Stable iridium dinuclear heterogeneous catalysts supported on metal-oxide substrate for solar water oxidation

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Atomically dispersed catalysts refer to substrate-supported heterogeneous catalysts featuring one or a few active metal atoms that are separated from one another. They represent an important class of materials ranging from single-atom catalysts (SACs) and nanoparticles (NPs). While SACs and NPs have been extensively reported, catalysts featuring a few atoms with well-defined structures are poorly studied. The difficulty in synthesizing such structures has been a critical challenge. Here we report a facile photochemical method that produces catalytic centers consisting of two Ir metal cations, bridged by O and stably bound to a support. Direct evidence unambiguously supporting the dinuclear nature of the catalysts anchored on α-Fe₂O₃ is obtained by aberration-corrected scanning transmission electron microscopy (AC-STEM). Experimental and computational results further reveal that the threefold hollow binding sites on the OH-terminated surface of α-Fe₂O₃ anchor the catalysts to provide outstanding stability against detachment or aggregation. The resulting catalysts exhibit high activities toward H₂O photooxidation.

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Atomically dispersed catalysts have recently gained significant attention for their advantages of high atomic efficiency, activity, and selectivity (1, 2). Different from bulk heterogeneous catalysts, the atomically dispersed ones feature well-defined structural moieties at the atomic level (3). This feature opens a window to understand mechanisms that underpin heterogeneous catalysts in a similar fashion to how homogeneous catalysts are studied (4). It also presents opportunities to tailor the functionalities of heterogeneous catalysts with unprecedented controls (5). New and unique reaction pathways have been discovered on these catalysts. For instance, when used for CO oxidation, a Pt single-atom catalyst (SAC) has been shown far less susceptible to poisoning by CO than Pt nanoparticles (2). Within this context, research on SACs has thrived, where significant progress in synthesis, characterization, and mechanistic understanding has been made. Notwithstanding, these prior efforts primarily focused on reactions that may require only a single metal-active site, such as CO oxidation (6) and water-gas shift (4). Atomically dispersed catalysts featuring more than one metal-active site are rare in the literature. Consequently, reactions dependent on multiple metal-active sites without organic ligands remain poorly studied. Pioneering works by Iwasawa and coworkers (7) and Gates and coworkers (8, 9) have demonstrated several systems featuring multiple metal atoms per active site for hydrogenolysis and hydrogenation reactions. Another class of catalysts featuring such structures is found in dinuclear Cu or Ni catalyst for methane oxidation reactions (10, 11). However, two key challenges remain in this field: (i) to prepare the dinuclear catalyst with a high yield by a facile and general approach and (ii) to achieve a direct observation to unambiguously support the dinuclear nature. To fill in this important knowledge gap, here we report the synthesis of dinuclear heterogeneous catalysts (DHCs) consisting of two Ir atoms per catalytic site stably bound to a support. The structure was enabled by a heterogenization method followed by a room-temperature photochemical ligand removal treatment. The resulting DHCs exhibit outstanding stability against detachment and aggregation yet are highly active toward water oxidation. Water oxidation was chosen for this present work because, as an essential step in natural and artificial photosynthesis, it has provided a material platform to study reactions that would require more than one active site.

Significance

While research on single-atom catalysts (SACs) is arguably mature, rare work has been done on atomically dispersed catalyst featuring two atoms. We synthesized Ir dinuclear heterogeneous catalyst in a facile photochemical way. It exhibits outstanding stability and high activity toward water oxidation. The significance of this work can also be appreciated from the catalysis perspective. A grand challenge in heterogeneous catalysis is how to understand the detailed mechanisms at the molecular level, because the most active heterogeneous catalysts are often poorly defined in their atomic structures. Our finding is built upon recent advances aimed at studying SACs but takes a crucial step forward. It provides a material platform to study reactions that would require more than one active site.


