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Tuning electronic states of catalytic sites enhances SCR activity of hexagonal WO$_3$ by Mo framework substitution

Catalytic sites of selective catalytic reduction (SCR) of NO with NH$_3$ are identified to be located at the tunnel openings of the Mo-doped hexagonal WO$_3$ catalyst. The framework substitution by Mo reduces the bandgap between the highest occupied molecular orbitals and the lowest unoccupied molecular orbitals, thus accelerating electron transfer in SCR redox cycles and enhancing catalytic activity.

As featured in:
1. Introduction

Nitrogen oxides (NOx), as typical atmospheric pollutants, have a great contribution to acid rain, photochemical smog, and ozone formation, and hence have attracted great attention in the past several decades. Selective catalytic reduction (SCR) of NOx with NH3 for controlling NOx emissions from stationary sources and mobile sources is a promising technology. V2O5–WO3–MoO3/TiO2 catalysts have been commercially used for SCR because of their high catalytic activity and strong SO2 tolerance. However, the toxicity of vanadium demands that V2O5–WO3–MoO3/TiO2 catalysts should be replaced by environmentally friendly catalysts in the long run. SCR essentially follows a reduction–oxidation mechanism, and thus the activity is intimately associated with the redox properties of catalysts. MnO2-based catalysts with good redox properties show high catalytic activity in SCR, especially at low temperatures, but their poor sulfur tolerance makes it impossible for them to be used under practical SCR conditions. Thus, it is highly desirable to develop environmentally benign and sulfur-tolerant SCR catalysts for practical applications.

Although WO3 and MoO3 in V2O5–WO3(MoO3)/TiO2 can promote the SCR activity,10,11 stabilize the TiO2 structure,12,13 and inhibit the oxidation of SO2, they often serve as promoters in SCR because their redox ability is much weaker than that of W2O5. An appealing alternative is to improve the redox ability of WO3 and MoO3, so they can act as active phases. In fact, WO3 and MoO3 have been used as active components of SCR catalysts after their redox properties are improved. For instance, Kobayashi et al. found that WO3/TiO2 prepared by coprecipitation is more active, in the presence of 10% H2O, than WO3/TiO2 prepared by conventional impregnation due to the enhanced redox ability of WO3 in the former catalyst. Giamello et al. found that surface Mo5+ centers are responsible for SCR via a Mo6+/Mo5+ redox cycle resulting from the interaction between the MoO3 phase and the TiO2 support. Thus, it is feasible for WO3 and MoO3 to become active phases by improving their redox ability.

WO3 has different crystalline structures, including hexagonal and monoclinic phases. Hexagonal WO3 (HWO), with a space group of P6/mmm, is constructed from corner-sharing WO6 octahedra. HWO has strong acidity and excellent reducibility due to the co-existence of W6+ and W5+, and thus it can be used as an active component of catalysts. Because
the electronic structures and the ionic sizes of Mo and W are similar, the hexagonal bronze structure can be maintained if WO₆ in the HWO framework is substituted by MoO₆ (i.e., to form Mo–HWO). After such framework substitution, the redox properties of HWO may be tuned. It was reported that the partial substitution of W in HWO by Mo can improve the sensing properties towards NH₃ or NOₓ, which are closely associated with the redox properties of Mo–HWO. The standard reduction potential (φEHWO) of Mo⁶⁺/⁵⁺ is ~0.40 V, close to that of W⁶⁺/⁵⁺ (~0.39 V), and hence the reaction between Mo and W cations is favorable in Mo–HWO. The Zener exchange may enhance the redox ability of Mo–HWO, i.e., the substitution of W in HWO by Mo improves the ability to activate lattice oxygen and gaseous oxygen. It is well known that the activation of NH₃ or NOₓ and oxygen plays crucial roles in SCR, which motivates us to investigate SCR over Mo–HWO.

