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# CoMoO<sub>4</sub> as a novel heterogeneous catalyst of peroxymonosulfate activation for the degradation of organic dyes†

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Sulfate radical-based advanced oxidation processes (SR-AOPs) are receiving more and more attention for the removal of recalcitrant organic pollutants. In this study, we employ  $CoMoO_4$  as a novel heterogeneous catalyst for peroxymonosulfate (PMS) activation to release powerful sulfate radicals for the first time. The  $CoMoO_4$ , prepared through a hydrothermal route and high-temperature calcination, displays a hierarchical microstructure assembled from ultrathin nanosheets and a large surface area (61.9 m² g⁻¹). Methylene blue (MB) is selected as a model organic pollutant, and it is found that the  $CoMoO_4$ /PMS system can realize 100% degradation of MB in 40 min and maintain its removal efficiency during three recycling experiments. Such a catalytic performance of  $CoMoO_4$  is indeed superior to those of conventional  $Co_3O_4$  and  $CoFe_2O_4$ . The effects of some potential influential factors, including reaction temperature, dosages of PMS and  $CoMoO_4$  and the initial pH value are systematically evaluated. More importantly, the  $CoMoO_4$ /PMS system not only shows its universality in the degradation of other organic dyes (e.g. orange II and rhodamine B), but also exhibits considerable degradation efficiency under some actual water background conditions. The quenching experiments confirm that sulfate radicals are the main active species for the degradation of dyes, and XPS spectra reveal that Co sites on the surface of  $CoMoO_4$  are the primary active sites for the generation of sulfate radicals.

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#### Introduction

Wastewater from textile, printing, and other industries usually contains various dyes in high-concentration, which can produce considerable adverse effects on eco-environments due to their toxicity and recalcitrance.1 In the past few decades, some conventional treatments, such as physical, chemical and biological techniques, have been utilized for environmental remediation, while their effectiveness in the removal of organic dyes is still far away from the expected performance.2,3 As an attractive alternative, advanced oxidation processes (AOPs) offer immense potential for overcoming the challenges related to the deep purification of wastewater owing to their excellent oxidation ability and mineralization efficiency. 4,5 Hydroxyl radicals ('OH) are the classic reactive species in AOPs, and their notable reactivity is almost applicable to all organic compounds.6 However, their practical applications always suffer from many limitations, e.g. sophisticated and cost-ineffective radical

generation.7-9 Recently, sulfate radicals ('SO<sub>4</sub>-) have received intensive research interests as another powerful reactive radicals, because they can not only compensate some critical inadequacies of 'OH, but also provide higher standard reduction potential (2.5-3.1 V) under neutral conditions than 'OH (1.8-2.7 V). It is well known that 'SO<sub>4</sub> can be released from peroxymonosulfate (PMS) or persulfate (PS) by heating, UV irradiation, ultrasonication, and catalytic activation, where catalytic activation has its own advantages without the assistance of necessary equipment and high-energy input. 6,11-13 Compared with PS, PMS is a preferable candidate for the generation of 'SO<sub>4</sub> in catalytic system, because its asymmetrical molecular structure and relatively large O-O bond length in free molecules will facilitate the activation process.14 Literature review reveals that homogeneous catalytic activation of PMS by some transition metal ions, especially Co<sup>2+</sup>, is very effective for the generation of 'SO<sub>4</sub>-, 15 while the secondary pollution caused by the high solubility and great toxicity of transition metal ions will severely hinder the application of homogeneous catalysis in water treatment.16 As a result, it is of great importance to develop high-performance heterogeneous catalysts to alleviate the potential secondary contamination from the homogeneous systems.

generation processes, pH adjustments and potential sludge

To date, many kinds of heterogeneous catalysts, including zero-valence metal, oxides, supported catalysts, and carbon

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 $\dagger$  Electronic supplementary information (ESI) available: SEM image of precursor;  $N_2$  adsorption/desorption isotherm of CoMoO4; MB degradation in recycling experiments, MeOH solution, and actual water bodies; RhB and OII degradations catalyzed by CoMoO4/PMS; the compositions of tap water and surface water. See DOI: 10.1039/c7ra04761d

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materials, have been continuously reported as promising PMS activators.8,17-20 Compared with free metal ions, immobilization of active sites in heterogeneous catalysts makes them lose the characteristic of single atom catalysis, resulting in inferior activities than those homogeneous catalysts. However, it is still found that some Co-containing heterogeneous catalysts can produce better catalytic performance in PMS activation than their analogous counterparts.21-23 For example, Lin and coworkers compared the catalytic activities of Prussian blue analogues (PBAs) with various metal ions (e.g. Co, Cu, Fe, Mn, Ni), and the results indicated that Co-containing PBAs exhibited excellent capability for activating PMS, while Cu, Fe and Ni based PBAs did not provide conspicuous catalytic activity to activate PMS;24 Ren et al. evaluated the performance of magnetic ferrospinel, including CoFe<sub>2</sub>O<sub>4</sub>, CuFe<sub>2</sub>O<sub>4</sub>, MnFe<sub>2</sub>O<sub>4</sub> and ZnFe<sub>2</sub>O<sub>4</sub>, in PMS activation, and they found that CoFe<sub>2</sub>O<sub>4</sub> had the highest catalytic performance among these ferrites towards PMS for the degradation of di-n-butyl phthalate.<sup>21</sup> In most previous studies on heterogeneous PMS activation, there was a consensus on the possible reaction mechanism, that was, electron transfer between M2+/M3+ and HSO5-/.SO4- would be responsible for the generation of 'SO<sub>4</sub>-. <sup>25,26</sup> Therefore, it may be an effective way to further enhance the effectiveness of PMS activation by employing a Co-containing heterogeneous catalyst with improved electron transfer features.

