Title

Crystalline salts of metal phthalocyanine radical anions [M(Pc˙3−)]˙− (M = CuII, PbII, VIVO, SnIVCl2) with cryptand(Na+) cations: structure, optical and magnetic properties

Author(s)

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Dmitri V. Konarev, Maxim A. Faraonov, Alexey V. Kuzmin, Salavat S. Khasanov, Yoshiaki Nakano, Semen I. Norko, Mikhail S. Batov, Akihiro Otsuka, Hideki Yamochi, Gunzi Saito and Rimma N. Lyubovskaya

Radical anion salts of metal phthalocyanines (cryptand\([2,2,2](\text{Na}^+)\))[\(M(Pc\cdot^3\cdot^\cdot^3\cdot)]\cdot^\cdot^\cdot\] \((M = \text{Cu}^{II}, \text{Pb}^{II}, \text{V}^{IVO}, \text{Sn}^{IVCl_2}\)) and (cryptand\([2,2,1](\text{Na}^+)\))[\(Sn^{IVCl_2}(Pc\cdot^3\cdot^\cdot^3\cdot)]\cdot^\cdot^\cdot\] \(C_6H_4Cl_2\) \((M = \text{Cu}^{II}\), \(\text{Sn}^{IVCl_2}\) have been obtained as single crystals. Phthalocyanine planes form channels to accommodate the cryptand\([2,2,2](\text{Na}^+)\) cations and solvent \(C_6H_4Cl_2\) molecules in \(1\)–\(3\). Nearly square phthalocyanine layers are formed in salts \(4\) and \(5\), which are parallel to the \(ac\) plane and have an effective \(\pi\)–\(\pi\) interaction between the Pc\(^{\cdot^3\cdot^\cdot^3\cdot}\) macrocycles. The using of smaller cryptand\([2,2,1](\text{Na}^+)\) cations in \(5\), instead of larger cryptand\([2,2,2](\text{Na}^+)\) cations in \(4\), allows one to obtain more closely packed Pc layers. Reduction in the salts was centered on the Pc macrocycles, providing alternation of the C–N (imine) bonds in Pc due to partial loss of aromaticity. New bands were also observed in the NIR spectra at 946–1006 nm and the Soret and Q-bands were essentially blue-shifted in the spectra of the salts. The \([\text{Cu}^{II}](Pc\cdot^3\cdot^\cdot^3\cdot)]\cdot^\cdot^\cdot\) and \([V^{IVO}(Pc\cdot^3\cdot^\cdot^3\cdot)]\cdot^\cdot^\cdot\) radical anions contain paramagnetic \(\text{Cu}^{II}\) and \(V^{IVO}\) centers and the \(Pc^{\cdot^3\cdot^\cdot^3\cdot}\) radical trianions, both having a \(S = 1/2\) spin state. These radical anions were nearly isolated in the salts. Weak antiferromagnetic coupling with a Weiss temperature \((\theta) = \approx 2^\circ\text{K}\) was observed in \(1\). The magnetic behavior of \(3\) was described by a modified singlet–triplet model with a ferromagnetic exchange coupling of \(J/k_B = 7.7^\circ\text{K}\) within \([V^{IVO}(Pc\cdot^3\cdot^\cdot^3\cdot)]\cdot^\cdot^\cdot\) and \(\theta = \approx 1^\circ\text{K}\). The stronger antiferromagnetic coupling with a Weiss temperature of \(\theta = \approx 64^\circ\text{K}\) was observed in \(4\) and was accompanied by the decrease of molar magnetic susceptibility below \(35^\circ\text{K}\) due to antiferromagnetic ordering of spins. The magnetic behavior of \(4\) was described using the Heisenberg model for quasi-square two-dimensional antiferromagnetic coupling between spins with an exchange interaction of \(J/k_B = \approx 18^\circ\text{K}\).

Introduction

Metal-free and metal-containing phthalocyanines (Pcs) in oxidized or reduced forms can have promising optical, conducting and magnetic properties. For example, oxidation of axially substituted phthalocyanine \([\text{M}^{III}L_2Pc]\) anions \((M = \text{Co, Fe}; L = \text{CN, Cl, Br})\) allows one to obtain crystalline salts with a \(\pi\)–\(\pi\) stacking columnar arrangement of the macrocycles showing quasi-one-dimensional metallic behavior down to liquid helium temperatures. Salts with Fe\(^{III}\) as a central metal atom show giant magnetoresistance. Chain compounds with ferrimagnetic ordering between spins were obtained with oxidized manganese phthalocyanine.

