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Spin-Triplet Cobalt Complexes of Biuret and Related Ligands

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A number of planar Co$_{111}$N$_4$ complexes have been prepared with biuret and N-substituted biurets. They are shown to have triplet electronic ground states in the temperature range 6.4–293°K. The zero-field splitting is estimated to be about 40 cm$^{-1}$. The triplet ground state is consistent with molecular orbital energies obtained from extended Hückel calculations. The $d_{x^2}$ and $d_{y^2}$ orbitals both have the same energy and are both occupied by one electron. Polarographic measurements show that these complexes can be reversibly reduced in a one-electron step. With amines, diamagnetic six-coordinated mixed-ligand species are obtained, whose uv–vis spectra can be interpreted as for Co(III) in an octahedral environment.

Introduction

In previous communications from our laboratory the rather unique properties of biuret as a ligand were discussed. It was shown that the well-known Cu(II) and Ni(II) complexes could be oxidized with various oxidizing agents yielding the corresponding Cu(III) and Ni(III) compounds.$^{1,2}$ With cobalt the very remarkable planar bis(biuretato)cobalt-

ate(III) ion could be synthesized having a spin-triplet ground state. The crystal structure of the 3-propyl-substituted biuret complex of Co(III) was recently published. We now report more details about the properties of a series of cobalt complexes with biuret and related ligands.

**Experimental Part**

**Abbreviations.** Abbreviations used in this paper are as follows: bi = [HINCONHCNH], 3-Ph(bi) = [HNCON(C6H5)CONH]2, 3-Pr(bi) = [HNCON(C3H7)CONH]2, 1-Pr(bi)H2 = H2NCONHCONHC3H7.

**DMSO** = dimethyl sulfoxide and Bu4NOH = tetra-n-butylammonium hydroxide.

**Syntheses.** o-Phenylenebis(biuret) and Ethylenebis(biuret).

These ligands were obtained in a way analogous to the method used for the preparation of 1-alkyl-substituted biuret starting with a 2:1 mixture of nitrobiuret and o-phenylenediamine or ethylenediamine, respectively.

KCo(bi). Cobalt acetate (25 g) was dissolved in 25 ml of water. Biuret (30 g) and KOH (36 g, dissolved in 50 ml of water) were added to this solution and the resulting blue slurry was oxidized with air during 24 hr. The insoluble material was then removed by filtration and upon careful neutralization with 6 N HCl to pH 7 KCo(bi) precipitated as a yellow compound. It was filtered off, washed with water, and dried in a vacuum desiccator. The compound is insoluble in any solvent.

**Anal.** Calculated for KCo(bi)2: K, 13.03; Co, 19.65; C, 16.00; H, 2.00; N, 28.00. Found: K, 13.4; Co, 19.5; C, 15.7; H, 2.2; N, 27.2.

KCo(3-Ph(bi))2DMSO. 3-Phenylbiuret was prepared according to Weith. Cobalt chloride (4 g) was dissolved in 50 ml of water. KOH (3 g dissolved in 5 ml of water) was added to this solution. The resulting blue slurry was oxidized with air during 24 hr. The insoluble material was removed by filtration. To the red filtrate about the same volume of alcohol was added and then sufficient ether was added to form red and colorless liquid layers. The upper colorless layer was re-filtered and the yellow complex was precipitated by adding a little alcohol and much ether. The precipitate was filtered off. It was dissolved in chloroform, the solution was filtered, and the complex was again precipitated by adding ether.

**Anal.** Calculated for Bu4NCo(3-Ph(bi))2: C, 58.61; H, 7.69; N, 14.95. Found: C, 58.2; H, 7.7; N, 14.9.

Bu4NCo(bi)H2. CoCl2-6H2O (5 g) was dissolved in 200 ml of water. Biuret (6 g) and 75 ml of a 40% Bu4NOH solution in water were added. The resulting mixture was oxidized with air during 4 hr. The insoluble material was filtered off and the red filtrate was concentrated as described above with alcohol and ether until a red precipitate, that was dissolved in DMSO. Dropwise aqueous 1 N HCl was added to this red solution until the color had turned to orange-yellow. The complex was then precipitated by adding a little alcohol and much ether. The precipitate was filtered off. It was dissolved in chloroform, the solution was filtered, and the complex was again precipitated by adding ether.