More recently, CoMoO<sub>4</sub> has appeared as a novel and promising pseudo-capacitor, and its unique advantages, such as great redox activity, stable crystalline structure, high electrical conductivity, as well as fast transport between ions and electrons, are highly desirable for the excellent electrochemical performance.27-29 It is very exciting that these characteristics also cater to PMS activation, and thus the catalytic performance of CoMoO<sub>4</sub> in the degradation of organic pollutants can be expected. Unfortunately, there are very few correlative literatures accessible. In this article, we employ CoMoO4 as a heterogeneous catalyst in PMS activation for the first time. The organic dye, methylene blue (MB), is selected as a model pollutant. It is found that CoMoO4 displays much better MB degradation than common CoFe<sub>2</sub>O<sub>4</sub> and Co<sub>3</sub>O<sub>4</sub>, and its good catalytic activity is also versatile for the degradation of other dyes [orange II (OII) and rhodamine B (RhB)], even under some actual water background conditions. We believe that CoMoO<sub>4</sub> can be an efficient and green heterogeneous catalyst for the degradation of organic pollutants, and this study provides new insights into the development of alternative catalysts for wastewater treatment.

### **Experimental**

#### Materials synthesis and characterization

CoMoO<sub>4</sub> was synthesized through a hydrothermal route according to previous literature.30 Briefly, 0.119 g (0.5 mmol) of CoCl<sub>2</sub>·6H<sub>2</sub>O, 0.877 g (0.071 mmol) of (NH<sub>4</sub>)<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub>·4H<sub>2</sub>O and 0.3003 g (5 mmol) of urea were dissolved in 36 mL of deionized (DI) water. This mixture was vigorously stirred at room temperature for 30 min to form a homogeneous solution. The transparent solution was then transferred into a Teflon-lined stainless steel autoclave and sealed to heat at 120 °C for 12 h.

The obtained purple particles were collected by centrifugation, and washed several times with DI water and ethanol till the filtrate became colourless. After drying at 60 °C for 12 h, the asprepared precursor was further calcined at 500 °C for 3 h with a heating rate of 1 °C min<sup>-1</sup> under air atmosphere to produce highly crystalline CoMoO<sub>4</sub>. For comparison, CoFe<sub>2</sub>O<sub>4</sub> and Co<sub>3</sub>O<sub>4</sub> were also prepared by co-precipitation method and hightemperature calcination under the same conditions.

Powder X-ray diffraction (XRD) pattern of the as-prepared samples was obtained at room temperature with a D/MAXRC X-ray diffractometer using Cu Kα radiation source which operated at 45 kV and 40 mA. The structure and morphology of the sample was performed by scanning electron microscopy (SEM, Quanta 200S) and transmission electron microscopy (TEM, JEM-3000F). Brunauer-Emmett-Teller (BET) surface area data were measured by a QUADRASORB SI-KR/MP (Quantachrome, USA). Samples were normally treated for porosity measurement by degassing at 200 °C for 4 h with a N<sub>2</sub> flow. X-ray photoelectron spectroscopy (XPS) was collected using PHI 5700 ESCA System with an Al Kα radiation as the X-ray source. The absorbance of different dye solution was determined by using a UV-Vis diffuse reflectance spectrophotometer (TU-1901). Total organic carbon (TOC) was determined by Analytik Jena AG MultiN/C 2100 TOC analyzer.

#### Catalytic test procedure

All the catalytic experiments were evaluated in 100 mL beaker containing 50 mL of pollutant solution at  $25 \pm 1$  °C. In a typical run, 5.0 mg of catalysts was dispersed into 50 mL MB solution  $(100 \text{ mg L}^{-1})$  under constant magnetic stirring (the agitation speed was fixed at 400 rpm) for about 30 min to achieve the adsorption-desorption balance. The oxidative process was initiated after the introduction of a certain amount of PMS (2.0 mM) [Note: PMS concentration was calculated from its actual content in commercially available oxone (KHSO<sub>5</sub>·0.5KHSO<sub>4</sub>-·0.5K<sub>2</sub>SO<sub>4</sub>]. At given time intervals, 0.4 mL of the reaction mixture was withdrawn and mixed with saturated Na2S2O3 solution to quench the reaction. The concentration of MB was analyzed using UV spectrophotometer at  $\lambda = 664$  nm, and the corresponding degradation efficiencies were obtained according to the following equation:

Degradation efficiency (%) = 
$$[(C_0 - C_t)/C_0] \times 100\%$$
 (1)

where  $C_t$  is the real-time concentration and  $C_0$  is the designed concentration. For the recycling test of CoMoO4, several identical reactions were performed simultaneously, and the used catalysts were separated by centrifugation, washed by deionized water and ethanol, and calcined at 500 °C for reuse. For studying the effect of initial pH value on MB degradation, NaOH (1.0 mM) and HCl (1.0 mM) were employed as the pH regulators. In addition, several water bodies were applied as actual water backgrounds to evaluate the practical application of CoMoO<sub>4</sub>, where tap water was from a drinking water treatment plant in Heilongjiang Province, China, and surface water was from Songhua River in Harbin, China.

