w, 2907 w, 2937 w, 3059 w, 3352 w, 3462 w.

Elemental analysis: 1, Mr = 550.4, C$_{18}$H$_{18}$CoN$_2$O$_4$: found: C, 56.28 H, 4.74; N, 7.25, requires C, 56.11; H, 4.71; N, 7.27%; 2, Mr = 567.9, C$_{24}$H$_{30}$CoN$_2$O$_4$: found: C, 61.38; H, 6.52; N, 5.90, requires C, 61.40; H, 6.44; N, 5.97%. 3 (composition calculated for [Co(neo)(4OH-
benz)\(\cdot 1.8\text{CH}_3\text{OH}\), \(M_r = 661.9\), \(C_{30}H_{30}CoN_2O_7\), found: C, 59.46; H, 4.56; N, 4.97, requires C, 59.74; H, 4.91; N, 4.68.

2.3. General Methods

Elemental analysis was performed by a Flash 2000 CHNS Elemental Analyzer (Thermo Scientific, Waltham, MA, USA). A Jasco FT/IR-4700 spectrometer (Jasco, Easton, MD, USA) was used for the collection of the infrared (IR) spectra in the range of 400–4000 cm\(^{-1}\) using the attenuated total reflection (ATR) technique on a diamond plate. The static magnetic data were measured on powdered samples pressed into pellets using a PPMS Dynacool (Quantum Design Inc., San Diego, CA, USA). The dynamic magnetic data were measured on powdered samples pressed into pellets stabilized by eicosane using a MPMS XL-7 Quantum Design SQUID magnetometer (Quantum Design Inc., San Diego, CA, USA).

2.4. X-ray Crystallography

Data collection for 1–3 was done using the standard rotational method on a D8 Quest diffractometer equipped with a Photon 100 CMOS detector (Bruker, Billerica, MA, USA) using the Mo-K\(\alpha\) radiation (\(\lambda = 0.71073\) Å). Data collection, data reduction, and cell parameters refinements were performed using the Bruker Apex III software package [35]. The structures were solved by SHELXT [36] and all non-hydrogen atoms were refined anisotropically on \(F^2\) using the full matrix least-squares procedure with Olex2.refine [37] in OLEX2 (version 1.5) [38]. All hydrogen atoms were found in differential Fourier maps and their parameters were refined using a riding model with \(U_{\text{iso}}(H) = 1.2(CH) \) or 1.5(–CH\(_3\), –OH) \(U_{\text{eq}}\). The molecular structures and packing diagram were drawn with MERCURY [39].

Powder diffraction data (Supplementary information, Figures S1–S3) were collected using a MiniFlex600 (Rigaku) equipped with the Bragg–Brentano geometry, and with iron-filtered Cu K\(\alpha_1,2\) radiation.

3. Results and Discussion

3.1. Synthesis and Crystal Structure

Compound 1 was prepared by a direct reaction between Co(ac)\(_2\)·4H\(_2\)O and neo (molar ratio 1:1) in methanol. Compounds 2 and 3 were prepared in a very similar way, but Co(NO\(_3\))\(_2\)·6H\(_2\)O was used as the starting Co(II) compound and the corresponding sodium salts (NaRCOO) were used as a source of the carboxylate ligands. The reaction mixtures were prepared by dissolving Co(NO\(_3\))\(_2\)·6H\(_2\)O, neo, and NaRCOO in methanol (molar ratio of 1:1:2). These procedures proved to be good for the preparation of the crystalline products including single crystals useful for X-ray diffraction; however, the reaction yields were relatively low (30–50%).

Compounds 1–3 were isolated as violet crystals, which diffracted rather well, and we were able to determine crystal structures by single-crystal X-ray diffraction. The basic crystallographic data are summarized in Table 1. All compounds consist of the [Co(neo)(RCOO)\(_2\)] complex molecules, only in 3 are two additional co-crystallized methanol molecules present in the asymmetric unit. The complex molecules are hexacoordinate, and all the ligands coordinate to the Co atom in a bidentate manner. The Co–N bonds adopt bond lengths between 2.08 and 2.13 Å, whereas the lengths of the Co–O bonds are more variable: 2.04–2.20 Å (Figure 1). The shapes of the coordination polyhedrons were evaluated using SHAPE algorithm and continuous shape measures (CSMs) [40]. We revealed that all the complex molecules possess very large distortions from the ideal geometries. The coordination polyhedrons of 2 and 3 are closer to regular octahedral (O\(_h\)) than trigonal prismatic geometry (D\(_{3h}\)) with the following CSMs (O\(_h\), D\(_{3h}\)): 7.650, 9.801, 2, 6.352, 10.110, 3. Complex 1 possesses the largest trigonal distortion and coordination geometry close to trigonal prism (11.893, 3.761, ESI Table S1).
Table 1. Crystallographic data for 1–3.

