Heterotrimetallic RuMnMn Species on a Hydrotalcite Surface as Highly Efficient Heterogeneous Catalysts for Liquid-Phase Oxidation of Alcohols with Molecular Oxygen**

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The creation of a well-defined, active metal site on a solid surface not only opens up an avenue to materials that boost catalytic performance but also aids an understanding of the molecular basis of heterogeneous catalysis.^[1] Modifying the coordination sphere of a metal species with other metals as ligands, which is a basic approach in organometallic and bioinorganic chemistry,^[2] is also being applied to heterogeneous catalysis^[3] because of the potential of performing unique catalytic reactions based on cooperation between diverse metals within a regular arrangement. In addition, this allows local compositional modeling for the strong metal–support interactions (SMSI) seen in supported metal catalysts.^[4]

Hydrotalcite (HT), which is a layered, mixed hydroxide of Mg and Al,^[5] has received attention as a material for advanced heterogeneous catalysts^[6,7] because of the cation-exchange ability of the Brucite layer, the anion-exchange ability of the interlayer, its adjustable surface basicity, and adsorption capacity. Recently, we created a monomeric Ru^{IV}-OH species on the HT (Ru/HT) surface by adsorption which turned out to be an excellent heterogeneous catalyst for the one-pot synthesis of a-alkylated nitriles by a Rucatalyzed alcohol oxidation coupled with a basepromoted aldol reaction.^[7d] As part of our ongoing project on the functionalization of supported Ru catalysts to achieve environmentally friendly ("green") organic syntheses, we present a novel HTbound heterotrimetallic Ru^{IV}Mn^{IV}Mn^{IV} species that has been structurally characterized on the atomic scale by X-ray absorption spectroscopy.^[8] This material is an

excellent solid catalyst for liquid-phase alcohol oxidation under mild conditions. This protocol to create a supported mixed-metal species can provide catalytically active compounds that are uniform in composition and distribution on a solid surface by applying concepts from coordination chemis-

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try, and bridges the gap between homogeneous and heterogeneous catalysis. $\ensuremath{^{[9]}}$

We obtained well-defined, heterotrimetallic RuMnMn species on the HT surface (RuMn₂/HT) by immobilizing Mn cations onto Ru/HT. The Ru/HT was prepared by treating HT with an aqueous solution of RuCl₃·*n* H₂O.^[7d] The retention of the HT interlayer distance (3.0 Å), as shown by XRD (X-ray diffraction),^[10] indicated that both metal species are accommodated on the HT surface. The K-edge XANES (X-ray absorption near-edge structure) spectrum of the Ru and Mn of RuMn₂/HT reveals that the surface Ru and Mn cations are in the oxidation state + IV. In the Fourier transformation of the Ru K-edge k^3 -weighted EXAFS (extended X-ray absorption fine structure) spectrum of RuMn₂/HT (Figure 1b) a peak near 3.5 Å, corresponding to the contiguous Ru sites (Ru–O–Ru),^[11] was barely detected.

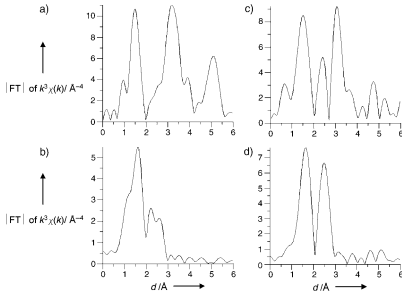


Figure 1. Fourier transformation (FT) of the k^3 -weighted K-edge EXAFS spectrum of a) RuO₂, b) RuMn₂/HT (Ru K-edge), c) β -MnO₂, and d) RuMn₂/HT (Mn K-edge). The phase shift was not corrected.

