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Unexpected cis-dioxido uranyl carboxylate compound: Synthesis, characterization and photocatalytic activity of uranyl-succinate complexes



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ABSTRACT

Two new complexes formed by uranyl ions with succinic acid and 4,4'-dipyridine have been investigated. Interestingly, the uranium(VI) coordination polymers possess two types dioxido unit, a cis-dioxido bond angle with 115.421(2)° and a trans-dioxido bond angle with 178.421(2)°, respectively. The complexes were characterized by elemental analysis, IR and UV-vis spectroscopy and X-ray crystal diffraction. In addition, the properties of luminescence and photocatalytic analysis were also investigated. The result revealed that the two complexes show different photocatalytic activity in the decomposition of RhB under visible light and UV light irradiation.

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Uranium is a potentially carcinogenic metal of growing importance, especially with the increased use of depleted uranium in munitions and armor plating by the military [1]. In the meantime, it may also lead to internal contamination of people worldwide by various compounds of uranium through alpha particle decay [2]. This constitutes an occupational risk and has led to questions about its potential health effects since contamination with uranium compounds may induce radiological and chemical toxicity [3].

Nowadays, the class of uranyl polycarboxylate has been developed as an important member of the uranium organic coordination polymer family due to their fantastic structural diversities and excellent physicochemical properties in potential applications of ion exchange [4,5], proton conductivity [6], chiral materials [7], and materials [8,9].

This field has aroused great interest with increasingly attractive structural architectures resulting from various coordination modes of the uranium atom and the modification of the organic residues of polycarboxylate [10–12]. For these uranyl polycarboxylates, the template species (H₃O⁺, NH₄⁺, metalions, organic linkers etc.), dimensions, configurations, and the coordination abilities play important roles in structure directing for constructing new structures.

However, the aim of obtaining desirable architectures and functional properties of coordination polymers constructed by organic ligands and uranyl ions is still a long-term challenge to chemists due to the difficult prediction of either the compositions or the structures and properties of the reaction products. Therefore, it is necessary to (i) explore the fundamental chemistry of 5f-elements [13], reactivity and coordination behavior; (ii) understand the bonding interactions between the metal center and ligand (ionic or covalent bonding); (iii) explore possible applications of new complexes, such as in catalysis or as sensors, molecular adsorption, separation, drug delivery. From the chemical point of view, a large number of studies have reported the reactivity of organic polycarboxylates with uranium [14–16], since its crystal chemistry reveals a wide variety of coordination states, which give rise to a large diversity of topologies. In this course, uranium was found to be a good candidate element for the generation of mixed organic-inorganic extended networks [17]. When it is associated with multidentate O-donor ligands, such as carboxylate-based solids, uranium (especially with oxidation state VI) exhibits an interesting chemical ability to successfully form a wide variety of frameworks showing different dimensionalities. In case of uranyl cation UO_2^{2+} , the possible coordination environments (4 + 2, square bipyramid; 5 + 2, pentagonal bipyramid; 6+2, hexagonal bipyramid) also contribute to the richness of the structural topologies of such solids. The so-called uranyl — organic frameworks (UOF) have now been described in many contributions with the use of a large range of organic aliphatic or aromatic carboxylates [18]. The $[O = U = O]^{2+}$ moiety is typically unreactive and terminates bonding along the axial plane, resulting in the widely recognized twodimensional sheets of uranyl polyhedra. The coordination chemistry of $[0 = U = 0]^{2+}$ is the most developed of the actinides, and the vast majority of complexes and solids exhibit the UO_2^{2+} uranyl ion.