#### Results and discussion

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The crystalline structure of the as-prepared CoMoO $_4$  is primarily studied by wide angle XRD (Fig. 1), and the diffraction peaks in the  $2\theta$  range of 10– $60^\circ$  can be well matched with  $\beta$ -CoMoO $_4$  phase (JCPDS 21-0868). Although CoMoO $_4$  can occur in the forms of  $\alpha$ -phase and  $\beta$ -phase, high temperature (330–410 °C) will induce the transformation from  $\alpha$ -CoMoO $_4$  to  $\beta$ -CoMoO $_4$ . In our case, the calcination temperature is 500 °C, and thus the characteristic peaks of  $\alpha$ -CoMoO $_4$  (JCPDS 25-1434) are not detected in final product. In addition, the absence of characteristic peaks of possible impurities, such as Co $_3$ O $_4$  (JCPDS 42-1467) and MoO $_3$  (JCPDS 47-1320 and 47-1081) further validates the formation of high-purity  $\beta$ -CoMoO $_4$  through hydrothermal treatment and high-temperature calcination.

SEM and TEM measurements were conducted in order to confirm the morphology and microstructures of the as-prepared product. Before calcination in the muffle furnace, the precursor holds the relatively uniform spheres with wrinkle surface and an average diameter of about 2.5 μm, as shown in Fig. S1.† It can be clearly seen that each flower-like microsphere is assembled by numerous ultrathin nanosheets, and these curved and corrugated nanosheets are connected to each other, forming a stable hierarchical microstructure. Very interestingly, SEM image reveals that this hierarchical microstructure can be well preserved during the high-temperature calcination, and the asprepared CoMoO4 still exhibits flower-like morphology except for a slight shrinkage in the average diameter (ca. 2.0 µm). It has to mention that this hierarchical microstructure in CoMoO<sub>4</sub> may provide abundant exposure of active sites, which will be quite beneficial to the catalytic process. TEM results are consistent with SEM data, further confirming that these microspheres are constituted by sequential nanosheets with a relatively thin thickness (Fig. 2b and c). The high-resolution TEM (HRTEM) image of the edge region identifies the lattice fringe of these nanosheets of about 0.336 nm, corresponding to the (002) plane of β-CoMoO<sub>4</sub>. Fig. S2† shows N<sub>2</sub> adsorptiondesorption isotherm of CoMoO<sub>4</sub>. As observed, this isotherm performs its profile between standard II-type and IV-type

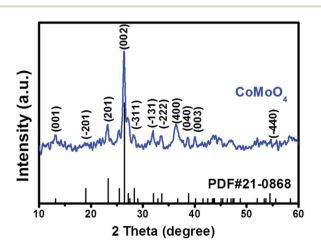


Fig. 1 XRD pattern of the as-prepared CoMoO<sub>4</sub>.

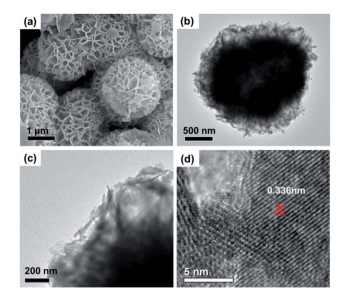


Fig. 2 SEM image (a), TEM images (b and c), and HR-TEM image (d) of  $CoMoO_4$ .

isotherms, characteristic of disordered mesoporous structure in the material according to the IUPAC classification.<sup>32</sup> The corresponding BET surface and pore volume are  $61.9 \text{ m}^2 \text{ g}^{-1}$  and  $0.44 \text{ cm}^3 \text{ g}^{-1}$ , respectively, and these superior structural parameters after high-temperature calcination may be associated with its unique hierarchical microstructure (Fig. 2).

Fig. 3 shows the degradation of MB *via* catalytic oxidation process with different catalysts. As observed, sole PMS can work for the decoloration of MB solution, and 32.2% of MB is removed after 60 min. CoMoO<sub>4</sub> fails to promise any MB removal in the absence of PMS, suggesting that the contribution from simple physical adsorption is negligible. It is very interesting that the removal of MB can be substantially reinforced with the presence of both PMS and CoMoO<sub>4</sub>, and complete degradation of MB can be achieved in 40 min. In previous reports, CoMoO<sub>4</sub>

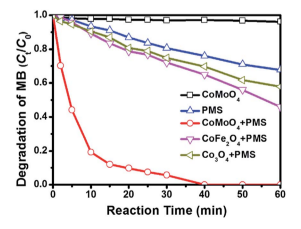


Fig. 3 MB degradation by various catalytic PMS systems. Conditions: [MB] $_0=100~{\rm mg~L^{-1}}$ , volume (MB) = 50 mL, [Oxone] = 2 mM, catalyst amount = 0.05 g (0.10 g L $^{-1}$ ), temperature = 25 °C, without pH adjustment.

