Manganese vacancy-confined single-atom Ag in cryptomelane nanorods for efficient Wacker oxidation of styrene derivatives†

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Single-atom catalysts provide a pathway to elucidate the nature of catalytically active sites. However, keeping them stabilized during operation proves to be challenging. Herein, we employ cryptomelane-type octahedral molecular sieve nanorods featuring abundant manganese vacancy defects as a support, to periodically anchor single-atom Ag. The doped Ag atoms with tetrahedral coordination are found to locate at cation substitution sites rather than being supported on the catalyst surface, thus effectively tuning the electronic structure of adjacent manganese atoms. The resulting unique Ag–O–MnO₆ unit functions as the active site. Its turnover frequency reaches 1038 h⁻¹, one order of magnitude higher than for previously reported catalysts, with 90% selectivity for anti-Markovnikov phenylacetaldehyde. Mechanistic studies reveal that the activation of styrene on the ensemble site of Ag–O–MnO₆ is significantly promoted, which can accelerate the oxidation of styrene and, in particular, the rate-determining step of forming the epoxide intermediate. Such an extraordinary electronic promotion can be extended to other single-atom catalysts and paves the way for their practical applications.

Introduction

Supported single-atom catalysts (SACs) have attracted considerable attention in the field of heterogeneous catalysis by virtue of their uniform active sites and coordination environment. It is critical to maintain the stability of SACs that have unambiguous active components under preparation and reaction conditions. Utilizing defects on supports to construct strong interaction between individual metal atoms and supports have become an effective approach to stabilizing SACs. Among them, metal cation vacancies on the surface of supports have great potential for anchoring single atoms, owing to their adjustable structure and multifarious electron and orbital distributions. Most recently, we reported a single-atom Pt catalyst anchored on the Ni²⁺ vacancy defects of Ni(OH)₂ nanoboard. Although the developed catalyst exhibited high efficiency in diboration reactions, this method has limitations for other metals. Cryptomelane-type octahedral molecular sieve (OMS) nanorods feature ordered arrays of MnO₆ moieties with tunable electronic structure. So far, the cation vacancies on OMS-2 have not been reported, and utilizing the defect-rich OMS-2 to achieve a high metal-loading SAC has not yet been realized.

The Wacker oxidation reaction, an atom-economic route for the formation of C=O bonds, is one of the most important processes in organic synthesis, because its anti-Markovnikov carbonyl products and their derivatives are value-added chemicals. The protocols developed thus far either depend on precious metals, but have low activities (turnover frequency <108 h⁻¹), or generate Markovnikov products with no regiocontrol. Moreover, the nature of the active sites in heterogeneous catalysts remains elusive, as the multiple sites involved during the reaction may lead to unexpected synergies. Therefore, there is an urgent need to develop a robust heterogeneous...
catalyst for aldehyde-selective Wacker oxidation reactions. Notably, the active components in these reported heterogeneous catalysts for Wacker oxidation of styrene are all metal nanoparticles and clusters. Downsizing metal particles or clusters to single atoms is highly desirable to improve their catalytic activity and alter the product distribution. Therefore, the rational design of SACs would provide unexpected opportunities to obtain an ideal heterogeneous catalyst for target-oriented synthesis of anti-Markovnikov carbonyl compounds.

Herein, we report a catalyst with single Ag atoms periodically anchored on defect-rich cryptomelane-type octahedral molecular sieve (Ag1/OMS-2) nanorods via a simple thermal diffusion method. Although various strategies have been developed for preparing SACs, the construction of Ag SAC on cation vacancies of OMS-2 has never been achieved owing to the easy aggregation of Ag species. Aberration-corrected annular dark-field scanning transmission electron microscope (AC-HAADF-STEM) image simulation and theoretical calculations identify that the single Ag atoms occupy manganese vacancy sites and form strong Ag–O bonds with the neighbour O atoms rather than being supported on the catalyst surface, which effectively modulates the Wacker oxidation activity. The turnover frequency (TOF) and the selectivity for phenylacetaldehyde can reach 1038 h⁻¹ and 90%, respectively. To the best of our knowledge, this catalytic performance outperforms those of previously reported heterogeneous catalysts. Density functional theory (DFT) results indicate that single Ag atoms play a crucial role in promoting styrene activation, and lower the energy barrier for the rate-limiting step of formation of the epoxide intermediate. By contrast, the Ag nanoparticles supported on OMS-2 tend to over-oxidize the epoxide into benzaldehyde, leading to a lowered TOF and selectivity for phenylacetaldehyde (173 h⁻¹ and 50%, respectively). Our method here is also applicable to other metals (Au, Ir, and Pd) and therefore is a general and robust fabrication route to SACs.