Herein, we investigate the SCR performance of Mo–HWO. Firstly, we synthesize Mo–HWO by a hydrothermal method, test its catalytic activity in SCR, and determine the structure of Mo–HWO by synchrotron X-ray diffraction (SXRD) and transmission electron microscopy (TEM). Next, the catalytically active sites (CASSs) of Mo–HWO are identified by SXRD, ³¹H magic angle spinning nuclear magnetic resonance (³¹H MAS NMR), and hydrogen temperature-programmed reduction (H₂-TPR). Finally, X-ray absorption near-edge structure (XANES) and X-ray photoelectron spectra (XPS) coupled with theoretical calculations are used to illustrate why the substituted Mo can dramatically enhance the SCR performance of HWO.

2. Experimental

2.1 Catalyst synthesis

Mo–HWO powders were synthesized via a hydrothermal method. Ammonium paratungstate [(NH₄)₆W₁₂O₄₀·13H₂O, 1.300 g], ammonium sulfate [(NH₄)₂SO₄, 8.600 g], ammonium molybdate [(NH₄)₆Mo₇O₂₄·4H₂O, 0.940 g], and oxalic acid (H₂C₂O₄, 2.100 g) were dissolved in deionized H₂O (60 mL). The resulting solution was transferred to a 100 mL autoclave and kept at 180 °C in an oven for 18 h. The final slurry was filtered, washed with deionized H₂O, and dried at 80 °C. The resulting solid was calcined at 350 °C for 4 h in air to give the Mo–HWO catalyst. HWO powders were prepared using the same method as Mo–HWO without the addition of (NH₄)₆Mo₇O₂₄ during the preparation process. To further load K⁺, Mo–HWO was impregnated with a K₂SO₄ solution. K/Mo–HWO was obtained after being dried at 105 °C for 12 h and then calcined at 350 °C for 12 h. The K⁺ loading was set to be 400 μmol gcat⁻¹.

2.2 Catalytic evaluation

SCR reactions were conducted in a fixed-bed quartz reactor (i.d. = 8 mm) under atmospheric pressure. A certain amount of catalyst (40–60 mesh) was charged for each run. For the activity tests, the feed gas was composed of 500 ppm NO, 500 ppm NH₃ and 3.0 vol% O₂, and balanced using N₂. The gas hourly space velocity (GHSV) was fixed at 192 000 h⁻¹. The gas flow rate was 1000 mL min⁻¹, and the amount of the catalyst was 0.5 g. For the water- and SO₂-resistance tests, the feed gas was composed of 1000 ppm NO, 1000 ppm NH₃, 3.0 vol% O₂, 2700 mg m⁻³ SO₂ and 10 vol% H₂O vapor, and balanced using N₂. The GHSV was fixed at 32 000 h⁻¹. The gas flow rate was 500 mL min⁻¹, and the amount of the catalyst was 1.5 g. The concentrations of NO and NO₂ in the inlet and outlet gas were measured using an on-line chemiluminescence NO–NO₂–NOx analyzer (42i-HL, Thermo Electron Corporation, USA). The data were recorded after the reactions reached a steady state. The conversion rate (XNO%) was defined as follows: XNO% = (1 - [NO]out/[NO]in) × 100%, where [NO]out and [NO]in represent the NO concentrations of the inlet and outlet gas, respectively. Arrhenius plots were obtained according to the SCR activity of the samples at 260–320 °C with XNO% less than 20%. We eliminated the effects from the external and internal diffusion by varying the linear velocity of the feed gas passing through the catalysts and the size of catalysts’ particles, respectively, for the reaction kinetics, as shown in Fig. S1 (ESI†).

