



Communication Synthesis of Catena-bis(μ-bromo)-(O-methyl-Nphenylthiocarbamate)-dicopper(I) and Its Reactivity towards PAr₃ (Ar = Ph, *p*-Tol)

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Abstract: In order to investigate the coordination chemistry of *O*-alkyl *N*-aryl thiocarbamate ligands toward coinage metals, CuBr was reacted with one equivalent of MeOC(=S)N(H)Ph L in MeCN solution to afford the 1D-polymeric title compound $[{Cu(\mu_2-Br)_2Cu}{\mu_2-MeOC(=S)N(H)Ph}_2]_n$ **CP1**. Compound **1** was characterized by IR spectroscopy and an elemental analysis. The formation of a polymeric 1D ribbon built upon μ_2 -bridging bromido and thione ligands via the C=S bond was ascertained by a single-crystal X-ray diffraction study performed at 100 K. In the presence of PAr₃ (Ar = Ph, *p*-Tol), the polymer chain was broken to yield the mononuclear complexes [(Ar₃P)₂Cu{MeOC(=S)N(H)Ph}Br] **C1** and **C2**.

Keywords: copper bromide; thiocarbamate; thione; crystal structure; hydrogen bonding; aryl phosphine

1. Introduction

Thiocarbamates, also called thiocarbamides, with the general formula ROC(=S)N(H)R' play not only a role in organic chemistry as building blocks for further transformations but also in applications in various domains, such as biological and pharmacological activities or catalysis [1–5]. Furthermore, they have attracted the attention of coordination chemists, since the soft C=S thione function (according to Pearson's HSAB principle) [6] readily coordinates to a wide range of transition metal complexes, forming complexes with Cu(I), Ag(I), Au(I), Hg(II), Ru(II), Rh(III), etc. [7–11]. Furthermore, they may act as potential ditopic ligands capable of coordinating to a metal center both via their soft C=S function or through the harder HNR₂ group as demonstrated by the crystal structure determination of the square-planar bis(O-methylphenylthiocarbamato)(triphenylphosphine)-palladium [11]. We have recently demonstrated that O-methyl-N-phenyl thiocarbamate L can even be coordinated to *p*-block metal salts, such as BiI₃ [12]. Some representative examples of crystallographically characterized mono- and dinuclear complexes ligated by O-methyl-N-phenyl thiocarbamate are presented in Scheme 1 [7,8,12].

With the aim to investigate putative arene-C–H··· π (quasi-chelate ring) interactions in copper(I) crystal structures, Tiekink et al. reported in 2014 on the synthesis of the complex [(Ph₃P)₂Cu{MeOC(=S)N(H)Ph}Cl] by the reaction of CuCl with L in the presence of two equivalents of PPh₃. [7], shown as complex A in Scheme 1. In a continuation of our investigations on the coordination chemistry of thione-type ligands on diverse metal centers [13–16], our group extended the preparation of this type of Cu(I) complex



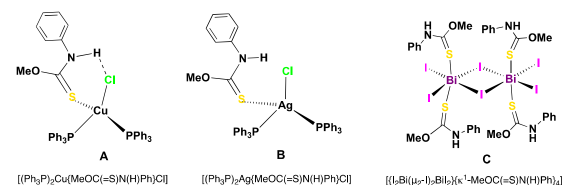
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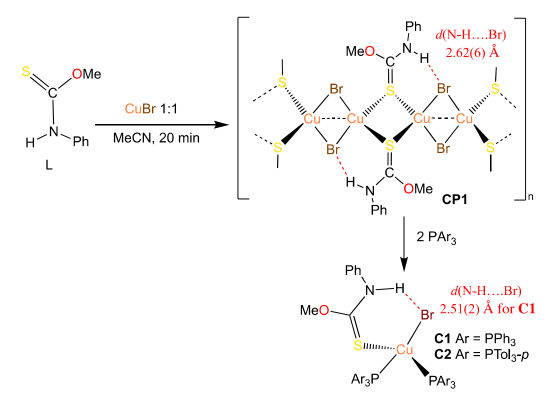
Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). to CuBr and CuI, systematically analyzed by crystal structure determinations a wide series of complexes $[(Ar_3P)_2Cu\{ROC(=S)N(H)Ph\}X)]$ (Ar = Ph, *p*-C₆H₄OMe; X = Br, I), and investigated the biological activity of these compounds featuring intramolecular N-H···Hal bonding, giving rise to six-membered cycles [17]. We also reported that the formation of $[(Ar_3P)_2Cu\{ROC(=S)N(H)Ph\}I)]$ is preceded in the initial phase of the reaction by the formation of mono-dimensional coordination polymers of type $[\{Cu(\mu_2-I)_2Cu\}(\mu_2-L)_2]_n$ [17]. We now investigate in more detail the outcome of the reaction of CuBr with L, describing herein the synthesis and crystal structure of the resulting coordination polymer $[\{Cu(\mu_2-Br)_2Cu\}(\mu_2-MeOC(=S)N(H)Ph\}_2]_n$ **CP1** and reporting on its reactivity towards PAr₃.