|          | 1                          | 2                          | 3                          |
|----------|----------------------------|----------------------------|----------------------------|
| Formula  | C_{18}H_{16}CoN_{2}O_{4}  | C_{24}H_{30}CoN_{2}O_{4}  | C_{30}H_{30}CoN_{2}O_{8}  |
| Formula weight | 385.27                     | 469.43                     | 605.49                     |
| Crystal system | monoclinic                 | triclinic                  | monoclinic                 |
| Space group   | \(C2/c\)                   | \(P\bar{T}\)               | \(P2_1/c\)                 |
| Cell parameters |
| \(a/\text{Å}\) | 14.0976(17)               | 9.5282(13)                 | 9.922(4)                   |
| \(b/\text{Å}\) | 9.4555(12)                 | 9.8422(13)                 | 18.083(7)                  |
| \(c/\text{Å}\) | 25.259(4)                  | 14.4559(16)                | 15.191(6)                  |
| \(a/\text{deg}\) | 90                         | 87.490(4)                  | 90                         |
| \(\beta/\text{deg}\) | 95.643(12)                | 80.364(4)                  | 91.472(14)                 |
| \(\gamma/\text{deg}\) | 90                         | 62.028(4)                  | 90                         |
| \(V/\text{Å}^3\) | 3350.1(8)                  | 1179.5(3)                  | 2675(2)                    |
| Density, \(\text{g cm}^{-3}\) | 1.52                        | 1.32                       | 1.399                      |
| Abs. coefficient/\(\text{mm}^{-1}\) | 1.050                      | 0.759                      | 0.650                      |
| Data/restraints/param |
| \(R1 \ a, wR2 \ b (all data)\) | 0.0412, 0.0730             | 0.0594/0.1068              | 0.1200/0.1847              |
| \(R1 \ a, wR2 \ b [I > 2 s(I)]\) | 0.0296, 0.0700             | 0.0411/0.0999              | 0.0611/0.1614              |
| Goodness of fit | 1.073                     | 1.034                      | 1.064                      |
| CSD number | 2,126,276                  | 2,126,278                  | 2,126,275                  |

\(R1 = \sum (|F_o| - |F_c|)/\sum |F_o|, \quad wR2 = \sqrt{\sum w(|F_o|^2 - |F_c|^2)^2)/\sum w(|F_c|^2)} \).  

Figure 1. Molecular structures of complex molecules in the crystal structure of 1 (a), 2 (b), and 3 (c). Selected bond lengths (in Å): Co1–N1 = 2.105(2) in 1, 2.133(2) in 2, 2.130(4) in 3; Co1–N2 = 2.125(2) in 1, 2.118(2) in 2, 2.078(3) in 3; Co–O1 = 2.175(2) in 1, 2.198(9) in 2, 2.102(3) in 3; Co–O2 = 2.163(2) in 1, 2.150(9) in 2, 2.160(4) in 3; Co–O3 = 2.160(2) in 1, 2.149(3) in 2, 2.196(4) in 3; Co–O4 = 2.104(2) in 1, 2.119(6) in 2, 2.131(3) in 3.

The non-covalent interactions in 1–3 are mainly of weak nature, including C–H···O, C–H···π, and π···π interactions of the neo aromatic rings. Of note here is the crystal structure of 3, involving OH groups of the 4OH-benz ligands and co-crystallized molecules of methanol. These formed 2D networks of the molecules are interconnected by rather strong O–H···O hydrogen bonds with the donor···acceptor distances ranging between 2.65 and 2.85 Å (Figure S4). Even such relatively strong non-covalent contacts did not sufficiently stabilize the crystal structure of 3. When the crystals were transferred outside of the mother-liquor, the solvent loss occurred upon drying and was accompanied by a loss of crystallinity and/or change in the unit cell parameters. Thus, at ambient conditions (\(T = 298\text{ K}, p = 1\text{ atm}\)), we were not able to unambiguously confirm the phase uniformity of 3, because the X-ray diffraction pattern of the dried 3 differed slightly from the pattern calculated from the single-crystal structure. Nevertheless, the experiments performed using grinded crystals of 3 immersed in a highly viscose crystallographic oil showed that, after
1 h, the diffraction pattern underwent significant changes (Figure S3), which indicated that the batches of 3 we prepared were phase pure.

