Synthesis and Molecular Structure of the First Metallamacrocyclic Bi(III)–Cu(II) 15–MC–5 Complex Derived from Pyrazinohydroxamic Acid

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Dedicated to Professor Dieter Wöhrle on the occasion of his 80th Birthday

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Polynuclear metallamacroyclic complexes named as metallacrowns are considered as a promising exceptional class of macroheterocyclic compounds that structurally resembles crown ethers. In this communication, we report a general synthetic approach and characterization of the first metallamacrocyclic Bi(III)-Cu(II) 15-MC-5 complex derived from pyrazinohydroxamic acid. The molecular and crystal structures of complex were determined from single-crystal X-ray diffraction studies. The single-crystal structure reveals the classic metallamacroyclic 15-MC-5 configuration. The Bi(III) ion is located at the center of the 15-MC-5 ring consisting of five [Cu(II)-N–O] repeat units. Detailed NBO and QTAIM studies based on DFT calculations were carried out for {Bi[15-MC^Cu(II)Pyzha-5]}^{3+} species.

Keywords: Metallacrown, metallamacrocyclic complex, bismuth, copper, pyrazinohydroxamic acid (Pyzha), X-ray structure, DFT.

Синтез и молекулярная структура первого металламакроциклического Bi(III)–Cu(II) комплекса на основе пиразингидроксамовой кислоты

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Полиядерные металламакроциклические комплексы, названные металлакраунами, представляют собой перспективный класс макрогетероциклических соединений, структурно напоминающих краун-эфиры. В данной статье приведены условия синтеза и характеристика первого металламакроциклического комплекса Bi(III)-Cu(II) 15-MC-5, полученного из пиразингидроксамовой кислоты. Молекулярная и кристаллическая структура комплекса были определены методом рентгеноструктурного исследования монокристаллов. Молекулярная структура представляет классическую металламакроциклическую конфигурацию 15-MC-5. Ион Bi(III) расположен в центре кольца 15-MC-5, состоящего из пяти повторяющихся [Cu(II)-N–O] звеньев. Подробные NBO- и QTAIM-исследования, основанные на DFT расчётах, были проведены для ионов {Bi[15-MC^Cu(II)Pyzha-5]}^{3+}.

Ключевые слова: Металлакраун, металламакроциклический комплекс, висмут, медь, пиразингидроксамовая кислота (Pyzha), PCA, DFT расчет.
Introduction

Investigations of polynuclear metallamacrocyclic complexes named as metallacrowns (MC) form a fascinating and rapidly growing branch of macroheterocyclic chemistry. First reported by Prof. Vincent L. Pecoraro in 1989, metallacrowns presented an exceptional class of supramolecular compounds with a metallamacrocyclic motif that structurally resembles crown ethers. Nowadays, the excellent and extensive reviews have described the structural and functional diversity of MCs inspiring interest to polynuclear metallamacrocyclic complexes due to their rich coordination chemistry and diverse properties combined with a range of applications. Among numerous structures including 8-MC-4, 9-MC-3, 10-MC-5, 12-MC-3, 12-MC-4, 12-MC-6, 14-MC-7, 5-MC-3, 15-MC-5, 15-MC-6, 16-MC-4, 16-MC-8, 18-MC-6, 18-MC-9, 20-MC-10, 22-MC-11, 24-MC-6, 24-MC-12, 28-MC-14, 30-MC-10, 32-MC-8, 32-MC-16, 36-MC-12, 36-MC-18, 40-MC-10, 48-MC-24, and 60-MC-20, heterometallic hydroximate complexes containing a 3d metal ion as a ring metal and a 4f ion as a core metal are the most abundant. Hydroxamic acids and their derivatives were extensively studied due to their synthetic and effective biological importance as constituents of antibacterial, antifungal, or anticancer agents, and as key functional groups of potential chemotherapeutics targeting cardiovascular diseases, HIV and Alzheimer’s disease or specific enzyme inhibitors. Their special (N,N)-(O,O) bridging bis-chelating ability towards transition metal ions such as Cu(II), Ni(II), Zn(II) and others results in the formation of the most common 15-MC-5 metallamacrocyclic compounds. The first recognized 15-MC-5 with M-N-O connectivity was a nonplanar system synthesized with salicylic acid.