Less attention was paid to the preparation of compounds with reduced metal phthalocyanines most likely due to their air and moisture sensitivity. However, some theoretical works predict high conductivity, including even superconductivity, for electron doped metal phthalocyanines. At present, powdered anionic Pc compounds can be prepared by doping of metal phthalocyanines by alkali metals in the gas phase. Several crystalline salts were obtained by metal phthalocyanine reduction with alkali metals or the KCl reductant in coordinating solvents.

**PAPER**


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like tetrahydrofuran or 1,2-dimethoxyethane. Recently, we have developed a method to prepare crystalline radical anion salts of metal-free and metal-containing phthalocyanines by neutral Pc reduction with sodium fluorenone ketyl in the presence of organic cations. As a result, salts of iron(I) and cobalt(I) phthalocyanine anions whose reduction is centered on metal atoms were obtained. A series of radical anion (cation \{M(Pc^{3-})\}^- and anions, \{cryptand[2,2,2](Na+)\}) were synthesized and structurally characterized. This allows one to study crystal structures, optical and magnetic properties of radical anion salts with different Pcs as well as to estimate the effect of reduction on their geometry, optical and magnetic properties.

To study how the geometry of metal Pc and size of cations affect the packing of Pc macrocycles in a crystal, we obtained a series of metal phthalocyanine salts with cryptand[2,2,2][Na+] and cryptand[2,2,1][Na+] counter cations, whose compositions are shown in Table 1. Until now, only several metal phthalocyanine salts with cryptand[2,2,2][Na+] cations are known. Those are the salts with iron phthalocyanine [Fe(Pc^{3-})]^- anions, \{cryptand[2,2,2][Na+]\}(\{H_2Pc\} ) \cdot 1.5C_6H_4Cl_2 and \{cryptand[2,2,2][Na+]\}(\{Sn^{VCl_2}(Pc^{3-})\}^- \cdot C_6H_4Cl_2). Obtained salts are categorized in two types according to the packing manner of Pcs: with channels formed by Pc macrocycles or with closely packed Pc layers. Pcs are nearly isolated in the structures of the first type (1–3) and magnetic interactions are realized only between the spins of the metal center and the Pc^{3-} macrocycle. At the same time, salt 4 with a layered structure shows strong coupling between spins with long-range antiferromagnetic ordering below 35 K. We report and discuss the syntheses, crystal structures, optical and magnetic properties of these salts.

### Results and discussion

#### 1. Synthesis

Syntheses of 1–3 were carried out in the presence of the organometallic compound [Cp*M(CO)3]2, which essentially improved the crystal growth and allows one to obtain high-quality single crystals but was not incorporated in the crystals. Interestingly, similar syntheses with Cu^2+Pc, Pb^2+Pc and V^5+Pc without [Cp*M(CO)3]2 afforded only polycrystalline products composed of small-sized crystals in low yield.

In the case of Sn^VCl_2Pc, direct reduction with sodium fluorenone ketyl does not allow the preparation of crystals probably due to possible substitution of fluorenone ketyl radical anions for chloride anions interfering the formation of good-quality crystals. Similar substitution of chloride anions by fluorenone ketyl radical anions is observed, for example, when boron subphthalocyanine chloride is reduced with sodium fluorenone ketyl. However, reduction of Sn^VCl_2Pc with Cao provides high-quality single crystals of radical anion salts 4 and 5. Since the (PPN)^+\{Sn^{VCl_2}(Pc^{3-})\}^+ salt was previously obtained by this method, it is seen that it is suitable for the preparation of crystalline radical anion salts of metal Pcs coordinated with chloroanions at the metal center.

#### 2. Spectra of phthalocyanine salts in the IR and UV-visible-NIR ranges

IR spectral data for the starting compounds and obtained salts are presented in Table S1 and Fig. S1–S4 (ESI†). The IR spectra of 1–4 show a superposition of the absorption bands of parent metal Pcs, cations and solvent molecules (see the ESI†), but some absorption bands of Pcs are essentially shifted or disappeared by the formation of the salts (see Table S1, ESI†).

Spectra of starting Pcs and salts 1–4 in KBr pellets in the UV-visible-NIR range are presented in Fig. 1, and Fig. S5 and S6 (ESI†) and are listed in Table 2. New intense bands are observed in the spectra of the salts in the NIR range of 946–1022 nm. These bands are characteristic of metal Pc radical anions, in which an additional electron is delocalized over the Pc macrocycle. The spectra of the salts based on Fe(II) and Co(II) Pc anions with the reduced metal centers do not show such bands. Thus, we can conclude that reduction is centered on the Pc macrocycles in 1–4.

