Article

Reactions of a Dioxidomolybdenum(VI) Complex with Thionation Reagents—Formation of Mo(IV) Species with Sulfur Donors

Esko Salojärvi, Anssi Peuronen, Narhari Sapkota and Ari Lehtonen *

Intelligent Materials Chemistry Research Group, Department of Chemistry, University of Turku, FI-20014 Turku, Finland
* Correspondence: ari.lehtonen@utu.fi

Abstract: Molecular molybdenum complexes with sulfur donor ligands are generally studied as soluble model compounds for molybdenum enzymes essential for life. The dioxidomolybdenum(VI) complex with tetradentate aminobisphenolate ligand undergoes a reaction with thionation reagent \( \text{P}_2\text{S}_5 \) or its organic derivative, Lawesson’s reagent, to yield stable Mo(IV) aminobisphenolate complexes, where pristine oxido ligands have been replaced by bidentate sulfur donors tetrasulfide, \( \text{S}_4^{2-} \) or (4-methoxyphenyl)phosphonotrithioate residue derived from Lawesson’s reagent. This is in contrast to the behaviour of analogous dioxidotungsten(VI) complex, which, under similar conditions, yields W(VI) \( \text{S}_2 \) systems. The overall \( \text{cis},\text{trans},\text{cis} \) geometry of the parent dioxidomolybdenum(VI) aminobisphenolate is retained, namely, the neutral nitrogen donors are in \( \text{cis} \) positions, phenolate oxygens are \( \text{trans} \) to each other and sulfur donors are \( \text{cis} \). Although formally Mo(IV), thus \( d^2 \) system, the studied complexes have diamagnetic singlet electron configurations as a result of the axially compressed octahedral structures.

Keywords: molybdenum; sulfur donor; Lawesson’s reagent; transition metal complexes

1. Introduction

Molybdenum sulfides are useful catalysts in a number of important reactions used to activate small molecules, e.g., hydrogen evolution reaction, \( \text{CO}_2 \) reduction and \( \text{N}_2 \) fixation [1]. Moreover, molybdenum–sulfur bonds are found in all molybdenum enzymes; therefore, molecular molybdenium sulfide complexes are investigated as soluble model compounds for the active sites of such catalysts as well as for metal enzymes [2–5]. In principle, diverse metal–organic sulfide complexes can be synthesized by the reaction of simple metal sulfides and ligand precursors, but in practice, they are mostly made of easily available oxido complexes by simple thionation reactions, i.e., oxido-to-sulfido substitutions. Typically, the thionations are run using \( \text{H}_2\text{S} \) [6–10], \( \text{B}_2\text{S}_3 \) [11–14], (\( \text{R}_3\text{Si} \))\( _2\text{S}/\text{R}_3\text{SiSH} \) [15–18], \( \text{P}_2\text{S}_5 \) [12], or Lawesson’s reagent (2,4-bis(4-methoxyphenyl)-1,3,2,4-dithiadiphosphetane-2,4-disulfide) [9,19]. We have earlier used the reaction of dioxidotungsten(VI) complexes with \( \text{P}_2\text{S}_5 \) and its soluble derivative, Lawesson’s reagent, to prepare stable disulfidotungsten(VI) complexes supported with tetradeutate aminobisphenolates [20]. As equivalent Mo and W dioxido compounds have, in many cases, very similar molecular and crystals structures and rather similar chemical properties, we might expect parallel reactivity in the thionation reactions as well. However, under identical reaction conditions, the dioxidomolybdenum(VI) aminobisphenolates did not yield the anticipated disulfido derivatives; instead, the reactions led to the reduction of the metals’ centres as the isolated products were identified as Mo(IV) complexes. In this article, we report the reactivity of dioxidomolybdenum(VI) aminobisphenolate with thionation reagents and structural characterization of two Mo(IV) species with sulfur donor chelate ligands.
2. Results and Discussion

The oxido-to-sulfido substitution of dioxidomolybdenum(VI) aminobisphenolates was studied, allowing complex 1 to react with either P2S5 or Lawesson’s reagent in 1,2-dichloroethane at reflux temperature following the substitution procedure used previously to synthesize corresponding disulfidotungsten(VI) compounds. As a result, the reactions yielded dark mixtures of several strongly coloured, poorly stable compounds (seen by thin-layer chromatography), which could not be isolated or further characterized. However, 1 reacted with P2S5 at room temperature in a toluene solution to form a dark solution, which allowed the isolation of green 2 and brownish-red 3 along with several unstable compounds. The isolated yields of major products in the reaction with Lawesson’s reagent depended on the reaction time—after a 20 h reaction, 3 was isolated in ca. 40% yield together with 5% of 2, whereas 2 was found in ca. 35% yield with only a minor amount of 3 after one-week reaction.

