Activation of Peroxymonosulfate Using Secondary Pyrolysis Oil-Based Drilling Cuttings Ash for Pollutant Removal

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ABSTRACT: In this study, the utilization of secondary pyrolysis oil-based drilling cuttings ash (OBDCA-sp) to activate peroxymonosulfate (PMS) for pollutant removal was investigated. The chemical and physical properties of OBDCA-sp were explicitly analyzed via multiple characterization. The activation efficiency of OBDCA-sp for PMS was tested using humic acid (HA) as the target pollutant. 92% of HA and 52% of total organic carbon in solution could be removed using OBDCA-sp-activated PMS under optimal conditions: OBDCA-sp dosage at 4 g/L, PMS concentration at 4 mmol/L, HA concentration at 10 mg/L, and pH value at 7. After four cycles, 84% removal rate of HA could still be achieved using OBDCA-sp to activate PMS. The main catalysis elements for PMS activation in OBDCA were postulated to be Fe(III), Co(III), and Mn(III), based on X-ray photoelectron spectroscopy and X-ray diffraction results. The results of the quenching experiment indicated that SO₄²⁻, *OH, and ¹O₂ were the main reactive oxygen species (ROS) and that ¹O₂ was the dominant ROS in the HA removal process. Radical trapping experiments indicated the presence of SO₄²⁻, *OH, and ¹O₂ in the reaction system. This study presented a novel utilization path of OBDCA in the field of environmental remediation.

1. INTRODUCTION

In China, oil-based drilling cuttings (OBDCx) are classified as a hazardous waste, which is mainly derived from the process of crude oil development and extraction. OBDC has a complex composition, including oil−water mixtures, heavy metals, soil components, and various surfactants. The discharge of untreated/inappropriately treated OBDC could cause severe adverse impact on the surrounding environment. At present, many treatment strategies for OBDC have been studied, including pyrolysis, oxidation, microbial treatment, and solvent extraction.

Pyrolysis is the most common treatment method for OBDC due to its advantages of high scalability, ease of operation, and relatively low cost. Large amounts of OBDC ash (OBDCA) are produced as the pyrolysis product of OBDC, which are industry pain points and should be further utilized. Usually, OBDCA was employed as a raw material for bricks, cement, and proppants. Nowadays, OBDCA is rarely used in the field of environmental remediation. In our previous study, OBDCA was used as an environment-friendly material for treating Cr(VI)-contaminated soil and water. There are only rare studies about the use of OBDCA for organic pollutant removal.

Persulfate activation is an important advanced oxidized technology that can rapidly and efficiently degrade organic pollutants in an environment without causing secondary pollution. Persulfate activation includes peroxymonosulfate and peroxydisulfate activation. Transition metals, especially Fe, Co, and Mn, are the most common persulfate catalysts. Interestingly, OBDCA is composed of complex composition, which contains certain transition metals. Therefore, it is interesting to explore whether OBDCA can potentially be used to activate persulfate for organic pollutant removal.

Humic acid (HA) is a common natural organic substance in natural water bodies. The presence of excessive HA could cause some adverse impact on the environment. For water bodies, HA could result in changes in color and produce unpleasant odors. It is reported that the accumulation of HA may cause the clogging of constructed wetlands. In the chlorine disinfection process, the presence of HA will produce some carcinogenic byproducts. In addition, HA has adsorption and complexation effects on some organics and heavy metals in the environment, which is not conducive to the removal of other pollutants. Therefore, the removal of HA in the water body is of significant importance, which can improve
the quality of the water body and reduce the migration and conversion of pollutants.

The aim of this paper is to: (1) study the efficiency of removal of HA using OBDCA-activated peroxymonosulfate (PMS); (2) investigate the effect of different experimental conditions of PMS activation on HA removal; (3) analyze the reusable performance of OBDCA and the main reactive oxygen species (ROS) produced in PMS activation; and (4) characterize fresh and used OBDCA using X-ray diffraction (XRD) and X-ray photoelectron spectroscopy (XPS).

