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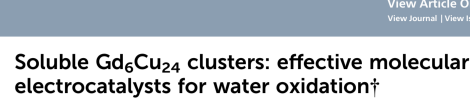
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Introduction

Hydrogen production by water splitting is a promising solution for future energy needs;¹ however, water oxidation is ratelimiting, primarily due to its four-electron transfer process with slow reaction kinetics.² Many researchers have been inspired by natural photosynthesis to develop catalytic systems to resolve this rate-limiting issue.3 Several studies have developed molecular water oxidation catalysts.⁴ In particular, significant efforts have focused on mimicking the structure and catalytic properties of the natural oxygen-evolving complex (OEC), a unique heterometallic-oxide Mn₄CaO₅-cluster, in photosystem II (PS II) of plants, algae, and cyanobacteria.5 Although several multinuclear clusters simulating OEC structures have been reported,6 their catalytic performance is still far from that of natural OEC, with the mechanism behind the O-O bond formation still not yet conclusively known.7 Catalysts usually should easily attain four electrons and contain effective synergistic catalytic structural motifs to achieve a similar catalytic activity.8 Previous studies on structural motifs have mainly focused on the Mn-O-Mn-H2O bimetallic structure.9,10 Recent studies have shown that the redox-inert Ca²⁺ ion in Mn₄CaO₅ may play an important role in maintaining the structural



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The water oxidation half reaction in water splitting for hydrogen production is extremely rate-limiting. This study reports the synthesis of two heterometallic clusters (Gd_6Cu_{24} -IM and Gd_6Cu_{24} -AC) for application as efficient water oxidation catalysts. Interestingly, the maximum turnover frequency of Gd₆Cu₂₄-IM in an NaAc solution of a weak acid (pH 6) was 319 s⁻¹. The trimetallic catalytic site, $H_2O-Gd^{III}Cu_2^{II}$ -H₂O, underwent two consecutive two-electron two-proton coupled transfer processes to form highvalent Gd^{III}-O-O-Cu^{III} intermediates. Furthermore, the O-O bond was formed via intramolecular interactions between the Cu^{III} and Gd^{III} centers. The results of this study revealed that synergistic catalytic water oxidation between polymetallic sites can be an effective strategy for regulating O-O bond formation.

> integrity of clusters and supporting ligand sites for water molecules or in assisting proton transfer.¹¹ However, the poor stability of Ca²⁺-containing Mn₄CaO₄ clusters in polar solvents has hindered their development.¹² Therefore, developing efficient and stable cluster catalysts for water oxidation remains challenging.

> Several studies have recently reported that replacing Ca²⁺ ions with lanthanide ions can stabilize the cluster structure in performance simulations.^{6c,13} Furthermore, lanthanide ions have high coordination numbers, rich geometries, and

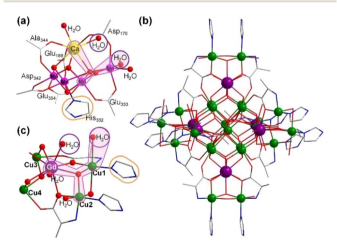


Fig. 1 (a) Structure of the Mn₄CaO₅ cluster. (b) Core of the Gd₆Cu₂₄-IM. (c) Asymmetric unit (GdCu₄) of Gd₆Cu₂₄-IM. H₂O-Ca-O-Mn₂-H₂O and H₂O-Gd-O-Cu₂-H₂O motifs are marked in pink and water molecules are marked by a purple circle. Color codes: Gd, purple; Cu, green; Mn, pink; Ca, yellow; C, gray; N, blue; O, red. All H atoms were omitted

