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Ru(IV)—Ru(IV), Ru(III)—Ru(IV), and Ru(III)—Ru(III) complexes having a sulfato bridging ligand on the doubly oxido-/hydroxido-bridged core, and their crystallographic and electronic structures†

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The Ru₂(IV,IV), Ru₂(III,IV), and Ru₂(III,III) complexes with the doubly oxido- and/or hydroxido-bridged diamond core $\{Ru_2(\mu-O(H))_2\}$, bridged by an $\eta^1:\eta^1:\mu^2$ -type bidentate sulfato ligand, $[\{Ru(L)\}_2(\mu-O)_2(\mu-O_2SO_2)]^{m+1}$ ($m=1: [III,IV]^+$; $m=2: [IV,IV]^{2+}$), $[\{Ru^{III,IV}(L)\}_2(\mu-O)(\mu-OH)(\mu-O_2SO_2)]^{2+1}$ ($[III,IV]^+$), and $[\{Ru^{III,III}(L)\}_2(\mu-OH)_2(\mu-O_2SO_2)]^{2+1}$ ($[III,III]^+$) (L=1) (L

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Introduction

Multi-nuclear transition metal complexes having a doubly oxido- and/or hydroxido-bridged core, $\{M_2(\mu\text{-O(H)})_2\}$, have attracted much attention as reaction centres in a number of natural metalloenzymes. For instance, the oxidation states of the active site of soluble methane monooxygenase (sMMO) range from $\text{Fe}_2(\Pi,\Pi)$ to $\text{Fe}_2(\Pi,\Pi)$ during the course of the catalytic reaction. Studies on the structural changes of/around the reaction centres, namely, the $\{M_2(\mu\text{-O(H)})_n\}$ (n=1, 2) core, accompanied by redox reactions coupled with proton transfers, are important for discussions on reaction mechanisms. For example, the doubly oxido- and/or hydroxido-bridged metal

core, $\{Fe_2^{IV,IV}(\mu-O)_2\}$, is the potential structure of the active species Q, but the debate over whether the core structure is the "open-type" or "closed-type" has long been continued regarding Q. Furthermore, the resting state (MMOHox) has been reported to have the $\{Fe_2^{III,III}(\mu\text{-OH})_2\}$ core. A number of studies on the electronic structures, catalytic oxidation reactions, and model reactions by sMMO analogues have been reported using diiron² or dinickel centres.³ As structural model compounds of such diiron-centred reaction sites, ruthenium complexes as the same 8th group element-containing compounds have also been intensively investigated; however, the number of studies is still limited. Although Ru₂(v,v), ^{6a,b} Ru₂(IV,IV), ⁶ Ru₂(III,IV), ⁴ and $Ru_2(III,III)^{6b,5}$ complexes with the $\{Ru_2(\mu-O(H))_2\}$ diamond core have been reported in different supporting ligand systems, only a few in the same supporting ligand system have been reported to date, making direct comparisons of the properties and reactivities of the reaction centres difficult. Bercaw et al. reported the isovalent species, Ru₂(v,v), Ru₂(iv,iv), and Ru₂(iii,iii) with the phosphate-donor oxygen tripod ligand [Co^{III}(Cp)(PO $(OR)_2)_3$ (Cp = cyclopentadienyl, R = Me, Et) system; 6a,b however, the mixed-valent species have not been isolated and characterised in their ligand system.

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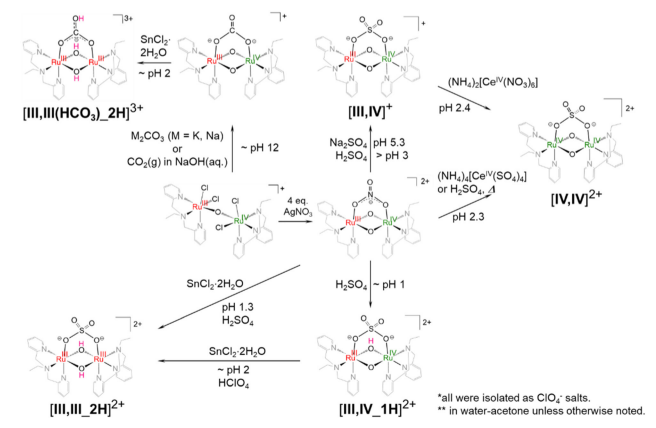
We reported diruthenium complexes with $\{Ru_2(\mu-O(H))_n\}$ (n = 1, 2) cores using the ethylbis(2-pyridylmethyl)amine (ebpma) tridentate ligand. For the $\{Ru_2(\mu-O(H))_2\}$ core, we synthesised carbonato-, 7e hydrogencarbonato-, 7e acetato-, 7c,d and nitrato-bridged^{7c} complexes in the $Ru_2(III,IV)$ states, in which S = 1/2 at 298 K, and compared their structures and properties. In addition, we recently reported the Ru₂(IV,IV) complexes using carbonato and hydrogenearbonato bridging ligands, and their reactivity in solution and application toward the oxidation of organic substrates were demonstrated.⁷f

In this study, sulfato-bridged complexes in the Ru₂(IV,IV), Ru₂(III, IV), and Ru₂(III,III) states and a hydrogencarbonato-bridged Ru₂(III,III) complex, for which only the higher-valent Ru₂(IV,IV) and Ru₂(III,IV) species with the corresponding frameworks have been identified, ^{7ef} were synthesised. The crystal structures and electronic structures of these complexes in one ligand system were investigated and directly compared. These complexes are significant as structural models for the series of reaction intermediates of sMMO.

Results and discussion

Synthesis of sulfato-bridged complexes in the Ru₂(IV,IV), Ru₂(III, IV), and Ru₂(III,III) states and a hydrogencarbonato-bridged $Ru_2(III,III)$ complex having the $\{Ru_2(\mu-O(H))_2\}$ cores

The synthetic procedure is summarised in Scheme 1. Sulfatobridged Ru₂(III,IV) complexes having the {Ru₂(μ-O)₂} or {Ru₂(μ-O)₂} O)(μ -OH)} core, [{Ru^{III,IV}(ebpma)}₂(μ -O)₂(μ -O₂SO₂)]ClO₄·0.25 $(CH_3)_2CO\cdot 4H_2O$ ([III,IV]ClO₄) and $[\{Ru^{III,IV}(ebpma)\}_2(\mu-O)(\mu-O)]$ OH)(μ -O₂SO₂)](ClO₄)₂·3H₂O ([**III,IV_1H**](ClO₄)₂), respectively, were synthesised through reactions of a nitrato-bridged $[{Ru^{III,IV}(ebpma)}_{2}(\mu-O)_{2}(\mu-O_{2}NO)](ClO_{4})_{2}\cdot 1.5H_{2}O$ complex, ([III,IV(NO₃)](ClO₄)₂), which was prepared by the reaction of a singly oxido-bridged complex ([{Ru^{III,IV}Cl₂(ebpma)}₂(μ-O)] ClO₄·0.75(CH₃)₂CO)^{7b} with four molar equivalents of AgNO₃.⁷⁰ Above pH 3 in the presence of sulfate sources such as Na₂SO₄, the doubly oxido-bridged complex ([III,IV]ClO₄) was synthesised, and at approximately pH 1, the hydroxido- and oxidobridged complex ([III,IV_1H](ClO₄)₂) was isolated. [III,IV_1H] (ClO₄)₂ could also be formed in a reaction of $[{Ru^{III,IV}Cl_2(ebpma)}_2(\mu-O)]^{+7b}$ with ten-fold Na₂SO₄ in water/ acetone if four molar equivalents of AgTfO (TfO = trifluoromethanesulfonate) were present. The corresponding Ru₂(IV,IV) complex, $[\{Ru^{IV,IV}(ebpma)\}_2(\mu-O_2SO_2)](ClO_4)_2 \cdot 0.5H_2O$ ([IV, IV[(ClO₄)₂), was isolated through the one-electron oxidation of [III,IV(NO₃)]²⁺ by cerium(IV) ammonium sulfate (CAS) at pH 2.3, owing to nitrato ligand substitution to sulfate in CAS. [IV, $[V]^{2+}$ was also afforded by the one-electron oxidation of $[III,IV]^{+}$ by cerium(IV) ammonium nitrate. The Ru₂(III,III) complex having the doubly hydroxido-bridged core $\{Ru_2(\mu\text{-OH})_2\}$, $\lceil \{Ru^{III,III}(ebpma)\}_2(\mu\text{-OH})_2(\mu\text{-O}_2SO_2)\rceil (ClO_4)_2 \cdot 3H_2O \quad \left(\begin{array}{cc} \textbf{[III,III}_2H \end{array} \right]$ $(ClO_4)_2$), was synthesised *via* a one-electron reduction reaction of [III,IV_1H]²⁺ by SnCl₂·2H₂O, coupled with a protonation, at approximately pH 2. [III,III_2H]²⁺ was also afforded directly



Synthesis of a series of sulfato-bridged diruthenium complexes and a hydrogencarbonato-bridged complex.