**Anal.** Calculated for Bu4NCo(3-Ph(bi))2: C, 58.61; H, 7.69; N, 14.95. Found: C, 58.2; H, 7.7; N, 14.9.

KCo(3-Pr(bi))2(1-Pr(bi)H2)2. When the method of Weith for the preparation of 3-phenylbiuret is used for the preparation of alkybiurets, a mixture of 1-alkylbiuret and 3-alkylbiuret is always obtained. So when a mixture of 0.1 mol of dry propyleneurea and 0.1 mol of PCl3 is refluxed during 4 hr, a acid solution containing 1-propylbiuret and 3-propylbiuret is obtained after hydrolysis of the reaction mixture with ice. In such a solution excess CoCl2-6H2O was dissolved. KOH was added until a blue slurry was formed, which was oxidized with air during 24 hr. The precipitate was filtered off and upon careful neutralization of the red solution with 6 N HCl to pH 7 the yellow complex precipitated. It was filtered off and washed with water. The complex is soluble in DMSO, alcohol, and acetone.

**Anal.** Calculated for KCo(3-Pr(bi))2(1-Pr(bi)H2)2: Co, 8.73; C, 35.51; H, 5.98; N, 24.92. Found: Co, 8.5; C, 35.2; H, 6.1; N, 25.0.

In analogous way other compounds of composition KCo(3-alkyl(bi))2(1-alkyl(bi)H2)2 could be prepared. KCo(3-Pr(bi))2H2O. From a solution of KCo(3-Pr(bi))(1-Pr(bi)H2)2 in moist DMSO the compound KCo(3-Pr(bi))(1-Pr(bi)H2)2H2O precipitated upon addition of chloroform and ether. It was filtered off and washed with chloroform. The precipitate was filtered off and washed with chloroform in dry air as it is sensitive to moisture.

**Anal.** Calculated for KCo(3-Pr(bi))(1-Pr(bi)H2)2H2O: Co, 14.0; C, 28.6; H, 5.3; N, 19.9. Found: Co, 13.9; C, 28.6; H, 4.9; N, 18.8.

KCo(3-Pr(bi))2DMSO. In dry DMSO KCo(3-Pr(bi))(1-Pr(bi)H2)2H2O was dissolved. After a few hours KCo(3-Pr(bi))2DMSO could be precipitated by adding dry chloroform. The precipitate was filtered off and washed with chloroform in dry air as it is sensitive to moisture.

**Anal.** Calculated for KCo(3-Pr(bi))2DMSO: C, 31.1; H, 5.59. Found: C, 30.9; H, 5.9.

KCo(en(bi))2. Ethylenebis(biuret) (1.2 g) and cobalt acetate (1.2 g) were dissolved in 40 ml of DMSO. To this solution 1.2 g of KOH dissolved in 4 ml of water was added. This mixture was oxidized with air during 3 hr. The precipitate of the red complex was filtered off and washed with alcohol. It was dissolved in a lot of water, and after cooling, the compound could be precipitated by adding a large amount of chloroform. The precipitate was filtered off and washed with chloroform.

**Anal.** Calculated for KCo(en(bi))2: C, 22.01; H, 2.46; N, 25.7. Found: C, 22.0; H, 2.7; N, 25.0.

KCo(o-phen(bi))2. CoCl2-6H2O (0.75 g) and o-phenylenebis(biuret) (0.85 g) were dissolved in 60 ml of DMSO. KOH (1 g dissolved in 6 ml of water) was added. This mixture was oxidized with air during 2 hr. The solution was filtered and the purple complex was precipitated from the filtrate by adding alcohol. The compound was filtered off and dissolved in DMSO. The solution was filtered, and the complex was again precipitated with alcohol and ether, filtered off, and washed with alcohol.

**Anal.** Calculated for KCo(o-phen(bi))2: C, 32.09; H, 2.15; N, 22.46. Found: C, 31.6; H, 2.3; N, 22.1.

Bu4NCo(en(bi)). Cobalt acetate (1.2 g) was dissolved in a mixture of 20 ml of water and 10 ml of DMSO. Ethylenebis(biuret) (1.2 g) and 12.5 ml of a 40% Bu4NOH solution in water were added and the resulting blue slurry was oxidized with air during 12 hr.