![Scheme 1](image)

Scheme 1. The reactions of a dioxidomolybdenum(VI) complex 1 with thionation reagents.

The molecular structures of 2 and 3 were studied by NMR spectroscopy and X-ray diffraction studies. The NMR spectra of 2 show the anticipated chemical shifts for the tetradentate aminobisphenolate ligand. For example, in the 1H NMR spectrum, the benzylic methylene protons are visible as two two-proton doublets at 5.66 and 4.30 ppm, respectively, accompanied by a two-proton singlet at 4.56 ppm for the methylene protons in the picoline arm. The XRD data shows that the product is virtually a C8 symmetric neutral molecule, where two oxides have been substituted by one S42− dianion to form a new Mo (IV) complex (Figure 1). The overall cis,trans,cis geometry of the parent complex 1 has been retained, i.e., the neutral nitrogen donors are in cis positions, phenolate oxygens are trans to each other and sulfur donors are cis (see Figure 1). Compared to 1, the metal–donor distances to the aminobisphenolate ligand are somewhat shorter in 2, whereas the O1-Mo-O2 and N8-Mo-N37 bite angles are slightly larger (see Table 1). The Mo-S distances and S-Mo-S angle are closely similar to the corresponding bonding parameters found previously for the dianionic tetrasulfido ligand in known Mo(IV) complexes [3,4,9,21,22]. In the same way, the 1H and 13C NMR spectra of 3 show chemical shifts for the coordinated aminobisphenolate ligand, while the number of the chemical shifts indicates a C1 symmetric structure. The spectra also show the presence of 4-methoxyphenyl group originated from Lawesson’s reagent. Correspondingly, the XRD studies show that the complex 3 is a neutral species, wherein one tetradentate aminobisphenolate and one bidentate, dianionic (4-methoxyphenyl) phosphonotrithioate surround the Mo(IV) ion (see Figure 2). The overall geometry around the metal centre is cis,trans,cis and the phenolate groups are in trans positions, i.e., the aminobisphenolate ligand is symmetrically coordinated. Therefore, the asymmetry seen in the NMR spectra rises from the phosphonotrithioate ligand. The metal–donor distances to the aminobisphenolate ligand are considerably shorter than in 2 (Table 1). The bite angles O1-Mo-O2 and N8-Mo-N37 are slightly larger and the S-Mo-S angle is clearly smaller than in 2, obviously due to the rigidity of the phosphonotrithioate ligand. Complexes 2 and 3 are
formally Mo(IV) complexes with d² electron configurations, and are therefore expected to show paramagnetic properties caused by the two odd electrons. However, both compounds are diamagnetic, as evidenced by the well-resolved NMR spectra. Principally, the d² metal centres may have either paramagnetic triplet or diamagnetic singlet electron configurations, depending on the coordination geometry. According to the visual inspection of the single crystal structures and further calculation of the continuous shape measures (CSHM’s, see Table S2 Supplementary Materials) [23], 2 and 3 are octahedral complexes with axially compressed structures with four long bonds in xy-planes (Mo-N8, Mo-N37, Mo-S1 and Mo-S2/S4) and two shorter ones (Mo-O1 and Mo-O2) along z axes. This compression causes the splitting of the dₓᵧ, dₓz, and dₓᵧ₂ orbital set, the first lower in energy compared to the other ones, which results singlet ground state system with both electrons on the dₓᵧ orbital. The diamagnetic ground states of 2 and 3 were also confirmed by DFT calculations using the respective crystal structures of the complexes as the basis of coordinates for computational analyses. The singlet-triplet gap of 2 is 103 kJ/mol, whereas in 3 the triplet lies 35 kJ/mol above the singlet ground state, which affirms the singlet ground states of the complexes evident from the experimental data. For 3, the lowest triplet state at the PBE0/def2-TZVP level shows Mo(IV) with d² electron configuration, whereas for the lowest triplet state of 2, the Mulliken spin densities (see ESI) in fact suggest a d¹ Mo centre with the other unpaired electron localized at the S₄-fragment while the d² triplet state could not be established even with an attempt of using fragment-based initial assignment of partial charges and spins separately for ligands and metal.