The first planar 15-MC-5 uranyl complex was obtained by self-assembly of pyrazinohydroxamic acid and the Mn(II) ion within the cavity. To achieve planarity in the 15-MC-5 structures the transition metal ions are not ideally suited for the central cavity due to their smaller ionic radii. Therefore, this gives an opportunity for the lanthanide and actinide ions. The first planar 15-MC-5 uranyl complex was obtained by self-assembly of picolinohydroxamic acid with Cu(II) and UO2(NO3)2 ions. A number of MCs were prepared in a similar way in the following few years, leading to the isolation of 15-MC-5 of copper and aminohydroxamic acids encapsulating Ln3+, Y3+, and non-transition metals Ca2+, Ag+, Pb2+ and Hg2+. Following our interest in Ln(III)-Cu(II) 15-MC-5 metallacrowns with aminohydroxamic ligands, we report now on a metallamacrocyclic complex constructed from Cu2+ and Bi3+ metal centers and aminohydroxamic moieties. The preparation of heterometallic complexes incorporating bismuth and copper metals is quite challenging due to the differences in electronic and coordination requirements for the metal centers. It is also known that bismuth binds to nitrogen donor macrocycles even in strongly acidic solutions. In this communication, we describe a general synthetic approach and characterization of the first metallamacrocyclic Bi(III)-Cu(II) 15-MC-5 complex derived from pyrazinohydroxamic acid.

Experimental

General Procedures

All chemicals were reagent-grade and were used as received from Sigma Aldrich without any additional purification. The pyrazinohydroxamic acid was synthesized via methyl pyrazinamide according to the literature method. The C, H, N elemental analyses were performed by the Microanalytical laboratory of G.A. Razuvaeve Institute of Organometallic Chemistry RAS (IOMC) on Euro EA 3000 Elemental Analyser. IR spectra were obtained on a Perkin Elmer 577 spectrometer and recorded from 4000 to 450 cm−1 as a Nujol mull on KBr plates. A part of experiments was performed using the instruments of the Collective Use Analytic Centre of IOMC.

Synthesis

Bi(Cl)(H2O)(15-MCCu(II)Pyzha-S)(NO3)(CH3OH)∙H2O (I). Pyrazinohydroxamic acid (0.139 g, 1 mmol) was added to a stirred solution of Bi(NO3)3∙5H2O (0.097 g, 0.2 mmol) and Cu(OAc)2∙H2O (0.199 g, 1 mmol) in 30 ml of methanol. The dark green clear solution was obtained. After stirring overnight and filtering the solution was left to evaporate slowly. Dark green needle crystals were formed. Yield 110 mg, 37 %. Found: C 22.38, H 2.05, N 15.80 %. C20H42BiCuN10O13∙2H2O∙3CH3OH∙2(NO3)2∙Cl− requires C 22.36, H 2.08, N 15.83 %. IR (KBr) νmax cm−1: 471m, 490 w, 513 m, 565m, 678m, 753m, 761w, 828w, 866w, 952s, 1046m, 1070s, 1160s, 1195m, 1388s, 1410w, 1443s, 1565s, 1573s, 1587s, 1617m, 339br.