Metal Pcs manifested intense absorption in the UV-visible region, in which the Soret and split Q-bands were located at 333–381 and 620–850 nm, respectively (see Table 2). The formation of 1–4 provided noticeable narrowing of Q-bands and the splitting into two or three bands was still observed in the spectra of the salts (Fig. 1). In all cases, the formation of the salts was accompanied by noticeable blue shifts of Q-bands by 0.04–0.15 eV (15–51 nm) and Soret bands by 0.01–0.31 eV (1–33 nm). Such shifts are generally observed for the metal Pc radical anions, whose Pc macrocycle is reduced. The largest shift of both Soret and Q-bands was observed during the formation of 4 with [Sn^{VCl_2}(Pc^{3-})]^+.

#### 3. Crystal structures of salts 1–5 and molecular structures of \([M(Pc^{3-})]^+\)

Geometric parameters of metal Pcs in neutral and radical anion states of 1–4 are shown in Table 3.

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**Table 1 Composition of salts 1–5**

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>1</td>
<td>{Cryptand[2,2,2][Na+]}(Cu^2+Pc^{3-})^- \cdot C_6H_4Cl_2</td>
</tr>
<tr>
<td>2</td>
<td>{Cryptand[2,2,2][Na+]}(Pb^2+Pc^{3-})^- \cdot C_6H_4Cl_2</td>
</tr>
<tr>
<td>3</td>
<td>{Cryptand[2,2,2][Na+]}(V^5+Pc^{3-})^- \cdot C_6H_4Cl_2</td>
</tr>
</tbody>
</table>
| 4  | {Cryptand[2,2,2][Na+]}(Sn^{VCl_2}(Pc^{3-})^- |}
| 5  | {Cryptand[2,2,2][Na+]}(Sn^{VCl_2}(Pc^{3-})^- \cdot C_6H_4Cl_2 |

\(\text{Cryptand[2,2,2]}\) \(\text{Na+}\) \(\text{SnIVCl2}\) \(\text{Pc}\) \(\text{VIVO}\) \(\text{CuII}\) \(\text{PbIIPc}\) \(\text{VIVOPc}\) \(\text{Cp*Mo(CO)2}\) \(\text{FeI}\) \(\text{CoI}\)
The reduction of metal phthalocyanines was accompanied by slight shortening of equatorial M–N bonds in 1–4 relative to parent Pcs (Table 3). Salts 1 and 3–5 contain Pc macrocycles with nearly planar conformation. In this case only phenylene groups slightly deviate from planarity (Table 3). However, only copper atoms are located exactly in the 24-atom Pc plane in 1. Tin atoms come out insignificantly from this plane (0.007 Å). Lead and vanadium atoms deviate significantly away from the 24-atom Pc plane, and the deviation was equal to 1.461 Å for the lead atoms, which is noticeably enhanced than 1.364 Å for octakis(hexylsulfanyl)phthalocyaninato lead(ii).

The Pc macrocycle has an aromatic 18 π-electron system. Reduction of this macrocycle yields an 19 π-electron system and should partially disrupt the aromaticity of the system. Indeed, Pc has 8 nearly equal C-N(imine) bonds in neutral form (Table 3) whereas alternation of these bonds in 1–3 supports partial loss of aromaticity in the radical trianion Pe(−3)− macrocycle. The differences between shorter and longer C-N(imine) bonds are in the range of 0.018–0.045 Å (Table 3). The bonds are positioned in such a way that shorter and longer bonds belong to the oppositely located isoidole units. Such changes in the molecular structure of the reduced Pc macrocycles are characteristic of the [M(Pe(−3)−)]− salts.18–21

Dependent on the geometry of parent metal Pcs, the formation of two types of salts was observed. In the case of the absence of axial ligands or the presence of only one axial ligand in 1–3, structures with channels are formed. The walls of these channels are built with four Pc planes and the channels are occupied by alternating cryptand[2,2,2](Na+) cations and solvent C6H4Cl2 molecules. Similar channel structures were previously found for the radical anion salts of free-base, iron and tin(ii) phthalocyanines with the cryptand[2,2,2](Na+) cations.14–17 In spite of the channel structure, Pcs form pairs in salts 2 and 3 with one side-by-side van der Waals (vdW) C ⋯ C contact between Pe(−3)− in 3 (Fig. 2b) and without such contacts in 2 (Fig. S7, ESI†). It was found that the [VIVO(Pe(−3)−)]− radical anions considerably tend to form π ⋯ π stacking dimers in the radical anion salts,16–27 but it is not the case for 3. Therefore, this is the first structure of [VIVO(Pe(−3)−)]− with nearly isolated Pc macrocycles. Chains formed by phthalocyanines can be outlined in 1 along the a axis due to the presence of vdW contacts of 3.39 Å length between them. However, those are only side-by-side vdW C ⋯ C contacts (Fig. 2a).