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In some UO₂²⁺ uranyl complexes, it was found that there was cation– cation interaction (CCI) [19,20] through the oxo group and the two terminal oxygen atoms located in trans-position, which is an important feature in the chemistry of 5f elements. On the other hand, carboxylates and polycarboxylates are particularly suited for uranyl complexation [6], and they are very commonly used in this field. Among them, the simplest of these ligands is the linear, aliphatic dicarboxylates of the general formula $[OOC-(CH_2)_n-COO]^{2-}$, where Cn is denoted in the following. The coordination polymers formed by uranyl ions with malonate (C3), succinate (C4), glutarate (C5), adipate (C6), pimelate (C7), suberate (C8), azelate(C9), and sebacate (C10) have been frequently investigated [7–9], and compared with lanthanide ion complexes. Some one- to three-dimensional uranyl-organic assemblies were isolated, and the effect of additional N-heterocyclic small molecules, which can assume the different roles of ligand or structure-directing species, was also examined. But, we have found that two terminal oxygen atoms of UO_2^{2+} uranyl structures are both in trans-position (O = U = O). More recently, during an investigation of succinate (C4) as a ligand for uranyl ions, it is worthy to note that we have obtained two types of uranyl ions; one is trans-dioxido unit succ complex and the other one is cis-dioxido unit succinate complex, which is not reserved in common uranyl complexes so far. In particular, the discovery of the unexpected cis-dioxido uranyl compound is an exciting event and a milestone in the history of actinide chemistry. Although actinide is situated beneath the lanthanides within the f block of the periodic table, the lighter members of the actinide series (that is, Th-Am) retain a closer affinity in the ability to access higher formal oxidation states and engage in metal-ligand multiple bonding. These chemical properties are perhaps best exemplified by the dioxide cations MO_2^{n+} (n = 1, 2), which represent a group of considerable environmental importance for the actinides [21].

However, while the geometry of the dioxido unit (Fig. 1) in transitionmetal complexes is highly dependent on the valence-electron count, this electronic distinction is not observed with the actinides, where, regardless of oxidation state or valence-electron count, the dioxido group always adopts a linear (trans) geometry, which imparts unique chemical behavior. Herein we describe the structures and spectroscopic characterization of the carboxylate-bridged uranium (VI) coordination polymers, $[(UO_2)(suc)(H_2O)]$ (1) and $[(UO_2)(suc)_{1.5}] \cdot (4,4'-bipy)_{0.5} \cdot (suc)_{0.5}$ \cdot 3H₂O (2), in which complex 1 possesses a bent (cis) dioxido unit with an extraordinarily acute (115.421(2)°) bond angle and complex 2 is a trans-dioxido unit with an extraordinarily acute (178.421(2)°). The corresponding two uranyl (VI) complexes 1 and 2 were prepared by the reaction of [UO₂(CH₃COO)₂] and succinate ligands in the system of aqueous at 100 °C [22]. The two complexes were fully characterized by elemental analysis, IR spectra, UV-vis spectroscopy, single-crystal X-ray diffraction, thermogravimetric analysis, luminescence and photocatalytic

X-ray single crystal analysis [23] indicates that complex 1 is crystallizing in orthorhombic, space group Ama2. The molecular structure contains one uranium atom, two terminal oxygen atoms, one succinate ligand, and one coordinated water molecule. The uranium (Fig. 2a) atom was seven-coordinated by two terminal oxygen atoms (O1, O1 $^{\#}$ from uranyl unit, $^{\#}1 = 0.5$ -x, y, z) with O = U = O angle of 115.421(2) $^{\circ}$ and bond length of 1.754(9)Å, four oxygen atoms (O3, O3 $^{\#}$, O4, O4 $^{\#}$) from the carboxyl group of four succinate ligands, respectively, with the corresponding bond distances in the range of 2.382(8)Å to 2.404(7) Å and one oxygen atom (O2) of from a water molecule with the bond length

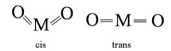


Fig. 1. Dioxido geometries of MO₂ⁿ⁺ complexes.

of 2.446(12)Å, forming a monocapped trigonal prismic geometry. The O = U = O bond angle is variously smaller from $178(1)^\circ$ in $UO_2(CH_2COO)_2 \cdot H_2O$ (3) [24] and $178.4(4)^\circ$ in $[UO_2(C_4H_4O_4)] \cdot H_2O$ (4) [25]. While, the bond length of U = O is slightly bigger than 1.74(2)Å of 3 [24] and 1.73(1)Å and 1.750(9)Å of 4 [25]. The adjacent uranium polyhedra were linked by two oxygen atoms from one carboxyl group of succinate acid ligand to form one-dimensional chain (Fig. 2b). These chains were linked by carbon atom of succinate acid ligand to form a two-dimensional layer (Fig. 2c). Furthermore, the three-dimensional network is formed by the bridging of the 2D layer through bidentate succinate groups (Fig. 2d). There is a kind of hydrogen bond in complex 1: O2–H2C---O4^{#1} (#1 = 1/2 - x, -1/2 + y, -1/2 + z), which enhance the molecular structural stability.