2.3 Catalyst characterization

SXRD patterns were recorded at BL14B of the Shanghai Synchrotron Radiation Facility (SSRF) at a wavelength of 0.6883 Å. Rietveld refinement analyses of the diffraction profiles were conducted using the Rietica v1.77 program. TEM, high-resolution TEM (HRTEM), and scanning TEM energy dispersive X-ray spectroscopy (STEM-EDX) mapping were conducted on a JEM 2100F transmission electron microscope. XPS experiments were carried out with an RBD 147 upgraded PerkinElmer PHI 5000C ESCA system equipped with a hemispherical electron energy analyzer. The Mg Kα (λ = 1253.6 eV) anode was operated at 14 kV and 20 mA. The spectra were recorded in constant pass energy mode with a value of 46.95 eV. The experimental errors were within ±0.2 eV. Data analysis and processing were undertaken using the XPSPeak4.1 software with the Shirley type background. H₂-TPR experiments were performed using a Micromeritics 2920 adsorption instrument with a thermal conductivity detector to monitor the consumed H₂. H₂-TPR was conducted at 10 °C min⁻¹ under a 50 mL min⁻¹ flow of 5 vol% H₂ in Ar.

³¹H MAS NMR spectra were recorded on a Bruker AVANCE III 400 WB spectrometer. The chemical shifts of ³¹H were referenced to TMS at 0 ppm. ³¹H MAS NMR experiments were performed at a spinning rate of 20 kHz. ³¹H MAS spectra were recorded in a spin echo pulse sequence (π/2–τ–π–τ-acquire), where τ is equal to one rotor period (rotor synchronized). The excitation pulse length was 2.3 μs (π/2), and typically ~40 scans were accumulated with a 5 s delay. XANES spectra were measured at the W L₃-edge at BL14W of the SSRF with an electron beam energy of 3.5 GeV and a ring current of 200–300 mA. Data were collected with a fixed exit monochromator using two flat Si(311) crystals. The XANES spectra were
acquired at an energy step of 0.5 eV. The raw data were analyzed using the IFEFFIT 1.2.11 software package. Raman spectra were obtained using a Horiba Jovin Yvon XploRA spectrometer. The measurements with an excitation laser of 532 nm were calibrated with the Raman peak of Si at 520 cm\(^{-1}\). The interchangeable holographic grating of the spectrometer is 1800 grooves per mm. The laser powers are less than 1 mW to avoid overheating of the samples.

### 2.4 Theoretical calculations

Density functional theory (DFT) calculations were performed with the Vienna ab initio Simulation Package (VASP).\(^{26,27}\) The exchange and correlation effects are described by the general gradient approximation (GGA) method.\(^{28}\) The projector-augmented wave (PAW) method describes the interactions between the core and valence electrons. The plane-wave energy cutoff was set at 450 eV. The Perdew–Burke–Ernzerhof (PBE) exchange-correlation functional method was used in our calculations.\(^{29}\) The lattice constant for a conventional hexagonal cell with a space group of \(P6/mmm\) is 7.319 Å \(\times\) 7.319 Å \(\times\) 3.881 Å. A super cell of 12 formula units of HWO was employed in the charge calculations. The (001) surface of Mo–HWO was cut from the stable supercell after the geometry was optimized.

### 3. Results and discussion

#### 3.1 Catalytic activity and catalyst structures

As shown in Fig. 1a, the activity of HWO is significantly low under the SCR conditions (GHSV = 192 000 h\(^{-1}\), in the absence of \(SO_2\) and \(H_2O\)). The NO conversion (\(X_{NO}\)) on HWO is lower than 40% even at 350 °C. However, the \(X_{NO}\) on Mo–HWO is much higher. The \(X_{NO}\) reaches ~75% at a high GHSV of 192 000 h\(^{-1}\) at 350 °C. In the chemical kinetic regime where the \(X_{NO}\) is lower than 15%, the \(X_{NO}\) of Mo–HWO (12%) is four times as that of HWO (3%) at 250 °C. Furthermore, we tested the stability of Mo–HWO and HWO under mimicked working conditions (2700 mg m\(^{-3}\) \(SO_2\), 10 vol% \(H_2O\), GHSV 32 000 h\(^{-1}\)). Mo–HWO shows higher activity and stability than HWO in the presence of high-concentration \(SO_2\) and \(H_2O\) in the feed gas, as shown in the stability test in Fig. 1a and Fig. S2 (ESI†).