In the case of [SnIVCl2(Pe(−3)−)]− with two axial ligands, closely packed Pc layers parallel to the ac plane are formed in 4 (Fig. 3a). The layers are separated by the cryptand[2,2,2](Na+) cations and solvent C6H4Cl2 molecules (Fig. 3b). Salt 5 is isostructural to 4, but has much more closely packed Pc layers due to the smaller size of the cryptand[2,2,1](Na+) cations (Fig. 4). As a result, the number of vdW contacts between [SnIVCl2(Pe(−3)−)]− in 5 (30 for each unit, Fig. 4) exceeds the number of such contacts in 4 (14 for each unit, Fig. 3a). These contacts shown by green dashed lines in Fig. 3a and 4 are in the range of 3.28–3.56 Å. That is a first example of such closely packed phthalocyanine layers in the radical anion salts. The distances between tin atoms along the a and c axes are also smaller for the layers in

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5 (9.975 and 10.580 Å, respectively) in comparison with those in the layers in 4 (10.217 and 10.702 Å, respectively). Thus, we can affect the packing of Pc layers by changing the size of the cations.

Pc layers in 4 and 5 have almost uniform square arrangement of the \([\text{Sn IV}Cl_2(Pc^\cdot/C_{15}/C_{0}/C_{15}/C_{0})]^-\) radical anions, but according to the number of vDW contacts more closely packed chains can be outlined in 4 along the \(a\) axis (Fig. 3a). At the same time, the layers are more uniform in 5 with nearly equal number of vDW contacts in both \(a\) and \(c\) directions (Fig. 4).

4. Magnetic properties

Magnetic measurements by SQUID and EPR techniques were carried out for polycrystalline samples of 1, 3 and 4 under strictly anaerobic conditions (Table 4).

Magnetic moments of 1 and 3 were 2.44 and 2.19 \(\mu_B\), respectively, at 300 K (Fig. S8a and Fig. 5a), which is in agreement with the contribution of two independent \(S = 1/2\) spins per formula unit (the calculated magnetic moment is 2.45 \(\mu_B\)).

Temperature dependences on the \(\chi M T\) values are shown in Fig. S9a and b (ESI†). Since both radical anions contain paramagnetic metal centers, we can suppose that one \(S = 1/2\) spin is localized on the metal atom and the other \(S = 1/2\) spin is delocalized over the Pc radical macrocycle. The Weiss temperature

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</table>
for 1 is only $-2 \text{ K}$, indicating weak antiferromagnetic coupling between Cu$^{II}$ and Pc$^{3-}$ spins within one radical anion (Fig. S8b, ESIT). Similarly the salts based on copper[n] phthalocyanine in oxidized Cu(Pc)I or reduced (Bu$_4$N$^+$)[Cu$^{II}$(Pc$^{3-}$)]$^+$ ($\text{Br}^-$) forms also have close Weiss temperatures of $-4.2$ and $-4 \text{ K}$, respectively, indicating weak antiferromagnetic coupling between spins localized on Cu$^{II}$ and delocalized over the Pc macrocycle.$^{4,16}$