2. EXPERIMENTAL SECTION

2.1. Chemicals and Reagents. The chemicals and reagents used in this study were of analytical grade and used as received without further purification. OBDCA was provided by the Agriculture and Forestry Department of Fuling Shale Gas Company. HA was obtained from the International Humic Acid Association. Peroxymonosulfate, sulfuric acid, sodium hydroxide, tert-butanol (TBA), ethanol (EtOH), dimethyl pyridine N-oxide (DMPO), 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO), phenyl methyl sulfoxide (PMSO), and tryptophan were purchased from Aladdin Company. The stock solutions of HA (100 mg/L) and PMS (100 mg/L) were prepared using ultrapure water.

2.2. Secondary Pyrolysis Procedure for OBDC. Secondary pyrolysis was used to remove the residual organic matter in OBDCA. According to our previous study, a modified pyrolysis procedure was employed. In brief, the secondary pyrolysis of OBDCA was conducted in a muffle at 750 °C for 90 min at a heating rate of 6 °C/min under N2 atmosphere. The secondary pyrolysis product of OBDCA was abbreviated as OBDCA-sp.

2.3. Experimental Work. A certain amount of OBDCA-sp and 100 mL of HA aqueous solution were added to a flask. The pH of the solution was adjusted using 1 mol/L H2SO4 and 1 mol/L NaOH. Then, the flask was put in a shaker to conduct experiments at 200 rpm and 25 °C. At predetermined time intervals, 5 mL of the solution was withdrawn and filtered using a syringe filter, and the concentration of HA was measured using an UV spectrometer at 254 nm.

2.4. Analysis for OBDCA and HA Solution. The crystalline structures of OBDCA-sp were analyzed using an X-ray diffractometer (D8 Advance). The surface compositions and oxidation states of OBDCA-sp were analyzed using an X-ray photoelectron spectrometer (EscaLab Xi+). The morphology of OBDCA-sp was identified using a scanning electron microscope (JSM-IT300) and a transmission electron microscope (Tecnai F20). The surface area and pore volume were calculated using a Brunauer–Emmett–Teller (BET) surface area analyzer (Mike 2020). The zeta potential of OBDCA-sp was analyzed using a zeta potential analyzer (ZetaPALS). Electronic paramagnetic resonance (EPR/ESR, Brook A300) was used to capture the radicals produced in solution. The removal rate of HA was calculated according to eq 1

\[
\text{Removal rate} = \frac{C_0 - C_t}{C_0}
\]

where \( C_t \) is the HA concentration at time \( t \) (mg/L), and \( C_0 \) is the initial HA concentration (mg/L).

3. RESULTS AND DISCUSSION

3.1. Effect of the Dosage of OBDCA-sp on HA Removal. Figure 1 shows the effect of OBDCA-sp dosage on HA removal. Experimental parameters: dosage of OBDCA-sp: 0–4 g/L; PMS: 1 mmol/L; HA: 10 mg/L; pH: 7.

3.2. Effect of PMS concentration on HA removal. Experimental parameters: dosage of OBDCA-sp: 0.4 g/L; PMS: 0–4 mmol/L; HA: 10 mg/L; pH: 7.

The removal rate of HA increased with the increasing dosage of OBDCA-sp from 0 to 4 g/L. With no addition of OBDCA-sp, the removal rate of HA reached 19% after 90 min of reaction, which was ascribed to the directly oxidized HA in solution by PMS. When the dosage of OBDCA-sp was 4 g/L, the removal rate of HA reached 89% after 90 min of reaction. A higher dosage of OBDCA-sp could provide more active sites to activate PMS and produce more ROS. However, excessive dosage of OBDCA-sp increased the
cost and may also cause the blockage of active sites, which was not conducive to the removal of HA. Therefore, the dosage of OBDCA-sp in the subsequent experiments was set as 4 g/L.