The crystal structures of HWO and Mo–HWO were studied using SXRD. As shown in Fig. 1b, all the diffraction peaks can be indexed to the hexagonal tungsten bronze structure with a space group of \(P6/mmm\) (JCPDS 33-1387).\(^{30}\) HWO has one-dimensional tunnels where different cations can insert themselves. Moreover, the framework \(W\) ions of HWO can exist as \(W^{6+}\), \(W^{5+}\), and \(W^{4+}\) with different oxidation states and ionic radii.\(^{30}\) These framework \(W\) ions can be readily substituted by Mo,\(^{18–22}\) V,\(^{31}\) and Nb.\(^{32}\) The SXRD pattern of Mo–HWO is also shown in Fig. 1b. The hexagonal tungsten bronze structure is preserved after the incorporation of Mo in HWO, and no MoO\(_2\) species appear. However, the structure of HWO is modified, as judged from the intensity of the peaks after the introduction of Mo. The intensity ratio of the (001) plane to the (100) plane is \(\sim 1.4\) for HWO, whereas it decreases to \(\sim 1.2\) for Mo–HWO, indicating that the incorporation of Mo suppressed the growth of HWO along the [001] direction. Similarly, the intensity ratio of the HWO(200) plane to the HWO(100) plane is \(\sim 1.6\) for HWO, whereas it decreases to \(1.2\) for Mo–HWO, indicating the lattice distortion of the HWO structure after the incorporation of Mo. Because Mo ions have almost the same ionic radius as \(W\) ions, for instance, the radius of Mo\(^{5+}\) (0.75 Å) is the same as that of \(W^{6+}\) (0.74 Å),\(^{33}\) no distinct shift in the diffraction peaks of HWO is observed after the incorporation of Mo.\(^{34}\) The results support the substitution of the HWO framework by Mo.

The effect of the Mo framework substitution on the morphologies and structures of HWO was studied by TEM, HRTEM, and STEM (Fig. 2). HWO has a rod-shaped morphology with a mean width of \(\sim 80\) nm and a mean length of \(\sim 370\) nm (Fig. 2a, S3 and S4, ESI†). The fringe of 3.8 Å, assigned to the HWO(001) spacing, can be clearly observed by HRTEM (Fig. 2b), implying that HWO rods grow along the [001] axis because the (001) plane has a higher surface energy than other low-index Mille planes such as the (100) plane.\(^{35}\) The growth direction of HWO is also in agreement with a geometrical prediction made by the Bravais–Friedel–Donnay–Harker method,\(^{36}\) as well as some experimental observations.\(^{37}\) After the Mo framework substitution shown in Fig. 2c–f, Mo–HWO still preserves the rod-shaped structure.
in the [001] growth direction, but the length (∼260 nm) decreases more distinctly than the width (∼60 nm) [Fig. S3 and S4, ESI†] in comparison with HWO, implying that the aspect ratio decreases. Hence, the framework substitution by Mo increases the fraction of the (001) planes exposed. Moreover, the relatively obscure fringe of the (001) plane in the HRTEM image (Fig. 2d) and the highly dispersed Mo species (Fig. 2e) further demonstrate the Mo framework substitution, as evidenced by the SXRD patterns (Fig. 1b) and Raman spectra (Fig. S5, ESI†).

3.2 Identification of catalytically active sites

As discussed above, the (001) top-facets of Mo–HWO, consisting of the tunnel openings, have higher surface energy than the (100) side-facets,35 and hence the CASs may be located at the (001) facets of Mo–HWO. To verify this hypothesis, we anchored K⁺ ions (a SCR catalyst poison) onto the tunnel openings. Fig. 3a shows the SXRD pattern with Rietveld refinement of Mo–HWO after loading K⁺ (K/Mo–HWO), and the crystallographic data and structure parameters obtained by the Rietveld refinement are listed in Tables S1 and S2,† respectively. Obviously, K⁺ ions are situated at the Wyckoff 1a sites or at the crystallographic (0,0,0) sites.16,38 According to our recent report,38 K⁺ ions occupy the tunnel openings of the (001) top-facets. After the addition of K⁺, the catalytic activity of Mo–HWO decreases drastically (Fig. S6, ESI†), demonstrating that the tunnel openings should be the CASs for SCR. Moreover, the tunnel openings on the (001) facets are suitable for NH₃ adsorption and subsequent activation.16,39