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W. Weith, Ber., 10, 1744 (1877).
The precipitate of the crude red complex was then filtered off. It was dissolved in DMSO, the solution was filtered, and the complex was precipitated by adding chloroform and ether. The presence of chloroform was confirmed by the mass spectrometric analysis of the compound at 120°.

All these complexes do not have good melting points but decompose in general above 170°.

Physical Measurements. Magnetic susceptibilities were measured with Gouy type equipment with HgCo(NCS)₄ as a reference over a temperature range of −150 to +20°. A vibrating-sample magnetometer (Princeton Applied Research Type 155) equipped with a standard PAR temperature control unit was used in the temperature range of −150 to +20°. A dropping mercury electrode was used as solvent with 0.1 M Bu₄NOH solution in water as supporting electrolyte. A saturated calomel electrode. Ultraviolet and visible spectra were measured with a Unicam SP 700C; transmission spectra were measured in solid KBr with a Cary Model 14.

Results and Discussion

Paramagnetic compounds in which Co(III) is coordinated with four atoms in an essentially planar geometry are very rare. Except for some cobalt(III) dithiolate complexes no good examples are known. Apparently the electron-donating capacity of the biuretato and the dithiolato ligands is so strong that the relatively large electron density on the central cobalt atom makes coordination of strong electron-donating ligands (e.g., amines) unfavorable, while there is in the case of the biuretato complexes no indication of coordination with weak electron donators (e.g., H₂O, DMSO, alcohol). Sometimes, however, the solid compounds contain DMSO, H₂O, or 1-alkylbiuret. It has been shown by crystallographic structure determination⁴ that the 1-alkylbiuret molecules are not coordinated with the cobalt atom and that the cobalt atom is in planar coordination with four nitrogen atoms of deprotonated 3-alkylbiurets. Through formation of H bridges the 1-alkylbiuret molecules are incorporated in the crystal lattice around the potassium ions, yielding magnetic compounds which can at best be formulated as [K{1-alkyl(bi)H₂}]₂[Co{3-alkyl(bi)}₃]. Compounds of this composition are insoluble in water, so the coincidental presence of 1-alkylbiuret in the reaction mixture from which these compounds are prepared (see Experimental Part) enabled us to prepare these compounds from aqueous solutions. The CoN₄ complexes that are soluble in water (e.g., KCo(3-Pr(bi))₂-2DMSO) are decomposed in neutral or acid solution, due to protonation of the ligand, so they have to be prepared in nonaqueous solvents such as DMSO. We believe that the cobalt atom in the solid compounds containing DMSO or H₂O is also four-coordinated, as the spectral and magnetic properties of these compounds are similar to those of the compound containing 1-alkylbiuret (see Table I and Figure 1).

These complexes could be studied in solutions of rather weak electron-donating solvents such as alcohol, DMSO, and acetone, as they do not take up weak electron donors as further ligands. The magnetic moments of KCo(3-Pr(bi))₂-2H₂O and KCo(3-Pr(bi))₂(1-Pr(bi)H₂)₂ in DMSO solution indicate that the triplet ground state is not affected and the positions of the uv-vis absorption bands of KCo(3-Pr(bi))₂(1-Pr(bi)H₂)₂ measured in DMSO, alcohol, and the solid compound are quite the same, indicating that the solvent molecules are not interacting with the planar CoN₄ species.

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![Figure 1. Electronic spectrum of KCo(3-Pr(bi))₂(1-Pr(bi)H₂)₂ in DMSO.](image-url)
Spin-Triplet Cobalt Complexes of Biuret