The coordination number (CN), distance (*R*), and Debye– Waller factor ($\Delta\sigma$) of the Ru–O, Ru–Mg, and Ru–Mn bonds, as estimated by a curve-fitting analysis,^[8] are listed in Table 1. The Ru^{IV} species of RuMn₂/HT are surrounded by six oxygen atoms with different bond distances. The shortest Ru–O bond was assigned to a Ru^{IV}–OH moiety.^[12] Two bonds of Ru–Mg and Ru–Mn shells, with CNs of 0.9 and 1.9, respectively, prove that the Ru^{IV} species is in the vicinity of the Mg cation within the Brucite-like sheets and two Mn cations on the HT surface. Furthermore, a Mn–Mn shell with a distance of 2.32 Å and CN of 1.1 shows the formation of dimeric Mn^{IV} cation species.^[13]

The above results show a Ru^{IV}Mn^{IV}Mn^{IV} trimetallic species on the HT surface, as shown in Figure 2, in which dimeric Mn–Mn species are connected to a single Ru^{IV} cation through OH groups and water. To our knowledge, this is the first report of the preparation of heterotrimetallic species

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Table 1	Curve-fitting	results for	K-edge	FXAFS	of Ru	and Mn
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Sample	Shell	CN ^[a]	R [Å] ^[b]	$\Delta\sigma [{ m \AA}^2]^{[c]}$
RuMn ₂ /HT	Ru-K			
_,	Ru-O(1)	2.1	1.97	-0.0079
	Ru-O(2)	2.1	2.07	-0.0045
	Ru-O(3)	1.1	1.83	-0.0088
	Ru-O(4)	0.6	2.61	-0.0087
	Ru-Mg	0.9	3.19	-0.1023
	Ru-Mn	1.9	3.28	0.0499
	Mn-K			
	Mn–O(1)	2.6	1.85	-0.0030
	Mn–O(2)	3.1	1.98	-0.0060
	Mn-Mn	1.1	2.32	-0.0017
RuO ₂ ^[d]	Ru-O(1)	2	1.94	_
	Ru-O(2)	4	1.98	_
β -MnO ₂ ^[e]	Mn-O	6	1.89	_
	Mn-Mn(1)	2	2.87	-
	Mn-Mn(2)	8	3.42	-

[a] Coordination number. [b] Interatomic distance. [c] $\Delta\sigma$ is the difference between the Debye–Waller factor of the sample and that of the reference sample. [d] Taken from the crystallographic data.^[11] [e] Taken from the crystallographic data.

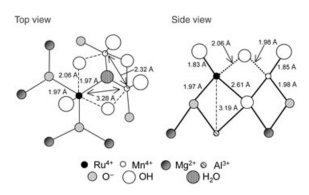


Figure 2. Proposed structure of the heterotrimetallic RuMnMn species on HT.

consisting of metal cations on a support involving metal oxide and metal hydroxides,^[14] whereas the preparation of supported heterobimetallic or metal alloy species have been reported.^[3] Adjusting the basicity of the hydroxy groups around the Ru^{IV} cation, produced by reaction of the surface OH groups of HT with RuCl₃ species,^[7d] brought about selective immobilization of Mn cations in the vicinity of the Ru species to give the unique Ru^{IV}Mn^{IV}Mn^{IV} sites.

The catalytic ability of the RuMnMn species was explored in the oxidation of alcohol with O_2 at atmospheric pressure (Table 2).^[15,16] The trimetallic RuMn₂/HT shows a higher catalytic activity than Ru/HT,^[7d] Ru/Al₂O₃,^[15c,e] and RuO₂,^[15a] which are typical heterogeneous Ru catalysts for the oxidation of benzyl alcohol (1).^[17] Benzyl alcohol was oxidized to benzal-

Table 2: Oxidation of benzyl alcohol (1) in the presence of various Ru catalysts under 1 atm $O_2.^{[a]}$

Entry	Catalyst	Conv. [%] ^[b]	Yield [%] ^[b]
1	RuMn ₂ /HT	100	99
2	reuse 1 ^[c]	100	99
3	reuse 2 ^[c]	100	98
4	reuse 3 ^[c]	100	99
5	Ru/HT	66	66
6	Ru/Al ₂ O ₃ ^[d]	56	55
7	$RuO_2 \cdot nH_2O$	5	4
8	Mn ₂ /HT ^[e]	0	0
9	HT	0	0

[a] Reaction conditions: catalyst (3 mol% Ru), benzyl alcohol (1 mmol), toluene (5 mL), 60 °C, 1 atm O_2 , 1 h. [b] Conversions and yields were determined by GC analysis using an internal standard method based on benzyl alcohol. [c] Catalyst recovered and reused. [d] Prepared by the reported procedure.^[15c] [e] 6 mol% Mn.

dehyde (2) quantitatively within 1 h in the presence of the $RuMn_2/HT$ catalyst.^[18]

The initial turnover frequency based on Ru for RuMn₂/HT (140 h⁻¹) is almost five times larger than that for Ru/HT. On a 10-mmol scale oxidation of **1** in the presence of 0.1 mol-% Ru, the turnover number based on Ru reached 840 at 90 °C. Moreover, the high catalytic ability of RuMn₂/HT was demonstrated by the quantitative oxidation of **1** within 10 h, even at 40 °C.