NH₃ adsorption and activation are commonly regarded as important steps for SCR. We studied NH₃ adsorption on Mo–HWO surfaces by using ¹H MAS NMR spectroscopy. As shown in Fig. 3b, before the adsorption of NH₃, a strong symmetric peak shows up at a chemical shift (δ) of ~6.8 ppm. This peak can be assigned to the hydrogen resonance of the lattice NH₄⁺ in the Mo–HWO tunnels. Here, we denote it as NH₄⁺ at a “static state” because NH₄⁺ ions are strictly confined in the tunnel lattice sites of Mo–HWO.39,40 Another relatively weak peak at ~4.8 ppm is assigned to the hydrogen resonance of NH₄⁺ adsorbed in the tunnels.40 This NH₄⁺ species might have a relatively weak interaction with the Mo–HWO tunnels.41 No resonant peaks due to H₂O can be observed because Mo–HWO was annealed at 350°C.

After the adsorption of NH₃ on Mo–HWO in an inert N₂ gas flow, two new resonant features appear at chemical shifts of ~1.2 and ~2.8 ppm (Fig. 3b), assigned to the OH⁻ groups and H₂O, respectively.40 The data indicate that NH₃ is oxidized by the lattice oxygen of Mo–HWO. The intensity of the peak due to the lattice NH₄⁺ at a chemical shift of 6.8 ppm increases, possibly originating from the reaction of the produced H₂O with adsorbed NH₃ (NH₃ + H₂O → NH₄⁺ + OH⁻), further indicating that NH₃ can be activated by Mo–HWO. The occurrence of this activation process is possibly associated with the tunnel structures, because the amount of NH₃ adsorbed on monoclinic WO₃ without any tunnel is significantly less than the amount of that adsorbed on HWO with the tunnel structure.39 Moreover, according to the curve-fitting of the ¹H NMR spectrum of Mo–HWO, the chemical shift at ~5.6 ppm can be ascribed to NH₃.40 Often, SCR does not occur in the HWO tunnels due to the big molecular sizes of the reactant O₂ and the produced N₂. Thus, SCR mainly occurs at the tunnel openings.
H₂-TPR experiments were conducted to further identify the CASs. The reduction of the bulk lattice oxygen species of pure WO₃ often occurs at temperatures higher than ~570 °C, and thus possible H₂ consumption at temperatures below ~570 °C should be assigned to the surface lattice oxygen species. Fig. S7 (ESI†) shows the H₂-TPR profiles of HWO, Mo–HWO, and K/Mo–HWO at 250–900 °C. A reduction peak at ~750 °C with an onset reduction temperature of ~600 °C can be ascribed to the reduction of the bulk lattice oxygen of HWO to form WO₂.⁴³ After the Mo framework substitution, this bulk reduction peak shifts to ~640 °C for Mo–HWO and the onset temperature is ~550 °C, while the reduction feature of this peak almost remains unchanged after the K⁺ loading. Thus, the weak reduction peaks at temperatures lower than ~550 °C can be due to the reduction of the surface lattice oxygen.