Figure 1. Molecular structure of 2, with H-atoms and disorder of the S-atoms omitted. Displacement ellipsoids are drawn at the 50% probability level.

Figure 2. Molecular structure of 3, with H-atoms and solvent molecules omitted. Displacement ellipsoids are drawn at the 50% probability level.
Table 1. Selected bond lengths (Å) and angles (°) for studied complexes.

|           | 1 [24]     | 2        | 3        |
|-----------|------------|----------|----------|
| Mo-O1     | 1.951 (3)  | 1.9370 (15) | 1.901 (6) |
| Mo-O2     | 1.9152 (15) | 2.237 (6)  |
| Mo-N8     | 2.372 (3)  | 2.3388 (19) | 2.290 (7) |
| Mo-N38    | 2.345 (4)  | 2.309 (2)  | 2.237 (6) |
| Mo-S1     | 2.378 (2)  | 2.378 (2)  | 2.378 (2) |
| Mo-S2/S4  | 2.2223 (6) | 2.373 (2)  |
| O1-Mo-O2  | 156.5 (1)  | 158.01 (7) | 161.5 (3) |
| N8-Mo-N38 | 68.9 (1)   | 73.89 (7)  | 75.9 (3)  |
| S1-Mo-S2/S4| 95.62 (2)  | 80.71 (7)  |
| N8-Mo-S1  | 167.29 (5) | 175.0 (3)  |
| N37-Mo-S2/S4| 94.05 (5)  | 101.76 (19) |

Based on our investigation, it is evident that the reaction of Mo(VI) species 1 with thionation reagents leads to the formation new Mo(IV) compounds. Similar reactivity leading to the formation of Mo(IV) tetrasulfide complexes (Me₃tacn)MoO(S₄) and (Bu₃tach)MoO(S₄) is seen in the reactions of Mo(VI) compounds (Me₃tacn)MoO₃ with S₈ and (Bu₃tach)MoO₃ with B₂S₃ (Me₃tacn = 1,4,7-trimethyl-1,4,7-triazacyclonane, Bu₃tach = 1,3,5-tri-tert-butyl-1,3,5-triazacyclohexane) [3,9]. Although the formation mechanism of the reduced products 2 and 3 is not clear, we may suppose that the initial step is the oxido-to-sulfido substitution to form MoS₂(L), similarly to the reactions of analogous W(VI) complexes. The unstable disulfido reduces then to eliminate elemental sulfur yielding a reactive Mo(IV) intermediate MoS(L). This intermediate reacts further with Lawesson’s reagent or other sulfur-containing species in the reaction mixture to form distinct isolatable products 2 and 3. Both 2 and 3 are stable in the solid state as well as in inert solvents under open atmosphere. Although 3 decomposes in a prolonged reaction while 2 is formed, isolated and purified 3 is stable and can be stored in a toluene solution for a week without any noticeable formation of 2. In general, the formation of these Mo(IV) complexes demonstrates the relative inclination of Mo(VI) toward reduction which is in stark contrast to the behaviour of W(VI), although these to M(VI) species often reflect each other’s reactivity.