Results and discussion

The synthetic procedure for the catalyst is illustrated in Fig. 1a. To begin with, OMS-2 nanorods (that is, z-MnO2) were synthesized on a gram scale via a one-pot hydrothermal method.3,14 Subsequently, Ag species were anchored on the surface of the defective OMS-2 via two simultaneous redox and hydrolysis reactions.3,14 Namely, the chemical is etched onto the surface of OMS-2 nanorods by H2O2,3,36 thereby promoting the hydrolysis of AgNO3 to Ag(OH), and is further reduced into Ag nanoparticles, and then anchored onto the surface manganese-deficit defect sites of the OMS-2 nanorods. Next, the solid was dried at 120 °C overnight to obtain Ag NPs/OMS-2. After calcination in air at 500 °C, Ag1/OMS-2 was obtained through a thermal diffusion strategy.3,36 The defects in the OMS-2 nanorods were confirmed by positron annihilation spectroscopy, a well-established technique to determine defects in solids.3,36 The positron lifetime spectra of Ag1/OMS-2 and OMS-2 are similar (Fig. S1†), indicating that these two samples possess similar defect types. After analysing the spectra, we can see that Ag1/OMS-2 and OMS-2 feature three lifetime components (Table S1†). According to the two-state trapping model, the shortest lifetime component (\(r_1\)) could be attributed to the bulk lifetime of the samples, the longest lifetime (\(r_3\)) is due to the large voids (defect clusters) in the samples, and \(r_2\) could be ascribed to the positron annihilation trapped at manganese vacancies,3,36 which implies the presence of manganese vacancies in these two samples. In addition to the lifetime of the positron, the relative intensity of positron lifetime provided more information on the concentration of the defects. Table S1† summarizes the lowered manganese vacancy concentration (\(I_0\)) in the Ag1/OMS-2 sample in comparison to that of OMS-2, directly indicating that the single Ag atoms occupy the manganese vacancy sites rather than support on the catalyst surface.

A uniform nanorod structure can be observed in the TEM and high-resolution TEM (HRTEM) images (Fig. 1b and its inset.
and S2†), which inherits the structure of OMS-2, with diameters of about 13 nm and lengths of 30–100 nm. HAADF-STEM and energy-dispersive X-ray spectroscopy (EDX) images reveal that Ag, O and Mn elements are evenly distributed over the entire OMS-2 structure, and no Ag nanoparticles are observed (Fig. 1c). The content of Ag is estimated to be approximately 2.5 wt% based on inductively coupled plasma optical emission spectrometry (ICP-OES) analysis. In the AC-HAADF-STEM images (Fig. 1d and g), high density bright dots (highlighted by red circles) corresponding to single Ag atoms are observed. As shown in the intensity profiles along the line (Fig. S3†), the single Ag atoms are much brighter than the manganese atoms. Moreover, these images also demonstrate that those small bright dots of Ag locate at the lattice walls of the manganese columns, rather than in the spacing in between. This indicates that the single Ag atoms are stabilized in the manganese-deficit defect sites by substitution instead of interstitial doping. This can be further verified by the following simulation. Fig. 1e and f, respectively, show the experimental HAADF-STEM image and the simulated HAADF-STEM image using a model with seven Ag atoms replacing seven surface manganese atoms. In the simulated image, the atom column containing Ag shows higher contrast than the nearby Mn–O atom columns, as Ag has a larger atomic number. The agreement between experiment and simulation provides clear evidence that the Ag atoms are anchored as substitutes of manganese. The detailed simulated procedure is summarized in the ESI† In contrast, the simulated HAADF-STEM image using the model in which Ag atoms are in the interval between the manganese columns does not match the experimental image at all (Fig. S4†). Moreover, the surface area and pore size of the Ag1/OMS-2 catalyst were nearly identical to those of the pure OMS-2 support, indicating that the prepared Ag1/OMS-2 and OMS-2 materials have similar textural properties (Fig. S5 and Table S2†). This thermal diffusion strategy can be readily generalized to other elements, such as Ir, Au, and Pd. In addition, the atomically dispersed nature of the metal atoms throughout the OMS-2 support was demonstrated by the AC-HAADF-STEM images (Fig. 1h–j and S6†).