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is rarely studied as a catalyst for heterogeneous PMS activation, and thus the degradation of MB is also performed over some common Co-containing catalysts, e.g. CoFe<sub>2</sub>O<sub>4</sub> and Co<sub>3</sub>O<sub>4</sub>, to evaluate the catalytic activity of CoMoO<sub>4</sub>. It is found that although these Co-containing catalysts can also accelerate the degradation of MB in the same system, the removal efficiencies in 60 min catalysed by CoFe<sub>2</sub>O<sub>4</sub> and Co<sub>3</sub>O<sub>4</sub> are only 54.1% and 42.2%, respectively, which are far behind that over CoMoO<sub>4</sub>. It is also observed that the degradation rate of MB in CoMoO<sub>4</sub>/ PMS system is extremely fast in the first 5 min, implying that the active radicals may be released as soon as the introduction of PMS. As mentioned above, PMS activation is always accompanied by the electron transfer between active sites and HSO<sub>5</sub><sup>-</sup>/ 'SO<sub>4</sub>-, and thus the good performance of CoMoO<sub>4</sub> should be linked with its better electrochemical properties as compared with CoFe<sub>2</sub>O<sub>4</sub> and Co<sub>3</sub>O<sub>4</sub>.<sup>28,33</sup> What merits a special note is that CoMoO<sub>4</sub> not only provides high MB degradation efficiency, but also performs good mineralization capacity (Fig. S3†). After 60 min, TOC removal can reach 79.6%. By considering that Co sites are generally taken as the primary active sites for PMS activation, we also evaluate the catalytic activities of Co sites in Co<sub>3</sub>O<sub>4</sub>, CoFe<sub>2</sub>O<sub>4</sub>, and CoMoO<sub>4</sub> with turn over number (TON, the number of chemical conversions of substrate molecules on a single catalytic site in unit time). In terms of the degradation efficiencies of MB in 40 min, the TON values for Co<sub>3</sub>O<sub>4</sub>,  $CoFe_2O_4$ , and  $CoMoO_4$  can be deduced as  $9.25 \times 10^{-3}$ ,  $3.14 \times 10^{-3}$ 

 $10^{-2}$ , and  $5.48 \times 10^{-2}$ , respectively (Table 1). It is undoubted

that Co sites in CoMoO<sub>4</sub> display the best catalytic ability. In

general, the degradation of organic dye obeys the pseudo-first

order kinetics, and thus the kinetics constant over different

catalysts can be calculated by the following equation,

$$ln(C_t/C_0) = -kt$$
(2)

where  $C_t$  and  $C_0$  are the real-time concentration and the initial concentration, respectively, and k is the apparent first order rate constant of MB removal, and t is the reaction time. The results show that the kinetic rate constant of  $CoMoO_4$  (0.144  $min^{-1}$ ) is at least 11.5 and 16.1 times higher than those of  $CoFe_2O_4$  (0.0125  $min^{-1}$ ) and  $Co_3O_4$  (0.00895  $min^{-1}$ ), respectively (Table 1). To further address the superiority of  $CoMoO_4$ , we also collect the performances of various catalysts in PMS activation for MB degradation (Table S1†). Although it is unlikely for a comparison of the catalytic performances from different experimental conditions,  $CoMoO_4$  can realize the complete removal of high-concentration MB in a short time with lower relative dosages of PMS and catalyst, which may be considered as an indirect

Table 1 TON and kinetic rate constants of MB degradation over various catalysts

Catalysts	$TON^a$	$k  (\mathrm{min^{-1}})$
${ m Co_3O_4} \ { m CoFe_2O_4} \ { m CoMoO_4}$	$9.25 \times 10^{-3}$ $3.43 \times 10^{-2}$ $5.48 \times 10^{-2}$	$8.95 \times 10^{-3}$ $1.25 \times 10^{-2}$ $1.44 \times 10^{-1}$

<sup>&</sup>lt;sup>a</sup> TON is calculated by the degradation efficiency of MB at 40 min.

evidence of highly catalytic activity of  $CoMoO_4$ . The repeated batch experiments are carried out to investigate the catalytic stability of  $CoMoO_4$  (Fig. S4†), and the results demonstrate that the regenerated catalyst can still present good MB degradation over 94.0% in the second and third runs. Very interestingly, the used catalyst still shows typical  $\beta$ -phase and flower-like morphology, especially its BET surface area (56.0 m² g⁻¹) is very close to that of the fresh  $CoMoO_4$  (Fig. S5†). The stable crystallinity and microstructure may provide a good platform for the recovery of used  $CoMoO_4$  catalyst.

It is well known that reaction temperature is a critical operating parameter in AOPs, which can affect the degradation rate of organic pollutants greatly. The degradation of the reaction temperature also plays an important role in determining the degradation rate of MB. When the reaction temperature is increased from 15 °C to 35 °C, the kinetic rate constants will be increased from 0.056 min<sup>-1</sup> to 0.430 min<sup>-1</sup> (Table S2†). Furthermore, the activation energy ( $E_a$ ) of the reaction on the surface of CoMoO<sub>4</sub> is evaluated by plotting  $\ln k$  versus 1/T based on the Arrhenius equation and is determined to be 69.89 kJ mol<sup>-1</sup> (Fig. 4b). The  $E_a$  value is much higher than that of diffusion-controlled reaction (10–13 kJ mol<sup>-1</sup>), indicating that the apparent reaction rate of this oxidation process is more dependent on the rate of intrinsic chemical reactions on the surface of CoMoO<sub>4</sub> rather than the rate of mass transfer.