A Weiss temperature of $+2 \text{ K}$ in 3 determined in the 50–300 K range indicates an apparent weak ferromagnetic coupling between spins (Fig. 5b). The effective magnetic moment of 3 increases below 60 K, and an abrupt decrease is observed below 10 K (Fig. 5a) showing a competition between ferro- and antiferromagnetic interactions in 3. Since the [V$^{IV}$(Pc$^{3-}$)]$^+$ radical anions are nearly isolated in 3, we can suppose that magnetic coupling is realized within the radical anions having two $S = 1/2$ spins, which can access the triplet and singlet states. Indeed, the magnetic behavior of 3 is described well (Fig. 5c) by the modified singlet–triplet (S–T) model$^{28}$ with a ferromagnetic exchange interaction between the spins of $J/k_B = 7.7 \text{ K}$, a Weiss temperature of $-1.3 \text{ K}$ (antiferromagnetic), a $g$ value of 2 (fixed) and a scaling factor of $f = 0.782$. Previously, it was shown that the exchange interaction between the V$^{IV}$ and Pc$^{3-}$ spins in [V$^{IV}$(Pc$^{3-}$)]$^+$ is also rather weak and has an antiferromagnetic nature. For example, a Weiss temperature of $-9.6 \text{ K}$ is observed in [Bu$_3$N$^+$][V$^{IV}$(Pc$^{3-}$)]$^+$ also with an isolated arrangement of the macrocycles. The exchange interaction between the V$^{IV}$ and Pc$^{3-}$ spins was estimated to be only $J/k_B = -15.2 \text{ K}$ in (Me$_3$P$^+$)[V$^{IV}$(Pc$^{3-}$)]$^+$ (TPC)$_{0.5}$C$_{6}$H$_{4}$Cl$_{2}$ (TPC is triptycene) with $[[V^{IV}O(Pc)^{3-}]^{+}]_2$ π-stacking dimers. In this case the intermolecular coupling between the spins delocalized over the Pc$^{3-}$ macrocycles is $J/k_B = -105 \text{ K}$ due to the effective π–π interaction between the macrocycles.$^{27}$ These data show that despite the two spins being located in one anion at close distances, magnetic coupling between them is rather ineffective and many examples support this supposition. Intermolecular coupling in the presence of π–π interaction between the macrocycles is generally several times greater.

At the same time in both cases (salts 1 and 3) intermolecular coupling is very weak due to the absence of π–π interaction between the macrocycles in the perpendicular arrangement. That is seen if we consider the magnetic behavior of the salt with the radical anions of metal-free phthalocyanine with cryptand[2,2,2][Na$^+$] cations. It has only one $S = 1/2$ spin delocalized over the Pc$^{3-}$ macrocycle and their perpendicular arrangement results in the absence of magnetic coupling (Weiss temperature is in the range of 0 to $-1 \text{ K}$).$^{17}$

Salt 4 has an effective magnetic moment of 1.83 $\mu_B$ at 300 K (Fig. 6a) from one non-interacting $S = 1/2$ spin per formula unit delocalized on the Pc$^{3-}$ macrocycle (the calculated value is 1.73 $\mu_B$). The temperature dependence of the $\chi_M/T$ value is shown in Fig. S9c (ESI†). A Weiss temperature for 4 of $-64 \text{ K}$ in the 70–300 K range (Fig. 6b) indicates strong antiferromagnetic coupling between spins in the Pc layers (Fig. 3a). The deviation from the Curie–Weiss law to the antiferromagnetic side was observed below 70 K and was accompanied by a decrease of molar magnetic susceptibility below 35 K (Fig. 6c). The temperature behavior of molar magnetic susceptibility of 4 can be described well by the Heisenberg model for quadratic-layer antiferromagnetic coupling between spins$^{29}$ with an estimated exchange interaction of $J/k_B = -18 \text{ K}$ (Fig. 6c). The use of this model is supported by a nearly square arrangement of [Sn$^{IV}Cl_2$(Pc$^{3-}$)]$^+$ in the Pc layers of 4 (Fig. 3a).

That is a first example of such strong magnetic coupling between spins in the anion-radical phthalocyanine layers which...
is provided by effective π-π interactions within the layers. Previously even stronger antiferromagnetic coupling was observed only for the isolated π-stacking phthalocyanine [(M(Pc)²⁺)]⁺ ₁₂ dimers which show an exchange interaction up to J/\(k_B\) = −183 K. ²¹,²²

The EPR spectrum of 1 with two weakly interacting S = 1/2 spins shows two lines with \(g_1 = 2.0522\) and \(\Delta H = 6.37\) mT, and \(g_2 = 2.0794\) and \(\Delta H = 13.32\) mT, respectively, at 301 K (Fig. 7a).

Previously, two separate EPR signals from Cu²⁺ (\(g = 2.194\)) and Pc³− (\(g = 1.9999–1.9974\)) were observed for the radical anion (Bu₄N⁺)₂[Cu(II)(Pc³−)]⁺ salt. ¹⁸ Since \(g\)-factors of the signals in 1 have intermediate values between those characteristic of Cu²⁺ and Pc³−, we can conclude that these signals originate from both paramagnetic Cu²⁺ and Pc³− species having weak exchange interaction. The \(g\)-factors in 1 shift to smaller values with almost constant linewidth below 12 K (Fig. S10, ESI†).