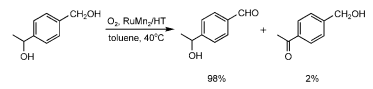
As displayed in Table 3, the RuMn₂/HT catalyst selectively oxidizes a wide variety of alcohols. Primary and secondary benzylic alcohols are converted into the corresponding carbonyl compounds in high yield (entries 1–3). The catalysis is also chemoselective; RuMn₂/HT preferentially oxidizes primary over secondary hydroxy groups, as shown by the selective oxidation of 1-[(4'-hydroxymethyl)phenyl]ethanol to 1-[(4'-formyl)phenyl]ethanol in 98 % yield (Scheme 1).

In the case of cyclopropyl(phenyl)methanol, the hydroxy group was oxidized without cleavage of the cyclopropyl ring

Table 3:	Oxidation	of various	alcohols	catalyzed by	RuMn ₂ /HT	in the presence of O_2 . ^[a]
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Entry	Alcohol	Product	<i>t</i> [h]	Conv. [%]	Yield [%] ^[b]
1	Benzyl alcohol (1)	Benzaldehyde (2)	1	100	99
2	1-Phenylethanol	Acetophenone	3.5	100	92
3	Benzhydrol	Benzophenone	2	94	93
4	Cyclopropyl (phenyl) methanol	Cyclopropyl phenyl ketone	6	100	99
5	Cinnamyl alcohol	Cinnamaldehyde	1.5	100	97
6	(2-Hydroxymethyl)thiophene	2-Thiophenecarboxaldehyde	2	100	89
7	2-Aminobenzyl alcohol	2-Aminobenzaldehyde	21	100	100

[a] Reaction conditions: catalyst (2 mol% Ru), alcohol (1 mmol), toluene (5 mL), 60 °C, O₂ flow. [b] Yields were calculated based on alcohols.



Scheme 1. Selective oxidation of primary hydroxy groups by RuMn_2/HT catalyst.

(entry 4). Oxidation of cinnamyl alcohol proceeded smoothly without affecting the carbon–carbon double bond (entry 5). Furthermore, RuMn₂/HT effectively catalyzes the oxidation of (2-hydroxymethyl)thiophene, a heteroaromatic alcohol, to 2-thiophenecarboxaldehyde, in high yield (entry 6), and the oxidation of 2-aminobenzyl alcohol to 2-aminobenzaldehyde, quantitatively (entry 7), in contrast to the homogeneous Pd and Ru complexes.^[16]

The spent RuMn₂/HT catalyst could be readily separated from the reaction mixture by filtration. The EXAFS spectrum of the recovered RuMn₂/HT catalyst confirmed retention of the original RuMnMn structure, and ICP (inductively coupled plasma) analysis of the filtrate indicated no leaching of Ru and Mn species during the oxidation. The RuMn₂/HT catalyst could be reused while maintaining the same high catalytic activity and selectivity (Table 2, entries 2–4). When the catalyst was removed at about 50% conversion of the alcohol, no further oxidation was detected in the filtrate after 3 h under 1 atm O₂ which shows that the present alcohol oxidations proceed at the interface between the catalyst surface and the liquid phase.

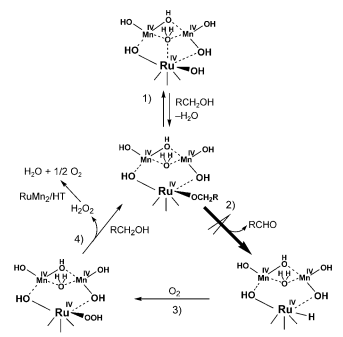
To the best of our knowledge, RuMn₂/HT is the most effective, environmentally benign Ru catalyst for the oxidation of benzylic and aromatic allylic alcohols in the liquid phase^[15,16c,d,19] with the advantages of 1) high catalytic activity, even with O₂ at 1 atm as the sole oxidant, 2) wide applicability to alcohols, including those containing heteroatoms, 3) simple preparation and work-up procedures, and 4) reusability as a solid catalyst.