Scheme 1. Examples of some *O*-methyl *N*-arylthiocarbamate complexes.

2. Results and Discussion

The hitherto unknown coordination polymer [$\{Cu(\mu_2-Br)_2Cu\}\{\mu_2-MeOC(=S)N(H)Ph\}_2]_n$ **CP1** was obtained by the addition of one equivalent of O-methyl N-phenyl thiocarbamate L to a solution of CuBr in MeCN (Scheme 2). After stirring for 20 min, the precipitation of a colorless microcrystalline material occurred, which upon heating for 5 min almost redissolved. Upon cooling, colorless crystals started to grow. According to the elemental analysis of this air-stable material, one L molecule is attached per CuBr motif. This polymeric material, whose characterization is described below, reacts straightforwardly upon the addition of two equivalents of PPh₃ per CuBr unit to form by the splitting of the μ_2 -Br bridges of **CP1** the mononuclear tetrahedral complex [(Ph₃P)₂Cu{MeOC(=S)N(H)Ph}Br] C1 (Scheme 2). Since this complex has been recently fully characterized, we refer for its spectroscopic data to ref. [17]. In an analogous manner, **CP1** reacts with tris(p-tolyl) phosphine PTol₃-p to yield the derivative [(Tol₃P)₂Cu{MeOC(=S)N(H)Ph}Br] C2 with over an 85% yield as an air-stable crystalline solid. Its ³¹P NMR spectrum in CDCl₃ shows a single broadened resonance at δ –6.34 with a $\Delta\delta$ of 0.92 ppm with respect to free PTol₃ [18]. The ¹H NMR spectrum displays a strong singlet at δ 2.33 ppm, attributed to the six equivalent methyl groups of the *p*-tolyl rings. The characteristic signals of the thiocarbamate moiety at δ 4.06 and 10.54 ppm are assigned to -OCH and the NH group while the aromatic protons are observed as a multiplet between δ 7.03 and 7.39 ppm (Figure S2). In the proton-decoupled ¹³C NMR spectrum, a single resonance at δ 21.4 ppm is seen for the six magnetically equivalent Me substituents. The C-atom of the ligated thione function resonates at δ 187.3 ppm (Figure S3). The ATR-IR spectrum features a broadened N-H vibration at 3160 cm⁻¹, indicative of an intramolecular N-H…Br interaction, as was crystallographically evidenced for C1 (Figure S4). Upon the treatment of **CP1** with an excess of PPh₃, the dissociation of the coordinated thione ligand occurs, yielding finally, among other non-identified products, the cluster compounds [(PPh₃)₂Cu(μ_2 -Br)₂CuPPh₃] and [Cu₄Br₄(PPh₃)₄], which are already known in the literature, as evidenced by X-ray diffraction [19].



Scheme 2. Synthesis of the title compound CP1 and its reactivity with PAr₃ yielding C1 and C2.

Since the compound **CP1** is not soluble enough to be characterized by NMR spectroscopy, we examined the material crystallizing in the triclinic space group $P\overline{1}$ by an X-ray diffraction study performed at 100 K. As shown in Figure 1, a mono-dimensional ribbon has formed, which is constituted of rhomboid-shaped dinuclear {Cu(μ_2 -Br)₂Cu} units that are interconnected through the bridging C=S groups of the L molecules. Each Cu(I) center of **CP1** has thus a tetrahedral coordination sphere (neglecting the Cu···Cu interaction). The Cu–Cu contact within the {Cu(μ_2 -Br)₂Cu} rhomboid of 2.8339(10) Å falls within the limit of the Van der Waals radii of two Cu atoms (2.8 Å) and can be considered as weakly bonding. However, without computation, it remains speculative to discuss the contribution of metallophilic interactions, which can occur both in molecular compounds or coordination polymers incorporating neighboring coinage metal centers [20–22].