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from [III,IV(NO₃)]²⁺ under similar reaction conditions via the formation of [III,IV_1H]²⁺ in solution. A corresponding Ru₂(III, III) complex having a hydrogencarbonato bridging ligand, [III, III(HCO₃)_2H](ClO₄)₃·H₂O, was newly prepared in the reaction of the corresponding $Ru_2(III,IV)$ complex, $[III,IV(CO_3)_2H]^{3+}$.

The sulfato ligand functions as a multidentate bridging ligand with versatile coordination modes to form multinuclear complexes (for example, with first, 8-16 second, 17-24 and third row²⁵⁻²⁹ d-block elements and lanthanoids³⁰⁻³³ actinoids^{34,35}), and it has been investigated as one of the building blocks of multinuclear frameworks or self-assembled structures. Among the coordination modes, there have been a limited number of Ru dimers, 20 and to the best of our knowledge, only a small number of crystal structures have been characterised or finely proposed in the $\eta^1:\eta^1:\eta^1:\mu^2$ -mode^{20f} or in the $\eta^1{:}\eta^1{:}\mu^2{-}mode^{20{\it a-c},g}$ which is the same type as the complexes described herein. Therefore, reporting the $\eta^1:\eta^1:\mu^2$ -O₂SO₂²⁻-bridged complexes in the Ru₂(IV,IV), Ru₂(III,IV), and Ru₂(III,III) states significantly expands the available structural data across a wide range of oxidation states.

X-ray crystallography

The crystal structures of all the species, [IV,IV]²⁺, [III,IV]⁺, [III, $IV_1H_1^{2+}$, $[III,III_2H_1^{2+}]$, and $[III,III(HCO_3)_2H_1^{3+}]$, were successfully characterised. Crystallographic data are summarised in Table S1,† and selected metric parameters are listed in Table 1. The crystal structures are shown in Fig. 1-3 and Fig. S1 and S2.† Each ruthenium centre had a distorted hexacoordinate octahedral geometry and was coordinated by tridentate ebpma in the facial manner; the other three coordination sites were occupied by two bridging oxido or hydroxido ligands and one of the sulfato oxygens. The sulfato ligand bridged the $\{Ru_2(\mu-O(H))_2\}$ core through two oxygens as a bidentate ligand. The hydrogencarbonato in [III,III(HCO₃)_2H]³⁺ bridged the {Ru₂(μ-OH)₂} core in a similar manner to a bidentate ligand. Both the bridging oxido or hydroxido ligands were located trans to the pyridyl (py) and amine nitrogen atoms, which was the same geometrical configuration as that of previously reported systems. 7c-f

The metric parameters around the $\{Ru_2(\mu-O(H))_2\}$ core differed depending on the electronic structure of the Ru centres and the number of protons on the core, whereas those around

the $\{\mu - O_2 S O_2\}^{2-}$ ligand were similar among the complexes (Table 1). Between $[\mathbf{IV}, \mathbf{IV}]^{2+}$ and $[\mathbf{III}, \mathbf{IV}]^{+}$ having the $\{Ru_2(\mu-O)_2\}$ core, the Ru-O and Ru...Ru distances were shorter for [IV,IV]2+ (average: 1.917 and 2.4470(3) Å) than those of [III,IV]⁺ (average: 1.939 and 2.4705(4) Å) because of the more cationic Ru(IV) centres, while the bond angles around the core were similar. The shorter Ru···Ru distance in [IV,IV]2+ indicated stronger interactions between the Ru centres. In [III,IV_1H]2+ having the $\{Ru_2(\mu-O)(\mu-OH)\}\$ core, the Ru-O2 distances in $[III,IV_1H]^{2+}$ (average: 2.029 Å) was elongated by 0.05-0.10 Å compared to the other Ru-O1 bond lengths in [III,IV_1H]2+ (average: 1.917 Å) and $[\mathbf{III},\mathbf{IV}]^+$ having the $\{Ru_2(\mu-O)_2\}$ core (average: 1.939 Å), suggesting that one of the bridging oxido ligands (O2), which was away from one of the terminal sulfato oxygens (O6), was singly protonated. Accordingly, the Ru...Ru distance was elongated (2.5342 (3) Å) compared to that of $[\mathbf{III}, \mathbf{IV}]^+$ (2.4705(3) Å), and the Ru-OH-Ru angle (77.28(8)°) was slightly smaller than the Ru-O-Ru angle (82.77(9)° in [III,IV_1H]²⁺). It is noteworthy that, according to the theoretical calculations, a geometrical isomer of [III, IV_1H|2+, in which O1 is singly protonated instead of O2 (Table S10† Model-B), could slightly be stabilized by 3.13 kcal mol⁻¹ in free energy (Table S11†) owing to the intramolecular hydrogen bonding interactions between O1-H and O6-SO₃²⁻ (1.677 Å). However, it was rather challenging to separate the isomers in the synthetic procedures.

For [III,III_2H]²⁺, the two Ru-OH bond lengths (average: 2.019 Å) were almost identical to those of [III,IV_1H]²⁺ (average: 2.019 Å); however, because both oxygen atoms were singly protonated, the Ru···Ru distance (2.5695(7) Å) was longer than that in [III,IV_1H]²⁺ (2.5342(3) Å) but comparable to that in $[III,III(HCO_3)_2H]^{3+}$ (2.5657(4) Å). The angles around the {Ru₂(μ-OH)₂} core differ slightly from those of [III, IV_1H]²⁺. The longer Ru···Ru distances in [III,III_2H]²⁺ and [III,III(HCO₃)_2H]³⁺ suggest weaker interactions between the $d\pi(Ru)$ centres. The closer orbital energies of the LUMO and HOMO may contribute to unpairing the electrons in the HOMO, resulting in the observation of a certain extent of a paramagnetic character of [III,III_2H]²⁺, as discussed later.

The Ru-O3,4(sulfato) bond lengths were longer in [III, III_2H^{2+} (average: 2.115 Å), followed by $[III,IV]^+$ (average: 2.091 Å) and $[III,IV_1H]^{2+}$ (average: 2.083 Å), and $[IV,IV]^{2+}$ (average:

Table 1 Selected metric parameters of the Ru₂(IV,IV), Ru₂(III,IV), and Ru₂(III,III) complexes

| Ru(III)-Ru(III) | | | | Ru(m)-Ru(v) | | | | Ru(w)-Ru(w) | | |
|----------------------------|---|------------|---|-------------|---|------------|--|------------------------------|--|------------|
| | [III,III(HCO ₃)_2H] (ClO ₄) ₃ ·0.25H ₂ O | | [III,III_2H] (ClO ₄) ₂ ·1.5H ₂ O | | [III,IV]ClO ₄ ·3.5H ₂ O | | [III,IV_1H](ClO ₄) ₂ ·0.75H ₂ O | | [IV,IV](ClO ₄) ₂ ·H ₂ O | |
| Bond distances/Å | | | | | | | | | | |
| Ru-O1 | 2.028(3) | 2.026(2) | 2.023(6) | 2.023(6) | 1.930(2) | 1.945(2) | 1.921(2) | 1.913(2) | 1.9080(19) | 1.9042(19) |
| Ru-O2 | 2.014(3) | 2.024(3) | 2.010(6) | 2.020(6) | 1.938(3) | 1.937(3) | 2.030(2) | 2.028(2) | 1.9173(19) | 1.926(2) |
| Ru1···Ru2 | 2.5657(4) | . , | 2.5695(7) |) | 2.4705(4) | . , | 2.5342(3) | | 2.4470(3) | |
| Ru-O(bidentate ligand) | 2.095(3) | 2.070(3) | 2.121(5) | 2.109(5) | 2.093(3) | 2.089(3) | 2.080(2) | 2.086(3) | 2.0520 (18) | 2.0419(18) |
| Bond angles/° | | . , | . , | . , | | . , | | | | , |
| Ru1-O1,2—Ru2 | 78.52(9) | 78.88(10) | 78.9(2) | 79.2(2) | 79.22(9) | 79.22(10) | O1 _{O2-} 82.77(9) | O2 _{OH-} : 77.28(8) | 79.87(7) | 79.08(7) |
| O1-Ru1,2-O2 | 101.03(10) | 100.74(10) | 101.0(2) | 100.6(2) | 100.92(10) | 100.46(10) | 99.51(10) | 99.87(10) | 100.49(8) | 100.30 (8) |
| Ru-O(bidentate ligand)-S/C | 120.0(2) | 121.7(3) | 119.7(3) | 117.9(3) | 122.89(16) | 123.53(15) | 119.93(15) | 118.31(14) | 121.76(11) | 121.71(11) |

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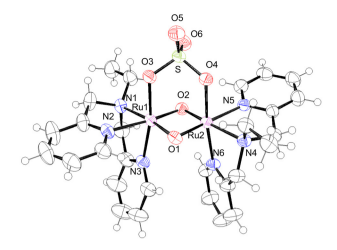


Fig. 1 ORTEP drawing of [IV,IV]²⁺ shown at the 50% thermal probability.