3.2. Effect of PMS Concentration on HA Removal. Figure 2 shows the effect of PMS concentration on the removal of HA. The concentration of PMS increased from 0 to 4 mmol/L. With no addition of PMS, the removal of HA was very low (<1%), which indicated the poor and neglected adsorption of HA on OBDCA-sp. The removal rate of HA slightly increased with the increasing PMS concentration. When the PMS concentration was 4 mmol/L, the removal rate of HA reached 92% after 90 min of reaction. The poor effect from the increased PMS concentration for HA removal was due to the following reasons: (i) although a higher concentration of PMS in the solution could produce more ROS during the activation process, according to formula 2, PMS was one of the scavengers of \( \text{SO}_4^{2-} \), which was not conducive to HA removal; (ii) due to the limitation of the catalyst (OBDCA-sp) quantity, it could provide only limited active sites for PMS activation, resulting in the incomplete activation of excessive PMS. To achieve the maximum HA removal, the PMS concentration in the subsequent experiments was set as 4 mmol/L.

\[
\text{HSO}_5^- + \text{SO}_4^{2-} \rightarrow \text{SO}_5^{2-} + \text{SO}_4^{2-} + \text{H}^+ \\
\]  

3.3. Effect of pH on HA Removal. Figure 3 shows the effect of pH on the removal of HA. The removal rate of HA decreased as the solution changed from neutral to acidic or alkaline. When the pH of the solution was 3, the removal rate of HA reached 69.5% after 90 min of the reaction; when the pH of the solution was 11, the removal rate of HA reached 18.7% after 90 min of the reaction. The results indicated that both acidic and alkaline solutions were not conducive to the removal of HA. PMS existed in different forms in solution at different pH (\( \text{pK}_{a1} \) of PMS <0, and \( \text{pK}_{a2} \) of PMS = 9.4). Under acidic conditions, the main forms of PMS in the solution were HSO_5^- and \( \text{SO}_4^{2-} \); in neutral solutions, the main form of PMS in the solution was HSO_5^-; while in alkaline solutions, according to formula 3, the main form of PMS in solution was \( \text{SO}_5^{2-} \). The \( \text{pK}_a \) value of OBDCA-sp was calculated as 2.01, indicating that when the pH of the solution was less than 2.01, the surface of OBDCA-sp in the

Figure 3. Effect of pH on HA removal. Experimental parameters: OBDCA-sp: 4 g/L dosage; PMS: 4 mmol/L; HA: 10 mg/L; pH: 3–11.

Figure 4. (a,b) SEM image of OBDCA-sp; (c,d) TEM image of OBDCA-sp.
solution was positively charged, and when the pH of the solution was greater than 2.01, the surface of OBDCA-sp in the solution was negatively charged. Therefore, the reasons for the poor removal effect of HA under acidic conditions were: (i) the complexation reaction of HA to the metal on the surface of OBDCA-sp; (ii) according to formula 4, the generated SO$_4^{2-}$ may be changed to form *OH. The reasons for the poor removal effect of HA under alkaline conditions were: (i) according to formula 5, OH$^-$ was also the scavenger of SO$_4^{2-}$, forming *OH with a lower redox potential; (ii) according to the above dissociation coefficient, the surface of OBDCA-sp had a significant electrostatic mutual repulsion effect with SO$_5^{2-}$, resulting in a decrease in the HA removal effect; (iii) in addition, the main composite of OBDCA-sp was SiO$_2$, which could be dissolved in alkaline conditions and was not conductive to the activation of PMS. Therefore, in the subsequent experiments, the pH value in the solution was adjusted to 7.

\[
\text{HSO}_3^- \rightarrow \text{SO}_3^{2-} + \text{H}^+ \\
\text{SO}_4^{2-} + \text{H}_2\text{O} \rightarrow \text{*OH} + \text{SO}_4^{2-} + \text{H}^+ \tag{4}
\]

\[
\text{SO}_4^{2-} + \text{OH}^- \rightarrow \text{SO}_4^{2-} + \text{*OH} \tag{5}
\]
3.4. Morphology, BET, and Surface Potential Analyses of OBDCA-sp.