Similar values have been reported for the 1D CPs [{Cu(μ_2 -Br)_2Cu}{ μ_2 -PhC(=S)NH_2}]_n and [{Cu(μ_2 -Br)_2Cu}(μ -PhS(CH₂)_3SPh)_2]_n, in which the metal centers within the rhomboid clusters are separated by 2.857(3) and 2.7851(12) Å, respectively [23,24]. For the molecular compounds [{Cu(μ_2 -Br})_2}(1-oxa-4,7-dithiacyclononane)₂] (2.852(1) Å), [{Cu(μ_2 -Br})_2{dibenzo[e,m][1,4,8,11]dioxadithia-cyclotetradecine-S,S'}2] (2.919(7) Å), and the supramolecular compound tetrakis(4,5-bis(methylsulfanyl)-1,3-dithiole-2-thione)-bis(μ_2 -bromo)dicopper (2.893(1) Å), the Cu…Cu interaction lies in a similar range [16,25,26]. A further example is the thione-bridged dinuclear complex [Cu₂Br₂(μ -S-mimzSH)₂(η^1 -S-mimzSH)₂] (2.877(6) Å), obtained by the reaction of CuBr with 1-methyl-1,3-imidazoline-2-thione (mimzSH) in a 1:2 molar ratio [27].

This is in contrast with the recently described iodido analog [$\{Cu(\mu_2-I)_2Cu\}\{\mu_2-MeOC (=S)N(H)Ph\}_2$]_n **CP2**, for which a much shorter Cu–Cu separation of only 2.6563(7) Å was evidenced [17]. The C=S bond is slightly elongated with respect to that of non-ligated MeOC(=S)N(H)Ph (1.711(5) vs. 1.6708(11) Å) [28].

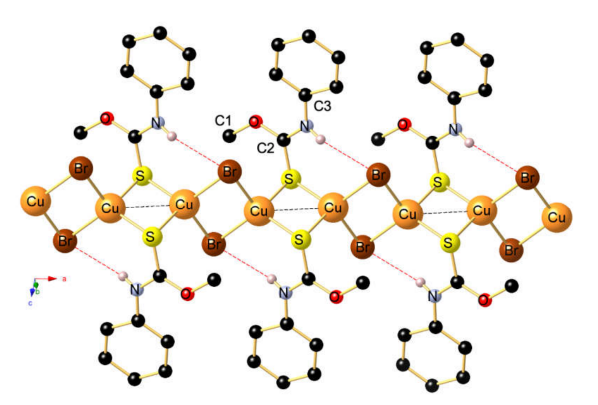


Figure 1. Crystal structure of the 1D ribbon of $[{Cu(\mu_2-Br)_2Cu}(\mu_2-L)_2]_n$ **CP1** running along the *a* axis. Selected bond lengths (Å) and angles (deg) of **CP1**: Cu···Cu¹ 2.8649(10), Cu–Cu² 2.8339(10), Br–Cu 2.4620(7), Cu–S 2.2966(12), Cu–S² 2.4157(12), S–C1 1.711(5), O–C1 1.313(5), O–C2 1.453(5), N–C1 1.335(5), N–C3 1.421(6); Cu²–Cu–Cu¹ 171.28(5), S–Cu–Br 117.68(4), S–Cu–Br¹ 107.64(4), S–Cu–S 106.11(4), Br–Cu–Br¹ 109.12(2), Br–Cu–Cu¹ 54.83(2), Br–Cu–Cu² 130.17(3), Br¹–Cu–Cu² 120.12(3), S–Cu–Cu¹ 131.40(4), S–Cu–Cu² 51.13(3), S–Cu–Cu¹ 131.30(4), N–C1–S 121.7(3), C8–C3–N 115.3(4), and C7–C8–C3 120.4(3). Symmetry transformation used to generate equivalent atoms: ¹ 2-*x*, 1-*y*, 2-*z*; ² 1-*x*, 1-*y*, 2-*z*.