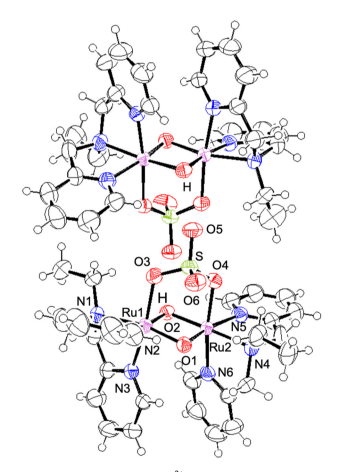


Fig. 2 ORTEP drawing of [III,IV_1H]²⁺ shown at the 50% thermal probability (notation of H is added to the bridging oxygen where the Ru-O bond lengths were elongated).

2.047 Å), indicating that the sulfato ligand coordinated more strongly to the more cationic $\{Ru_2(\mu-O(H))_2\}$ core. Furthermore, the Ru-N2,5(py, trans-O(oxido/hydroxido)) bond lengths were longer in $[III,IV]^+$ and $[IV,IV]^{2+}$ having the $\{Ru_2(\mu-O)_2\}$ core

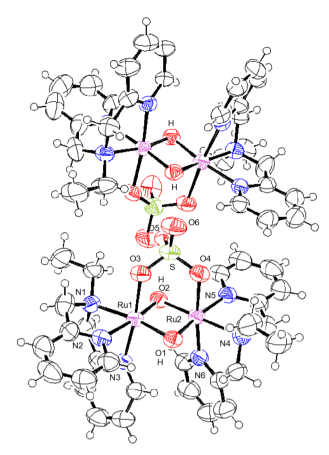


Fig. 3 ORTEP drawing of [III,III_2H]²⁺ shown at the 50% thermal probability (notation of H is added for the bridging oxygen).

(2.0912.103 Å) compared to those in $[III,IV_1H]^{2+}$ and $[III,IV_1H]^{2+}$ III_2H $^{2+}$ having the {Ru₂(μ -O)(μ -OH)} or {Ru₂(μ -OH)₂} cores (2.062–2.078 Å), respectively. Upon protonation to the bridging oxido ligand, the $p\pi(O)$ - $d\pi(Ru)$ interactions became stronger by decreasing the number of lone pair electrons on oxygen.

The metric parameters were also compared with those of complexes having different bridging ligands, $\{\mu\text{-O}_2\text{CO}\}^{2-}$ ([IV, $IV(CO_3)]^{2+7f}$ or $[III,IV(CO_3)]^{+7e}$), $\{\mu\text{-O}_2COH\}^ ([IV,IV(HCO_3)]^{3+,7f}$ $[III,IV(HCO_3)]^{2+}$, 7e $[III,IV(HCO_3)_1H]^{3+}$, 7e or $[III,III(HCO_3)$ $[2H]^{3+}$), $\{\mu\text{-O}_2\text{CCH}_3\}^-$ ([III,IV(CH₃O₂)]^{2+7c} or [III,IV(CH₃O₂) $[III,IV(NO_3)]^{2+7c}$, and $[\mu-O_2NO]^-$ ($[III,IV(NO_3)]^{2+7c}$). The order of the Ru···Ru distances was consistent among the Ru₂(IV,IV) and $Ru_2(III,IV)$ complexes; $[IV,IV]^{2+}$ (2.4470(3) Å) > $[IV,IV(HCO_3)]^{3+}$ $(2.4389(3) \text{ Å}) > [IV,IV(CO_3)]^{2+} (2.4261(3) \text{ Å}) \approx [III,IV]^+ (2.4705(4))$ \mathring{A}) > $[III,IV(HCO_3)]^{2+}$ (2.4547(2) \mathring{A}) > $[III,IV(CH_3O_2)]^{2+}$ (2.4463) $(10) \text{ Å} > [III,IV(CO_3)]^+ (2.4434(8) \text{ Å}) > [III,IV(NO_3)]^{2+} (2.4334(4))$ Å). These results were in accordance with the order of the angles of Ru–O–X (X = $S(sp^3) > C(sp^2) > N(sp^2)$) of the respective $\{\mu - O_2 X O_n\}^{m-}$ (n = 1, 2), in which the central X atom has a different geometry and flexibility upon coordinating to the Ru centres. Notably, the electrostatic nature of the $\{\mu$ -O₂XO_n $\}^{m-}$ ligand does not seem to influence the metric parameters. The same reasoning regarding the Ru...Ru distance is applicable to complexes having the asymmetric $\{Ru_2(\mu-O)(\mu-OH)\}\$ core and

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the Ru₂(III,III) complexes having the {Ru₂(μ -OH)₂} core; [III, IV_1H^{2+} (2.5342(3) Å) > [III,IV(HCO₃)_1H]³⁺ (2.5332(5) Å) > [III, $IV(CH_3CO_2)_1H^{2+}$ (2.5201(6) Å) and $[III,III_2H]^{2+}$ (2.5695(7) Å) $> [III,III(HCO_3)_2H]^{3+} (2.5657(4) \text{ Å}).$

Electronic structures

The ¹H NMR spectra of [IV,IV]²⁺, [III,IV]⁺, and [III,IV_1H]²⁺ showed some broad signals, which were observed only in the diamagnetic region and were not assignable (Fig. S4 and S5†). Therefore, magnetic measurements were performed at 298 K. The effective magnetic moment (μ_{eff}) of $[\text{IV,IV}](\text{ClO}_4)_2 \cdot 0.5 \text{H}_2\text{O}$ was $0.94\mu_{\rm B}$, indicating the paramagnetic character of the d⁴-d⁴ system, almost consistent with those of the corresponding carbonato- and hydrogenearbonato-bridged complexes, [IV,IV $(CO_3)^{2+}$ and $[IV,IV(HCO_3)]^{3+}$ (1.04 and 0.85 μ_B , respectively).^{7f} We newly performed the theoretical calculations of [IV,IV]²⁺ for the quintet, open-shell singlet, and closed-shell singlet states to testify which best describes the electronic structure of [IV, $[V]^{2+}$. The free energy was lower for the open-shell singlet state than those of other states (Table S8†). Specifically, the paramagnetic features experimentally studied could be supported by the theoretical result that the free energy of the open-shell singlet was lower than that of the closed-shell singlet state by -6.27 kcal mol⁻¹. The metric parameters of the optimised structure for the open-shell singlet state were closer to those observed in the single crystal XRD analysis than those obtained from the calculation for the quintet state (Table S7†). Therefore the MO energy diagram of the open-shell singlet state is shown in Fig. S35.† The spin density was clearly localised on the $\{Ru_2(\mu-O)_2\}$ core (Fig. S36†), showing the stronger anti-ferromagnetic interactions between the Ru (S = 1) centers.

The μ_{eff} values of [III,IV]ClO₄·0.25(CH₃)₂CO·4H₂O and [III, IV_1H](ClO₄)₂·3H₂O were 1.79 (296 K) and 1.69 $\mu_{\rm B}$ (298 K), respectively, suggesting one electronic spin on the {Ru₂^{III,IV}(μ- O_{2} core (S = 1/2). The theoretical calculations were performed assuming the doublet state for both [III,IV]⁺ and [III,IV_1H]²⁺ and the optimised structures (Fig. S37 and S39†) were wellconsistent with the crystal structures (Tables S9 and S10†). The spin density plot supported the paramagnetic character of the $\{Ru_2(\mu-O(H))_2\}$ core (Fig. S38 and S40†). ESR signals centred at g = 2.07 - 2.09 were observed for [III,IV]⁺ and [III,IV_1H]²⁺ at both 298 (powder samples) and 93 K (frozen acetonitrile) (Fig. S8, S9 and Table S3†), which supported the paramagnetic characters of the $\{Ru_2(\mu-O(H))_2\}$ core.

For [III,III_2H]²⁺ and [III,III(HCO₃)_2H]³⁺, the ¹H and ¹³C NMR spectra confirmed the diamagnetic character of the $\{Ru_2^{III,III}(\mu\text{-OH})_2\}$ core (Fig. S6 and S7†). The proton or carbon atoms of the ebpma ligand could be assigned in the respective spectrum. The protons at the bridging oxido ligands were observed at 16.60 and 15.75 ppm for [III,III_2H]²⁺ and [III,III (HCO₃)_2H³⁺, respectively, and were reasonably quantified as 2H (Fig. S6†). The more downshifted hydroxido proton signals for [III,III_2H]²⁺ suggested higher electron density, which was consistent with the p K_a values of the $\{Ru_2(\mu\text{-OH})_2\}$ (5.7) and $\{Ru_2(\mu-O)(\mu-OH)\}\$ cores (4.2) (eqn (3)), determined from spectroscopic investigations (Fig. S14 and S15†). Although the NMR spectra of [III,III_2H](ClO₄)₂·3H₂O were reasonably assigned, its paramagnetic character was also suggested in the magnetic measurements ($\mu_{\rm eff} = 1.15 \mu_{\rm B}$ at 298 K). Based on the theoretical calculations, which were performed for both closed-shell singlet and triplet states, the singlet state was stabilised by only -0.168 kcal mol⁻¹ in free energy (298 K) compared to the triplet state (Table S13†). The spin state transition from the closed-shell singlet to triplet state could occur by such a lower thermal barrier. These computational results could explain the magnetic characters of [III,III 2H]²⁺.