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been a topic of intense research (12). A large number of catalysts have been extensively studied, and their performance has been benchmarked (13). Despite these successes, gaps in the understanding of catalytic water oxidation exist (14). For instance, the mechanisms of heterogeneous catalysts for water oxidation, especially at the molecular level, are poorly understood (15). An important reason has been that the high-performance heterogeneous catalysts are often poorly defined in their structures, especially at the molecular and atomic levels. Despite recent successes in this area (16), the complex interactions between water-oxidation catalysts and light absorbers for integrated solar-to-fuel conversion are still poorly understood (17). Controlled synthesis of high-performance heterogeneous catalysts with well-defined structures and integration with light absorbers are therefore of great importance. Taking an important step aimed at addressing these issues, here we report a facile method for direct synthesis of heterogeneous catalysts consisting of two Ir atoms per catalytic site on the α-Fe$_2$O$_3$ substrate. The resulting catalysts preserve the atomic arrangements of the active two metal centers (Ir-O-Ir) but present two key features. First, the Ir DHCs are tightly bound to the supporting substrate (α-Fe$_2$O$_3$) and show outstanding stability against aggregation or detachment. Second, the catalysts are highly active toward water oxidation even without the organic ligands, which are critical to the functionality of the homogeneous analog (18). Significantly better performance than Ir SACs or Ir NPs was measured on Ir DHCs.

**Results**

**Synthesis Strategy for Ir DHCs.** Building upon our previous successes in immobilizing homogeneous water oxidation catalysts (Ir homodimer) (19), we designed the synthesis of Ir DHCs as shown in Fig. 1. The key step added to this work is the photochemical treatment, which was implemented to remove the organic ligands. The characterization of the α-Fe$_2$O$_3$ substrate and the resulting catalysts at each stage are presented in SI Appendix, Figs. S1–S3. In addition to structure results to be presented below, X-ray photoelectron spectroscopy (XPS) (SI Appendix, Fig. S2) and electron energy-loss spectroscopy (EELS) (SI Appendix, Fig. S3) clearly support that there is no N signal after the photochemical treatment. This provides strong evidence that the pyridyl ligands in the precursor have been removed.

**Direct Structure Characterization of Ir DHCs.** The most direct evidence that proves we have obtained a dinuclear structure of Ir atoms that are not in direct contact but connected by an oxygen bridge is from the high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) data (Fig. 2 A and B and SI Appendix, Figs. S4–S11). The bright two-point features in Fig. 2 A were identified as Ir DHCs, and the lighter gray points in Fig. 2 B were identified as Ir SACs, and the darker gray points in Fig. 2 C were ascribed to Fe atoms. Tilting experiments further supported that there were no large aggregates of Ir clusters. The lack of Ir SACs or clusters was proven by spectroscopic characterization to be discussed below. Taken as a whole, the unique two-atom arrangement and the lack of aggregated byproducts set the stage for detailed studies of DHCs as a new class of water-oxidation catalyst.

Our previous work has proven that the Ir–O–Ir arrangement within the Ir homodimer (Fig. 1) is key to its high activity toward water oxidation (18, 19). We therefore expect that Ir–O–Ir rather than Ir–Ir arrangement is preserved in the Ir DHC. Evidence from at least two complementary experimental techniques strongly supports our expectation. First, we performed in situ CO diffuse reflectance infrared Fourier transform (DRIFT) spectroscopic characterization, and the purpose was to use CO as a probe to study the Ir atomic arrangement within the DHC. While the dinuclear nature of the structure is unambiguously confirmed by the HAADF data, an alternative arrangement to the proposed one could feature Ir–Ir bonding. In the in situ CO DRIFT experiment, the two arrangements are expected to be readily distinguished by the characteristic absorption features at ∼1,850 cm$^{-1}$, which is indicative of the bridge adsorption of CO at the Ir–Ir site (20). Indeed, the broad peak at ∼1,850 cm$^{-1}$ was observed for Ir NPs containing a known Ir–Ir bonding (Fig. 2 D) but absent from Ir DHCs (Fig. 2 H). Additionally, a strong dipole–dipole coupling effect between adjacent CO probe molecules was evident in the control sample with Ir aggregates but was absent in Ir DHCs, further confirming the atomic dispersity of the latter. When the oxygen bridge of Ir DHC was broken to yield Ir SAC, we observed a redshift of the peaks characteristic of dicarbonyl species (SI Appendix, Fig. S13). Second, X-ray absorption spectroscopy data were acquired to reveal the coordination environment of the Ir center. We see from SI Appendix, Figs. S14–S16 and Table S1 that the first shell coordination of Ir in DHCs is dominated by Ir–O (21), in excellent agreement with the proposed bonding of Ir–O–Ir but not Ir–Ir. Taken together, the microscopic and spectroscopic characterization unambiguously confirmed that we have obtained a dinuclear Ir catalyst that features discrete Ir–O–Ir rather than Ir–Ir structures without organic ligands. Here we further highlight the uniqueness of the synthesis method. Our previous experiments show that other treatments such as thermal annealing (19) or electrochemical decomposition (22) lead to obvious Ir aggregation. Only by photochemical treatments could we achieve the DHC structure in the high yield (>80%).