3.2. DC Magnetic Properties

The magnetic properties for 1–3 measured in a static magnetic field are depicted in Figure 2 as the temperature dependence of the effective magnetic moment and isothermal magnetization. Evidently, the profile of $\mu_{\text{eff}}$ versus $T$ is varied within the series owing to the variation of the geometry of the coordination polyhedra, and hence the ligand field. As there are negligible intermolecular interactions, the decrease in $\mu_{\text{eff}}$ is attributed to the zero-field splitting/large magnetic anisotropy of these compounds. Indeed, this is also confirmed by the saturation values of $M_{\text{mol}}$ of the isothermal magnetization curves well below the theoretical limit $g \cdot S$.
with the Hamiltonian parameters in Table 2, and the red and blue lines correspond to the negative and positive value of $\Delta_{ax}$, respectively.

Table 2. The parameters of the Hamiltonian in Equation (1) derived from the experimental and calculated data.

| Parameters                  | 1   | 2 \(^{a}\) | 3   |
|-----------------------------|-----|-----------|-----|
| $\Delta_{ax}$ (cm\(^{-1}\)) | -3317 | -523/810 | -1051 |
| $\Delta_{rh}$ (cm\(^{-1}\))  | -133  | -23.6/32.5 | -39.6 |
| $\alpha$                    | 1.66  | 1.04/1.25 | 1.21 |
| $\lambda$ (cm\(^{-1}\))    | -75.8 | -167/ -180 | -151 |

\(^{a}\) The parameters corresponds to the best-fit for the negative and positive value of $\Delta_{ax}$.

Usually, the magnetic anisotropy is treated with the spin Hamiltonian comprising the zero-field splitting and Zeeman terms; however, hexacoordinate Co\(^{III}\) complexes with the shape of the chromophore close to octahedron or trigonal prism possess orbital angular momentum, hence the spin Hamiltonian is inappropriate. This was confirmed by ab initio calculations that predicted magnetic behavior based on E ground state for 1 and low lying (below 1000 cm\(^{-1}\)) excited states for 2 and 3, both contradicting the use of spin Hamiltonian formalism [41] \((vide infra)\).

Therefore, the DC magnetic data were analysed with the L-S Hamiltonian based on Griffith and Figgis [42-44], which describes the splitting of the \(^4\)T\(_{1g}\) term originating from the \(^4\)F atomic term in lower symmetries than O\(_h\), as follows:

$$\hat{H} = -\alpha \cdot \lambda \left( \vec{S} \cdot \vec{L} \right) + \Delta_{ax} \left( \hat{L}_{x}^{2} - \hat{L}_{y}^{2} / 3 \right) + \Delta_{rh} \left( \hat{L}_{x}^{2} - \hat{L}_{y}^{2} \right) + \mu_{B} B \left( g_{e} \vec{S} - \alpha \vec{L} \right) \quad (1)$$

The splitting of the \(^4\)T\(_{1g}\) term is described by $\Delta_{ax}$ and $\Delta_{rh}$ parameters; $\alpha$ is an orbital reduction factor, $\lambda$ is a spin-orbit coupling parameter, and $g_{e} = 2.0023$. Owing to the utilization of T\(_{1g}\)-P isomorphism, the angular orbital momentum $L$ adopts the value of 1 with the effective Lande $g$-factor, $g_{L} = -1$. The Hamiltonian acts on $1\vec{S}, 1\vec{L}, M_{S}, M_{L} > \text{functions}$
with $M_L = 0, \pm 1$ and $M_S = \pm 1/2, \pm 3/2$ [45]. Next, the orbital reduction factor embodies two parameters, $\alpha = A\kappa$, where $A$ is the Figgis coefficient of the configuration interaction resulting from the admixture of the excited terms reflecting the ligand field strength, and $\kappa$ describes the lowering orbital contribution due to covalency of the metal–ligand bond. Moreover, the spin-orbit coupling parameter $\lambda$ can be reduced in comparison with its free-ion value $\lambda_0 = -180 \text{ cm}^{-1}$, which is attributable to the covalent character of the donor–acceptor bond.