The MO energy diagram of the closed-shell singlet state is shown in Fig. S41†. The metric parameters of the optimized structure obtained for a singlet state calculation perfectly agreed with those experimentally confirmed (Table S12†). Based on the theoretical results for [III,III_2H]²⁺, the calculations were performed for [III,III(HCO3)_2H]3+ as the closedshell singlet state (Fig. S42 and Table S14†).

UV-vis-NIR spectroscopic properties and electrochemical behaviours in CH₃CN

The electron transition spectra in acetonitrile of [IV,IV]²⁺, [III, IV]⁺, and [III,IV_1H]²⁺ (Fig. 4 and Table 2) exhibited maximum absorption wavelengths (λ_{max}) of 699, 945, and 1077 nm, respectively, which could be assigned to the characteristic electron transitions from and to the $\{Ru_2(\mu-O(H))_2\}$ diamond core. These bands indicated that the SOMO of the $\{Ru_2(\mu-O)_2\}$ core was more stabilised in the order of $[IV,IV]^{2+} > [III,IV]^{+} > [III,$ IV_1H]²⁺. [III,III_2H]²⁺ showed an intense band at 328 nm, assigned to the metal-to-ligand charge transfer (MLCT) band from the $\{Ru_2^{III,III}(\mu\text{-OH})_2\}$ core to the $\pi^*(py)$ of the ebpma ligand. On comparing the spectrum of [IV,IV]²⁺ (699 nm) with those of $[IV,IV(CO_3)]^{2+}$ (762 nm) and $[IV,IV(HCO_3)]^{3+}$ (669 nm), ⁷ it is found that the order of the λ_{max} corresponds to that of the electron-donating character of the bidentate ligand on the $\{Ru_2(\mu-O)_2\}$ core $\{\{\mu-O_2CO\}^{2-7f} > \{\mu-O_2SO_2\}^{2-} > \{\mu-O_2SO_2\}^{2-7f} > \{\mu O_2COH$; Table 2 and Fig. S10†), indicating the destabilisation of the core due to the coordination of a more electrondonating ligand. Time-dependent DFT calculations were performed for all the complexes and the wavelengths of character-

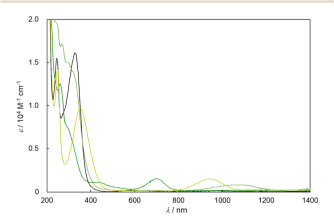


Fig. 4 UV-vis-NIR spectra of [IV,IV]²⁺ (green line), [III,IV]⁺ (olive line), [III,IV_1H]²⁺ (yellow-green line), and [III,III_2H]²⁺ (black line) in CH₃CN.

Table 2 Electrochemical and spectroscopic properties of complexes with the {Ru₂(µ-O)₂} core in CH₃CN at 298 K

| (E vs. Ag 0.01 M AgNO | O ₃)/V | $\frac{\lambda_{\rm max}/{\rm nm} \left(\varepsilon/{\rm M}^{-1} \ {\rm cm}^{-1}\right)}{\{{\rm Ru}_2(\mu\text{-O})_2\}\text{-}\{{\rm Ru}_2(\mu\text{-O})_2\}\text{+}}$ | | |
|----------------------------------|--|---|--|--|
| - () - () | - () - () | | | |
| Ru(ɪv)–Ru(ɪv)/ Ru(ɪɪɪ)–Ru(ɪv) | Ru(m)=Ru(v)/ Ru(m)=Ru(m) | Ru(ıv)-Ru(ıv) | Ru(III)-Ru(IV) | |
| 0.15 | -1.10 | 762 (1440) | 1067 (660) | |
| 0.31 | -1.02 | 699 (1710) | 945 (1440) | |
| 0.57 | -0.75 | 677 ^a | 977 (1440) | |
| 0.60 | | 669 (1970) | 972 (1490) | |
| 0.93 | -0.52 | 586 ^a , | 762 (1300) | |
| | Ru(iv)-Ru(iv)/ Ru(iii)-Ru(iv) 0.15 0.31 0.57 0.60 | Ru(m)-Ru(rv) Ru(m)-Ru(m) 0.15 -1.10 0.31 -1.02 0.57 -0.75 0.60 | $ \begin{array}{c ccccccccccccccccccccccccccccccccccc$ | |

^a Not isolated, confirmed only in solution.

Table 3 Electrochemical and spectroscopic properties of complexes with the $\{Ru_2(\mu-O)(\mu-OH)\}$ and $\{Ru_2(\mu-OH)_2\}$ cores in CH_3CN and $\{Ru_2(\mu-OH)_2\}$ cores in CH_3CN and $\{Ru_2(\mu-OH)_2\}$ cores in $\{Ru_2(\mu-OH)_2\}$ cores

| | (E vs. Ag 0.01 M AgN | | | |
|---|---------------------------------|------------------------------------|------------------------------------|---|
| | Ru(ɪv)–Ru(ɪv)/ Ru(ɪɪ)–Ru(ɪv) | Ru(III)-Ru(IV)/ Ru(III)-Ru(III) | Ru(III)-Ru(III)/ Ru(II)-Ru(III) | $\lambda_{ m max}/{ m nm} \left(\varepsilon/{ m M}^{-1} \ { m cm}^{-1} ight)$ |
| [III,IV_1H] ^{2+ this work} | 1.07 | -0.29 | -0.98 | 294 (15 000), 1077 (840) |
| [III.IV(CH ₂ CO ₂) 1H] ^{3+7c} | 1.33 | -0.22 | -0.90 | 269 (16 400), 1161 (841) |
| [III.IV(HCO ₂) 1H] ³⁺⁷¹ | 1.32 | -0.20 | -0.86 | 336 (11 600), 964 (1330) |
| III.III 2H] ^{2+ this work} | 1.14 | -0.30 | -0.98 | 328 (16 100) |
| [III,III(HCO ₃)_2H] ^{3+ this work} | 1.25 | -0.22 | -0.88 | 331 (15 200) (in CH ₃ OH) |

istic absorption bands in the visible and near-IR regions relatively matched the experimentally derived spectra (Fig. S43–S46 and Table S15†).

Furthermore, a relationship was found between λ_{max} and the redox potentials, which will be discussed in the following section. These results were consistent with those previously reported for $Ru_2(III,IV)$ complexes (Table 2). The effect of the electron-donating character of the bidentate bridging ligands in complexes having the asymmetric $\{Ru_2^{III,IV}(\mu\text{-O})(\mu\text{-OH})\}$ core was evident from their electrochemical properties $(\{\mu\text{-O}_2SO_2\}^{2-} > \{\mu\text{-O}_2CCH_3\}^{-7d} > \{\mu\text{-O}_2COH\}^-;^{7e,f}$ Table 3 and Fig. S11†). However, the effect was much smaller than that on the corresponding complexes with the $\{Ru_2^{III,IV}(\mu\text{-O})_2\}$ core. This would be because the impact of protonation of the $\{Ru_2(\mu\text{-O})_2\}$ core on the destabilization of MO energies of frontier orbitals was much larger than that of the electrostatic differences of bidentate bridging ligands $\{\mu\text{-O}_2XY\}^{n-}$.

The absorption wavelengths of $[III,III_2H]^{2+}$ (331 nm) and $[III,III(HCO_3)_2H]^{2+}$ (328 nm), which could be attributed to the MLCT bands, and molar coefficients ($\varepsilon = 1.61 \times 10^4$ and 1.52×10^4 M⁻¹ cm⁻¹, respectively) were comparable (Fig. 4 and Table 3), indicating that the electron transition energies and probabilities are approximately independent of the type of bidentate bridging ligand.