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structural flexibility.¹⁴ This study reports the synthesis of two high-nuclearity clusters— $[Gd_6Cu_{24}(IM)_{12}(L-Al)_{12}(\mu_3-OH)_{30}(\mu_2-OH)_6(CO_3)(H_2O)_{24}]\cdot(ClO_4)_{16}\cdot(H_2O)_6$ $(Gd_6Cu_{24}-IM)$ and $[Gd_6-Cu_{24}(Ac)_6(L-Al)_{12}(\mu_3-OH)_{30}(\mu_2-OH)_6Cl(H_2O)_{24}]\cdot(ClO_4)_{11}\cdot(H_2O)_{19}$ $(Gd_6Cu_{24}-AC)$ (IM = imidazole, L-Al = L-alanine, Ac = acetate)— possessing the same metal core frames (Fig. 1 and S1†). Similar to H_2O-Ca-O-Mn_2-H_2O in OEC, the trimetallic structural motif of H_2O-Gd-O-Cu_2-H_2O in Gd_6Cu_{24} may be a potentially efficient synergistic catalytic site for water oxidation.

Results and discussion

Clusters Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC were obtained by the reaction of Gd(ClO₄)₃, Cu(ClO₄)₂·6H₂O, and L-alanine with imidazole and NaAc, respectively. The asymmetric structural unit [GdCu₄] of Gd₆Cu₂₄ can be considered as two cornerdeficient [GdCu₂O₃] and [GdCu₂O₄] cubes linked together by sharing a Gd³⁺ ion (Fig. 1c). Six [GdCu₄] units were connected by four μ_3 -O²⁻, resulting in an octahedral conformational arrangement. The metal core of Gd₆Cu₂₄ displayed an octahedral inner core of Gd₆Cu₁₂ connected to six outer Cu₂ units. Each [GdCu₄] unit contained five metal ions, in which Gd, Cu₁, and Cu2 were bound to water molecules. Furthermore, the Gd₆Cu₂₄-IM cluster was protected by 12 amino acids and 12 imidazole ligands, highly resembling the peripheral ligands of the OEC. Furthermore, each cluster core unit of Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC had 16 and 11 positive charges in addition to 12 sets of water-binding sites, which can promote water oxidation catalysis.

The catalytic properties were characterized in an aqueous solution, and the stability was verified in the high-resolution electrospray ionization mass spectrometry (HRESI-MS) spectra (Fig. S6[†]). For Gd₆Cu₂₄-IM, the peak at 1978.73 can be attributed $[Gd_6Cu_{24}(IM)_{12}(L-Al)_{12}(\mu_3-OH)_{28}(O)_8(CO_3)_1(H_2O)_{24}(ClO_4)_5]^{3+}$ to (Fig. S6a^{\dagger}); for Gd₆Cu₂₄-AC, these peaks are observed and can be attributed $[Gd_6Cu_{24}(Ac)_x(L-Al)_{10}(\mu_3-OH)_{28}(O)_8(H_2O)_2$ to $(ClO_4)_{8-x}^{\uparrow^{4+}}$ (X = 0-6) (Fig. S6b[†]). The redox properties of Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC were analyzed by cyclic voltammetry (CV) and differential pulse voltammetry (DPV) in an NaAc/HAc buffer solution (0.5 M, pH = 6) (Fig. 2, S7 and S8^{\dagger}). The redox current of Gd_6Cu_{24} -IM at $E_{1/2}$ (0.15 V, all potentials were compared to the normal hydrogen electrode, NHE) depended linearly on the square root of the scan rate and corresponded to diffusion control. Moreover, the difference between the cathodic and anodic peak potentials of 30 mV (ΔE_p , Fig. 2a, b and S7[†]) indicated a two-electron quasi-reversible process.15 However, the reduction current of Gd₆Cu₂₄-AC (Fig. 2c, d and S8[†]) at 0.13 V was not reversible for the reaction, although it linearly depended on the square root of the scan rate owing to the dissociation of the charged acetate during the reduction process.¹⁶ By performing DPV at a low scan rate, the reduction current corresponded to the splitting of the oxidation current peak into two peaks attributed to two electron transfers as well as the relocation of the acetate ligand. As determining the anodic shoulder peak potential (E_p) of approximately 1.35 V is difficult, the half-peak potential $E_{p/2}$ *versus* $\ln(v)$ was fitted using the Laviron equation (Fig. 3a, b, S9 and S10[†]).¹⁷ The fitted slope was equal to $RT/(1 - \alpha)nF$ for the