The analysis encompasses both temperature- and field-dependent magnetic data and was done both for positive and negative values of $\Delta_{\text{ax}}$ with the help of a program POLYMAGNET [46]. However, only in the case of compound 2 were reasonably good fits achieved for both signs of $\Delta_{\text{ax}}$, whereas the negative sign of $\Delta_{\text{ax}}$ was found for 1 and 3—Figure 2. The values of the fitted parameters are listed in Table 2. The negative values of the fitted $\Delta_{\text{ax}}$ parameters resulted in the easy axis type of magnetic anisotropy, as visualized in the three-dimensional plots of molar magnetization—Figure S5. Such a type of magnetic anisotropy is essential for the formation of the spin reversal barrier needed for the observation of the Orbach type mechanism of the slow relaxation of magnetization. Moreover, the respective energy levels in the zero magnetic field are also plotted for 1–3 in Figure S5.

### 3.3. AC Magnetic Properties

The SMMs are generally characterized by the AC susceptibility measurements, evidencing the slow relaxation of magnetization. Therefore, first, the AC data were measured in the zero static magnetic field for 1–3, but only in the case of 1 did we observed a very weak signal of the imaginary susceptibility ($\chi''$). Thus, the data were also measured for a varying static magnetic field, which resulted in a clear observation of non-zero $\chi''$ susceptibility—Figures S6–S8. Thus, the temperature and frequency AC susceptibility was measured at small $B_{\text{DC}} = 0.1 \text{ T}$ to suppress the quantum tunneling of magnetization for 1–3, which revealed frequency-dependent maxima of the imaginary susceptibility, thus confirming the slow relaxation of magnetization—Figures 3–5. The experimental data were analyzed with a program MIF&FIT [47] to the one component Debye’s model based on Equation (2):

$$\chi(\omega) = \frac{\chi_T - \chi_S}{1 + (i\omega\tau)^{1-\alpha}} + \chi_S$$

Such an analysis resulted in the values of isothermal ($\chi_T$) and adiabatic ($\chi_S$) susceptibilities, relaxation times ($\tau$), and distribution parameters ($\alpha$) for 1–3 (Tables S2–S4). For further analysis, only the data for which the fitted parameters were calculated with the standard deviation two times smaller than the value of the fitted parameter were considered. Afterwards, the temperature dependences of the relaxation times were analyzed with a model comprising the direct and Orbach mechanism:

$$\frac{1}{\tau} = AT + \frac{1}{\tau_0} \exp(U_{\text{eff}}/kT)$$

The fitted data are displayed in Figures 3–5 and the best-fit parameters are as follows: $A = (6.52 \pm 0.62) \text{ K}^{-1}\text{s}^{-1}$, $\tau_0 = (1.361 \pm 0.074) \times 10^{-7} \text{ s}$, and $U_{\text{eff}} = (37.7 \pm 0.25) \text{ K}$ for 1; $A = (471 \pm 29) \text{ K}^{-1}\text{s}^{-1}$, $\tau_0 = (6.2 \pm 2.0) \times 10^{-6} \text{ s}$, and $U_{\text{eff}} = (19.0 \pm 0.74) \text{ K}$ for 2; $A = (2053 \pm 49) \text{ K}^{-1}\text{s}^{-1}$, $\tau_0 = (1.04 \pm 0.31) \times 10^{-6} \text{ s}$, and $U_{\text{eff}} = (17.5 \pm 1.1) \text{ K}$ for 3. The fitted values of $U_{\text{eff}} = 26.2 \text{ cm}^{-1}$ for 1, $13.2 \text{ cm}^{-1}$ for 2, and $12.2 \text{ cm}^{-1}$ for 3 are smaller than the energy gaps between the first and the second Kramers doublets of $\Delta = 82 \text{ cm}^{-1}$ for 1, $\Delta = 187/145 \text{ cm}^{-1}$ for 2, and $\Delta = 181 \text{ cm}^{-1}$ for 3 (Figure S5), but such a feature is typical for Co$^{III}$ SMMs.
Figure 3. The AC magnetic data for 1. Temperature dependence of the real (\(\chi'\)) and imaginary (\(\chi''\)) components of the AC susceptibility at the applied external magnetic field \(B_{DC} = 0.1\) T for frequencies from 1 to 1500 Hz (full lines are only guides for eyes) (top). Frequency dependence of \(\chi'\) and \(\chi''\) molar susceptibilities fitted with one-component Debye’s model using Equation (2) (full lines) (middle). The Argand (Cole-Cole) plot with full lines fitted with Equation (2) and, on the right, the fit of resulting relaxation times \(\tau\) with the direct + Orbach relaxation processes (red line) using Equation (3) (bottom).
Figure 4. The AC magnetic data for 2. Temperature dependence of the real ($\chi'$) and imaginary ($\chi''$) components of the AC susceptibility at the applied external magnetic field $B_{DC} = 0.1$ T for frequencies from 1 to 1500 Hz (full lines are only guides for eyes) (top). Frequency dependence of $\chi'$ and $\chi''$ molar susceptibilities fitted with one-component Debye’s model using Equation (2) (full lines) (middle). The Argand (Cole-Cole) plot with full lines fitted with Equation (2) and, on the right, the fit of resulting relaxation times $\tau$ with the direct + Orbach relaxation processes (red line) using Equation (3) (bottom).
3.4. Theoretical Calculations