The Mn cations in the RuMnMn species evidently play a pivotal role in improving the Ru-catalyzed alcohol oxidation since Mn₂/HT does not catalyze the reaction (Table 2, entry 8). We have proposed a catalytic cycle for the alcohol oxidation (Scheme 2)^[20] that proceeds via a ruthenium alkoxide intermediate, which undergoes β -hydrogen elimination to produce the carbonyl compound and a ruthenium hydride species, as observed by IR spectroscopy.^[10] Reaction of this hydride with O₂ and subsequent ligand-exchange with the alcohol completes the catalytic cycle.^[15bc]

A rate equation based on a Michaelis–Menten-type $model^{[21]}$ for this proposed mechanism [Eq. (1)] agrees well with the kinetic data.

$$\mathbf{Ru}-\mathbf{OH}+\mathbf{RCH}_{2}\mathbf{OH}\underset{k_{1}}{\overset{k_{1}}{\longrightarrow}}[\mathbf{Ru}-\mathbf{OCH}_{2}\mathbf{R}]\overset{k_{2}}{\longrightarrow}\mathbf{Ru}-\mathbf{H}+\mathbf{RCHO}$$
(1)

For the RuMn₂/HT-catalyzed oxidation of **1**, $K_{\rm M}$ and k_2 were calculated to be 2.96 mM and 0.047 s⁻¹, respectively, at 60 °C. The rate constant (k_2) of β -hydrogen elimination from the ruthenium alkoxide intermediate is therefore almost twice that for the Mn-free Ru/HT. The β -hydrogen elimination is considered as the rate-determining step in the overall alcohol oxidation from the primary kinetic isotope effect in the competitive oxidation of **1** and C₆D₅CD₂OH (4.2).^[15b,e] Thus, the Mn cations in the heterotrimetallic sites facilitate β hydrogen elimination from the ruthenium alkoxide intermediate. Removing the water molecule that binds the Ru and Mn cations improves the situation of the Ru species during β hydrogen elimination by, for example, producing a coordinately unsaturated Ru site.^[16c]



Scheme 2. A proposed mechanism for the oxidation of a primary alcohol by trimetallic RuMnMn sites.

For the secondary alcohol 1-phenylethanol, $K_{\rm M}$ and k_2 were found to be 133 mM and 0.026 s⁻¹, respectively, at 60 °C. Importantly, the $K_{\rm M}$ value is significantly greater than that for the oxidation of **1**, which is reflected in the preferential oxidation of primary hydroxy groups by this species.^[22] The formation of metal alkoxide intermediates of primary alcohols is favored over secondary alcohols in the ligand-exchange step.^[23]

In summary, a robust heterotrimetallic Ru^{IV}Mn^{IV}Mn^{IV} species coordinated to a hydrotalcite surface as a macroligand facilitates the highly efficient aerobic oxidation of alcohols. Such cooperative action among high-valence metal cations in a heterometallic species on a solid surface provides a unique protocol for the preparation of functionalized heterogeneous catalysts for environmentally benign organic syntheses.^[24] Ongoing efforts are focused on creating high-valence Ru cation species, that is, Ru^{VI} and Ru^{VII}, based on the redox interaction within a heterometallic site for highly efficient epoxidation and dihydroxylation reactions of alkenes with molecular oxygen.

Experimental Section

 $Ru/HT^{[7d]}$ was treated with an aqueous solution of $MnCl_2\cdot 4\,H_2O$ to afford $RuMn_2/HT.^{[10]}$ Elemental analysis (%) calcd for $Ru_{0.071}Mn_{0.142}/Mg_6Al_2(OH)_{16}CO_3$: Ru 1.03, Mn 1.12, Mg 21.0, Al, 8.14; found: Ru 1.05, Mn 1.20, Mg 21.4, Al 7.9.

The X-ray absorption spectra were recorded at the BL01B1 beamline in SPring-8 of JASRI, Japan (2003B0944-UXa-np and 2004A489-NXa-np) and at the BL-10B beamline of PF at KEK, Japan (2001G143 and 2002G102). The data were reduced using computer

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systems at the Data Processing Center of Kyoto University according to a previously reported procedure.^[25]

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