Salt {cryptand[2,2,2](Na⁺)}[VIVO(Pc³−)]⁻·C₆H₄Cl₂ (3) shows two lines at RT with \(g_1 = 1.9923\) and \(\Delta H = 8.91\) mT, and \(g_2 = 2.0217\) and \(\Delta H = 1.44\) mT, respectively (Fig. 7b). The integral intensity of the narrower line is only 0.1% from that of the broad line and can be attributed to impurities. The broader main line according to the \(g\)-factor value and linewidth can be attributed to paramagnetic V¹⁺ and Pc³− species having weak exchange interactions since \(g\)-factors of the signals are intermediate between those characteristic of the main signal from V¹⁺ in neutral V¹⁺O(Pc³−) (\(g = 1.9858\) and \(\Delta H = 24.9\) mT) and the signal from the (H₂Pc)⁺⁺ salts (\(g = 2.0033–2.0037\) and \(\Delta H = 0.08–0.24\) mT). ¹⁷,¹⁸ A similar signal from both V¹⁺ and Pc³− were observed in (Bu₄N⁺)[V¹⁺O(Pc³−)]⁺ (\(g = 1.9921\) and \(\Delta H = 7.2\) mT at RT) ¹⁸ and in (Me₄P⁺)[V¹⁺O(Pc³−)]⁺ (TPC)₀.₅·C₆H₄Cl₂ (\(g_1 = 1.9954\) and \(\Delta H = 25.32\) mT, and \(g_2 = 2.0002\) and \(\Delta H = 10.49\) mT RT) ²⁷. The \(g\)-factor of the broad signal shifts to smaller values and narrows with the temperature decrease down to 40 K (\(g = 1.9901\) and \(\Delta H = 3.98\) mT at 40 K) (Fig. S11, ESI†). Below this temperature, the \(g\)-factor shifts slightly to higher values (\(g = 1.9910\) at 4 K) without noticeable broadening (\(\Delta H = 4.37\) mT at 4.1 K) (Fig. S11, ESI†).

Salt 4 manifests a broad Lorentzian line with \(g_1 = 1.9933\) and \(\Delta H = 29.9\) mT at 299 K (Fig. 7c) which strongly narrows with the temperature decrease and shifts to smaller \(g\)-factors (Fig. S12, ESI†). A similar broad signal is observed in [PPN⁺][SnCl₄(II)(Pc³−)]⁺. ⁹,¹⁹ The central metal atom even in the cases of diamagnetic salts, affects the EPR spectra of the radical anions salts. Instead of narrow signals characteristic of (H₂Pc)⁺⁺ (\(g\)-factor of 2.0033–2.0037, linewidth of 0.08–0.24 mT), ¹⁷,¹⁸ broad temperature-dependent signals are observed in the EPR spectra of the radical anion salts based on Ni³⁺Pc, Pb³⁺Pc, Sn³⁺Pc and SnIVCl₂Pc. ¹⁸,¹⁹ An increase in the linewidth of the EPR signals is also observed during the transition from the radical anion salt of aluminium(III)phthalocyanines to gallium(III)phthalocyanine and finally to indium(III)-phthalocyanine. ³⁰ Most probably this effect could be explained by weak transfer of spin density from the Pc³− macrocycle to the central metal atom accompanied by the formation of paramagnetic metal centers affecting the EPR spectra of the salts. Similar effects were previously reported for the oxidized Ni³⁺Pc species. ³¹,³²

**Experimental**

**Materials**

Cu³⁺Pc (Aldrich, sublimed grade, 99%), Pb³⁺Pc (TCI, 95%), V¹⁺O(Pc) (Acros, 85%), and SnIVCl₂Pc (TCI) were used as received. Cryptand[2,2,2] and cryptand[2,2,1] were purchased from Aldrich. C₆H₄ of 99.9% purity was received from MTR Ltd. The pentamethylcyclopentadienylmolybdenum dicarbonyl ([C₅⁺Mo(CO)₂]₂) was purchased from Strem. Sodium fluorenone ketyl was obtained as described. ²³ Solvents were purified under an argon atmosphere. o-Dichlorobenzene (C₆H₄Cl₂) was distilled over CaH₂ under reduced pressure. n-Hexane was distilled over Na/benzophenone. The solvents were degassed and stored in a glove box. The crystals of 1–5 were stored in the glove box. KBr pellets for IR- and UV-visible-NIR measurements were also prepared in the glove box. EPR and SQUID magnetic measurements were performed on polycrystalline samples of 1, 3 and 4 sealed in 2 mm quartz tubes with ambient pressure argon.