Cyclic voltammetry (CV) was performed in acetonitrile containing tetra-*n*-butylammonium perchlorate (TBAP) at 298 K (Fig. 5) and electrochemical analyses were carried out (Fig. S16–S33†). [**IV,IV**]²⁺ and [**III,IV**]⁺ showed four distinct redox waves: a quasi-reversible oxidation wave at 2.03 V, reversible one-electron

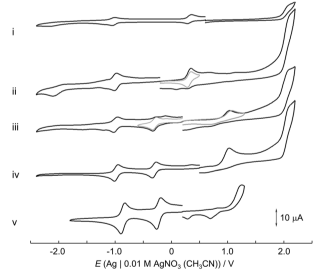


Fig. 5 Cyclic voltammograms of (i) $[IV,IV]^{2+}$ (0.66 mM), (ii) $[III,IV]^{+}$, (iii) $[III,IV]^{+}$, (iv) $[III,III]^{2+}$, and (v) $[III,III]^{2+}$ (HCO₃)]³⁺ in CH₃CN (c = 1.0 mM unless otherwise noted; scan rate, 100 mV s⁻¹; 298 K).

oxidation and reduction waves at 0.31 and -1.02 V, respectively, and an irreversible reduction wave at -2.16 V. Based on the hydrodynamic voltammetry (HDV) analysis of [III,IV]⁺ (Fig. S19–S21†), the waves at 0.31 and -1.02 V were attributed to the one-electron couples of $Ru_2(w,w)/Ru_2(w,w)$ and $Ru_2(w,w)/Ru_2(w,w)$, and the reversibility was confirmed using different scan rates (Fig. S16, S17 and Table S4†). The irreversible waves at 2.03 and

-2.16 V were attributed to the one-electron couples of Ru₂(iv,v)/ Ru₂(IV,IV) and Ru₂(III,III)/Ru₂(II,III), respectively, in line with our previous reports. 7b,c For the reduction wave at -2.16 V, the electrode reaction could be assigned to a one-electron process involving an electrode reaction-chemical reaction-electrode reaction (ECE) according to our previous reports. 7b-e In the cyclic voltammogram of [III,IV]⁺, after the second oxidation of Ru₂(IV,V)/Ru₂(IV, IV), no cathodic peak current corresponding to the Ru₂(IV,IV)/ Ru₂(III,IV) redox couple was observed, suggesting the occurrence of a structural change upon the second oxidation and a high reactivity for the Ru₂(IV,V) species. The variations in the redox potentials of complexes having different bridging ligands indicated that the electron-donating nature is in the order of {µ- $O_2CO_1^{2-7f} > {\mu - O_2SO_2}^{2-} > {\mu - O_2CCH_3}^{-7c} \approx {\mu - O_2COH}^{-7e} > {\mu - O_2COH}^{-7e}$ O_2NO ^{-.7c} The λ_{max} values of the $Ru_2(v,v)$ complexes were related to the order of their redox potentials, which reflect the SOMO, namely, the $\{Ru_2(\mu-O)_2\}$ core energies. The cyclic voltammogram of [III,IV_1H]²⁺ exhibited irreversible oxidation waves at 1.07 and 2.02 V and reversible reduction waves at -0.29 and -0.98 V. The reversible waves were assigned to the Ru₂(III,IV)/ $Ru_2(\Pi,\Pi)$ (-0.29 V) and $Ru_2(\Pi,\Pi)/Ru_2(\Pi,\Pi)$ couples (-0.98 V), which were confirmed by HDV analysis (Fig. S25-S27†), and the reversibility was confirmed using different scan rates (Fig. S22, S23 and Table S5†). The irreversible waves were attributed to the $Ru_2(rv,rv)/Ru_2(rr,rv)$ (1.07 V) and $Ru_2(rv,v)/Ru_2(rv,rv)$ couples (2.02 V). The oxidation wave at 2.02 V is identical to those of [III,IV] and $[\mathbf{IV},\mathbf{IV}]^{2+}$, indicating that deprotonation from the $\{\mathrm{Ru}_2(\mu-0)\}$ (μ-OH)} core was accompanied by the one-electron oxidation of [III,IV_1H]²⁺. The redox waves of [III,IV_1H]²⁺ having the $\{Ru_2(\mu - \mu)\}$ O)(μ -OH)} core were more positive than those of [III,IV]⁺ having the $\{Ru_2(\mu-O)_2\}$ core.

The comproportionation constants (K_c) of the mixed-valence state in [III,IV]⁺ and [III,IV_1H]²⁺ were calculated to be 3.1×10^{22} and 4.5×10^{22} , respectively, according to the Nernst equation (eqn (S4)–(S6)†), corresponding to Class III of the Robin–Day classification.³⁶ The evaluation on the mixed-valency revealed a stable mixed-valent state of Ru^{3.5}–Ru^{3.5},

which is described as $Ru_2(III,IV)$ for the clarity of redox reactions. These values were in line with those of other $Ru_2(III,IV)$ complexes in our Rbpma ligand system (R = ethyl or benzyl; $10^{22}-10^{24}$).^{7d}

[III,III_2H]²⁺ exhibited two irreversible oxidation waves at 1.14 and 2.05 V, corresponding to the Ru₂(IV,IV)/Ru₂(III,IV) and Ru₂(III,IV)/Ru₂(III,III) couples, respectively. The reversible redox waves at -0.30 and -0.98 V were assigned to the one-electron reduction couples of Ru₂(III,III)/Ru₂(II,III) and Ru₂(II,III)/Ru₂(II,III), respectively (Fig. S31–S33†). These redox behaviours indicated that upon the one-electron oxidation of [III,III_2H]²⁺, one-deprotonation occurs to form [III,IV_1H]²⁺, and the second oxidation involves further one-deprotonation to eventually form [IV,IV]²⁺. The reversible reduction processes (Fig. S28, S29 and Table S6†) indicated that both {Ru₂^{II,III}(μ-OH)₂} and {Ru₂^{II,II}(μ-OH)₂} cores were formed without structural changes under the measurement conditions. [III,III(HCO₃)_2H]³⁺ exhibited similar redox behaviours with potentials at 1.25 ($E_{\rm pa}$), -0.22 ($E_{\rm 1/2}$) and -0.88 V ($E_{\rm 1/2}$), which were slightly more positive than those of [III,III_2H]²⁺.

Electrochemical and spectroscopic behaviours in aqueous solutions

The electrochemical behaviours of [III,IV_1H]²⁺ were investigated in aqueous solution (Fig. S34†) and a Pourbaix diagram was obtained by plotting the redox potentials as a function of pH between pH 0.5 and 8.3 (Fig. 6). The UV-vis-NIR spectrum measured at pH 10.95 showed much lower intensity compared to those obtained at lower pH (Fig. S13†), indicating that [III, IV_1H]²⁺ underwent a structural change, and was thus excluded from the Pourbaix diagram. In all pH regions, one reversible oxidation wave, corresponding to the Ru₂(IV,IV)/Ru₂(III,IV) couple, and multiple reversible or irreversible reduction waves were observed.

The protonation/deprotonation equilibria and electrode reactions are summarised in eqn (1)–(15). In the $Ru_2(III,IV)$ region, the pK_a between $[III,IV_1H]^{2+}$ and $[III,IV]^+$ was at pH 2.1 (eqn (1)), consistent with the spectroscopic investigations (Fig. S13†). A pK_a was not observed in the $Ru_2(IV,IV)$ region

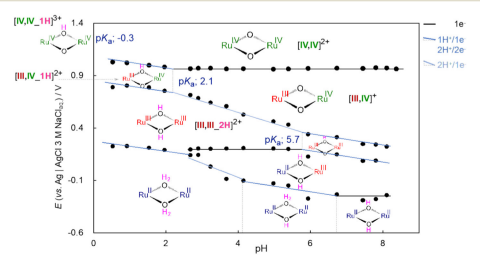


Fig. 6 Pourbaix diagram of sulfato-bridged complexes (the $\{\mu-O_2SO_2\}^{2-}$ moiety is omitted for clarity in the figure).

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between pH 0.5 and 8.3, indicating the presence of only [IV, $[V]^{2^+}$. The spectroscopic data suggested that the p K_a between $\{[IV,IV_1H]^{3+}\}$ and $[IV,IV]^{2+}$ was as low as -0.3 (eqn (2) and Fig. S12†), although it was out of scope of the accurate pH evaluation with the glass electrode, therefore, {[IV,IV_1H]³⁺} is only present under strongly acidic conditions. [IV,IV]²⁺ seemed to be unstable above pH 5, as evidenced by the UV-vis-NIR spectra above pH 5.03, likely changing into [III,IV 1H]²⁺ (Fig. S12 and S13†). In the Ru₂(III,III) region, the isolated species $[III,III_2H]^{2+}$ is present below pH 5.7 (p K_a), consistent with the spectroscopic results (Fig. S14†). The correponding pK_a of [III,III(HCO₃)_2H]³⁺was observed at 4.2 (Fig. S15,† eqn (3), eqn (S1) \dagger). Above the p K_a and below approximately pH 8, a $Ru_2(III,III)$ species having the $\{Ru_2(\mu-O)(\mu-OH)\}$ core, $\{[III,$ **III_1H**]⁺}, was present, but it started to decompose at pH 11 (grey line in Fig. S14†). The unstable character of {[III,III_1H]⁺} in strongly basic aqueous media was similar to that of {{III,III (CO₃)]. Finally, in the Ru₂(II,II) region, complexes having the $\{Ru_2(\mu-OH)_2\}, \{Ru_2(\mu-O)(\mu-OH)\}, \text{ or } \{Ru_2(\mu-O)_2\} \text{ cores would be}$ present owing to the two pK_as approximately at 4.1 and 6.6, although they were not isolated.