To further confirm that the single Ag atoms are dispersed in Ag1/OMS-2, and to unveil the changes in atomic and electronic structure of Ag1/OMS-2, we performed X-ray absorption near-edge structure spectroscopy (XANES), extended X-ray absorption fine structure spectroscopy (EXAFS), X-ray photoelectron spectroscopy (XPS), and X-ray diffraction (XRD). The manganese K-edge XANES spectra of OMS-2 and Ag1/OMS-2 are quite similar (Fig. S7†), except that the absorption edge of Ag1/OMS-2 was shifted by about 0.5 eV to higher energy, indicating a slightly higher oxidation state of manganese in Ag1/OMS-2 compared with that in OMS-2, which can be ascribed to the occupation of manganese defects by Ag atoms.41,42 The oxidation state of manganese was further analysed by manganese 3s transfer from the support to the Ag atoms, thus enhancing the rearrangement of the atoms in these planes induced by the addition of Ag, as Ag has a larger atomic radius than manganese.

Fig. 2a shows the XANES spectra of the Ag K-edge. The position of the absorption edge of Ag1/OMS-2 is located between those of Ag foil and AgO (Fig. 2a, inset), suggesting that the single Ag atom is positively charged and its oxidation state is between 0 and +2. The oxidation state of Ag was further analysed by XPS. As shown in Fig. 2b, the Ag 3d spectrum can be deconvoluted into two peaks centred at 367.8 and 373.8 eV, corresponding to 3d5/2 and 3d3/2 levels, respectively.43,44 The peak positions are between those of Ag(0) and Ag(0), indicating Ag atoms carry partially positive charge through electron transfer from the support to the Ag atoms, thus enhancing the metal–support interaction. Fig. 2c shows the FT k3-weighted EXAFS spectra of Ag1/OMS-2 and reference samples. The profile for Ag1/OMS-2 shows two main peaks at about 1.5 and 3.0 Å, associated with the first shell of Ag–O scattering and the higher shell of Ag–Mn, respectively. Compared with the FT profile of Ag foil, no scattering from the Ag–Ag contribution is observed, indicating the sole existence of atomically dispersed Ag in Ag1/OMS-2. The corresponding fits of the EXAFS spectra of Ag1/OMS-2 in R space and K space are exhibited in Fig. S11†. To further confirm this inference, a wavelet transform (WT) analysis of the Ag K-edge EXAFS oscillations was performed. In Fig. 2d, the WT contour plot of Ag1/OMS-2 in the first coordination shell shows one intensity maximum at around 4 Å−1, which is associated with the Ag–O contribution. However, for the higher coordination shell, the WT contour plot shows an additional maximum at around 5 Å−1, owing to the Ag–Mn contribution. In comparison to the WT contour plot of the Ag foil, there is no intensity maximum detected near 8.5 Å−1 (indexed to the Ag–Ag path), further confirming that Ag atoms are atomically dispersed.

To get the precise atomic structure and explore the interaction between single-atom Ag and OMS-2 support, we carried out quantitative EXAFS curve fitting, and the results for Ag1/OMS-2...
are listed in Table S4.† According to the fitting results, the Ag–O coordination number for Ag1/OMS-2 is 4, with a bond length of 2.06 Å, implying that each Ag atom is coordinated with four oxygen atoms in the OMS-2 nanorods. Moreover, the coordination number for the higher shell of Ag–Mn is 4.6 ± 0.7, with a bond length of 3.72 ± 0.02 Å. The possible structure of Ag1/OMS-2 was further elucidated by first-principles calculations, which can provide more insight into the interaction between manganese vacancies and single Ag atoms. The O-terminated (110) surface of OMS-2 was selected for the simulation model (Fig. S12†). For the OMS-2 with manganese vacancies, the manganese vacancy site and the fourfold hollow site of the oxygen atoms were confirmed to be the most stable adsorption sites for the single Ag atom. According to the results of charge density difference, single Ag atoms are fixed by the four top oxygen atoms near the manganese vacancy (Fig. 2e). In the simulation model, the bond lengths of Ag–Mn and Ag–O are 3.70 and 2.08 Å, respectively, which are in accord with the EXAFS results. The cyan region represents charge density depletion around a single Ag atom, which illustrates that the Ag is positively charged. The charge accumulation on Ag is [1.09e] based on the Bader charge analysis.