X-Ray Crystallographic Studies

The X-ray diffraction data for I were collected on a Bruker D8 Quest diffractometer with a CMOS area detector and a microfocus tube source (Mo-Kα radiation, λ=0.71073 Å, ω- and φ-scanning). The semi-empirical method SADABS(37) was applied for the absorption correction. The structure was solved by dual-space method using the SHELXT(38) software and refined by the full-matrix least-squares technique against F2 with the anisotropic displacement parameters for all non-hydrogen atoms. The hydrogen atoms in the MeOH molecules were placed geometrically and included in the structure factors calculation in the riding motion approximation. The H atoms in water molecules were found from Fourier synthoses of electron density and were refined isotropically with Uiso(H) = 1.2Ueq. Two methanol molecules and one NO3− moiety are disordered over two positions. All the data reduction and further calculations were performed using the SAIINT(39) and SHELXL-97(40) packages. The figures of structure I were drawn using the OLEX2 software. The details of crystallographic, collection and refinement data are shown in Table 1.

CCDC reference number is 1937326. The data can be obtained free of charge from the Cambridge Crystallographic Data Centre at www.ccdc.cam.ac.uk/data_request/cif.

Computational Details

The quantum chemical calculations were performed with the Gaussian09 software. The hybrid meta-GGA exchange-correlation functional M06(43) and the triple-c–plus-polarization Def2TZVP basis set(44) (with relativistic pseudopotential for Bi atom(45)) were employed for the DFT calculations. Full geometry optimization of the [Bi15-MC-Cu(II)Py3ha-5]- cationic complex (bearing 3285 primitive gaussians) was carried out with use of the ultrafine integration grid without molecular symmetry (Int=UltraFine and NoSymm keywords). The complex was treated as a high-spin sextet system. The wave function of the complex was proven to be stable by the tests for the SCF solutions stability.
The polynuclear metallamacrocyclic Bi(III)-Cu(II) pyrazinohydroximate complex Bi(Cl)(H$_2$O)[15-MC$_{Cu(II)Pyzha-5}$(NO$_3$)$_2$(CH$_3$OH)$_2$(H$_2$O)] (I) was prepared in methanol in the presence of HCl under ambient conditions by one-step reaction of pyrazinohydroxamic acid with copper acetate and bismuth(III) nitrate (Scheme 1).

The characterization was performed using elemental analysis and IR spectroscopy. The elemental analysis data are in agreement with the proposed structure. The IR spectrum of the complex shows a broad band in the region of 3534–3166 cm$^{-1}$, which is due to the water molecules in the lattice. The band at 1617 cm$^{-1}$ caused by water molecules overlaps the absorption band of the C-O group of the hydroximate moiety. In the IR spectra of complex I the vibrations assigned to the characteristic bands $\nu$(C=N) of pyrazine and $\nu$(N-O) of the Pyzha ligand participating in the formation of the metallacrown ring were located at 1573 cm$^{-1}$ and 1443 cm$^{-1}$, respectively. The bands at 513 and 565 cm$^{-1}$ can be associated with the $\nu$(Cu–O) mode.

The crystal and molecular structures of complex I were determined by single-crystal X-ray diffraction (SC-XRD) studies (Figure 1, Table 2). According to the XRD investigations, the molecule of complex I consists of the bismuthic 15-MC$_{Cu(II)Pyzha-5}$ moiety, three coordinated anions (one chlorine and two nitrate groups) as well as two methanol and two water molecules. Bismuth atom Bi(1) is disposed at the center of the metallacrown and bound to five oxygen atoms O(1,3,5,7,9). The Bi(1) cation additionally binds the chlorine anion Cl(1) and a water molecule which are located at the apical positions of the bismuth coordination environment. Thus, the coordination number of Bi(1) is 7 and the coordination polyhedron around bismuth has the geometry of a distorted pentagonal bipyramid. The angle between the apical atoms O(water)-Bi(1)-Cl(1) is 170.1(2)$^\circ$. The sum of angles O-Bi(1)-O in the basal plane equals to 360.1$^\circ$. Displacement of the Bi(1) atom from the least-square plane of five hydroxamate oxygen atoms is 0.02 Å.