Protonation/deprotonation equilibria:

$$\begin{split} &\left\{Ru_2^{III,IV}(\mu\text{-O})(\mu\text{-OH})\right\}^{4+}\left([\textbf{III},\textbf{IV}_\textbf{1H}]^{\textit{m}+}\right) \\ &\rightleftharpoons \left\{Ru_2^{III,IV}(\mu\text{-O})_2\right\}^{3+}\left([\textbf{III},\textbf{IV}]^{(\textit{m}-1)+}\right) + H^+ \end{split} \tag{1}$$

$$\begin{aligned} & \left\{ Ru_2^{\text{IV,IV}}(\mu\text{-O})(\mu\text{-OH}) \right\}^{5+} \left(\left[\textbf{IV}, \textbf{IV} \right]^{2+} \right) \\ & \rightleftharpoons \left\{ Ru_2^{\text{IV,IV}}(\mu\text{-O})_2 \right\}^{4+} \left(\left\{ \left[\textbf{IV}, \textbf{IV}\text{-1H} \right]^{3+} \right\} \right) + H^+ \end{aligned} \tag{2}$$

$$\begin{split} & \left\{ Ru_2^{\text{III},\text{III}} (\mu\text{-OH})_2 \right\}^{4+} (\left[\textbf{III},\textbf{III}_2\textbf{H}\right]^{2+}) \\ & \rightleftharpoons \left\{ Ru_2^{\text{III},\text{III}} (\mu\text{-O}) (\mu\text{-OH}) \right\}^{3+} \left(\left\{ \left[\textbf{III},\textbf{III}_1\textbf{H}\right]^+ \right\} \right) + \textbf{H}^+ \quad (3) \end{split}$$

$$\left\{Ru_{2}^{II,II}\big(\text{μ-OH}_{2}\big)_{2}\right\}^{4+} \rightleftharpoons \left\{Ru_{2}^{II,II}\big(\text{μ-OH}_{2}\big)\big(\text{μ-OH}\big)\right\}^{3+} + H^{+}$$

$${Ru_2^{II,II}(\mu\text{-OH}_2)(\mu\text{-OH})}^{3+} \rightleftharpoons {Ru_2^{II,II}(\mu\text{-OH})_2}^{2+} + H^+$$
 (5)

Redox reactions:

between Ru₂(III,IV) and Ru₂(IV,IV) above pH 2.1:

$$\left\{ Ru_{2}^{III,IV}(\mu\text{-O})_{2}\right\} ^{3+}\rightleftharpoons\left\{ Ru_{2}^{IV,IV}(\mu\text{-O})_{2}\right\} ^{4+}+e^{-} \tag{6}$$

below pH 2.1:

$${Ru_2^{III,IV}(\mu\text{-O})(\mu\text{-OH})}^{4+} \rightleftharpoons {Ru_2^{IV,IV}(\mu\text{-O})_2}^{4+} + e^- + H^+$$
 (2)

between Ru2(III,IV) and Ru2(III,III) above pH 5.7:

$$\left\{Ru_2^{III,IV}(\mu\text{-O})_2\right\}^{3+} + e^- + H^+ \rightleftharpoons \left\{Ru_2^{III,III}(\mu\text{-O})(\mu\text{-OH})\right\}^{3+} \quad (8$$

between pH 2.1 and 5.7:

$$\left\{Ru_{2}^{III,IV}(\text{μ-O})_{2}\right\}^{3+}+e^{-}+2H^{+}\rightleftharpoons\left\{Ru_{2}^{III,III}(\text{μ-OH})_{2}\right\}^{4+} \qquad (9)$$

$$\left\{Ru_{2}^{III,IV}(\text{μ-O)}(\text{μ-OH})\right\}^{4+} + e^{-} + H^{+} \rightleftharpoons \left\{Ru_{2}^{III,III}(\text{μ-OH})_{2}\right\}^{4+} \quad (10)$$

between Ru₂(III,III) and Ru₂(II,III)

above pH 5.7:

$$\{Ru_{2}^{III,III}(\mu\text{-O})(\mu\text{-OH})\}^{3+} + e^{-} + H^{+} \rightleftharpoons \{Ru_{2}^{II,III}(\mu\text{-OH})_{2}\}^{3+} \tag{11}$$

between pH 2.0 and 5.7:

$${Ru_2^{III,III}(\mu\text{-OH})_2}^{4+} + e^- \rightleftharpoons {Ru_2^{II,III}(\mu\text{-OH})_2}^{3+}$$
 (12)

between Ru₂(III,III) and Ru₂(II,II) below pH 2.0:

$$\{Ru_{2}^{III,III}(\mu\text{-OH})_{2}\}^{4+} + 2e^{-} + 2H^{+} \rightleftharpoons \{Ru_{2}^{II,II}(\mu\text{-OH}_{2})_{2}\}^{3+} \ \ (13)$$

between Ru₂(II,III) and Ru₂(II,II) above pH 6.6:

$${Ru_2^{II,III}(\mu\text{-OH})_2}^{3+} + e^- \rightleftharpoons {Ru_2^{II,II}(\mu\text{-OH})_2}^{2+}$$
 (14)

between pH 4.1 and 6.6:

$$\left\{Ru_{2}^{II,III}(\mu\text{-OH})_{2}\right\}^{3+} + e^{-} + H^{+} \rightleftharpoons \left\{Ru_{2}^{II,II}(\mu\text{-OH})(\mu\text{-OH}_{2})\right\}^{3+} \tag{15}$$

The $pK_a(s)$ in every oxidation state was(were) observed at a higher pH for the sulfato-bridged complex than that of the acetato-bridged complex in the Pourbaix diagram, indicating the higher basicity of the $(\mu$ -O(H))₂ core of the present system: $pK_a = 1.5$ between $[III,IV(CH_3CO_2)_1H]^{3+}$ and [III,IV] $(CH_3CO_2)^{2+}$ (eqn (1)); 5.6 between {[III,III(CH_3CO_2)_2H]^{3+}} and $[III,III(CH_3CO_2)_1H]^{2+}$ (eqn (3)); and 3.7 and 6.6 between $\{[II,II]$ $(CH_3CO_2)_4H^{3+}$ and $\{[II,II(CH_3CO_2)_3H^{2+}\}\ (eqn\ (4))\ and$ $\{[II,II(CH_3CO_2)_3H]^{2+}\}$ and $\{[II,II(CH_3CO_2)_2H]^+\}$ (eqn (5)), respectively. The p K_a s of singly protonated species in the $Ru_2(III,IV)$ state were 2.4, 2.1, and 1.5 for [III,IV(HCO₃)]²⁺, ^{7e} $[III,IV]^+$, and $[III,IV(CH_3CO_2)]^{2+}$, respectively, indicating that the order of increasing acidity is hydrogenearbonato > sulfato > acetato, which is not necessarily correlated with the electrostatic nature of the bridging ligand. For the Ru₂(III,III) state, p $K_a = 5.7$ and 4.2 for [III,III_2H]²⁺ and [III,III(HCO₃)_2H]³⁺, respectively, indicating that the basicity of the bridging oxygen is higher in [III,III(HCO₃)_2H]³⁺ owing to the weaker electron-donating hydrogenearbonato compared to sulfato.

Conclusions

(4)

A series of Ru(IV)-Ru(IV), Ru(III)-Ru(IV), and Ru(III)-Ru(III) complexes, in which the bidentate sulfato ligand bridged the doubly oxido- and/or hydroxido-bridged diruthenium core $\{Ru_2(\mu-O(H))_2\}$, were synthesised all using the same tridentate ligand, ebpma. The hydrogencarbonato-bridged Ru₂(III,III) complex was also synthesised. The electronic structures of these complexes were systematically investigated experimentally and theoretically as structural analogues of the reaction intermediates proposed in iron-containing metalloenzymes such as sMMO.

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The single-crystal X-ray analyses of all complexes revealed the structural differences among the $\{Ru_2(\mu-O)_2\}$, $\{Ru_2(\mu-O)(\mu-O)\}$ OH)} and {Ru₂(µ-OH)₂} cores, which depended on the oxidation state of the Ru centres and the number of protons on the oxido ligand(s). Interesting paramagnetic characteristics were revealed for [IV,IV]2+ and [III,III_2H]2+ in addition to those of $[\mathbf{III}, \mathbf{IV}]^+$ and $[\mathbf{III}, \mathbf{IV}_{\mathbf{1}}\mathbf{H}]^{2+}$ with $S = \frac{1}{2}$ through magnetic measurements and NMR spectroscopy, which could be rationalised by the quantum mechanical calculations that the openshell singlet state was more stabilized for [IV,IV]2+ and a possible spin transition from a closed-shell singlet state to triplet state could occur for [III,III_2H]²⁺. Notably, all the complexes isolated could be confirmed in the Pourbaix diagram. The basicity of the bridging oxido ligand(s) of the {Ru₂(μ-O(H))₂} core were discussed in each electronic state; the pK_a values of $\{[IV,IV_1H]^{3+}\}, [III,IV_1H]^{2+}, \text{ and } [III,III_2H]^{2+} \text{ were } -0.3, 2.1,$ and 5.7, respectively. Under more strongly acidic conditions, the sulfato ligand may be protonated and utilised as a proton accepting site in future applications. In addition to the Ru₂(IV, IV), $Ru_2(III,IV)$, and $Ru_2(III,III)$ states, the $Ru_2(II,III)$ or $Ru_2(II,III)$ state can be accessed using the ebpma ligand system. These oxidation states are analogous to the intermediate or transient species expected in the sMMO cycle.