In addition, the Ag atom adsorbed onto the OMS-2 with manganese vacancies has a formation energy of −0.9 eV, indicating that this stable structure is readily available (Fig. S13† inset). By contrast, for the Ag atom supported on the OMS-2 without manganese vacancies, it tends to locate at the site that has a slight deviation from the oxygen atoms locating at the fourfold hollow site. This is caused by the competition between the interaction between the Ag atom and the four top oxygen atoms and the electrostatic repulsion of the Ag and manganese atoms with positive charge (Fig. S14†). Based on these DFT calculation results and the XANES data, it can be inferred that the manganese vacancies play a critical role in stabilizing the isolated Ag atoms supported on OMS-2, through removing the spatial isolation between Ag atoms and O atom and thus promoting the charge transfer from OMS-2 to Ag. The total density of states and partial density of states (PDOS) of Ag1/OMS-2 were calculated to reveal the energy contribution from this atomic orbital hybridization. The PDOS of manganese atoms has more contributions, compared with those of O atoms, near the Fermi energy (Fig. S13†). The PDOS of the Ag atom mainly occupies the energy below the Fermi energy, thus the ensemble active site of Ag–O–MnO is favourable for the activation of reactants.† Fig. 2f shows the charge density difference of adsorbing styrene over an Ag atom in the substitution site of a manganese atom and supported on the surface. After adsorbing the styrene molecule, the charge is more accumulated on the O atom, which was adsorbed by the Ag atom in the substitution site of manganese, indicating that this configuration promotes the rate-determining step of formation of styrene oxide.

The selective oxidation of styrene to aldehydes is one of the most important processes in organic synthesis. When we used Ag1/OMS-2 to catalyse the Wacker oxidation of styrene as a representative substrate (Fig. 3a), we found that the catalyst achieved a nearly 100% conversion of styrene, and produced almost exclusively the anti-Markovnikov oxidation product, phenylacetaldehyde (detected by gas chromatography-mass spectrometer (GC-MS), 1H nuclear magnetic resonance (NMR) and 13C NMR; Fig. S15–S19†). The calculated TOF of Ag1/OMS-2 reached as high as 1038 h⁻¹. There is no evident metal Ag leaching in the solution, based on the ICP-OES result. The yield of phenylacetaldehyde catalysed by other SACs (Ru1/OMS-2, Rh1/OMS-2, Pd1/OMS-2, Ir1/OMS-2, Pt1/OMS-2, and Au1/OMS-
2) are shown in Fig. S20. Under the identical reaction conditions, no product was detected without any catalyst (Fig. 3b). We also assessed the catalytic performance of the pure OMS-2, and it gave a nearly 10% yield of phenylacetaldehyde, indicating that OMS-2 itself plays an important role in Wacker oxidation of styrene, which is consistent with previous reports. For comparison, a sample of Ag NPs/OMS-2 with an Ag loading of 2.5 wt% was prepared and tested, which offered a phenylacetaldehyde yield of merely 15% under the same conditions. Based on the reported literature, we conducted a controlled experiment to determine the reaction intermediates. The controlled experiment revealed that when we used styrene oxide as the reaction substrate, the complete transformation of styrene oxide to phenylacetaldehyde was realized under identical reaction conditions (Fig. S21†), indicating that styrene oxide is the intermediate for Wacker oxidation of styrene.

In practical production, catalyst recycling is an important issue in heterogeneous catalytic reactions. Therefore, we investigated the recyclability of our catalyst for the Wacker oxidation reaction by using styrene as a model substrate. The spent catalyst was separated, and then used after only washing with ethanol and drying, with no extra fresh catalyst being added. As shown in Fig. 3c, after five cycles, the activity and selectivity of Ag1/OMS-2 were well retained, suggesting a good recycling capability and potential for the Wacker oxidation of styrene in practical applications. Besides, the structures were unchanged with respect to those of the fresh samples, as identified by XRD and AC-HAADF-STEM (Fig. S22 and S23†). For the spent Ag1/OMS-2 catalyst, no Ag particles formed after the Wacker oxidation of styrene, which has been confirmed by EXAFS (Fig. S24†). Next, we examined the scope of alkenes for this Wacker oxidation reaction catalysed by Ag1/OMS-2, and the results are presented in Table S5.† When styrene substituted with electron-donating (Table S5, entries 1–4) or electron-withdrawing groups (Table S5, entries 5–9) were used, the desired anti-Markovnikov products were obtained in high yields (>90%). Furthermore, the catalysis comparison with reported works in the literature is summarized in Table S6.† Our Ag1/OMS-2 has both an outstanding TOF and a remarkable selectivity for phenylacetaldehyde, with low catalyst usage and simple preparation method.