Structural analysis shows that complex 2 is crystallizing in monoclinic, space group C2/c. The molecular structure of complex 2 consist of one uranium atom, two terminal oxygen atoms, one and a half succinate ligand, half a lattice 4,4'-bipy molecule, half a lattice succinate acid molecular and three lattice water molecules. The uranium (Fig. 3a) atom was eight-coordinated by two oxygen atoms (O1, O2 from uranyl unit) with O = U = O angle of 178.942(6)° and U-O bond length in the range of 1.760(4) Å to 1.769(4) Å and six oxygen atoms (O3–O8) from the carboxyl group of three succinate ligands with the bond distances in the range of 2.443(4) Å to 2.470(4) Å, forming a hexagonal bipyramid. The adjacent uranium polyhedra were linked by oxygen atoms from two carboxyl group of succinate acid ligand to form an infinite one-dimensional chain (Fig. S4a). The uranium polyhedra form a layer in ab plane by the dicarboxylate group (Fig. S4b). There are two kinds of hydrogen bonds in complex 2: (i) hydrogen bond (O-H---O) between the oxygen (donor) and oxygen (acceptor); (ii) hydrogen bonds (C-H---O) of carbon (donor) and the oxygen (acceptor). The two adjacent layers were connected to form a 3D network supermolecular structure (Fig. S4) by the hydrogen bond of C6–H6A---O2. The interlayer region includes lattice 4,4'-dipy and water molecules which are connected also by the hydrogen bonds (Fig. 3b).

The structural analysis indicates that the complexes 1 and 2 contain different geometries of uranyl moiety, 1 is a cis-dioxo uranyl moiety $(O = U = O \text{ band angle of } 115.421(2)^{\circ})$ and bond length of 1.754(9)Å. The complex 2 is a trans-dioxido unit (O = U = O band angle of 178.942(6)°). Although the synthesis method is similar, the obtained results are different. And the particularly easy and reversible formation of this coordination polymeric product from uranyl acetate and succinate acid, occurring at 100 °C is comparable to those found in many uranyl(VI) complexes. It is surprising to find that the O-U-O angle (115.421(2)°) of complex 1 is much more than that reported $([UO_2(fcdc)(thf) \cdot (Fc)]_n$, fcdc = 1,2-ferrocenedicarboxylate, Fc = ferrocene) by Duval et al.(70.8°), which was considerably the smallest than the O-M-O angles in cis-dioxido transition metal complexes, to be associated with U-O distances equal to those in trans-uranyl complexes. However, the result reported by Duval et al. is not confirmed furthermore so far and it failed to reproduce the product [21,26]. So, we think that the O-U-O angle (115.421(2)°) of the complex 1 should be more reasonable and reproducible.

It is worthy to note that in the synthetic process of complex 1, we introduced lanthanide element (Ln) and urea or thiourea to the reaction system, however, the reaction result did not contain the species, unexpectedly, a cis-dioxido uranyl succinate complex (1) was obtained. If in the reaction system, reagent of lanthanide element (Ln) and urea or thiourea was not added, reaction result is the same as that of reported in literature [24,25], namely, trans-dioxido uranyl succinate complex. So, we have thought that reagent of lanthanide element (Ln) and urea or thiourea plays important role probably in the directional synthesis course.