Experimental

Materials

All chemical reagents were used as purchased without further purification. RuCl₃·nH₂O (Ru content, 40.11 wt%) was purchased from Furuya Metal Inc. Co., Ltd Japan or Tanaka Holdings Co., Ltd Japan. The ebpma ligand and fac-[Ru^{III}Cl₃(ebpma)] were prepared according to the literature procedures.^{7a} $[{Ru^{III,IV}Cl_2(ebpma)}_2(\mu-O)]ClO_4 \cdot 0.75(CH_3)_2CO$ was prepared according to the procedure reported in ref. 7f and the synthesis of $[{Ru^{III,IV}(ebpma)}_2(\mu-O_2(\mu-O_2NO))](ClO_4)_2 \cdot 1.5H_2O$ ([III,IV(NO₃)](ClO₄)₂·1.5H₂O) followed the procedure of the corresponding PF₆-salt reported in ref. 7c.

Measurements

The $\mu_{\rm eff}$ was measured at 296–298 K using a Sherwood Scientific Magway MSB MK1 balance (UK) with Hg[Co(SCN)4] as the calibrant. X-band ESR spectra were recorded using a JEOL (Japan) JESFA300 ESR spectrometer at 93 and 298 K. ¹H and ¹³C NMR spectra were measured using a JEOL JNM-ECA (500 MHz) spectrometer. IR spectra were recorded using a Shimadzu (Japan) IR Affinity-1 spectrophotometer for samples prepared as KBr disks. Elemental analyses were carried out with a PerkinElmer (US) 2400-II or at the Institute of Physical and Chemical Research (RIKEN, Japan). Fast atom bombardment mass spectrometry (FAB MS) was conducted using a JEOL JMS-SX700 V with an m-nitrobenzyl alcohol matrix. UVvis-NIR spectra were measured using a Shimadzu (Japan) UV-3600 Plus spectrophotometer in acetonitrile, methanol, or aqueous solutions using a quartz cell with a 1 cm path length. Cyclic voltammetry, Osteryoung square wave voltammetry, and

differential pulse voltammetry were performed in acetonitrile containing 0.1 M TBAP (Nakarai Tesque Ltd, Japan) as a supporting electrolyte with a glassy carbon disk working electrode $(\phi = 1.0 \text{ or } 1.6 \text{ mm})$, Ag|0.01 M AgNO₃(CH₃CN) reference electrode, and platinum counter electrode using an ALS Model 630E electrochemical analyser (BAS Inc., Japan.). At the end of each measurement, ferrocene ([Fe^{III}Cp₂]⁺/[Fe^{II}Cp₂] (Cp = cyclopentadienyl), 0.074 V in CH₃CN vs. Ag | 0.01 M AgNO₃(CH₃CN)) was added as the internal standard to scale the redox potentials. For the measurements in 0.1 M NaOH-0.1 M H₃PO₄ aqueous buffer solutions, a glassy carbon working electrode (ϕ = 1.0 mm), Ag | 3.0 M AgCl reference electrode, and platinum wire counter electrode were used. Hydrodynamic voltammetry (HDV) was carried out with a rotating platinum disk or a rotating glassy carbon disk working electrode (ϕ = 3.0 mm) and a BAS RDE-2 rotating electrode.

Theoretical calculations

The computational calculations were performed with (U) B3LYP-D3BJ³⁷ as a functional and LANL08(f)(Ru), aug-cc-pVDZ (N, O, S), and cc-pVDZ(H, C) as basis sets after the self-consistent field (SCF) calculations, using the Gaussian 16 program³⁸ for $[IV,IV]^{2+}$, $[III,IV]^{+}$, $[III,IV_1H]^{2+}$, and $[III,III_2H]^{2+}$. The Grimme dispersion correction (D3BJ) was employed in addition to B3LYP as a better solution in our system in comparison with the ωB97XD and the sole B3LYP functions. The initial electronic structures were estimated by the fragment method, dividing the complex cations into four moieties; the $\{Ru_2(\mu-O(H))_2\}$ core, $\mu-O_2SO_2^{2-}$ ligand, and two ebpma ligands. The structures were optimized with full-optimization (no symmetric restrains) and the polarizable continuum model (PCM) was applied, in which acetonitrile was selected as a representative solvent used in experiments reported herein. For the assignment of electron transition spectra, time-dependent (TD)-DFT calculations were performed in similar methods. For [III,III(HCO₃)_2H]³⁺, costless simpler calculations were performed with B3LYP/LANL2DZ levels.

Single-crystal X-ray diffraction analysis

Single crystals suitable for X-ray diffraction analysis were obtained at 298 K by slow evaporation of a water/acetone reaction solution, with pH 2.7 for [IV,IV](ClO₄)₂·H₂O, pH 5.4 for [III, IV]ClO₄·3.5H₂O, pH 2.3 for [III,IV_1H](ClO₄)₂·0.75H₂O, pH 1.9 for $[III,III_2H](ClO_4)_2 \cdot 1.5H_2O$, and pH 1.9 for $[III,III(HCO_3)_2H]$ (ClO₄)₃·0.25H₂O. The intensity data were collected on a Rigaku (Japan) Model Synergy-S system, using multilayer mirror monochromated $CuK\alpha$ radiation (1.54184 Å). All the calculations were performed using CrysAlis^{Pro} data collection and processing software (Rigaku Corporation (2015)). Structures were solved using direct methods, expanded using Fourier techniques, and refined using full-matrix least-squares techniques on F2 using the SHELXT³⁹ implementation in Olex2 1.5.⁴⁰ For the disorder modelling of perchlorate and solvent water, parts 1 and 2 were used, restraints (DFIX, DANG, RIGU, SIMU, and ISOR) were applied where necessary to make more sensible models, and constraint **Paper**

(EADP) was utilized only if the split atoms were located too close to each other. The structures are represented by ORTEP-3 for windows⁴¹ (CCDC 2355836, 2355837, 2355838, 2355839 and 2355840† for [IV,IV](ClO₄)₂·H₂O, [III,IV]ClO₄·3.5H₂O, [III,IV_1H] (ClO₄)₂·0.75H₂O, [III,III_2H](ClO₄)₂·1.5H₂O, and [III,III(HCO₃) 2H](ClO₄)₃·0.25H₂O, respectively).

Synthesis of [{Ru^{IV,IV}(ebpma)}₂(μ-O)₂(μ-O₂SO₂)](ClO₄)₂·0.5H₂O ([IV,IV](ClO₄)₂·0.5H₂O). [{Ru^{III,IV}(ebpma)(μ -O)₂(μ -O₂NO)](ClO₄)₂· $1.5H_2O$ ([III,IV(NO₃)](ClO₄)₂· $1.5H_2O$; 75 mg, 0.077 mmol) was dissolved in water/acetone (7 mL, v/v = 4/3) and H₂SO₄ and $(NH_4)_4[Ce^{IV}(SO_4)_4]\cdot 2H_2O$ (50 mg, 0.079 mmol), which had been dissolved into a least amount of water, were added (pH 2.28). An emerald-green precipitate was obtained immediately and NaClO₄ (27 mg, 0.22 mmol) was added. The volume of the reaction mixture was reduced by a slow evaporation at 50 °C. The precipitate was collected by filtration and dried in vacuo. Yield: 44 mg, 58%. Elemental analysis/%; Calcd for C₂₈H₃₅N₆O_{14.5}SCl₂Ru₂; C, 33.87 H, 3.53 N. 8.47. Found C, 33.57 H, 3.40 N, 8.27. FAB MS(+) (m/z); 885.5 (30%, {M - ClO_4^-)⁺). ¹H NMR (500 MHz, acetonitrile- d_3) was measured at 298 K and only broadened signals with paramagnetic shifts were observed (Fig. S4†). ¹³C NMR (500 MHz, acetonitrile-d₃) measured at 298 K showed no specific signals. $\mu_{\rm eff}$ = 0.94 $\mu_{\rm B}$ (298 K). IR spectrum (KBr method); $\nu(SO_4)$: 1148, 931 cm⁻¹; $\nu(\text{ClO}_4)$: 1089 cm⁻¹ (Fig. S3†).

Synthesis of [{Ru^{III,IV}(ebpma)}₂(μ-O₂SO₂)]ClO₄·0.25 (CH₃)₂CO·4H₂O ([III,IV]ClO₄·0.25(CH₃)₂CO·4H₂O). [{Ru^{III,IV} (ebpma)}{(μ-O)₂(μ-O₂NO)](ClO₄)₂·1.5H₂O ([III,IV(NO₃)]-(ClO₄)₂·1.5H₂O 101 mg, 0.106 mmol) was dissolved in acetone-water (11.5 mL, $\nu/\nu = 13/10$) and Na₂SO₄ (149 mg, 1.05 mmol) dissolved in 3.5 mL water was added (pH 5.32). The volume of the solution was reduced to *ca.* 1 mL and the yellow-brown crystals were collected by filtration and dried *in vacuo*. Yield: 80 mg, 78%. Elemental analysis/%; calcd for C_{28.75}H_{43.5}N₆O_{14.25}SClRu₂; C, 35.57 H, 4.48 N, 8.66. Found C, 35.54 H, 4.21 N, 8.53. FAB MS(+) (m/z); 785.3 (40%, {M − ClO₄⁻}⁺). $\mu_{\rm eff} = 1.79\mu_{\rm B}$ (296 K). IR spectrum (KBr method); ν (SO₄): 1128, 949 cm⁻¹; ν (ClO₄): 1088.9 cm⁻¹ (Fig. S3†).