The nitrate anions are coordinated at the Cu(1) and Cu(2) atoms in a monodentate fashion. All copper atoms are square-pyramidal wherein the coordinated two nitrate anions, two methanol and one water molecules occupy the apical positions. Only one molecule of methanol at

Results and Discussion

The polynuclear metallamacrocyclic Bi(III)-Cu(II) pyrazinohydroximate complex Bi(Cl)(H$_2$O)[15-MC$_{Cu(II)Pyzha-5}$(NO$_3$)$_2$(CH$_3$OH)$_2$(H$_2$O)] (I) was prepared in methanol in the presence of HCl under ambient conditions by one-step reaction of pyrazinohydroxamic acid with copper acetate and bismuth(III) nitrate (Scheme 1).

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The copper metallacrown is almost planar and has a barely zigzag shape. The mean deviation of non-hydrogen atoms of the copper-crown moiety from the plane is 0.12 Å.

The nitrate anions are coordinated at the Cu(1) and Cu(2) atoms in a monodentate fashion. All copper atoms are square-pyramidal wherein the coordinated two nitrate anions, two methanol and one water molecules occupy the apical positions. Only one molecule of methanol at
the Cu(5) atom is located on the same side of the metallacrown as the Bi(1)-coordinated chlorine anion. There are numerous intermolecular interactions with the participation of a solvate MeOH molecule in the crystal of I.

Table 2. Selected bond lengths ($d$) for complex 1.

| Bond                  | $d$, Å |
|-----------------------|--------|
| Bi(1)-oxime           | 2.429(8)-2.451(7) |
| Bi(1)-O(water)        | 2.555(10)     |
| Bi(1)-Cl(1)           | 2.550(4)      |
| Cu-O (15-MC-5)        | 1.912(8)-1.951(7) |
| Cu-N (15-MC-5)        | 1.906(8)-2.024(10) |
| Cu-O(NO$_3$)          | 2.317(17), 2.404(8) |
| Cu-O(MeOH)            | 2.354(8), 2.16(2)     |
| Cu-O(water)           | 2.366(11)     |

Detailed analysis of crystal packing of I revealed that the ions form two-dimensional chains as zigzag ladders (molecules ABC and DEF) along the axis c (Figure 2). In these chains pairs of the hydrogen bonds N2...H19B and N5...H20B are realized with the distances 2.05 and 2.09, respectively. The chains of molecules ABC and DEF are linked via parallel stacking with some displacements of pirazine heterocycles (atoms N14, N11 and N8). The distances between centres of the pirazines with the atoms N14...N11 and N11...N8 are 3.65(1) and 3.73(1) Å, correspondingly. These distances may indicate the presence of $p$-$π$ interactions between the pyrazine rings.

To shed light on electronic structure of I we undertook quantum chemical calculations at the M06/Def2-TZVP level of DFT. Optimized geometry of the cationic form [Bi[15-MC$_{Cu1}[15-MC_{Cu1(Im}(Pyz)]^{5-}$] agrees well with the experimental X-ray structure of I and reveals the same trends in bond lengths (Table 3). Interatomic distances C-N(imine), C-N(imine-Ar), and C-O in the pyrazinohydroximate ligands indicate unsaturated character which is confirmed by the Wiberg Bond Indices (WBI) and ellipticity values at the Bond Critical Points (BPC) (Table 3). Thus, among O(carbonyl), C, and N(imine) atoms of the hydroximate groups a $π$-delocalization takes place, the $π$-contribution for the C-N(imine) bonds being higher (WBI 1.415, $ε$ = 0.267) than that for C-O (WBI 1.348–1.349, $ε$ = 0.079). For the N-O...
Table 3. Interatomic distances (d), Wiberg Bond Indices (WBI), and ellipticities (ε) at the Bond Critical Points (BCP) of [Bi15-MC$_{Cu(II)}$Pyza-5]$_{3+}$ cationic complex calculated at the M06/Def2-TZVP level.