In addition, the effects of catalyst dosage and PMS dosage are further studied in CoMoO<sub>4</sub>/PMS system. As shown in Fig. 4c, with increased dosage of PMS, the degradation efficiency of MB will rise from 69.5% at 0.5 mM PMS to 96.3% at 2.5 mM in 30 min. It is reliable that increasing PMS concentration can make more HSO<sub>5</sub><sup>-</sup> attach to the active sites of CoMoO<sub>4</sub>, which facilitates the generation of radical species. Interestingly, MB degradation shows a similar dependence on the dosage of CoMoO<sub>4</sub> (Fig. 4d). When the dosage of CoMoO<sub>4</sub> is set at 0.02 g  $L^{-1}$ , the degradation efficiency of MB will be 95.3% in 60 min. For the dosage of 0.40 g L<sup>-1</sup>, MB can be completely eliminated in 25 min, and its kinetic rate constant (0.1597 min<sup>-1</sup>) is about 2.36 times higher than that at  $0.02 \text{ g L}^{-1}$  (0.0677 min<sup>-1</sup>). However, it is also found that the stimulation effect will be strongly restrained once the dosage of CoMoO4 is beyond 0.40 g L<sup>-1</sup>. According to previous literature, <sup>38-40</sup> this phenomenon can be explained by the diffusion limitation in heterogeneous reactions. If the dosage of the catalysts exceeds the optimum value, the ineffective oxidant consumption on its surface will be accelerated and become dominant before radical species can react with organic pollutant molecules, leading to the constant or decreased degradation efficiency. In the studied interval of dosage, the diffusion limitation of CoMoO<sub>4</sub> is determined as  $0.40 \mathrm{~g~L^{-1}}$ , which is higher than those reported in some oxides and ferrites, 41,42 suggesting that CoMoO4 may be a better candidate as heterogeneous catalysts for PMS activation.

As reported in many previous studies, sulfate radical-based AOPs can work in a wider pH range as compared to conventional Fenton reaction. This deduction is also applicable to CoMoO<sub>4</sub>/PMS system (Fig. 5a). It is observed that the degradation efficiency of MB increases sharply as the initial pH value rises from 3.0 to 11.0. However, when the pH value further

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(a) (b) In (C/C<sub>0</sub>) 25 °C 30 °C 35 °C ᆂᄰ E\_=69.89KJ/mol 10 12 0.0032 0.0033 0.0034 0.0035 Reaction Time (min) 1/T (d)1 Degradation of MB(C,/C,) Degradation of MB(C,/C) [PMS]=0.5mM = catalyst 0.02g/L [PMS]=1.0mM [PMS]=1.5mM [PMS]=2.0mM catalyst 0.60g/L IPMS1=2.5mM

Fig. 4 Kinetic curves (a) and the Arrhenius curve (b) of MB removal at different temperature in  $CoMoO_4/PMS$  system. Conditions:  $[MB]_0 = 100 \text{ mg L}^{-1}$ , volume (MB) = 50 mL, [Oxone] = 2 mM, catalyst amount = 0.05 g  $(0.10 \text{ g L}^{-1})$ , temperature  $= 25 \,^{\circ}\text{C}$ , without pH adjustment. Effects of oxone concentration (c) and  $CoMoO_4$  dosage (d) on MB removal catalyzed by  $CoMoO_4/PMS$  system.

Reaction Time (min)

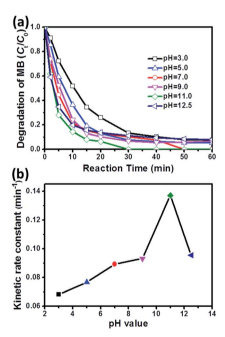


Fig. 5 Effect of initial pH value on MB degradation (a) and kinetic rate constant (b) in CoMoO<sub>4</sub>/PMS system. Conditions: [MB] $_0$  = 100 mg L $^{-1}$ , volume (MB) = 50 mL, [Oxone] = 2 mM, catalyst amount = 0.05 g (0.10 g L $^{-1}$ ), temperature = 25 °C.

increases to 12.50, the degradation rate is decreased reversely. The corresponding kinetic rate constants (k) at different initial pH values are deduced in Fig. 5b, which clearly demonstrates the effect of initial pH value on MB degradation catalyzed by  $CoMoO_4/PMS$  system. This phenomenon can be explained from

two aspects: (1) pH value has a great impact on the speciation of PMS, and acidic condition can induce the formation of strong hydrogen bond between H<sup>+</sup> and O–O bond in PMS, <sup>6,8</sup> which will obviously inhibit the interaction between PMS and CoMoO<sub>4</sub>; (2) the hydrogen bond will be continuously weakened with increased pH value, and meanwhile, hydroxyl radicals will be inevitably introduced into the catalytic system under alkaline condition, which can not only attack organic pollutants, but also react with other species to stimulate PMS activation. <sup>45,46</sup> As a result, the degradation of MB will be reinforced in the pH range of 3.0–11.0. However, excessive high basicity (*e.g.* pH = 12.5) will produce less oxidative  $SO_5^{2-}$  through the deprotonation of  $HSO_5^{-}$ , <sup>47</sup> leading to an inferior MB degradation.

Reaction Time (min)

It has been reported that both 'SO<sub>4</sub> and 'OH can be identified in conventional PMS activation system, because 'OH is a typical product from the reaction between 'SO<sub>4</sub> and H<sub>2</sub>O.<sup>43</sup> Therefore, the classical quenching tests are carried out to figure out the specific reaction mechanism in the system of MB degradation, where tert-butyl alcohol (TBA) is selected as a particular scavenger for 'OH and MeOH is utilized as a universal scavenger for both 'OH and 'SO4-.40,48 As shown in Fig. 6a and b, the degradation efficiency of MB displays an anticipated decrease with increasing the concentration of TBA or MeOH, and more importantly, MeOH always plays a more aggressive role in inhibiting MB degradation as compared with TBA. For example, when the concentration of TBA increases from 0 to 300 mM, the degradation efficiency of MB will decrease from 100% to 90.3% in 40 min. In contrast, the degradation efficiency of MB decreases from 100% to 77.2% in the presence of MeOH (300 mM). It is worth noting that the extremely high MeOH concentration (300 mM) does not **RSC Advances** 