Synthesis of [$\{Ru^{III,IV}(ebpma)\}_2(\mu-O)(\mu-OH)(\mu-O_2SO_2)]$ -(ClO_4)₂·3H₂O ([III,IV_1H](ClO_4)₂·3H₂O). [$\{Ru^{III,IV}(ebpma)\}(\mu-O)_2(\mu-O_2NO)\}$](ClO_4)₂·1.5H₂O ([III,IV(NO₃)]²⁺; 70 mg, 0.072 mmol) was dissolved in water/acetone mixed solution (13 mL, $\nu/\nu = 8/5$) and H₂SO₄ (0.070 mL) was added (pH 0.98). After the colour of the reaction mixture changed from ochre to deep green, NaClO₄ was added as a precipitant and the volume of the solution reduced by slow evaporation at 55 °C until deep-green crystals were obtained. The crystals were collected by filtration and dried *in vacuo*. Yield: 51 mg, 68%. Elemental analysis/%; calcd for $C_{28}H_{41}N_6O_{17}SCl_2Ru_2$; C, 32.37 H, 3.94 N, 8.09. Found C, 32.20 H, 3.55 N, 8.10. FAB MS(+) (m/z); 784.3 (86%, $\{M-H^+\}^+$). $\mu_{eff} = 1.69\mu_B$ (298 K). IR spectrum (KBr method); $\nu(SO_4)$: 1162.1 (weak), 949 cm⁻¹; $\nu(ClO_4)$: 1089 cm⁻¹ (Fig. S3†).

Synthesis of [{Ru^{III,III}(ebpma)}₂(μ -OH)₂(μ -O₂SO₂)](ClO₄)₂·3H₂O ([III,III_2H](ClO₄)₂·3H₂O). [{Ru^{III,IV}(ebpma)}₂(μ -O)₂(μ -O₂NO)]ClO₄·1.5H₂O (100 mg, 0.10 mmol) was dissolved in water/acetone (6.5 mL, ν/ν = 3/10) and H₂SO₄ was added (pH

1.28). The solution was degassed using argon and Sn^{II}Cl₂·2H₂O (14 mg, 0.062 mmol) dissolved in 3.5 mL water was added. After 1.5 h, when the colour of the reaction mixture changed from ochre to yellow green, NaClO₄ (18 mg, 0.15 mmol) was added, and the volume of the solution was reduced by slow evaporation at 50 °C until a pale yellow-green precipitate was obtained. The solid was collected by filtration and dried in vacuo (80 mg). The solid (64 mg) was purified in water/acetone (8 mL, v/v = 5/3), where the pH was adjusted to pH 1.37 by adding H₂SO₄. Yield: 51 mg, 49%. Elemental analysis/%; calcd for C₂₈H₄₂N₆O₁₇SCl₂Ru₂; C, 32.34 H, 4.04 N. 8.08. Found C, 32.14 H, 3.64 N, 7.97. FAB MS(+) (m/z); 785.0 $(100\%, \{M - 2ClO_4^- - H^+\}^+)$. $\mu_{eff} = 1.15\mu_B (298 \text{ K})$. ¹H NMR (500 MHz, acetonitrile- d_3) δ /ppm; 16.60 (s, 2H OH), 9.55 (d, J = 5.5 Hz, 2H py-6 or 6'), 8.03 (t, J = 7.3 Hz, 2H py-6' or 6), 7.60 (t, J = 7.3 Hz, 4 H py-4,4'), 7.34 (d, J = 8.0 Hz, 2 H, py-3' or 3),7.25 (t, J = 8.0 Hz, 2H, py-5' or 5), 6.38 (d, 2H, J = 5.5 Hz, 2H,py-3 or 3'), 6.14 (t, J = 6.5 Hz, 2H, py-5 or 5'), 5.59 (d, J = 16.0 Hz, 2H, $-CH_2$ -(methylene)), 5.47 (td, J = 58.7, 16.0 Hz, 2H, $-CH_2$ -(methylene)), 5.35 (d, J = 17.0 Hz, 2H, $-CH_2$ -(methylene)), 4.95 (d, J = 17.0 Hz, 2H, $-CH_2$ -(methylene)), 4.52 (sext, J= 6.6 Hz, 2H, $-CH_2$ -(ethyl)), 4.00 (sext, J = 6.9 Hz, 2H, $-CH_2$ -(ethyl)), 1.70 (t, J = 7.5 Hz, 6H, $-CH_3$ (ethyl)) (Fig. S6†). ¹³C NMR (500 MHz, acetonitrile- d_3) δ /ppm; 162.71, 160.35, 153.33, 151.18, 143.34, 141.00, 128.25, 125.77, 124.93, 124.70, 72.04, 67.03, 65.59, 10.68 (Fig. S7†). IR spectrum (KBr method); $\nu(SO_4)$: 1138, 954 cm⁻¹; $\nu(ClO_4)$: 1114, 1088 cm⁻¹ (Fig. S3†).

Synthesis of $[{Ru^{III,III}(ebpma)}_2(\mu-OH)_2(\mu-O_2COH)](ClO_4)_3\cdot H_2O$ ([III,III(HCO₃)_2H](ClO₄)₃·H₂O). Na[{Ru^{III,IV}(ebpma)}₂(μ -O)₂(μ -O₂CO)]₂(ClO₄)₃·17H₂O (50 mg, 0.047 mmol) was dissolved in water/acetone (6.5 mL, v/v = 3/10) and perchloric acid was added (pH 1.9). The solution was degassed using argon, and Sn^{II}Cl₂·2H₂O (13 mg, 0.058 mmol) was added. The solution was stirred at room temperature for 1 d, and after filtration and degassing, the volume of the yellow solution was reduced to ca. 0.5 mL using a rotary evaporator. NaClO₄ (29 mg, 0.24 mmol) was added to the solution and the mixture was cooled at 4 °C in the fridge for 2 h. The pale green precipitate was collected by filtration, washed with cooled water, and dried in vacuo. Yield: 21 mg, 42%. Elemental analysis/%; calcd for C₂₉H₃₉N₆O₁₈Cl₃Ru₂; C, 32.60 H, 3.65 N, 7.87. Found C, 32.66 H, 3.80 N, 7.96. ¹H NMR (500 MHz, acetonitrile- d_3) δ / ppm; 15.75 (s, 2H OH), 9.32 (d, J = 6.0 Hz, 2H py-6'), 8.08 (t, J =6.8 Hz, 2H, py-6), 7.72 (d, J = 8.0 Hz, 2H py-4'), 7.61 (t, J = 6.3 Hz, 2H, py-4), 7.38-7.32 (m, 4H, py-3,3'), 6.42 (d, J = 6.0 Hz, 2H, py-5'), 6.24 (t, J = 6.3 Hz, 2H, py-5), 5.54 (q, J = 38.0, 16.0 Hz, 4H, $-CH_2$ -(methylene)), 5.40 (d, J = 18.5 Hz, 2H, $-CH_2$ -(methylene)), 5.18 (d, J = 17.0 Hz, 2H, $-CH_2$ -(methylene)), 4.45 (sext, J = 7.0 Hz, 2H, $-CH_2$ -(ethyl)), 4.02 (sext, J = 7.1 Hz, 2H, $-CH_2$ -(ethyl)), 1.75 (t, J = 7.5 Hz, 6H, $-CH_3$ (ethyl)) (Fig. S6†). ¹³C NMR (500 MHz, acetonitrile- d_3) δ/ppm; 163.93, 161.75, 158.38, 151.88, 149.68, 139.80, 124.19, 123.55, 117.02, 70.31, 65.35, 65.34, 9.22 (Fig. S7†). IR spectrum (KBr method); $\nu(COO)$: 1449 cm⁻¹; $\nu(ClO_4)$: 1143, 1116, 1088 cm⁻¹ (Fig. S3†).

T. M.-S.: conceptualization, supervision, data curation, visualization, writing – original draft preparation, and writing – reviewing and editing; Y. K.: data curation, visualization, and writing – reviewing and editing. S. M. and T. N.: data curation and visualization. H. N.: supervision and writing – reviewing and editing.

Data availability

The data supporting this article have been included as part of the ESI. \dagger

Crystallographic data for $[[IV,IV](ClO_4)_2 \cdot H_2O$, [III,IV] $ClO_4 \cdot 3.5H_2O$, $[III,IV_1H](ClO_4)_2 \cdot 0.75H_2O$, $[III,III_2H]$ $(ClO_4)_2 \cdot 1.5H_2O$, and $[III,III(HCO_3)_2H](ClO_4)_3 \cdot 0.25H_2O]$ have been deposited at the CCDC under 2355836, 2355837, 2355838 2355839, and 2355840,† respectively, and can be obtained from [https://doi.org/10.1039/D4DT02833C].

Conflicts of interest

There are no conflicts to declare.

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