Fluorescence of uranyl complexes typically has a characteristic of six peak spectrum relating to the $S_{11} \rightarrow S_{00}$ and $S_{10} \rightarrow S_{0\nu}$, where $\nu = 0$ –4, electronic transitions [27], and the most intense peak (S10 \rightarrow S00) is positioned at 511 nm for UO₂(CH₃COO)₂ \cdot 2H₂O. Comparison of the

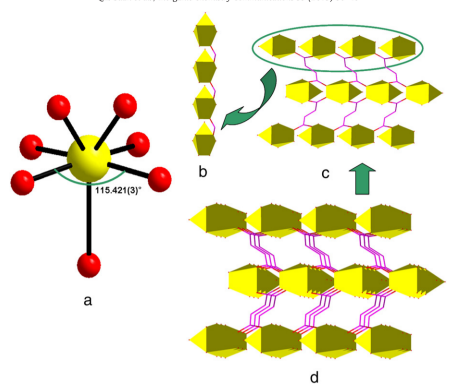


Fig. 2. (a) the coordination mode of complex 1; (b) a view of one-dimensional chain of complex 1; (c) a view of two-dimensional layer of complex 1; (d) a view of three-dimensional framework of complex 1.

spectra observed of the complex 1 displayed a red shift ranging from 10 to 15 nm. For complex 2, it was composed of broad signals with maximum at 547 nm and 573 nm but without resolution. It should be attributed to the coordination environment and space filling.

In order to further study the photocatalytic activity of complexes 1 and 2 in detail, we selected rhodamine B (RhB), a typical dye, as the model substrate to evaluate the photocatalytic activity. To our knowledge, photocatalytic materials have attracted much attention for solving the pollution problems because of their potential applications in the green degradation of organic pollutants [28]. Uranyl units have the potentialactivity for photocatalytically organic substrates [29,30], due to the thoroughly decomposing of organic ligand which completed by the uranyl double bond [29,31].

The absorption in the UV region should be attributed to the ligand to metal charge transfer (LMCT) between the U=0 bond which probably is responsible for the photocatalytic activities of the uranyl complexes

[29]. By analyzing the UV-vis spectra of complexes 1 and 2, it is revealed that the two complexes exhibit strong absorption in the UV and visible region, indicating that the complexes should be the photocatalytic behavior under the UV or visible irradiation. The concentration change of RhB was characterized by the variation of the absorbance at 554 nm. As illustrated in Fig. 4, the addition of complex 1 can greatly accelerate the degradation of RhB under UV light irradiation, however the addition of complex 2 much more accelerates the degradation of RhB under visible light irradiation. Approximately 84.1% and 10.9% of RhB were decomposed after 100 min of irradiation, respectively. It is worthy to note that the extent of degradation for complexes 1 and 2 is different, the reason may be due to the cis/trans-dioxido uranyl moiety and lattice molecules, which likely provides a value clue for further investigations. The results show that the two complexes possess potential photocatalytic activity in the decomposition of RhB under UV and visible light irradiation.

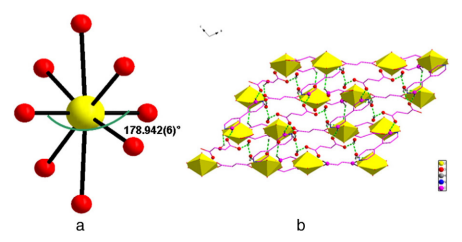


Fig. 3. (a) the coordination mode of complex 2; (b) a view of three-dimensional framework of complex 2.

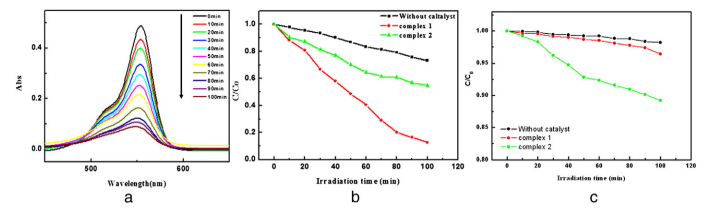


Fig. 4. (a) The UV-vis absorption spectra of the RhB solution during the decomposition reaction for complex 1. (b) Photocatalytic degradation of RhB solution under UV with the use of complexes; the black curve is the control experiment without any catalyst. (c) Photocatalytic degradation of RhB solution under visible light with the use of complexes; the black curve is the control experiment without any catalyst.