In conclusion, the oxido-to-sulfido substitution of a dioxidomolybdenum(VI) amino-bisphenolate 1 was studied using different thionation reagents. The reaction with P₂S₅ leads to the formation of Mo(IV) aminobisphenolate complex 2 with a bidentate sulfur chelate, i.e., tetrasulfide, S₄²⁻. The parallel reaction with Lawesson’s reagent, the organic derivative of P₂S₅, yields also complex 2 together with complex 3 as the initial major product, of which the latter has a (4-methoxyphenyl)phosphonotrithioate residue derived from Lawesson’s reagent coordinated to the Mo(IV) centre. Both compounds 2 and 3 have an overall geometry of cis,trans,cis, that is, the neutral nitrogen donors are in cis positions, phenolate oxygen atoms are trans to each other and sulfur donors are cis in the axially compressed octahedral structures. Both 2 and 3 are stable in the solid state as well as in inert solvents under ambient atmosphere. The studied complexes are formally Mo(IV) with d² metal centres. However, they are diamagnetic due to the singlet electron configurations as a result of the axially compressed octahedral structures. Similar thionation reactions of comparable dioxidotungsten(VI) aminobisphenolates are known to yield disulfidotungsten(VI) complexes, so these results demonstrate the higher inclination.

3. Materials and Methods

Complex 1 was prepared as reported previously [25]. Toluene was dried over 3A molecular sieves. Other chemicals were from commercial sources and were used as purchased. The synthetic reactions were run under a nitrogen atmosphere, whereas all isolations and analyses were conducted under open atmosphere. The IR spectra were measured using a Bruker VERTEX 70 FTIR instrument in transmittance mode, and peaks are reported in wavenumbers (cm⁻¹) and intensities (b = broad, w = weak, m = medium, s = strong,
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vs = very strong). All NMR spectra were recorded on a Bruker Avance III 500 MHz instrument (1H: 500.08 MHz, 13C: 125.75 MHz) equipped with a broad-band smart probe and were referenced to residual CHCL3 solvent signals (1H: δ 7.26, 13C: δ 77.16). The NMR samples were kept under vacuum prior to the measurements to remove possible solvate molecules. The samples were dissolved in acetonitrile for mass spectrometric analysis. The mass spectra were obtained by quadrupole–Orbitrap mass spectrometer (QExactiveTM, Thermo Fisher Scientific GmbH, Bremen, Germany) using direct infusion, negative electrospray ionization and full scan at m/z 150–2000 with the resolution of 140,000. The calibration was performed by Pierce ESI Negative Ion Calibration Solution (Thermo Fisher Scientific Inc., Waltham, MA, USA). The data was processed with Thermo Xcalibur Qual Browser software (Version 3.0.63, Thermo Fisher Scientific Inc., Waltham, MA, USA). Single crystal X-ray diffraction data were collected with Rigaku Oxford Diffraction custom system consisting of microfocus MicroMax™-007 HF rotating anode generator producing monochromatized Cu Kα1 radiation and HyPix-6000HE detector. Data collection and reduction were done using the CrysAlisPro software [26], whereas crystal structures were solved and refined using SHELXS and SHELXL programs within the Olex2 interface. DFT calculations were conducted using Gaussian 16 software [27]. The geometries were taken from single crystal X-ray structures and the C–H bond lengths were normalized. The relative energies of singlet and triplet spin states were determined as single point calculations at the PBE0/def2-TZVP (effective core potential for Mo) level using these geometries [28,29].

2: Method a: First, 0.35 g (0.50 mmol) of 1 and 0.15 g (0.68 mmol) of P2S5 were mixed in 20 mL of dry toluene. The dark mixture was stirred for 24 h, the solvent was evaporated and the green product (60 mg, 15%) was isolated by a silica column chromatography using CH2Cl2 as an eluent. Method b: First, 0.50 g (0.75 mmol) of 1 and 0.61 g (1.50 mmol) of Lawesson’s reagent were mixed in 20 mL of dry toluene and the solution was kept for week at room temperature. Afterwards, 132 mg (35%) of 2 as well as 33 mg (5%) of 3 were isolated by column chromatography using CH2Cl2 as an eluent. The crystals of 2 for XRD analyses were grown from acetonitrile at room temperature. IR: 3440 s, 2958 s, 2868 s, 1725 s, 1589 s, 1470 s, 1443 s, 1414 m, 1392 m, 1362 m, 1304 m, 1259 s, 1240 m, 1203 m, 1171 m, 1128 m, 1057 w, 1022 w, 976 w, 916 m, 850 s, 808 m, 760 s, 727 w, 648 w, 590 w, 557 s, 509 w, 474 m cm⁻¹. UV-Vis: λ = 425 nm, ε = 7062 cm⁻¹ M⁻¹, 1H NMR (CDCl₃): 10.11 (1H, d, J = 5.3 Hz, ArH), 7.69 (1H, t, J = 8.0 Hz, ArH), 7.38 (1H, t, J = 6.2 Hz, ArH), 7.09 (1H, d, J = 7.8 Hz, ArH), 7.02 (2H, d, J = 2.0 Hz, ArH), 6.87 (2H, d, J = 2.1 Hz, ArH), 5.66 (2H, d, J = 13.4 Hz, CH₂), 4.56 (2H, CH₃), 4.30 (2H, d, J = 13.4 Hz, CH₂), 1.22 (18H, tBu), 0.56 (18H, tBu). 13C NMR (CDCl₃): 158.28, 157.92, 154.54, 141.15, 138.43, 135.02, 130.51, 122.44, 121.19, 120.57, 66.41, 61.59, 33.23, 33.15, 30.54, 28.68, 28.13. ESI-(−)MS: (MeCN): m/z = 767.17367 [M-H]⁻ (calcld. m/z = 767.17310).