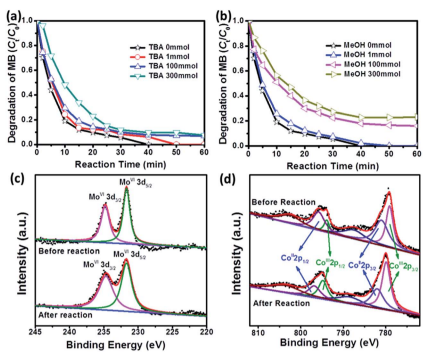


Fig. 6 Effect of (a) TBA and (b) MeOH on MB degradation by CoMoO<sub>4</sub>/PMS system. Conditions:  $[MB]_0 = 100 \text{ mg L}^{-1}$ , volume  $[MB]_0 = 100 \text{ mg}$ [Oxone] = 2 mM, catalyst amount  $= 0.05 \text{ g} (0.10 \text{ g L}^{-1})$ , temperature  $= 25 ^{\circ}\text{C}$ , without pH adjustment. XPS patterns of Mo 3d (c), Co 2p (d) in fresh and used CoMoO<sub>4</sub> sample.

suppress MB degradation completely, implying that CoMoO<sub>4</sub>/ PMS can release sufficient radicals to react with MB. To rule out the contribution of radicals completely, we further perform the degradation of MB in absolute MeOH solution, and only 11.4% of MB can be removed in 60 min (Fig. S6†), which suggests that the degradation of MB through non-radical pathway is rather limited.49,50

The existing states of elements Co and Mo in fresh and used CoMoO<sub>4</sub> are characterized by XPS to discern the primary functional sites for PMS activation. As shown in Fig. 6c, Mo 3d spectrum of fresh CoMoO<sub>4</sub> exhibits two peaks with binding energy at 234.7 eV and 231.0 eV, which can be assigned to Mo 3d<sub>3/2</sub> and Mo 3d<sub>5/2</sub>, respectively.<sup>51,52</sup> After degradation, the profile of Mo 3d in used CoMoO<sub>4</sub> is quite similar to that of fresh CoMoO<sub>4</sub>, and no peak shift or additional signal can be detected, indicating that Mo sites keep their initial state during the catalytic process. In contrast, the profiles of Co 2p in fresh and used CoMoO4 are a little different. In order to better illustrate the variation, the fitting curves of Co  $2p_{1/2}$  and Co  $2p_{3/2}$ , as well as their satellite peaks in fresh and used CoMoO4 are presented in Fig. 6d, where the peaks at binding energies of 779.9 eV and 794.8 eV with the satellite signal at 788.5 eV are due to Co(III), and peaks at binding energies of 782.0 eV and 796.8 eV with the satellite signal at 803.5 eV are characteristic of Co(II).53,54 It is clear that the proportion of Co(III) increases from 39.9% in fresh catalyst to 59.8% in used catalyst. This phenomenon means that Co sites in CoMoO<sub>4</sub> still play the primary functional sites for PMS activation.

According to the above experimental results, the possible mechanism of radicals generation in

decomposition of MB can be depicted as follows. 55,56 Firstly, the complex Co(II)-(OH)OSO<sub>3</sub> is generated by the complexation between PMS and Co sites on the surface of CoMoO4 as shown in eqn (3). By means of one-electron transfer inside the complex, 'SO<sub>4</sub>" is generated and new hydroxyl groups are formed (eqn (4)). Co(III) then oxidized PMS to  $SO_5$  that further attaches to the hydroxyl groups, producing the surface peroxo species, and reaching a lower transient valence (eqn (5)). Then, the surface peroxo species combine with each other, and recycled hydroxyl groups bonded to Co(II) are obtained (eqn (6) and (7)).

$$\equiv \text{Co(II)} - \text{OH} + \text{HSO}_5^- \rightarrow \equiv \text{Co(II)} - \text{(OH)OSO}_3^- + \text{OH}^-$$
 (3)

$$\equiv \text{Co(II)-(OH)OSO}_3^- \rightarrow \equiv \text{Co(III)--OH + SO}_4^{\bullet-}$$
 (4)

$$\equiv \text{Co(III)} - \text{OH} + \text{HSO}_5^- \rightarrow \equiv \text{Co(II)} - \text{OOSO}_3^- + \text{H}_2\text{O}$$
 (5)

$$2 \equiv \text{Co(II)} - \text{OOSO}_3^- + 2\text{H}_2\text{O} \rightarrow 2 \equiv \text{Co(II)} - \text{OH} + \text{O}_2 + \text{S}_2\text{O}_8^{2-} + 2\text{H}^+$$
 (7)

To test the generality of CoMoO<sub>4</sub>/PMS system, two additional dyes, RhB and OII, are further selected as model pollutants. As shown in Fig. S7,† sole PMS does not account for the obvious degradation for both RhB and OII, while CoMoO4/PMS removes over 80% of dyes in 30 min, confirming that this heterogeneous activation system can work for the degradation of different types of organic dyes. In addition, we also perform the degradation of MB in two actual water bodies (tap water and surface water), whose compositions are listed in Table S3.† It is very interesting that CoMoO<sub>4</sub>/PMS system can maintain its high MB removal efficiency in these actual water bodies, and its performance is always superior to that of CoFe<sub>2</sub>O<sub>4</sub>/PMS (Fig. S8†). In view of the results in this study, we believe that CoMoO<sub>4</sub> can be a promising heterogeneous catalyst for PMS activation with a bright prospect in practical application.