Noteworthy is also the occurrence of the secondary N-H…Br interaction of 2.62(6) Å (dN …Br 3.440(4) Å, indicated by the red dashed lines in Figure 1. This interaction is noticeably shorter and in consequence stronger than the weak intramolecular N-H…I contacts of 2.748 Å and 2.803 Å encountered for **CP2** [17]. A similar systematic shortening of the intramolecular N-H…X bonding occurring in [(Ph₃P)₂Cu{MeOC(=S)N(H)Ph}X] (X = I, Br, Cl) and giving rise to six-member cycles has also been evidenced recently by us (see Scheme 1) [17]. Note that in the latter case, this intramolecular hydrogen bonding is even stronger for the molecular bromido complex **C1** compared to **CP1** (2.51(2) vs. 2.62(6)). Furthermore, a loose intramolecular S…H contact of 2.672 Å occurs between the sulfur atom and the H atom of the NH group.

In the packing, the ribbons of **CP1** are aligned in a co-parallel arrangement and are running along the *a* axis (Figure 2). There are also weak supramolecular interchain contacts between Br and the H atom of C7 of the aryl cycle (C7-H7… Br 2.919 Å).

To check the phase purity of this material, a grounded sample has been analyzed by powder X-ray diffraction (PXRD). The experimental PXRD pattern matches well with the simulated ones, confirming the homogeneity of the phases. A comparison of the PXRD patterns of **CP1** is illustrated in Figure 3.

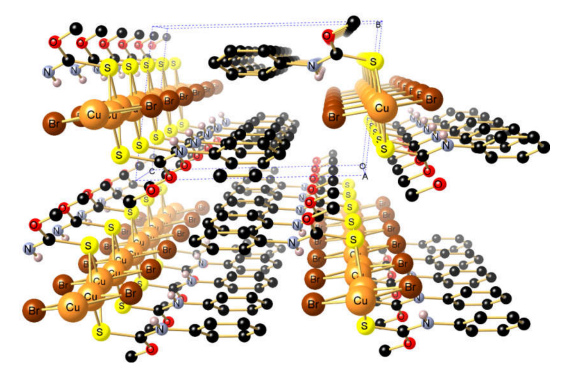


Figure 2. Perspective view of the packing, showing several ribbons of **CP1** running along the *a* axis in a parallel arrangement.

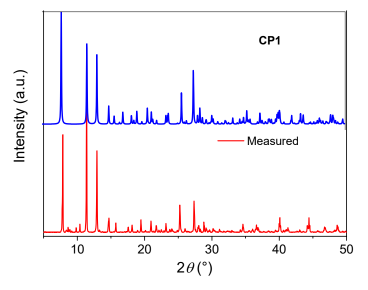


Figure 3. Comparison of the experimental PXRD pattern of CP1 with the simulated one (blue).

3. Experimental Section

Synthesis of **CP1**—We added 1 eq. of *O*-methyl *N*-phenylthiocarbamate L (334 mg, 2 mmol) to a solution of CuBr (286 mg, 2 mmol) in 12 mL of acetonitrile. A colorless precipitation was immediately noticed. The mixture was stirred 20 min at room temperature and then shortly heated till almost all of the precipitates had redissolved. After 1 d, pale yellowish crystals of **CP1** were formed and then collected by filtration. Yield: 79%. Anal. Calc. for C₈H₉BrCuNOS (M.W = 310.67 g·mol⁻¹). C, 33.09; H, 4.01; N, 4.29; S 9.81%. Found: C, 32.95; H, 4.00; N, 4.15; S 9.75. IR-ATR: 1026 (C–O), 1215 (C=S), 1440 v(C–N), 3199 v(N–H…Br), 3276 v(N–H) cm⁻¹.

Synthesis of **C1**—We added two equivalents of PPh₃ (52 mg, 0.2 mmol) to a suspension of **CP1** (31 mg, 0.1 mmol) in 4 mL of MeCN, prepared as described above. The mixture

was then heated for 20 min to 60 °C till all of the product was dissolved. Upon cooling, **C1** crystallized. The characterization data are identical to those described previously [18].