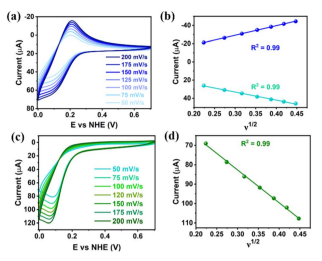


Fig. 2 (a) CV of 0.25 mM Gd₆Cu₂₄-IM in 0.5 M NaAc/HAc (pH = 6) buffer solution using a GC electrode with different scan rates (0.05–0.2 V s⁻¹) at 0–0.7 V. (b) Plots of i_p (μ A) vs. $\nu^{1/2}$ (V^{1/2} s^{-1/2}) for Gd₆Cu₂₄-IM. (c) CV of 0.25 mM Gd₆Cu₂₄-AC in 0.5 M NaAc/HAc (pH = 6) buffer solution using the GC electrode with different scan rates (0.05–0.2 V s⁻¹) at 0–0.7 V. (d) Plots of i_p (μ A) vs. $\nu^{1/2}$ (V^{1/2} s^{-1/2}) for Gd₆Cu₂₄-AC.

anodic shoulder peak, where α is the transfer coefficient (0.5) and n was calculated to be 2. These results indicated that the two peripheral Cu²⁺ ions in the [GdCu₄] unit could be considered equivalent groups for the single-electron reaction, *i.e.*, the quasi-reversible process near 0.15 V can be attributed to the reaction pair of two-electron reduction of **Gd₆Cu^{II}₂₄**/Gd₆Cu^{II}₂₂Cu^I₂, while the anodic shoulder peak around 1.35 V can be attributed to the two-electron oxidation of **Gd₆Cu^{II}₂₄**/Gd₆Cu^{II}₂₂Cu^{II}₂.

The DPVs of **Gd**₆**Cu**₂₄**-IM** and **Gd**₆**Cu**₂₄**-AC** transformed significantly by changing the solvent from superdry acetonitrile

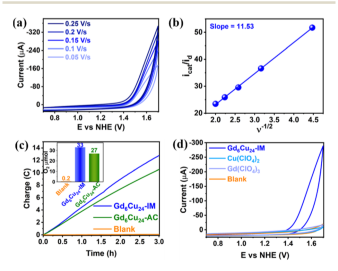


Fig. 3 (a) CVs of Gd₆Cu₂₄-IM (0.25 mM) in NaAc/HAc (0.5 M, pH = 6) buffer solution at different scan rates: $50-250 \text{ mV s}^{-1}$ at 0.7-1.7 V. (b) Plot of i_{cat}/i_d versus $\nu^{-1/2}$ at 1.7 V. (c) CPE data showing charge versus time for a 20 mL solution containing NaAc/HAc buffer solution (0.5 M, pH = 6) and 0.25 mM Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC. (d) CVs of Gd₆Cu₂₄-IM (0.25 mM), Gd(ClO₄)₃ (1.5 mM), and Cu(ClO₄)₂ (6 mM).

to an aqueous NaAc/HAc buffer solution (pH = 6; Fig. S11 and $S12^{\dagger}$).^{6e,13d} In acetonitrile, the oxidation peak at 1.72 V can be attributed to the two-electron oxidation of the Gd₆Cu^{II}₂₄/Gd₆Cu^{II}₂₂Cu^{III} reaction pair. In the NaAc/HAc buffer, the anodic current was significantly enhanced after 1.35 V, thus indicating the occurrence of aqueous oxidation. A linear relationship was observed in the plot of the catalytic peak current versus Gd₆Cu₂₄-IM concentration (Fig. S13[†]), thus indicating that the rate law for Gd₆Cu₂₄-IM catalyzed water oxidation can be expressed in terms of a pseudo-first-order rate reaction (eqn (1)-(3), ESI[†]).¹⁸ The catalytic peak currents were directly proportional to the square root of the scan rate, thus indicating a diffusion-controlled catalytic redox process.19 Linear scan voltammograms (LSV) of Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC were recorded on an indium tin oxide (ITO) electrode. The significantly enhanced catalytic currents indicated water oxidation (Fig. S14[†]). The overpotentials for Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC required to reach 1 mA cm⁻² were 598 and 689 mV, respectively. In comparison to previous reports (Table S2[†]), the overpotentials represent a relatively lower level. The difference in the overpotentials may be attributed to the different peripheral coordination groups between Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC (imidazole and acetate, respectively), and the conjugation effect of imidazole ligands may help to reduce the reaction energy barrier.²⁰ The apparent rate constants (k_{cat}) , called turnover frequency (TOF), for Gd₆Cu₂₄-IM and Gd₆Cu₂₄-AC at 1.7 V were 319 and 169 s⁻¹, respectively (Fig. 3b and S9[†]),²¹ much higher than those of previously reported mono- and di-nuclear catalysts $(0.4-100 \text{ s}^{-1}).^{22}$