**Synthesis**

Crystals of 1–5 were obtained using the diffusion technique. The reaction solutions were filtered into a 50 mL glass tube of 1.8 cm diameter with a ground glass plug, and 30 mL of n-hexane was layered over the solution. Slow mixing of two solvents during 1–2 months provided precipitation of crystals on the walls of the tube. The solvent was decanted from the crystals and they were washed with n-hexane.

Salts: {cryptand[2,2,2](Na⁺)}[Cu³⁺Pc]C₆H₄Cl₂ (1), {cryptand[2,2,2](Na⁺)}[Pb³⁺Pc]C₆H₄Cl₂ (2), and {cryptand[2,2,2](Na⁺)}[VIVO(Pc³−)]⁻·C₆H₄Cl₂ (3) were obtained as described. ²³
[V^VOPc]_2C_6H_4Cl_2 (3) were obtained by the reduction of C_60 (30 mg, 0.042 mmol) with an excess of sodium fluorenone ketyl (16 mg, 0.0789 mmol) in the presence of a stoichiometric amount of cryptand[2,2,2] (16 mg, 0.042 mmol) for two hours at 100 °C until complete dissolution of metal Pc and the formation of a deep-blue solution. After that, the [Cp*Mo(CO)_2]_2 dimer (12 mg, 0.0209 mmol) was added to the solutions, and they were additionally stirred at 80 °C for two hours. Then the solutions were cooled down to room temperature, and filtered into the tube for diffusion. Their diffusion into the n-hexane layer produced crystals as black blocks (1, 2) and black prisms (3) in 56–75% yield. All crystals had copper luster characteristic of metal Pc salts. The organometallic compound [Cp*Mo(CO)_2]_2 was added in the synthesis of the salts to study the possibility of the formation of coordination complexes of metal Pcs with C_60(MoPC), which were previously described for (PPN +)[SnIVCl_2(Pc^V)] and [SnIVCl_2(Pc^V)]⁻ radical anions. In the case of Cu^IPc, Pb^IPc and V^IVOPc, coordination complexes were not formed. However, instead of that, high-quality and large single crystals of metal Pc salts were obtained.

(Cryptand[2,2,2][Na])[[Sn^IVCl_2Pc]C_6H_4Cl_2 (4) and {cryptand-[2,2,1][Na]}[[Sn^IVCl_2Pc]C_6H_4Cl_2 (5) were synthesized as previously described for ([PPN]^[Sn^IVCl_2(Pc^V)]^+)^9,19. In the case of Cu^IPc, Pb^IPc and V^IVOPc, coordination complexes were not formed. However, instead of that, high-quality and large single crystals of metal Pc salts were obtained.

Table 5 X-ray diffraction data for 1–5

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This solution was cooled down to room temperature and filtered into a flask containing 29.2 mg [0.042 mmol] of Sn^IVCl_2(Pc^V). The solution was stirred at 80 °C for 2 hours and overnight at room temperature to produce a deep blue solution, which was filtered into a glass tube for diffusion. The diffusion of the solution into the n-hexane layer produced the crystals as well-shaped black blocks of 4 in 72% yield and 5 in 32% yield. The addition of a stoichiometric amount of [Cp*Mo(CO)_2]_2 to the [Cp*Mo(CO)_2]_2/[Sn^IVCl_2(Pc^V)]⁻ salt in o-dichlorobenzene solution was accompanied by an immediate color change from deep blue to green, and a previously described coordination compound [Cp*MoBr_3(CO)_3]Sn^IVCl_2 was isolated from this solution, whose structure and properties were described.21

Compositions of 1–5 were determined from X-ray structural analysis on single crystals. The optical microscopic observation of the obtained crystals in a glove box, as well as a sample check of single crystals from each synthesis by X-ray diffraction, showed that only one crystal phase was formed. Elemental analysis cannot be used to determine the composition of the obtained crystals due to their high air sensitivity (i.e., addition of oxygen to the samples during the procedure of elemental analysis).

General

UV-visible-NIR spectra were recorded in KBr pellets on a PerkinElmer Lambda 1050 spectrometer in the 250–2500 nm range. FT-IR spectra were obtained in KBr pellets with a PerkinElmer Spectrum 400 spectrometer (400–7800 cm⁻¹). EPR spectra were recorded for sealed polycrystalline samples of 1, 3 and 4 from 4 up to 295 K with a JEOL JES-TE 200 X-band ESR spectrometer.