In summary, in this work we presented the synthesis and structural characterization of two uranyl complexes constructed from succinic acid ligand. These results indicate that the cis-dioxido uranyl complex demonstrates different spectroscopy, luminescence and photocatalytic behavior in comparison to trans-dioxido uranyl complex. Further investigations will be considered studied for the cis-dioxido geometry.

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Appendix A. Supplementary material

Experimental materials and methods, X-ray Crystallographic Determination, Spectra studies, XRD analysis, Thermal properties, Photoluminescent Properties, Table S1-S3 and Figure S1-S12 are presented in Supplementary material. Copies of this information may be obtained free of charge, by quoting the publication citation and deposition numbers CCDC 968660 (1) and 968661 (2) from the Director, CCDC, 12 Union Road. Cambridge, CB2 1EZ, UK (fax +44- 1223-336033; e-mail deposit@ccdc.cam.ac.uk; http://www.ccdc.cam.ac.uk). Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.inoche.2015.06.027.

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- Synthesis of [(UO₂)(suc)(μ_1 -OH₂)] (1): UO₂(CH₃COO)₂ \cdot 2H₂O and succinic acid are available commercially and were used without any further purification. UO₂(CH₃COO)₂ · 2H₂O (0.039 g, 0.1 mmol), succinic acid (0.012 g, 0.1 mmol) $Sm(NO_3)_3 \cdot 6H_2O(0.034 \text{ g}, 0.1 \text{ mmol})$ and urea (0.006 g, 0.1 mmol) were dissolved in demineralized water (10 mL). The solution was stirred for 4 h at room temperature, and heated at 120 °C for 3 days. Yellow crystals are obtained with a yield of 69% based on UO₂(CH₃COO)₂ · 2H₂O. Yield: 55%. Anal.Calc. for C4H6O7U(404.12): C, 11.89; H, 1.48. Found: C, 11.86; H, 1.53. IR data (KBr, cm⁻¹): 3407, 1532, 1438, 1293, 940. Synthesis of $[(UO_2)(suc)_{1.5}] \cdot (4,4'-bipy)_{0.5} \cdot (suc)_{0.5} \cdot 3H_2O(2)$: A mixture of 0.040 g (0.1 mmol) UO₂(CH₃COO)₂ · 2H₂O, 0.016 g (0.15 mmol) succinic acid, 0.032 g (0.15 mmol) 4,4'-Dipyridine and 10 mL of demineralized water was put in a glass vessel and stirred for 4 h at room temperature. Then the solution was kept at 120 °C for 3 day. After filtration and wash with demineralized water, light yellow crystals were obtained with a yield of 42% based on UO₂(CH₃COO)₂ · 2H₂O. Anal. Calc. for C13H18NO13U (634.31): C, 24.6; H, 2.84; N, 2.21. Found: C, 24.55; H, 2.86; N, 2.19%. IR data (KBr, cm⁻¹): 3442, 1628, 1538, 1449, 1417, 1237, 809, 916.
- [23] Crystal data for [(UO₂)(suc)(H₂O)]: C4H6O7U, Mr = 404.12, orthorhombic, Ama2, a = 10.8569(14) Å, b = 9.5114(12) Å, c = 7.5892(10) Å, b = 90°, V = 783.69(18) Å3, Z = 4, ρ_{calcd} = 3.425 g cm⁻³, T = 296 K, Rint = 0.0244, R1 = 0.0264 (1> 2 σ (I)), wR2 = 0.0649 (all data), GOF = 1.144, CCDC 968660; [(UO₂)(suc)_{1.5}] · (4,4'-bipy)_{0.5} · (suc)_{0.5} · 3H₂O: C13H18NO13U, Mr = 634.31, monoclinic, C2/c, a = 23.843(2) Å, b = 14.4772(9) Å, c = 12.4312(10) Å, b = 106.735(10)°, V = 4109.2(6) Å3, Z = 8, ρ_{calcd} = 2.051 g cm⁻³, T = 293 K, Rint = 0.0318, R1 = 0.0276 (1> 2 σ (I)), wR2 = 0.0753 (all data), GOF = 1.059, CCDC 968661.
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