The reduction of the surface lattice oxygen species of the samples is shown in the H₂-TPR profiles in Fig. 4. Similar reduction features for these three samples are observed. These features can be fitted into two reduction peaks, i.e., the low-temperature (LT) and high-temperature (HT) peaks. The LT and HT peaks can be assigned to the reduction of the high surface energy (001) top-facets and the relatively low surface energy (100) side-facets, respectively, because these three samples expose only two kinds of facets on the surfaces.¹⁶ The LT peaks of these three samples are almost located at the same reduction temperature of ~400 °C, while the calculated amounts of consumed H₂ are distinctly different from each other. The Mo framework substitution leads to a considerable increase of the H₂ consumption based on the LT peaks, in line with the increasing fraction of the (001) top-facets due to the higher aspect ratio of Mo–HWO to HWO. Note that the amount of H₂ consumption determined from the LT peak of Mo–HWO dramatically decreases from ~780 to ~110 μmol gcat⁻¹ after K⁺ ions occupied the tunnel openings on the (001) facets. Hence, the tunnel openings of Mo–HWO are the CASs. When K⁺ ions occupy the CASs, the redox ability of the CASs drastically decreases, leading to much lower SCR activity (Fig. S6, ESI†).

Both the amount of the H₂ consumption and the reduction temperature of the HT peak of Mo–HWO (due to the reduction of the surface lattice oxygen on the (100) side-facets) remain almost unchanged after K⁺ ions occupy the (001) top-facets. This further demonstrates that the CASs are not situated on the (100) facets, but located only on the (001) facets. Note that the onset of the reduction of Mo–HWO occurs at a much lower temperature than that of HWO, allowing Mo–HWO to exhibit an activation energy (~37 kJ mol⁻¹) much lower than that of HWO (~67 kJ mol⁻¹) in SCR at low reaction temperature (Fig. S8, ESI†). Thus, the Mo framework substitution not only increases the number of CASs but also enhances the redox ability of the CASs.

### 3.3 Electronic states of catalytically active sites

Both the catalytic activity and the redox ability are intimately associated with the electronic states of the CASs.⁴⁴ Thus, the effect of the Mo framework substitution on the electronic structures of W was investigated by using XANES and XPS. Fig. 5 shows the W XANES spectra and the corresponding second-derivative spectra of HWO and Mo–HWO at the W L₃-edge. Two overlapping absorption peaks attributed to the 2p½ → ε₅ and 2p½ → ε₃g transitions appear in the XANES spectrum of HWO. These peaks shift to the low absorption energies after Mo substitution, indicating that the lowest unoccupied orbitals of W decrease in energy. The crystal field splitting energy (10Dq) of HWO is calculated to be ~4.5 eV according to the second-derivative spectra, and it decreases to 4.2 eV after the Mo framework substitution, indicating the decrease of the oxidation states of W.⁴⁵

The oxidation states of Mo–HWO surfaces were studied by XPS (Fig. 5b). The W⁶⁺ and W⁵⁺ species with a ratio of 5:1

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**Fig. 4** H₂-TPR profiles of HWO, Mo–HWO and K/Mo–HWO. The red shades show the reduction of the (001) top-facets, and the numbers in the shades are the corresponding amounts of the H₂ consumption during the reduction process. The purple ball represents the K⁺ ion.

**Fig. 5** (a) W L₃-edge XANES and the corresponding second-derivative spectra of HWO and Mo–HWO. (b) W 4f XPS of HWO and Mo–HWO.
are present in HWO, which is consistent with the hexagonal tungsten oxide with a formula of \( \text{AW}^{6+} \text{W}^{5+} \text{O}_{18} \), where A represents H\(^+\) and/or NH\(_4^+\) in the tunnels for charge balance.\(^{39}\) After the Mo framework substitution, the average oxidation states of W slightly decrease, in line with the decrease of the 10\(Dq\) value obtained from the XANES spectra at the W L\(_2\)-edge (Fig. 5a). Note that an extra XPS peak appears at a low binding energy of \( \sim 3.31 \) eV. After curve-fitting, the peak can be assigned to the W\(^{4+}\) 4f\(_{7/2}\) XPS line.\(^{39}\) Consequently, the W species in Mo–HWO include W\(^{6+}\), W\(^{5+}\), and W\(^{4+}\) (see Table S3† for more details), i.e., the electronic states of W in Mo–HWO are tuned by the Mo framework substitution.