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equipped with a JEOL ES-CT470 cryostat. A quantum design MPMS-XL SQUID magnetometer was used to measure the static magnetic susceptibility of 1, 3 and 4 under a magnetic field of 100 mT in the range of 300–1.9 K upon cooling and heating processes. A sample holder contribution and core temperature independent diamagnetic susceptibility ($\chi_d$) were subtracted from the experimental values. The $\chi_d$ values for salts 1, 3, and 4 were obtained by the extrapolation of the data (10–300 K, 1), (50–300 K, 3) and (70–300 K, 4) in the high-temperature range by fitting the data with the following expression: $\chi_M = C(T - \theta) + \chi_d$, where $C$ is the Curie constant and $\theta$ is the Weiss temperature. Effective magnetic moments ($\mu_{\text{eff}}$) were calculated with the formula: $\mu_{\text{eff}} = (B/3M^2)^{1/2}$.

**Crystal structure determination**

X-ray diffraction data for 1–5 are listed in Table 5. The data were collected on an Oxford diffraction “Gemini-R” CCD diffractometer with graphite monochromated MoKα radiation using an Oxford Instrument Cryojet system. Raw data reduction to $F^2$ was carried out using CrysAlisPro, Oxford Diffraction Ltd. The structures were solved by direct methods and refined by the full-matrix least-squares method against $F^2$ using SHELX 2016 and Olex2. Non-hydrogen atoms were refined in the anisotropic approximation. The positions of hydrogen were calculated geometrically.

The crystal structure of 2 contains cryptand[$2,2,2$][Na+] cations severely disordered between three orientations with the occupancies of 0.541(4), 0.263(4) and 0.196(4). Crystal structures of 4 and 5 were solved with rather low C–C bond length precision (0.012 and 0.018 Å, respectively) which prohibited the discussion on the macrocycle geometries. To keep the anisotropic thermal parameters of the disordered atoms within reasonable estimation on the macrocycle geometries. To keep the anisotropic thermal parameters of the disordered atoms within reasonable limits, the displacement components were restrained using SHELXL instructions of ISOR, SIMU and DELU. That resulted in great numbers of restraints for the refinement of crystal structures of 2, 4 and 5 (Table 5).

**Conclusion**

A series of radical anion salts with [M(Pc$^{3-}$)]$^-$, where M = Cu$^{II}$, Pb$^{III}$, V$^{IV}$O, and Sn$^{IV}$Cl$_2$ with cryptand(Na$^+$) counter cations, was obtained for the first time. The formation of these salts was accompanied by the reduction of Pc macrocycles. That results on the geometry of parent metal Pcs (M = Cu$^{II}$, Pb$^{II}$, V$^{IV}$O), structures with channels accommodating the cryptand[$2,2,2$] (Na$^+$) cations and solvent C$_6$H$_4$Cl$_2$ molecules were formed. No effective $\pi$–$\pi$ interaction was observed between Pc macrocycles in these crystal structures due to the perpendicular arrangement of the Pc$^{3-}$ macrocycles. Salts 1 and 3 demonstrated weak magnetic coupling between spins due to the absence of $\pi$–$\pi$ interactions between Pc$^{3-}$. There is mainly antiferromagnetic coupling between the Cu$^{II}$ and Pc$^{3-}$ ions in 1 and ferromagnetic coupling between the V$^{IV}$ and Pc$^{3-}$ ions in 3. The weakness of this coupling shows ineffective magnetic coupling between two spins localized on the metal centers and delocalized over the Pc macrocycle. There are several examples of such behavior. Tin(IV) dichloride phthalocyanine with two axial ligands formed closely packed Pc layers in 4 and 5. The decrease of the cation size allows one to obtain densely packed Pc layers. The salts with the [Sn$^{IV}$Cl$_2$(Pc$^{3-}$)]$^-$ radical anions containing one S = 1/2 spin per formula unit delocalized over the Pc macrocycle. A layered structure together with effective $\pi$–$\pi$ interactions results in strong antiferromagnetic coupling between spins in 4 with an exchange interaction of $J/k_B = -18$ K. That is a first example of such strong magnetic coupling between spins in the anion-radical phthalocyanine layers. Further work on studying the effect of small cations on the structure and magnetic properties of the salts is in progress.

**Acknowledgements**

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**Notes and references**