#### Conclusions

Paper

In this study, CoMoO<sub>4</sub> is employed as a novel heterogeneous catalyst for PMS activation for the first time. The as-prepared CoMoO<sub>4</sub> through hydrothermal route and high-temperature calcination displays hierarchical microstructure that is assembled by ultrathin nanosheets and high surface area (61.9 m<sup>2</sup>) g<sup>-1</sup>). CoMoO<sub>4</sub> exhibits excellent catalytic performance in activating PMS for MB degradation, which is much better than conventional Co<sub>3</sub>O<sub>4</sub> and CoFe<sub>2</sub>O<sub>4</sub>. Radical quenching tests reveal that radical pathway is dominant in the degradation of MB, and both sulfate and hydroxyl radicals will be released from PMS activation. The results of XPS spectra confirm that Co sites on the surface of CoMoO4 are primarily active sites. More importantly, it is found that the excellent catalytic performance of CoMoO<sub>4</sub> is not only versatile for the degradation of different dyes, but also available in some actual water bodies. We believe CoMoO<sub>4</sub> may be an advanced alternative to conventional heterogeneous catalysts for PMS activation in advanced oxidation processes.

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#### References

- 1 S. K. Ling, S. Wang and Y. Peng, *J. Hazard. Mater.*, 2010, **178**, 385–389.
- 2 T. Robinson, G. McMullan, R. Marchant and P. Nigam, *Bioresour. Technol.*, 2001, 77, 247–255.
- 3 K. Pazdzior, A. Klepacz-Smolka, S. Ledakowicz, J. Sojka-Ledakowicz, Z. Mrozinska and R. Zylla, *Chemosphere*, 2009, 75, 250–255.
- 4 X. G. Duan, H. Q. Sun, Y. X. Wang, J. Kang and S. B. Wang, *ACS Catal.*, 2015, 5, 553–559.
- 5 Y. J. Yao, H. Chen, C. Lian, F. Y. Wei, D. W. Zhang, G. D. Wu, B. J. Chen and S. B. Wang, *J. Hazard. Mater.*, 2016, 314, 129– 139.
- 6 Z. Wang, Y. Du, Y. Liu, B. Zou, J. Xiao and J. Ma, *RSC Adv.*, 2016, **6**, 11040–11048.
- 7 J. A. Zazo, J. A. Casas, A. F. Mohedano, M. A. Gilarranz and J. J. Rodriguez, *Environ. Sci. Technol.*, 2005, **39**, 9295–9302.
- 8 Y. Du, W. Ma, P. Liu, B. Zou and J. Ma, *J. Hazard. Mater.*, 2016, 308, 58–66.

- 9 Y.-P. Zhu, T.-Z. Ren and Z.-Y. Yuan, *RSC Adv.*, 2015, 5, 7628–7636.
- 10 Z. Huang, H. Bao, Y. Yao, J. Lu, W. Lu and W. Chen, *J. Chem. Technol. Biotechnol.*, 2016, **91**, 1257–1265.
- 11 Y.-H. Guan, J. Ma, X.-C. Li, J.-Y. Fang and L.-W. Chen, Environ. Sci. Technol., 2011, 45, 9308–9314.
- 12 M. Feng, R. Qu, X. Zhang, P. Sun, Y. Sui, L. Wang and Z. Wang, *Water Res.*, 2015, **85**, 1–10.
- 13 Z. Fang, X. Qiu, J. Chen and X. Qiu, *Appl. Catal., B*, 2010, **100**, 221–228.
- 14 X. Duan, Z. Ao, H. Sun, L. Zhou, G. Wang and S. Wang, *Chem. Commun.*, 2015, **51**, 15249–15252.
- 15 G. P. Anipsitakis and D. D. Dionysiou, *Environ. Sci. Technol.*, 2004, **38**, 3705–3712.
- 16 Z. Xu, J. Lu, Q. Liu, L. Duan, A. Xu, Q. Wang and Y. Li, *RSC Adv.*, 2015, **5**, 76862–76874.
- 17 Y. B. Ding, L. H. Zhu, A. Z. Huang, X. R. Zhao, X. Y. Zhang and H. Q. Tang, *Catal. Sci. Technol.*, 2012, **2**, 1977–1984.
- 18 W.-D. Oh, S.-K. Lua, Z. Dong and T.-T. Lim, *J. Hazard. Mater.*, 2015, **284**, 1–9.
- 19 Q. Yang, H. Choi, Y. Chen and D. D. Dionysiou, *Appl. Catal.*, *B*, 2008, 77, 300–307.
- 20 P. R. Shukla, S. Wang, H. Sun, H. M. Ang and M. Tade, *Appl. Catal.*, B, 2010, 100, 529–534.
- 21 Y. M. Ren, L. Q. Lin, J. Ma, J. Yang, J. Feng and Z. J. Fan, *Appl. Catal.*, *B*, 2015, **165**, 572–578.
- 22 P. Nfodzo and H. Choi, Chem. Eng. J., 2011, 174, 629-634.
- 23 Q. Yang, H. Choi, S. R. Al-Abed and D. D. Dionysiou, *Appl. Catal.*, B, 2009, 88, 462–469.
- 24 K.-Y. A. Lin, B.-J. Chen and C.-K. Chen, RSC Adv., 2016, 6, 92923–92933.
- 25 S. Indrawirawan, H. Sun, X. Duan and S. Wang, *J. Mater. Chem. A*, 2015, **3**, 3432–3440.
- 26 Y. Ji, C. Dong, D. Kong and J. Lu, *J. Hazard. Mater.*, 2015, **285**, 491–500.
- 27 C. T. Cherian, M. V. Reddy, S. C. Haur and B. V. R. Chowdari, *ACS Appl. Mater. Interfaces*, 2013, 5, 918–923.
- 28 D. Guo, H. Zhang, X. Yu, M. Zhang, P. Zhang, Q. Li and T. Wang, *J. Mater. Chem. A*, 2013, 1, 7247–7254.
- 29 M.-C. Liu, L.-B. Kong, C. Lu, X.-M. Li, Y.-C. Luo and L. Kang, *Mater. Lett.*, 2013, **94**, 197–200.
- 30 M. Q. Yu, L. X. Jiang and H. G. Yang, *Chem. Commun.*, 2015, 51, 14361–14364.
- 31 X. Xu, J. Shen, N. Li and M. Ye, *J. Alloys Compd.*, 2014, **616**, 58–65.
- 32 Y. Du, T. Liu, B. Yu, H. Gao, P. Xu, J. Wang, X. Wang and X. Han, *Mater. Chem. Phys.*, 2012, **135**, 884–891.
- 33 M.-C. Liu, L.-B. Kong, X.-J. Ma, C. Lu, X.-M. Li, Y.-C. Luo and L. Kang, *New J. Chem.*, 2012, **36**, 1713–1716.
- 34 H. Sun, S. Liu, G. Zhou, H. M. Ang, M. O. Tade and S. Wang, *ACS Appl. Mater. Interfaces*, 2012, 4, 5466–5471.
- 35 B. Yang, Z. Tian, B. Wang, Z. Sun, L. Zhang, Y. Guo, H. Li and S. Yan, *RSC Adv.*, 2015, 5, 20674–20683.
- 36 S. Zhang, Q. Fan, H. Gao, Y. Huang, X. Liu, J. Li, X. Xu and X. Wang, *J. Mater. Chem. A*, 2016, **4**, 1414–1422.
- 37 L. Xu and J. Wang, Environ. Sci. Technol., 2012, 46, 10145-10153.