To assess the activity of surface oxygen that participates in the styrene oxidation reaction, the electronic properties and the amount of adsorbed oxygen on OMS-2 and Ag1/OMS-2 were studied via XPS, temperature-programmed desorption of oxygen (O2-TPD), XANES, and Raman spectroscopy. The electronic density of oxygen was determined by the O 1s XPS, and an intensive peak with a shoulder at the high binding energy side is observed for OMS-2 and Ag1/OMS-2 catalysts in Fig. 4a. The main peak at about 528.5–530.5 eV and the broad one at 531.6 eV are assigned to surface lattice oxygen (Olat., O2−) and adsorbed oxygen (Oads.), respectively. In addition, a discernible shift to lower binding energy for Ag1/OMS-2 is observed in comparison to OMS-2, which is commonly assigned to an increase of its negative charge. As displayed in Fig. 4a, the content of adsorbed oxygen on the surface of Ag1/OMS-2 (20.4%) is higher than that of OMS-2 (18.9%), suggesting that Ag1/OMS-2 is more conducive to generating active oxygen species, thereby promoting the oxidation of styrene. O2-TPD was conducted to identify the evolution of oxygen from the prepared materials (Fig. 4b). The desorption peak below 350 °C can be ascribed to the release of chemisorbed oxygen molecules and active surface oxygen, whereas the peak in the range 400–650 °C is attributed to the release of sub-surface lattice oxygen. The peak above 700 °C is due to the evolution of bulk lattice oxygen. The oxygen released at low temperature was considered as labile oxygen species, which were highly active for styrene oxidation. A lower oxygen-releasing temperature indicates a looser binding between the oxygen species and manganese atoms. In the case of Ag1/OMS-2, the desorption temperature was like that for OMS-2, and the amount of labile oxygen was higher than that of OMS-2, indicating that Ag1/OMS-2 contains much more loosely bound labile oxygen species than does OMS-2, which is in accordance with the O 1s XPS result. Therefore, the Ag1/OMS-2 deserves to give a better catalytic activity for styrene oxidation.

Fig. 4c shows the O K-edge XANES spectra for OMS-2, fresh and used Ag1/OMS-2, and Ag NPs/OMS-2. The broad peaks at $\gtrsim 535$ eV are attributed to the transition from O 1s to O 2p and the manganese 4sp orbital, whereas the characteristic peak at around 530 eV corresponds to the transition from O 1s to O 2p and the manganese 3d orbital. The peak feature in the O K-edge XANES for Ag NPs/OMS-2 is almost the same as that for the OMS-2 support, implying that the electronic structure of oxygen in the sample is unchanged by impregnating Ag nanoparticles. By contrast, the absorption peak intensity of fresh Ag1/OMS-2 decreases notably with respect to that of OMS-2, indicating that the introduction of Ag atoms induced electronic structural changes around oxygen ions but barely any change around the manganese ions, which might be the main origin for the increased mobility of active oxygen species after the introduction of Ag. In addition, the absorption peak intensities of the fresh Ag1/OMS-2 are significantly changed after styrene oxidation reaction, which further confirms that the active oxygen species are involved in the reaction. As shown in Fig. 4d,
the Raman scattering spectra of OMS-2 and Ag₁/OMS-2 both feature four main contributions at 180, 383, 573, and 639 cm⁻¹. The Raman bands at around 180 and 383 cm⁻¹ are assigned to the deformation vibration of Mn–O–Mn. The bands at around 639 and 573 cm⁻¹ are attributed to the symmetric stretching vibration (Mn–O), which is perpendicular to the MnO₆ group direction. It is very interesting to note that, compared with the Mn–O stretching vibration mode (639 cm⁻¹) of OMS-2, a much weaker and a considerable downshift of the Mn–O stretching vibration can be observed for Ag₁/OMS-2 (627 cm⁻¹), suggesting that Ag incorporation increases the disorder of OMS-2 and induces more O vacancies in OMS-2, thus favouring the activity of the catalyst.