A gas-tight double-compartment cell with the cathode and anode separated by a Nafion membrane was utilized to verify the release of oxygen. Gd₆Cu₂₄-IM (0.25 mM) was subjected to controlled potential electrolysis (CPE) at 1.70 V in an NaAc/HAc buffer solution (0.5 M, pH = 6) using an ITO electrode (1.00 cm²; Fig. 3c). Electrolysis was performed for 3 h and oxygen production was determined by gas chromatography, where approximately 33 µmol of O2 were formed with a faradaic efficiency of 94%. The ratio of H_2 to O_2 was measured (2:1). For Gd₆Cu₂₄-IM, neither the blank nor the inorganic salt catalytic tests showed a significant catalytic current. The reproducibility of the 50 times cycle scan, the almost identical UV-Vis spectra, and the CV curves before and after 3 h of controlled potential electrolysis (Fig. 3d and S15-S18†) indicated that water oxidation occurred under homogeneous catalytic conditions in addition to the high activity and stability of Gd₆Cu₂₄-IM.²³

The DPVs of Gd_6Cu_{24} clusters in a mixture of superdry CH_3CN and H_2O at different concentrations were recorded to elucidate the water oxidation mechanism. The results (Fig. 4a) revealed that the DPV curves of Gd_6Cu_{24} -AC changed significantly with the addition of water, where the oxidation peak current increased dramatically and the oxidation onset potential shifted positively, indicating that the coordination of water can reduce the oxidation onset potential. The DPV curves of Gd_6Cu_{24} -IM changed in a pattern similar to that of Gd_6Cu_{24} -AC, although its two oxidation peaks did not show significant splitting (Fig. S19[†]). Changes in the oxidation onset potential

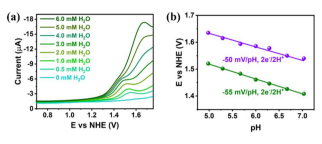


Fig. 4 (a) DPVs of Gd_6Cu_{24} -AC (0.25 mM) in CH₃CN with Et₄NClO₄ (0.1 M) before and after addition of different amounts of H₂O. (b) Pourbaix diagram of Gd_6Cu_{24} -AC (0.25 mM) in NaAc/HAc (0.5 M, pH 4.98–7.03) buffer solution. E_p values are cited rather than $E_{1/2}$ due to chemical irreversibility.

were observed by changing the pH of the NaAc/HAc buffer solution, indicating that the oxidation process involved a proton-coupled electron transfer (PCET). The Pourbaix diagram revealed that (Fig. 4b) in an NaAc/HAc buffered aqueous solution (0.5 M, pH 4.98–7), two consecutive oxidation peaks of **Gd₆Cu₂₄-AC** exhibited Nernstian responses at approximately -55 and -50 mV pH⁻¹, consistent with the $2e^-/2H^+$ PCET (-59 mV pH⁻¹, Nernstian ideal),²⁴ while the responses of **Gd₆Cu₂₄-IM** were -54 and -57 mV pH⁻¹ (Fig. S22†). In addition, the catalytic processes of **Gd₆Cu₂₄-IM** and **Gd₆Cu₂₄-AC** showed low solvent kinetic isotope effects of 1.23 and 1.25, respectively, (KIE = k_{H_2O}/k_{D_2O} , Fig. S23†),²⁵ significantly different from the hydrophilic nucleophilic attack mechanism of mononuclear catalysts but consistent with the intramolecular O–O bond formation route for Cu₃ catalysts.²⁶