Table II. Uv Spectral Data of Some Mixed-Ligand Complexes

<table>
<thead>
<tr>
<th>Compd</th>
<th>(\nu_1), kK</th>
<th>(\nu_2), kK</th>
<th>10D(\epsilon), kK</th>
<th>B, kK</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co(3-Pr(bi))(_2)(NH(_3))(_2)(^{-})</td>
<td>20.4 (70)</td>
<td>27.4 sh</td>
<td>21.4</td>
<td>500</td>
</tr>
<tr>
<td>Co(3-Pr(bi))(_2)(C(_3)H(_7)NH(_2))(^{-})</td>
<td>18.5 (45)</td>
<td>27.8 sh</td>
<td>19.0</td>
<td>730</td>
</tr>
<tr>
<td>Co(3-Pr(bi))(_2)(pyrrolidine)(^{-})</td>
<td>17.4 (42)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co(3-Pr(bi))(_2)(py)(^{-})</td>
<td>18.2 (47)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Writing square root terms of the type
\[12\sqrt{-(D + E)^2 + 4g_x^2\beta H_x^2} + 4g_y^2\beta H_y^2} + 4g_z^2\beta H_z^2} = (D + E)^2\]
all symbols having their usual significance, the energies of these levels can be written as a constant plus a function of \(H^2\), so in the general form\(^9\)

\[U_n = U_n(0) + U_n(1)H + U_n(2)H^2\]

\(U_n(1)\) is always zero in these cases. Now inserting these values into van Vleck's formula

\[X_M = \frac{N\Sigma_n \left[ U_n(1)^2 \right]}{kT} \exp\left(-\frac{U_n(0)}{kT}\right)\]

it is possible to write down expressions for \(X_M\), \(X_M_x\), and \(X_M_y\), \(X_M_z\) (i.e., \(X_M\) when all the molecules have their \(x\), \(y\), or \(z\) axis, respectively, parallel with the magnetic field). The bulk susceptibility of the powdered sample will then be

\[X_M = X_M_x + X_M_y + X_M_z\]

This results in

\[X_M = \frac{3N\beta^2}{(1 + \exp(-D/E))) + \exp(-D/E))\left(\frac{g_x^2}{E} - \frac{g_x^2}{E} - \frac{g_y^2}{E} - \frac{g_z^2}{E}\right)}\]

At low temperatures this expression for \(X_M\) approximates

\[X_M = \frac{2N\beta^2}{3(3kT - 2D)}(\frac{g_x^2}{D} + \frac{g_y^2}{D} + \frac{g_z^2}{D})\]

This means that at low temperatures \(\chi\) is independent of the temperature which is in agreement with our measurements. In order to obtain a rough estimate for the value of \(D\), we set \(E\) equal to zero and take for \(g_x\) and \(g_y\) the value of 2. The measured constant low-temperature \(1/X_M\) equals 29.3 so \(D = 81.4 \times 10^{16}\) erg = 40.9 cm\(^{-1}\).

This zero-field splitting value is of the same order of magnitude as found by Dale\(^{10}\) for iron(II) phthalocyanine where \(D = 69.9\) cm\(^{-1}\). At higher temperatures the expression for \(X_M\) approximates

\[X_M = \frac{2N\beta^2}{3(3kT - 2D)}(\frac{g_x^2}{D} + \frac{g_y^2}{D} + \frac{g_z^2}{D})\]

This formula has the Curie-Weiss form \(\chi = C(T - \Theta)\) which is also in agreement with our measurements.

Extended Huckel molecular orbital calculations are consistent with a triplet ground state for these planar Co \(_4\) complexes. The calculations were carried out using the VSIP's published by Basch, et al.,\(^{11}\) as diagonal elements \(H_{ii} = \langle a_i | H | a_i \rangle\) in the Hamiltonian matrix and using the Wolsberg-Helmholz\(^{12}\) approximation for the estimation of \(g^2\).

Figure 4. Axis system chosen for the extended Hückel calculations.

The off-diagonal elements \( \langle \alpha_0 | H | \alpha_0 \rangle = \frac{1}{2} (H_{ii} + H_{jj}) \) \( S_{ij} \).