| Interaction          | d, Å (DFT) | WBI   | ε     |
|----------------------|------------|-------|-------|
| C-C                  | 1.478      | 0.993 | 0.127 |
| C-N(imine)           | 1.312      | 1.415 | 0.267 |
| C-N(imine-Ar)        | 1.345      | 1.309 | 0.154 |
| C-O                  | 1.263      | 1.348–1.349 | 0.079 |
| N-O                  | 1.359–1.360| 1.037 | 0.057 |
| Bi-O(oxime)          | 2.402–2.403| 0.256–0.257|       |
| Cu-O(oxime)          | 1.969–1.970| 0.277–0.278|       |
| Cu-O(carbonyl)       | 1.922–1.923| 0.335 |       |
| Cu-N(imine)          | 1.903–1.904| 0.388 |       |
| Cu-N(imine-Ar)       | 2.008      | 0.304 |       |

The Bi-O(oxime) contacts are described by the lower WBI slightly exceed 1 and the ellipticity is 0.057. Taking into account the N-O interatomic distances close to those typical for a single bond, the obtained data are indicative of a weak $p,\pi$-conjugation between the oxime oxygen lone pair and the delocalized $\pi$-system. The bonding situation in the hydroximate groups is similar to that in the 15-MC-5 complexes of Ca$^{2+}$ and Y$^{3+}$ bearing simple glycinehydroximate ligands.[32,33] Notably, the $\pi$-contribution to the C-N(imine) bonds is higher than that to C-N(imine-Ar) in the aromatic rings (WBI 1.309, $\varepsilon = 0.154$). For the corresponding interactions with copper QTAIM predicts the following values of the electron density $\rho(r)$ and Laplacian $\nabla^2 \rho(r)$ at the BCP: $\rho(r) = 0.110$ a.u., $\nabla^2 \rho(r) = 0.433$ a.u. for Cu-N(imine) and $\rho(r) = 0.088$ a.u., $\nabla^2 \rho(r) = 0.321$ a.u. for Cu-N(imine-Ar). This allows us to consider Cu-N(imine) interactions to be stronger than Cu-N(imine-Ar). In addition, QTAIM reveals ellipticity values of 0.127 for the C-C bonds located between the two donor N atoms which testifies for a $\pi,\pi$-conjugation between the hydroximate and aromatic systems. For comparison: $\varepsilon$ values of the C-C bonds in the aromatic rings are 0.180–0.221.

The Bi-O(oxime) contacts are described by the lower WBI than Cu-O(oxime) (0.256–0.257 and 0.277–0.278, respectively) that indicates the more polar character of the bismuth interactions. Indeed, WBI values correlate well with the QTAIM atomic charges: +1.896 e for Bi, +1.122 e for Cu, and -0.774 e for O(oxime). Moreover, the BCP parameters for the Bi-O(oxime) bonds ($\rho(r) = 0.052$ a.u., $\nabla^2 \rho(r) = 0.173–0.174$ a.u.) are also lower than for Cu-O(oxime) ($\rho(r) = 0.083–0.084$ a.u., $\nabla^2 \rho(r) = 0.396$ a.u. Total energies $\sum E^{(2)}$ of donor-acceptor interactions between the oxygen lone pairs and the metal vacant orbitals are 35.3–35.5 kcal/mol for the Bi-O(oxime) contacts. In summary, interactions of the Bi$^{3+}$ cation with five oxime oxygen atoms give 177.0 kcal/mol that provides stability of the 15-MC-5 architecture. The natural orbitals providing the most significant contributions to the O(oxime) $\rightarrow$ Bi donation are described as hybrids: $\alpha$($\rho(72.6–72.7 \%) \rho^{p,\pi}(27.3–27.4 \%)$ with occupancies of 0.9506–0.9507 and $\beta$($\rho(67.3 \%) \rho^{p,\pi}(32.7 \%)$ with occupancies of 0.9488–0.9489. Corresponding acceptor orbitals of Bi possess pure $p$-character: $\alpha$-$6p^*(99.8 \%)$ with occupancies of 0.1141 and $\beta$-$6p^*(99.8 \%)$ with occupancies of 0.1134. The $E^{(2)}$ values corresponding to these interactions lie in the ranges of 8.1–12.9 kcal/mol for $\alpha$-orbitals and 8.5–14.2 kcal/mol for $\beta$-orbitals (Table 4). In Figure 3 the donor and acceptor $\beta$-orbitals are presented. The natural orbitals agree well with maps of the Deformation Electron Density (DED) and Electron Localization Function (ELF) which reveal domains of the oxygen lone pairs directed toward the Bi$^{3+}$ cation (Figure 4). The Cu-O(oxime) $\sum E^{(2)}$ values are drastically higher (99.8–100.0 kcal/mol).