**Determination of Binding Sites by Density-Functional Theory and HAADF Simulation.** The strong metal–support interaction is crucial to maintaining the catalytic structure under catalytic operation conditions for many systems (4). How SACs can be anchored onto supporting substrates by defect sites has been demonstrated on reducible metal oxides (23). Nevertheless, it would be a significant challenge to rely on the randomly distributed defect sites as the binding sites for DHCs that feature well-defined and uniform M–O–M arrangements. To understand how our Ir DHCs bind to the α-Fe$_2$O$_3$ support, we carried out density-functional theory (DFT) calculations. Our strategy was to construct hydroxyl-terminated α-Fe$_2$O$_3$ (001) surfaces, as shown in Fig. 1. Synthesis procedure of Ir DHC. It starts with the immobilization of molecular Ir catalysts, followed by photochemical removal of organic ligands. The binding details of H$_2$O and OH groups are proposed.
Direct structural characterization of Ir DHCs represents the linescan to Ir DHCs. The dotted line between the two stars in B represents the linescan for HAADF intensity linescan data reproduced the experimental data in terms of the Ir atomic location (Fig. 3). Importantly, the model built here allowed us to simulate the expected HAADF image under the experimental condition (Fig. 3E), which is in excellent agreement with the experimental data in terms of the Ir atomic location (Fig. 3A). Similarly, the simulated intensity linescan data reproduced the experimental ones as well, illustrating the accuracy of the interatomic distance in Ir–O–Ir site (Fig. 3B and F). We note that the data presented here represent a typical example of one facet of α-Fe₂O₃ (001), which provides a basis for more comprehensive future studies of other facets.

Catalytic Characterization of Ir DHCs. Photoelectrochemical (PEC) characterization was next carried out to study the catalytic activity of the Ir DHC following previously developed protocols. The Ir-based catalysts are known to be most active under acidic conditions (19). As such, it is desirable to study them at low pHs. The support (α-Fe₂O₃), however, would be dissolved by acids, leading to possible detachment of Ir that would undermine efforts to study its inherent catalytic activity and stability. For these reasons, we optimized the test conditions at a pH of 6.0. As shown in Fig. 4A, compared with control samples with only α-Fe₂O₃ support (bare in Fig. 4A), the addition of Ir catalysts improved the overall performance by shifting the polarization curves toward the cathodic direction. Previous thermodynamic and kinetic studies have revealed that such a shift may be due to either improved charge separation within the support or better charge-transfer kinetics or a combination of both (25, 26). Given that the surface coverage of Ir DHCs on α-Fe₂O₃ is sparse, we strongly believe faster kinetics is the true reason for such a shift. The understanding is consistent with our previous kinetic studies (25). The per-atom turnover frequency (TOF) of Ir DHCs at 1.23 V (vs. reversible hydrogen electrode, RHE) is 2.6 and 5 times higher than the corresponding Ir SACs (SI Appendix, Fig. S20) and Ir NPs (SI Appendix, Fig. S21), respectively, at the same potential. In addition, compared with similar systems that feature Ir-based water photoredox reactions (22, 25, 27–29), the Ir DHC on α-Fe₂O₃ features the higher normalized TOFs (SI Appendix, Table S3). That the sample with Ir DHCs exhibited the high activity attests to its functionality as an effective water-oxidation catalyst despite the fact that the organic ligands of the molecular precursor have been removed.