Further DFT calculations were carried out to unravel the reaction mechanism and to elucidate the fact that the Ag SAC has higher catalytic performance than Ag NPs. The atomic structure of Ag NPs is shown in Fig. S25. The configuration of Ag₁/OMS-2 has single Ag atoms surrounded by manganese atoms and four oxygen atoms with bond lengths of 3.70 and 2.08 Å, respectively. The styrene oxidation reaction catalysed by both Ag₁/OMS-2 and Ag NPs/OMS-2 follows the E₁ (epoxidation–isomerization) mechanism. As shown in Fig. 5, the styrene molecule first approaches the Ag₁/OMS-2 catalyst via a non-planar configuration, where single Ag atoms play an important role in the adsorption and activation of styrene molecules (first step). Then, the styrene molecule connects to the active oxygen species derived from tert-butyl hydroperoxide (TBHP), forming an intermediate state of styrene oxide (second step). Subsequently, the hydride migration and rearrangement lead to the formation of the desired product, phenylacetaldehyde (third step). The energy barriers in the styrene oxide formation step are the highest, suggesting that the second step is the rate-determining step of this reaction. For the rate-limiting step, the Ag–O–MnO₂ moieties acting as active sites play a critical role in facilitating the formation of epoxide. Overall, in the first two elementary steps, the energy barriers over Ag₁/OMS-2 are thermodynamically lower than those over Ag NPs/OMS-2. The energy barriers of the third step for Ag₁/OMS-2 and Ag NPs/OMS-2 are similar, because the Lewis acidity of OMS-2 can make styrene oxide spontaneously isomerize into phenylacetaldehyde. The Lewis acidity has been verified by pyridine infrared spectroscopy (Fig. S26†). Besides, it is relatively difficult to break the epoxy ring to form benzaldehyde over Ag₁/OMS-2, as the energy barrier of the third step is relatively high (Fig. S27†). Furthermore, the single Ag atoms substituting the manganese sites have lower energy barriers for Wacker oxidation than that of single Ag atoms supported on OMS-2, and pristine OMS-2 (Fig. S28 and S29†). These calculated results are in agreement with the superior catalytic performance for Ag₁/OMS-2 observed in experiment.

**Conclusions**

In summary, we report a defect-rich OMS-2 nanorod-supported Ag SAC with remarkable performance towards anti-Markovnikov Wacker oxidation. This OMS-2 support with abundant manganese vacancy defects plays a key role in stabilizing the single Ag atoms via an enhanced charge-transfer mechanism. The TOF of the as-fabricated Ag₁/OMS-2 reaches up to ~1038 h⁻¹, which surpasses those of conventional heterogeneous catalysts based on nanoparticles and clusters. In addition, no evident metal leaching or aggregation were found after the reaction. Combined with surface science, theoretical, and experimental data, the enhanced catalytic properties could be explained at the molecular level, which reveals the critical role of the location of single Ag atoms. Specifically, the single Ag atoms occupy the manganese vacancy sites of OMS-2 rather than being supported on the catalyst surface, and thus facilitate the activation of styrene and lower the energy barrier for the rate-limiting step of formation of the epoxide intermediate. Moreover, this is the first report of an SAC being used for the aldehyde-selective Wacker oxidation of styrene. Using the cation vacancy defects in supports to anchor single metal atoms could become an efficient approach to preparing these SACs on a large scale.

**Author contributions**

H.Y. performed the experiments, collected and analysed the data, and wrote the paper. X.C., D.Q., and H.X. conducted the DFT calculations and analysis. X.Z. and Y.Y. obtained the AC-HAADF-STEM image simulation. Z.C., Q.L., and Y.L. discussed the catalytic process and gave useful suggestions. W.C. helped with the HRTEM characterization and analysis. Z.Z. helped with the X-ray structure characterization and analysis of the XAS data. Q.P. and Y.L. helped with data analyses and discussions. C.C. conceived the experiments, planned the synthesis, analysed results, and wrote the paper.

**Conflicts of interest**

There are no conflicts to declare.

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