Synthesis of C2—We added two equivalents of PTol₃ (608 mg, 2 mmol) in several portions to an in situ prepared suspension of **CP1** (1 mmol). The mixture was then heated for 20 min to 60 °C till all product was dissolved. Upon cooling, **C2** crystallized in form of colorless plates and was isolated with 82% yield. Anal. Calc. for C₅₀H₅₁CuBrNOP₂S (M.W. = 919.43 g·mol⁻¹): C, 65.32; H, 5.59; N, 1.52; S, 3.49%. Found C, 65.20; H, 5.53; N, 1.59; S, 3.42%. ¹H NMR: δ 2.33 (s, 18H, CH₃), 4.08 (s, 3H, OCH₃), 7.04 (d, 12H, PC₆H₄, ³JHH = 7.9 Hz), 7.15–7.39 (m, 17H, Ph and PC₆H₄, ³JHH = 7.9, ³JPH = 9.7 Hz), 10.54 (s, br, 1H, NH). ¹³C{¹H} NMR: δ 21.4 (CH₃Ar), 58.4 (CH₃O), 121.9 (CH, Ph, Cmeta), 125.3 (CH, Ph, Cpara), 128.9 (CH, Ph, Cortho), 129.2 (d, CH, m-PC₆H₄, ³JCP = 9.2 Hz), 130.4 (d, Cq, CipsoPC₆H₄, ¹JCP = 27.7 Hz), 133.9 (d, CH, o-PC₆H₄, ²JCP = 14.9 Hz), 137.2 (s, Cq, Cipso), 139.4 (Cq, *p*-PC₆H₄), 187.3 (Cq, C=S). ³¹P{¹H}: δ – 6.34. IR-ATR: 1093 v(C–O), 1222 v(C=S), 1444 v(C–N), 3160, 3110 v(N–H) cm⁻¹.

Crystal data for C₈H₉BrCuNOS were as follows: M = 310.67 g·mol⁻¹, pale yellow crystals, crystal size $0.362 \times 0.102 \times 0.047$ mm³, Triclinic, space group $P\overline{1}$, a = 5.6823(2) Å, b = 7.6782(4) Å, c = 11.4019(5) Å, $\alpha = 97.689(2)^{\circ}$, $\beta = 91.877(2)^{\circ}$, $\gamma = 91.143(2)^{\circ}$, V = 492.58(4) Å³, Z = 2, $D_{calc} = 2.095$ g/cm³, T = 100 K, $R_1 = 0.0475$, $Rw_2 = 0.0976$ for 5163 reflections with I $\geq 2\sigma$ (I) and 2995 independent reflections. Largest diff. peak/hole e/Å⁻³ 1.04/-1.34. Data were collected on *Bruker D8 Venture* four-circle diffractometer equipped with a PHOTON II CPAD detector by Bruker AXS GmbH, using graphite-monochromated MoK_{α} radiation l = 0.71073 Å and were deposited at the Cambridge Crystallographic Data Centre as CCDC 2252606 (Supplementary Materials). The data can be obtained free of charge from the Cambridge Crystallographic Data Centre via http://www.ccdc.cam.ac.uk/getstructures, accessed on 3 February 2023. The structure was solved by direct methods and refined by full-matrix least-squares calculation against F² (SHELXL, 2015) [29–31].

4. Conclusions

We have crystallographically evidenced that CuBr reacts like CuI with a stoichiometric amount of *O*-methyl-*N*-phenylthiocarbamate to yield a halide-bridged mono-dimensional coordination polymer. We are currently extending this facile preparation of polymeric materials to other *O*-alkyl *N*-arylthiocarbamates, such as MeOC(=S)N(H)Cy, and are investigating the splitting of the polymer chain by other P and As donor ligands.

Supplementary Materials: The following supporting information can be downloaded online: CIF file, Check-CIF report, and IR and NMR spectra.

Author Contributions: W.A. and M.K. prepared the compounds; C.S., R.S. and L.V. collected the X-ray data and solved the structure; I.J., L.V. and M.K. designed the study and analyzed the data and M.K. wrote the paper. A.B.A., I.J. and M.K. contributed with the conceptualization. All authors have read and agreed to the published version of the manuscript.

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Conflicts of Interest: The authors declare no conflict of interest.

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