The electronic structure and magnetic properties of 1–3 were also studied by theoretical methods suitable for complexes with a multireference character. Therefore, the multireference calculations based on the state average complete active space self-consistent field (SA-CASSCF) [48] wave function method complemented by N-electron valence second-order perturbation theory (NEVPT2) [49,50] were conducted with an ORCA 5.0 computational package [51,52]. The experimental molecular structures were used, and just the positions of hydrogen atoms were normalized with Mercury software. The triple-ζ basis set def2-TZVP was used for all atoms except for carbon and hydrogen atoms, for which def2-SVP was applied [53]. The speed of the calculations was increased by using def2/J and def2-TZVP/C auxiliary basis sets [54,55], together with the chain-of-spheres (RIJCOSX) approximation to exact exchange [56,57] as implemented in ORCA. The active space was defined by seven
electrons in five d-orbitals of Co\textsuperscript{II} (CAS(7e,5o)), and all possible multiplets, 10 quartets and 40 doublets, were involved in the calculations. Subsequently, the ab initio ligand field theory (AILFT) \cite{58,59} was applied to calculate the splitting of d-orbitals, as shown in Figure 6. It is evident that splitting of d-orbitals for 1 is close to the pattern typical for a trigonal prism ligand field, whereas the splitting for 3 resembles a more typical octahedral ligand field. This nicely demonstrates the gradual change in the ligand field symmetry within the series of 1–3. Next, the 4T\textsubscript{1g} term is split within 0–2000 cm\textsuperscript{-1} for 2 and 3 owing to their deviations from ideal O\textsubscript{h} symmetry, but, in the case of 1, there is evidently a split 4E ground term (0–930 cm\textsuperscript{-1}) belonging to D\textsubscript{3} pseudosymmetry (Figure 6, middle). It can be also interpreted as a large distortion of O\textsubscript{h} symmetry, evidenced by the large splitting of the 4T\textsubscript{1g} term spanning energy interval of 0–5000 cm\textsuperscript{-1}. This is also mirrored in the splitting of the six lowest Kramers doublets (Figure 6, right), which is largest for 1.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure6.png}
\caption{The outcome of the CASSCF/NEVPT2 calculations for complexes of 1–3. Plot of the d-orbitals splitting calculated by ab initio ligand field theory (AILFT) (left), low-lying ligand-field terms (LFT) (middle), and ligand-field multiplets (LFM) (right). Note: different multiplicities of LFT are shown in a different color.}
\end{figure}