Proposed Catalytic Mechanism by DFT Calculations. Inspired by our previous studies on the Ir homogeneous catalysts (18) (SI Appendix, Fig. S22), we propose a mechanism that involves multiple proton-coupled electron transfer (PCET) steps by DFT calculations using the model built in Fig. 3C and D. Ir DHCs and homodimer molecules (18) share similar Ir⁴⁺(OH₂⁻)–O–Ir⁴⁺(OH₂⁻) structural motifs (x = 2 for Ir homodimer and x = 1 for Ir DHC). Thus, it is reasonable to assume that the water-oxidation mechanism of Ir DHCs is similar to their homogeneous molecular analogs, albeit in the absence of the organic supporting ligands. As shown in Fig. 5A, Ir DHCs are proposed to undergo three PCET processes to build enough oxidation potential to oxidize H₂O and produce Ir-OOH (steps E to F). The free-energy changes of the
three PCET steps are 1.06, 1.37, and 1.21 eV, respectively, at $U = 0$ V ($U$ is the applied potential; Fig. 5A). A substrate $\text{H}_2\text{O}$ molecule binds to the Ir DHC with a free-energy increase of 0.44 eV, which then undergoes nucleophilic attack to form hydrogen peroxide with a free-energy increase of 0.36 eV. The oxidation of Ir-OOH intermediate $F$ is easy to proceed through a PCET, requiring only 0.75 eV. The subsequent $\text{O}_2$ release is driven by the following substrate $\text{H}_2\text{O}$ binding. By comparing the free-energy changes under different applied potentials, an overpotential as low as 0.14 V is enough for the overall four-electron oxidation to be thermodynamically downhill. The calculated overpotential indicates that the Ir DHC is an efficient catalyst for water oxidation, consistent with the experimentally observed low onset potential of the Ir DHC/α-Fe$_2$O$_3$ ($V_{on} = 0.55$ V, corresponding to an applied potential of ~1.35 V by assuming a photovoltage of 0.8 V by α-Fe$_2$O$_3$) (26).

As an important control experiment, we carried out DFT calculations to compare the catalytic steps between the Ir DHC and SAC (Fig. 5 and SI Appendix, Figs. S23 and S24). The atomic structure of Ir SAC was built by placing one Ir atom in one threefold O site. The resting state of Ir SAC was identified as an Ir (IV) bound with an $\text{OH}^-$ and two $\text{H}_2\text{O}$ ligands (Fig. 5B). For water oxidation, Ir SAC first undergoes PCET processes twice, which is similar to the first two steps of water oxidation by Ir DHC. The key difference between the two catalysts lies in the next step. Whereas Ir DHC undergoes another PCET (steps $C$ to $D$ in Fig. 5A) before forming the $\text{O}–\text{O}$ bond, Ir SAC has to bind to a $\text{H}_2\text{O}$ substrate and form the $\text{O}–\text{O}$ bond first (steps $C$ to $E$ in Fig. 5B) before the third PCET. This is because direct oxidation of Ir SAC after the second PCET step would require too high an energy (more than 1.8 eV). By comparison, the availability of another Ir atom nearby in Ir DHC reduces the energy need to

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**Fig. 3.** Binding sites of Ir DHC on α-Fe$_2$O$_3$. Atomic resolution experimental (A) and simulated (E) HAADF-STEM micrographs of Ir DHC. The basis for the HAADF-STEM simulation is the atomic structure as shown in C and D. (Scale bars: A, 1 nm, E, 0.5 nm.) Proposed atomic structure of Ir DHC (support: α-Fe$_2$O$_3$). Top view along [001] zone axis (C) and side view (D). In the atomic model, green ball is Ir, gold ball is Fe, red ball is O, and white ball is H. The corresponding experimental (B) and simulated (F) HAADF line-intensity profiles show excellent match.