3: First, 3.9 g (5.8 mmol) of 1 and 2.4 g (5.9 mmol) of Lawesson’s reagent were mixed in 35 mL of dry toluene and the reaction mixture was stirred under a nitrogen atmosphere for 20 h at room temperature. Next, 3.55 g of brown precipitate was isolated and recrystallized from hot acetonitrile to obtain 2.08 g (39%) of brownish-red crystals. The filtrate was evaporated and 0.22 g (5%) of complex 2 was isolated by column chromatography. IR: 3440 s, 2958 s, 2868 m, 1725 s, 1589 s, 1470 s, 1443 s, 1414 m, 1392 m, 1362 m, 1302 w, 1288 w, 1250 s, 1230 m, 1171 m, 1126 m, 1095 m, 1055 w, 1022 m, 916 m, 868 s, 854 s, 808 w, 798 w, 762 s, 667 m, 619 m, 594 w, 575 m, 559 w, 532 w, 507 w cm⁻¹. UV-Vis: λ = 408 nm, ε = 4408 cm⁻¹ M⁻¹, 1H NMR (CDCl₃): 8.90 (1H, d, J = 5.1 Hz, ArH), 8.43 (2H, dd, J = 12.1 Hz, J’ = 11.7 Hz, ArH), 7.62 (1H, t, J = 7.8 Hz, ArH), 7.29 (1H, t, J = 7.8 Hz, ArH), 7.09 (1H, d, J = 6.2 Hz, ArH), 7.08 (1H, d, J = 6.3 Hz, ArH), 7.04 (2H, t, J = 2.5 Hz, ArH), 6.95 (1H, d, J = 7.9 Hz, ArH), 6.91 (2H, d, J = 6.7 Hz, ArH), 4.83 (1H, d, J = 13.1 Hz, CH), 4.66 (1H, d, J = 12.7 Hz, CH), 4.54 (1H, d, J = 17.0 Hz, CH), 4.36 (1H, d, J = 17.2 Hz, CH), 4.22 (1H, d, J = 13.2 Hz, CH), 4.12 (1H, d, J = 12.7 Hz, CH), 3.81 (3H, OCH₃), 1.24 (18H, tBu), 1.05 (9H, tBuH), 0.86 (9H, tBuH) ppm. 13C NMR (CDCl₃): 161.30, 161.27, 160.61, 157.19, 156.97, 154.47, 144.46, 144.30, 139.91, 136.75, 135.95, 132.33 132.20, 125.10, 124.75, 124.65, 124.53, 123.84, 123.40, 123.17, 120.55, 113.46, 113.34, 70.07, 66.65, 60.90, 55.32, 34.92,
34.68, 34.60, 34.53, 31.45, 31.44, 29.90, 29.83 ppm. ESI(−)-MS: (MeCN): \( m/z = 873.22504 \) [M−H]− (calcd. \( m/z = 873.22448 \)).

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27217154/s1, Electronic Supplementary Information (ESI) available: Details of computational methods and crystallographic data.

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