The energies of the six lowest Kramers doublets were used for the analysis of the parameters of the Hamiltonian in Equation (1). Such a procedure we applied for the first time in the investigation of the above mentioned [Co(neo)(PhCOO)\textsubscript{2}] polymorphs \cite{34} and then also for other Co\textsuperscript{II} complexes \cite{60,62}. This procedure resulted in the values of \(\alpha \cdot \lambda, \Delta_{ax}, \Delta_{rh}\), which are listed in Table 2 and graphically presented in Figure S9. However, this procedure has one drawback, as it is not possible to determine the values of \(\alpha\) and \(\lambda\) separately. To overcome this problem, we calculated temperature- and field-dependent magnetization data directly in an ORCA package resulting from CASSCF/NEVPT2 calculations. Subsequently, these magnetic data were fitted to Equation (1), but with fixed values of \(\alpha \cdot \lambda, \Delta_{ax}, \Delta_{rh}\), which enabled the determination of \(\alpha\) and \(\lambda\), as listed in Table 2. The respective fits are depicted in Figure S10. There are clear trends visible from the calculated values: \(\alpha\) and \(\Delta_{ax}\) are much larger for 1 than for 2–3, evidencing the impact of the geometry change from the trigonal prism to the octahedral shape. We can also comment on the possible source of discrepancies between the values of the Hamiltonian parameters derived from fitting of the experimental data and from CASSCF/NEVPT2 calculations. Firstly, the fitting of magnetic data is limited only to the temperature interval of 1.9–300 K, whereas the energy levels from Equation (1) span an interval of up to several thousand cm\textsuperscript{-1}/K, which means that the Boltzmann population of energetically higher Kramers doublets is negligible, and hence does not affect magnetic data. Secondly, the active space of theoretical calculations was limited to only five d-orbitals, thus the ligand-based orbitals are missing;
has the largest occupied and the ab initio magnetization blocking barriers were computed for magnets, where (temperature-assisted) quantum tunneling (blue), Orbach/Raman mechanisms (red), and direct/Raman mechanisms (green) [64].

The outcome of SINGLE_ANISO CASSCF/NEVPT2 calculations for 1–3 are close to trigonal prismatic geometry, whereas 2 and 3 adopt very trigonally distorted octahedral coordination environments. The magnetic properties were studied by DC and AC magnetometry, and it was revealed that very large magnetic anisotropy dominates the magnetic behavior of 1–3. CASSCF/NEVPT2 calculations revealed that the origin of the large anisotropy in 1 is in the $^4E$ ground state, whereas in 2 and 3, the large magnetic anisotropy arises from low-lying excited ligand field terms owing to the distorted hexacoordinate coordination geometry. The largest intra-Kramers doublet splitting was calculated for 3, whereas the smallest was calculated for 1. This agrees rather well with the distortion from the regular octahedral geometry of the coordination polyhedron in these complexes (CSMs): 6.352 (in 1) < 7.650 (in 2) < 11.893 (in 1). AC susceptibility measurements revealed that 1 exhibits the slow relaxation of magnetization even in the zero static magnetic field. However, the observed signal was rather weak, and this prevented us from performing further analysis. Thus, for all three compounds, AC susceptibility was measured in a static magnetic field ($B_{dc} = 0.1$ T) and the slow relaxation of magnetization was confirmed for all of them. Thus, 1–3 behave as field-induced single-ion magnets, where 1 has the largest $U_{dd} = 38$ K. SINGLE_ANISO calculations revealed that the probability of
13/2,+3/2> ↔ 13/2,−3/2> quantum tunneling is the lowest for 1, whereas in 2 and 3, it should be the dominant relaxation process in the absence of the external magnetic field. This agrees rather well with the experimental observations presented in this report and, furthermore, it underlines the importance of trigonal coordination geometry (as in 1) for the preparation of Co(II) zero-field SIMs.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/ma15031064/s1, Figure S1: XRPD data for (1); Figure S2: XRPD data for (2); Figure S3: XRPD data for (3); Figure S4: A perspective view on a 2D network; Figure S5: The three-dimensional plot of molar magnetization; Figure S6: In-phase \( \chi_{\text{in}} \) and out-of-phase \( \chi_{\text{out}} \) molar susceptibilities for 1; Figure S7: In-phase \( \chi_{\text{in}} \) and out-of-phase \( \chi_{\text{out}} \) molar susceptibilities for 2; Figure S8: In-phase \( \chi_{\text{in}} \) and out-of-phase \( \chi_{\text{out}} \) molar susceptibilities for 3; Figure S9: The low-lying ligand-field multiplets originating from the \( ^4T_{1g} \) state of 1–3; Figure S10: The analysis of the magnetic data calculated from CASSCF/NEVPT2 calculations; Table S1: The results of SHAPE calculation for 1–3; Table S2: Parameters of one-component Debye’s model for 1; Table S3: Parameters of one-component Debye’s model for 2; Table S4: Parameters of one-component Debye’s model for 3.

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