Table 4. Electronic configurations of the natural orbitals providing the most significant contributions to the Bi-O(oxime) interactions.

| Interaction | $E^{(2)}$, kcal/mol | Donor orbitals | Acceptor orbitals |
|------------|---------------------|---------------|-------------------|
| $\alpha$-orbitals: | | | |
| O(1)$\rightarrow$Bi | 8.8 | $\sigma(72.6 \%) \rho^{p,\pi}(27.4 \%)$ | $6p^*(99.8 \%)$ |
| O(3)$\rightarrow$Bi | 11.9 | $\sigma(72.6 \%) \rho^{p,\pi}(27.4 \%)$ | $6p^*(99.8 \%)$ |
| O(5)$\rightarrow$Bi | 12.9 | $\sigma(72.7 \%) \rho^{p,\pi}(27.3 \%)$ | $6p^*(99.8 \%)$ |
| O(7)$\rightarrow$Bi | 11.5 | $\sigma(72.6 \%) \rho^{p,\pi}(27.4 \%)$ | $6p^*(99.8 \%)$ |
| O(9)$\rightarrow$Bi | 8.1 | $\sigma(72.7 \%) \rho^{p,\pi}(27.3 \%)$ | $6p^*(99.8 \%)$ |

| $\beta$-orbitals: | | | |
| O(1)$\rightarrow$Bi | 14.2 | $\sigma(67.3 \%) \rho^{p,\pi}(32.7 \%)$ | $6p^*(99.8 \%)$ |
| O(3)$\rightarrow$Bi | 12.3 | $\sigma(67.3 \%) \rho^{p,\pi}(32.7 \%)$ | $6p^*(99.8 \%)$ |
| O(5)$\rightarrow$Bi | 8.5 | $\sigma(67.3 \%) \rho^{p,\pi}(32.7 \%)$ | $6p^*(99.8 \%)$ |
| O(7)$\rightarrow$Bi | 10.2 | $\sigma(67.3 \%) \rho^{p,\pi}(32.7 \%)$ | $6p^*(99.8 \%)$ |
| O(9)$\rightarrow$Bi | 13.4 | $\sigma(67.3 \%) \rho^{p,\pi}(32.7 \%)$ | $6p^*(99.8 \%)$ |
mol) than those for the Bi-O(oxime) interactions which is associated with an increased covalent contribution in the copper contacts. The Cu-O(carbonyl) interactions are characterized by slightly higher $\sum E(2)$ values of the 106.3–106.6 kcal/mol.

Conclusions

The first polynuclear metallamacrocyclic Bi(III)-Cu(II) 15-MC-5 complex bearing the pyrazinohydroximate ligands has been synthesized. The single-crystal structure determined by SC XRD studies revealed the classic metallamacro cyclic 15-MC-5 configuration. The NBO and QTAIM calculations provided quantitative description of $\pi$-interactions in the pyrazinohydroximate ligands and parameters of the donor-acceptor contacts involving the Cu$^{2+}$ and Bi$^{3+}$ cations. The nature of bonding in the metallacrow was studied in details.

As discussed, the Bi(III)-Cu(II) and Ln(III)-Cu(II) 15-MC-5 compounds appear to be structurally very similar. This structural resemblance of metallamacro cyclic complexes bearing a $p$-block metal ion and 4f metal ions represents another interesting outcome of this investigation.

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