The co-presence of W\(^{6+}\), W\(^{5+}\), and W\(^{4+}\) may be important for improving the SCR activity. Similar standard reduction potentials, such as \( \phi^0 \), \( \phi^0(\text{W}^{6+}/\text{W}^{5+}) = -0.03 \) V and \( \phi^0(\text{W}^{5+}/\text{W}^{4+}) = -0.04 \) V,\(^{23}\) allow W to transfer between different oxidation states, thus enhancing the redox ability, as evidenced by the LT onset of the H\(_2\)-TPR profile of Mo–HWO (Fig. 4). Meanwhile, Mo\(^{6+}\) and Mo\(^{5+}\) co-exist in Mo–HWO as shown in the Mo 3d XPS in Fig. S9 and Table S3†. Likewise, similar \( \phi^0 \) values of Mo\(^{5+}/\text{Mo}^{6+}(0.40 \text{ V})\)\(^{23}\) and W\(^{6+}/\text{W}^{4+}(0.39 \text{ V})\)\(^{24}\) might make the reaction \( \text{W}^{6+} + 2\text{Mo}^{5+} = \text{W}^{4+} + 2\text{Mo}^{6+} \) occur more easily, thus accelerating the electron transfer via the Mo–O–W bridge in Mo–HWO. Although the reduction potentials are obtained in aqueous solution, Owen \textit{et al.} suggested that the reduction potentials of many cations in the solid states also follow a similar sequence.\(^{46}\)

To shed light on the electronic states of the CASs of Mo–HWO, the projected density of states (DOS) of HWO and Mo–HWO are calculated by DFT (Fig. 6). For HWO, the DOS of O 2p and W 5d appear in the same energy regimes, implying the presence of W–O hybridization. Thus, the bandgap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) is \( \sim 2.5 \) eV. The bandgap is then reduced to \( \sim 1.2 \) eV after the Mo framework substitution mainly caused by a downshift in energy of the LUMO, which is consistent with the experimental data that the absorption edge shifts to the low energy region after the incorporation of Mo (Fig. 5a). Likewise, the hybridization between the W and Mo cations through interactions with the bridging O atoms implicitly occurs because the DOS of W, Mo, and O in Mo–HWO are almost in the same energy regimes. Thus, owing to the Mo–O–W hybridization and the narrow bandgap between HOMO–LUMO, the electron transfer between the W and Mo \textit{via} the bridging O is expected to become relatively easy,\(^{47}\) leading to the co-existence of several oxidation states of W and Mo. In all, the Mo framework substitution reduces the bandgap between the HOMO and LUMO and increases the redox ability of the CASs, thus being favorable for SCR.

4. Conclusions

In conclusion, we tuned the electronic states of the catalytically active sites of HWO by Mo framework substitution. The Mo–HWO catalyst showed high activity in SCR, even in the presence of high-concentration SO\(_2\) (2700 mg m\(^{-3}\)) and 10 vol\% H\(_2\)O. The SXRD, TEM, \(^1\)H MAS NMR, and H\(_2\)-TPR data demonstrated that the tunnel openings on the (001) facets of the Mo–HWO rods are the catalytically active sites, \textit{i.e.}, they consist of hexagonal hollow sites surrounded by six W\(_6\) and Mo\(_6\) octahedra. XANES and XPS data in conjunction with DFT calculations exhibited that the Mo framework substitution reduces the bandgap between the HOMO and LUMO, leading to the co-existence of multivalent W and Mo cations in Mo–HWO by hybridizing W and Mo with the bridging oxygen ions of W–O–Mo. The decrease in bandgap and the co-existence of multivalent metal cations may make the electrons produced from the SCR redox cycles transfer between the reactants and the CASs more easily, thus enhancing the activity.

Acknowledgements

This work was financially supported by the NSFC (21477023) and the STCSM (14JC1400400). The SXRD measurements were conducted at the SSRF.

Notes and references

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