**Fig. 4.** Catalytic performance of Ir DHC in water photooxidation. (A) The polarization curve of the Ir DHC (pink) in comparison with three control samples, the Ir SAC (green), Ir NP (gray), and bare α-Fe$_2$O$_3$ support (black). The electrolyte was 0.1 M KNO$_3$ (pH 6.0), and the illumination intensity was 100 mW cm$^{-2}$ with AM 1.5 filters. (B) Chronoamperometry showing the stability of Ir DHC over 10 h with negligible decay. Evolution of Ir DHC on α-Fe$_2$O$_3$ during electron beam exposure for the following times: (C) 0 s, (D) 35 s, (E) 75 s. The electron dose received by the sample was ~100 eVÅ$^2$/s. (Scale bar: 2 nm.)
1.2 eV. We note that more accurate calculations of the energy need for catalytic steps would require the inclusion of the supporting substrate ($\alpha$-Fe$_2$O$_3$) and solvation effects, which is beyond the scope of the present work. Our results nonetheless highlight the benefits of the synergistic effects between a dinuclear catalytic site (Ir DHC) over a mononuclear site (Ir SAC). The understanding aligns with recent experiment observations that a dinuclear site may be more active toward water oxidation (30, 31). We emphasize that the well-defined structure of the Ir DHC is key to our understanding reported here, which permits the construction of mechanistic models for detailed studies of the processes.

Stability Characterization of Ir DHCs. We characterized the stability of Ir DHCs in two different settings. First, the chronoamperometry data of Ir DHCs on $\alpha$-Fe$_2$O$_3$ for the first 10 h were compared with other Ir catalysts. No measurable degradation was observed (Fig. 4B), suggesting that under PEC conditions the Ir DHC on $\alpha$-Fe$_2$O$_3$ is stable. The stability is better than Ir SACs and Ir NPs (SI Appendix, Fig. S25). Inductively coupled plasma optical emission spectroscopy (ICP-OES) characterization indicated that no Ir was detected in the electrolytes of the Ir DHCs. In addition, the key features in the data of STEM-EDS, in situ DRIFTS, and XPS by Ir DHCs after PEC test are similar to those by fresh samples, indicating that there was no aggregation (SI Appendix, Figs. S26–S28). Second, the Ir DHCs were subjected to focused electron beam irradiation under HAADF-STEM conditions; the purpose was to observe the inherent thermal stability of Ir DHCs. As seen in Fig. 4 C–E, no aggregation or detachment was observed for up to ~1 min of irradiation, which is longer than the reported SAC systems under the similar condition (32). The O$_3$-binding site offered by the $\alpha$-Fe$_2$O$_3$ substrate has proven critical to the formation and stability of Ir DHCs (Fig. 3 C and D and SI Appendix, Fig. S18). We envision, however, that the binding is not unique to the chemical nature of the $\alpha$-Fe$_2$O$_3$ substrate. As long as similar binding sites are available, Ir DHC structures with comparable stabilities and catalytic activities are expected on other supporting substrates.

Conclusion

Heterogeneous catalysts with active moieties that are well defined in their atomic and/or molecular structures are expected to play important roles in the development of catalysis. The main challenge of this research direction lies in the synthesis and characterization of such catalysts. Within this context, the Ir DHCs reported here represent a significant advance. Not only are Ir DHCs a heterogeneous catalyst featuring two active-metal atoms, their structures are also well characterized at the atomic level. The atomic arrangement of the catalytic center (Ir–O–Ir), which is key to the water-oxidation activities, is preserved by the strong binding provided by the supporting substrate. The resulting Ir DHC exhibits outstanding stability against aggregation or detachment. It shows high activity toward water oxidation. The finding is built upon recent advances aimed at synthesizing and studying SACs but takes a crucial step forward. It presents opportunities to study the detailed mechanisms of heterogeneous catalysis involving multiple but individually separated active atoms, which was carried out using DFT in this article. The insights are expected to contribute to the design and optimization of heterogeneous catalysts and electrocatalysts.

Materials and Methods

Preparation of Ir Dinuclear Catalyst on $\alpha$-Fe$_2$O$_3$ (Ir DHC) and Loading Estimate. Step 1. An $\alpha$-Fe$_2$O$_3$ substrate was soaked in the Ir homodimer solution for 10 h and then thoroughly rinsed with deionized water to form the Ir intermediate. The loading amount was estimated by comparing the change of the UV-vis absorbance of the Ir homodimer solution. We used the average absorbance decrease for a 32-time loading procedure, the overall geometry and solvation effects, which is beyond.