Based on the typical mechanisms reported for natural OECcatalyzed water oxidation,12b,27 it was inferred that the two molecules of water (W1 and W2) that coordinated on the heterometallic centers Cu^{II} and Gd^{III} can be used as the oxygen source for O-O bond formation (Fig. S24⁺). W1 and W2 had a distance of 2.81 Å, close to the distance between the two water molecules coordinated on Ca^{II} and Mn^{III} in the OEC (3.26 Å).²⁸ Based on the experimental results of previous studies and previously reported catalytic mechanisms for water oxidation by multinuclear molecular catalysts,^{6e,13d} a plausible and reasonable mechanism for water oxidation catalyzed by Gd₆Cu₂₄ was proposed (Fig. 5). Under electrolytic catalytic conditions, the trimetallic reaction site is oxidized from the initial $Gd^{III}(OH_2)$ $Cu_2^{III}(OH_2)$ species 1 through a $2e^{-}/2H^{+}$ PCET process to give Gd^{III}(OH)Cu^{III}(OH) species 2, which can be further oxidized by the $2e^{-}/2H^{+}$ PCET process to $Gd^{III}(O^{\bullet})Cu_{2}^{III}(O^{\bullet})$ species 3. The redox potentials of these two steps were 1.35 and 1.55 V, respectively. Species 3 forms species 4, containing O-O bonds through intramolecular interactions, while species 4 releases O2 and completes the cycle by coordinating with two water molecules to form starting cluster 1. The M₃-O₂ intermediate is stabilized by the trimetallic site of Gd^{III}Cu₂^{III}, which may facilitate the formation of O-O bonds. This pathway differs from the common mechanism of O-O bond formation by the nucleophilic water attack on the highly oxidized state of M^{n+} =O. The formation of the highly oxidized state of Cu^{IV}=O and ligand hydrocarbon oxidation are also avoided.29

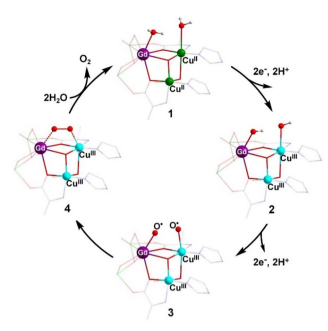


Fig. 5 Proposed catalytic mechanism for the water oxidation based on Gd_6Cu_{24} -IM. Color codes: Gd, purple; Cu^{II}, green; Cu^{III}, turquoise; H, white; C, gray; N, blue; O, red.

Conclusions

In conclusion, this study reported an efficient catalytic water oxidation using water-soluble Gd_6Cu_{24} clusters in aqueous solutions of weak acids. The trimetallic catalytic site synergistically catalyzed water oxidation by promoting O–O bond formation, while simple amino acid ligands provided both water solubility and stability, which assisted the catalytic process. Interestingly, the TOFs of Gd_6Cu_{24} -IM and Gd_6Cu_{24} -AC at 1.7 V reached 319 and 169 s⁻¹, respectively. This work offers a possibility to understand the synergistic effect of multiple metals in the water oxidation mechanism. Furthermore, Gd_6Cu_{24} has a tunable molecular structure and metal sites, which is beneficial for designing highly active and stable catalysts in the future.

Data availability

The crystal data and structure of the clusters, experimental details and experimental data for this article are available in the ESI.†

Author contributions

J.-N. C., Z.-H. P. and X.-J. K. conceived and designed the research; J.-N. C. and Z.-H. P. synthesized and characterized the compounds. X.-J. K., J.-N. C., Z.-H. P., Q.-H. Q., C. W., L.-S. L., and L.-S. Z. analyzed the data. J.-N. C., Z.-H. P. and X.-J. K. wrote the manuscript with contributions from all authors. All authors analyzed the data and commented on the manuscript. J.-N. C. and Z.-H. P. contributed equally to this work.

Conflicts of interest

The authors declare no competing financial interest.

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