- 38 X. Duan, K. O'Donnell, H. Sun, Y. Wang and S. Wang, *Small*, 2015, 11, 3036–3044.
- 39 S. L. Luo, L. Duan, B. Z. Sun, M. Y. Wei, X. X. Li and A. H. Xu, *Appl. Catal.*, *B*, 2015, **164**, 92–99.
- 40 L. J. Xu, W. Chu and L. Gan, *Chem. Eng. J.*, 2015, **263**, 435-443.
- 41 K.-Y. A. Lin and B.-J. Chen, *Chemosphere*, 2017, **166**, 146–156.
- 42 P. H. Shi, R. J. Su, S. B. Zhu, M. C. Zhu, D. X. Li and S. H. Xu, *J. Hazard. Mater.*, 2012, **229**, 331–339.
- 43 G. P. Anipsitakis and D. D. Dionysiou, *Environ. Sci. Technol.*, 2003, 37, 4790–4797.
- 44 M. G. Antoniou, A. A. de la Cruz and D. D. Dionysiou, *Appl. Catal.*, B, 2010, 96, 290–298.
- 45 S. Yuan, P. Liao and A. N. Alshawabkeh, *Environ. Sci. Technol.*, 2014, **48**, 656-663.
- 46 J. Zou, J. Ma and J. Zhang, Environ. Sci. Technol., 2014, 48, 4630-4631.
- 47 C. Tan, N. Gao, Y. Deng, J. Deng, S. Zhou, J. Li and X. Xin, *J. Hazard. Mater.*, 2014, **276**, 452–460.

- 48 Y.-H. Guan, J. Ma, Y.-M. Ren, Y.-L. Liu, J.-Y. Xiao, L.-q. Lin and C. Zhang, *Water Res.*, 2013, 47, 5431–5438.
- 49 T. Zhang, W. Li and J.-P. Croue, *Environ. Sci. Technol.*, 2011, 45, 9339–9346.
- 50 X. Duan, H. Sun, Y. Wang, J. Kang and S. Wang, *ACS Catal.*, 2015, 5, 553–559.
- 51 N. S. McIntyre, D. D. Johnston, L. L. Coatsworth, R. D. Davidson and J. R. Brown, *Surf. Interface Anal.*, 1990, 15, 265–272.
- 52 X. Xia, W. Lei, Q. Hao, W. Wang and X. Wang, *Electrochim. Acta*, 2013, **99**, 253–261.
- 53 G. Chen, X. Si, J. Yu, H. Bai and X. Zhang, *Appl. Surf. Sci.*, 2015, 330, 191–199.
- 54 Y. Ding, L. Zhu, A. Huang, X. Zhao, X. Zhang and H. Tang, *Catal. Sci. Technol.*, 2012, 2, 1977–1984.
- 55 Y. Yao, Y. Cai, F. Lu, F. Wei, X. Wang and S. Wang, *J. Hazard. Mater.*, 2014, **270**, 61–70.
- 56 T. Zhang, H. Zhu and J.-P. Croue, Environ. Sci. Technol., 2013, 47, 2784–2791.