Atomic orbitals were described by single-\( \xi \) Slater-type orbitals for \( s \) and \( p \) orbitals with \( \xi \) values taken from Clementi and Raimondi(12) and from Richardson, et al.,(14) and double-\( \xi \) functions were used for the description of the Co \( 3d \) orbitals taken from Richardson, et al.(15) The MO diagram of the unsubstituted Co(bi)\(_4\)\(^-\) ion was calculated using interatomic distances as in the alkyl-substituted complex.(4) Assuming \( D_{2h} \) symmetry average values were used for the bond lengths. The cartesian axes system used is given in Figure 4. Iterative calculations were performed using a charge criterion of 0.002. Three different values for the Wölfserg-Helmholz parameter \( K \) were used (2.50, 2.00, 1.75). The \( d \) orbitals show little overlap with ligand orbitals. There are five molecular orbitals which have very large coefficients for the Co \( 3d \) atomic orbitals and they can be considered as almost pure \( d \) orbitals (see Table III). The energy difference between these molecular orbitals appeared to be strongly dependent on the value of \( K \). Quantitative correlation between observed spectra and calculated \( d \)-orbital splittings cannot be expected to exist due to the inherent inaccuracy in the Hückel approximation. The most important result of these calculations is that the \( d_{xz} \) and \( d_{x^2-y^2} \) orbitals have almost the same energy for all values of \( K \) (see Table III). The configuration \((\frac{1}{2}d_{xz})^3(d_{x^2-y^2})^3\) is in excellent agreement with the observed magnetic triplet ground state which is retained even at temperatures as low as 6°K. The calculated d-orbital sequence is the same as found by Shupack,(11) et al.,(16) and used by Baker-Hawkes, et al.,(17) to explain the paramagnetism of Co(dt)\(_2\)^+.

One of the molecular orbitals which is occupied by an unpaired electron has \( xz \) symmetry. It is largely built up from the Co \( d_{xz} \) atomic orbital but there is also a contribution from all the \( p_z \) orbitals of the ligand atoms. The \( p_z \) orbital of the nitrogen atom on the 3 position has a coefficient of about 0.1 (see Table IV). In view of this fact it is not surprising that the value of the polarographic half-wave potential is in part determined by the nature of the substituent on the remote 3 position as its influence can be transferred through the delocalized \( \pi \)-electron system up to the cobalt atom (see Table V). The influence of a \( \pi \)-electron-withdrawing substituent on the 1 position has a much more pronounced effect as can be seen in Table V comparing the half-wave potentials of the ethylene(biuret) complex with that of the o-phenylene(biuret) complex.

As the molecular orbitals in which the unpaired electrons are present have \( xz \) and \( x^2-y^2 \) symmetry, the electronic ground state in \( D_{2h} \) symmetry is \( ^3B_2g \). All compounds have \( D_{2h} \) symmetry except the \( o \)-phenylene(biuret) and the ethylene(biuret) complex. In the latter no extensive delocalization over the ethylene bridge is possible. In the \( o \)-phenylene(biuret) complex the \( xz \) molecular orbital, consisting of the \( 3d_{xz} \) atomic orbital and all ligand \( p_z \) orbitals, has no center of symmetry in contrast with the situation in any other of the compounds reported here. So some transitions involving this orbital will no longer be Laporte forbidden. This accounts for the larger extinction coefficients found for some of the d-\( d \) transitions in the \( o \)-phenylene(biuret) complex.

**Conclusions**

The paramagnetic (\( S = 1 \)) cobalt(III)-bis(biuretato) complexes show a normal Curie-Weiss magnetic behavior from room temperature down to \(-150^\circ\). Below 50°K deviation from the Curie-Weiss behavior occurs due to zero-field splitting but the triplet ground state is not affected. The unpaired electrons are largely localized in the Co \( 3d_{xz} \) and \( 3d_{x^2-y^2} \) atomic orbitals. The polarographic half-wave potential is

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**Table III. Coefficients of Co 3d Atomic Orbitals in the Four Highest Filled and the First Empty Molecular Orbitals, together with the Energies of These Orbitals for Different Values of the Wölfserg-Helmholz Parameter**