Preparation of Ir Single-Atom Catalysts on $\alpha$-Fe$_2$O$_3$ (Ir SAC). Similar to the preparation of Ir DHC, Ir SAC was obtained by carrying out step 2, the photochemical treatment. The loading of Ir DHC sample, bec...
time of 16 μs per pixel were chosen for desirable signal-to-noise ratios. EDS was performed on a Super-X EDS system consisting of four Bruker silicon drift detector. During EDS mapping, sample current for imaging was chosen to be a simultaneous image collector. The Fe K edge, O K edge, and Ir K edge were used for elemental mapping. EELS was collected with a Gif quantum EELS spectrometer, a standard power-law background subtraction was utilized to remove the background. Figs. 2 A and B, 3A, and 4 C-E and SI Appendix, Figs. S4–S6 and S9 were collected on an FEI Titan G2 80–200 CryoN “ChemSTEM” microscope, which was equipped with a high-brightness field-emission gun, a probe spherical aberration (Cₐ) corrector, and an EDs mapping system (33). The semiconvergence angle for imaging was ~24.7 mrad, while the collection semiangle was ~70–200 mrad. A higher beam current of 280 pA was used with a longer dwell time of 0.1 ms per pixel and repeated 200× for atomic-resolution EDS mapping. Spatial drifting was corrected using a simultaneous image collector. All high-resolution HAADF images shown in this work are raw data without any postfiltering.

In Situ DRIFTS. In situ DRIFTS measurements were performed using a Nicolet iS50 FTIR spectrometer equipped with a DTGS KBr detector and a Harrick-praying mantis HVC-DRM cell equipped with KBr windows (34). After each sample was placed in the cell, it was heated to 105 °C under pure He for at least 15 min to remove physically adsorbed H₂O and was then gradually cooled to room temperature. If Nₐ-vFe₃O₄ prepared by method 1 was further treated under 20% H₂ at 150 °C for 3 h to remove potential thin oxide surface coatings, Background spectra were then recorded at room temperature. Subsequently, the gas flow was changed to CO for 15 min and was changed back to He to avoid possible interference of gaseous CO to the spectra. DRIFT spectra were recorded in the CO absorption region, 2,200–1,800 cm⁻¹ after varying the He purging time (5, 10, and 40, and 70 min). The resolution was selected at 4 cm⁻¹. Sixty-four scans were recorded for each spectrum shown in the work.

PEC Characterization. PEC measurements were carried out using a potentiostat (Modulab XM equipped with Modulab XM ECS software). The light source was a solar simulator (100 mW/cm², Solarlight model 165–300-M Air Mass Solar Simulator, AM 1.5). Ir (H/C)=Fe₃O₄, Ir SACₓFe₃O₄, Ir Nₐ-vFe₃O₄, and Fe₃O₄ were used as the working electrodes, with an SCE electrode as the reference electrode, and a Pt wire used as the counter electrode. The electrolyte was a 0.1 M KNO₃ solution with the pH adjusted to 6.0. The potential was corrected to the RHE scale by the Nernst equation (E_RHE = E_SCE + 0.599 pH + 0.241). For linear sweep voltammetry data, the scan rate was 20 mV/s.

Computational Details. We used the Vienna ab initio simulation package (VASP) for all periodic boundary calculations (35). Projector augmented plane-wave method (36) together with the Perdew-Burke-Ernzerhof exchange-correlation functional (37) were employed to describe the electron–ion interactions. A cutoff of 450 eV was chosen for the plane-wave basis set in all calculations. A 5 × 5 × 5 Monkhorst-Pack k-point grid was chosen for the optimization of bulk Fe₃O₄. The Gaussian smear method was used for Fe₃O₄. The value used was 0.1 eV. The energy convergence criterion was set to be 10⁻⁶ eV per unit cell and the geometry convergence criterion was set to be 10⁻³ eV per unit cell for energy difference between two consecutive ionic steps. Additional details of the calculations are available in SI Appendix.

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