<table>
<thead>
<tr>
<th>( K )</th>
<th>( d_{xy} )</th>
<th>( d_{x^2-y^2} )</th>
<th>( d_{xx} )</th>
<th>( d_{yy} )</th>
<th>( d_{zz} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.50</td>
<td>0.93382</td>
<td>-0.39514</td>
<td>0.98489</td>
<td>0.97602</td>
<td>0.94288</td>
</tr>
<tr>
<td>2.00</td>
<td>0.94451</td>
<td>-0.39514</td>
<td>0.98588</td>
<td>0.97607</td>
<td>0.92285</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>( K )</th>
<th>Energy, eV</th>
<th>( K )</th>
<th>Energy, eV</th>
<th>( K )</th>
<th>Energy, eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.50</td>
<td>-0.94919</td>
<td>2.00</td>
<td>-10.86347</td>
<td>1.75</td>
<td>-13.76943</td>
</tr>
<tr>
<td>2.00</td>
<td>-0.94919</td>
<td>2.00</td>
<td>-10.86347</td>
<td>1.75</td>
<td>-13.76943</td>
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<tr>
<td>1.75</td>
<td>-10.86347</td>
<td>2.00</td>
<td>-10.86347</td>
<td>1.75</td>
<td>-13.76943</td>
</tr>
</tbody>
</table>

**Table IV. Coefficients of \( p_z \) Atomic Orbitals of Ligand Atoms in the MO that Largely Consists of Co 3d\(_{xz}\)**

<table>
<thead>
<tr>
<th>Compd</th>
<th>( K )</th>
<th>1/4 ( E_{1/2} )</th>
<th>( E_{1/4} - E_{1/2} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>KCo(3-Pr(bi)), ( \cdot )H(_2)O</td>
<td>2.50</td>
<td>-0.71</td>
<td>60</td>
</tr>
<tr>
<td>KCo(3-Ph(bi)), ( \cdot )DMSO</td>
<td>2.00</td>
<td>-0.64</td>
<td>58</td>
</tr>
<tr>
<td>Bu(_4)Co(bi)(_3)</td>
<td>1.75</td>
<td>-0.65</td>
<td>64</td>
</tr>
<tr>
<td>Bu(_4)Co(en(bi))(_3)</td>
<td>2.50</td>
<td>-0.63</td>
<td>60</td>
</tr>
<tr>
<td>KCo(o-phen(bi))(_3)</td>
<td>2.00</td>
<td>-0.47</td>
<td>60</td>
</tr>
</tbody>
</table>

\( a \) All potentials were measured with 0.1 \( N \) Bu\(_4\)NClO\(_4\) as supporting electrolyte. All waves were of the reduction type.

\( b \) Theoretical value: \((R/\gamma|^2|\Pi)\) in 9. At 20° this value is 55.5 mV for a reversible one-electron transfer.

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tentials are dependent on the nature of substituents on the biuret ligand probably due to electron donating or withdrawing through the delocalized π-electron system which overlaps with the Co 3d\textsubscript{zz} orbital.

**Registry No.** KCo(bi)\textsubscript{2}, 38637-46-6; KCo(3-Ph(bi))\textsubscript{2}·2DMSO, 38637-47-7; Bu\textsubscript{4}NCo(3-Ph(bi))\textsubscript{2}, 38637-48-8; Bu\textsubscript{4}NCo(bi)\textsubscript{2}·H\textsubscript{2}O, 38637-49-9; KCo(3-Pr(bi))\textsubscript{2}(1-Pr(bi)H\textsubscript{2})\textsubscript{2}, 36472-84-1; KCo(3-Pr(bi))\textsubscript{2}·2H\textsubscript{2}O, 38637-51-3; KCo(3-Pr(bi))\textsubscript{2}·2DMSO, 38637-52-4; KCo(en(bi))\textsubscript{2}, 38637-53-5; KCo(o-phen(bi))\textsubscript{2}, 38637-54-6; Bu\textsubscript{4}NCo(en(bi))\textsubscript{2}, 38637-55-7; Bu\textsubscript{4}NCo(o-phen(bi))\textsubscript{2}·CHCl\textsubscript{3}, 38637-56-8; KCo(3-Pr(bi))\textsubscript{2}·(NH\textsubscript{3})\textsubscript{2}, 31282-33-4; KCo(3-Pr(bi))\textsubscript{2}(C\textsubscript{3}H\textsubscript{7}NH\textsubscript{2})\textsubscript{2}, 31282-34-5; KCo(3-Pr(bi))\textsubscript{2}(pyrrolidine)\textsubscript{2}, 31249-56-6; KCo(3-Pr(bi))\textsubscript{2}(pyridine)\textsubscript{2}, 31240-71-8.

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