The morphology analysis of OBDCA-sp was observed using SEM and TEM (Figure 4). OBDCA-sp was irregularly shaped with a coarse surface covered by corundum-like crushed particles. The BET and zeta potential analyses of OBDCA-sp were also carried out (Figure 5). The N2 adsorption/desorption isotherms for OBDCA-sp were classified as type III with H3 hysteresis, loops based on the Brunauer−Deming−Deming−Teller classification, which indicated that the surface of OBDCA-sp exhibited a slit hole formed by the accumulation of flake particles. The specific surface area and pore volume of OBDCA-sp (Table 1) were 1.09 m2/g and 0.0013 cm3/g, respectively. The pHpzc value of OBDCA-sp was obtained according to the calculation of zeta potentials, which indicated that the surface charge of OBDCA-sp was negative, with the pH exceeding 2.01.

3.5. Recycled Utilization and Characterization of OBDCA-sp for HA Solution.

The recycled utilization of OBDCA-sp to activate PMS for HA removal was tested (Figure 6). After four rounds of recycling, 84% HA removal rate could still be achieved using OBDCA to activate PMS. The reduced removal efficiency was owing to the weight loss of OBDCA-sp. The HA solution before and after the reaction was measured using EEM (Figure 7), which evidently indicated the removal of HA in the characteristic regions (λex = 200−250 nm).

Table 2. Leachate Concentration of OBDCA-sp

| element | concentration (mg/L) |
|---------|----------------------|
| Co      | 0.0001               |
| Cr      | 0.0062               |
| Fe      | 0.0036               |
| Mn      | 0.0043               |

Figure 8. XRD pattern of OBDCA-sp.
nm; $\lambda_{em} = 380–540$ nm and $\lambda_{ex} = 250–400$ nm, $\lambda_{em} = 380–540$ nm). The TOC removal rate under optimal conditions was calculated as 52.7%, which indicated the uncompleted minimization of HA. In addition, the leachate concentration of the main metal element in OBDCA-sp was analyzed using ICP-OES (Table 2). The leachate concentration of heavy metals was very low, guaranteeing the safe use of OBDCA-sp in environmental remediation.
To postulate the potential metal components and elements in OBDCA-sp for PMS activation, characterization by XPS and XRD was done for structural analysis. The XRD pattern of OBDCA-sp (Figure 8) exhibited complex and mass peaks. The diffraction peaks at 2θ = 20.8, 26.6, 36.5, 39.4, and 50.1° were assigned to quartz (SiO2, PDF 79-1906). The 2θ = 32.9, 55.2, 23.1, 38.2, and 65.8° corresponded to bixbyte (FeMnO3, PDF 75-0894). The characteristic diffraction peaks at 2θ = 23.1, 32.9, 38.2, and 55.1° were ascribed to manganese oxide (Mn2O3, PDF 71-0636). The characteristic diffraction peaks at 2θ = 32.9, 38.2, 55.2, and 65.8° were ascribed to iron oxide (Fe2O3, PDF 39-0238). The results indicated that the main components of OBDCA-sp were SiO2, FeMnO3, Mn2O3, and Fe2O3. The surface element state of OBDCA-sp was analyzed using XPS (Figure 9). Co was detected in the XPS spectra of OBDCA-sp. The corresponding peaks of Co were 780.1 and 782.3 eV. No obvious spin orbital was detected for Mn 2p, and two spin orbitals were detected for Mn 3s. The distance for spin energy separation of the Mn 3s level was 3 eV, which indicated that the possible value state for the Mn element was Mn3+. According to a previous report, the existing form of Mn was speculated to be Co2O3. Two spin orbitals of core-level Fe 2p were detected. Fe 2p3/2 and Fe 2p1/2 were located at 711.1 and 724.7 eV, respectively. The spin energy separation between the two spin orbitals was 13.6 eV, indicating that the possible value state for the Fe element was Fe3+. According to the XRD results, the existing forms of Mn were speculated to be Co2O3, FeMnO3, Mn2O3, and Fe2O3. The surface element state of OBDCA-sp was analyzed using XPS (Figure 9). Co was detected in the XPS spectra of OBDCA-sp. The corresponding peaks of Co were 780.1 and 782.3 eV. No obvious spin orbital was detected for Mn 2p, and two spin orbitals were detected for Mn 3s. The distance for spin energy separation of the Mn 3s level was 3 eV, which indicated that the possible value state for Mn was Mn3+. Co-containing component was not detected in XRD, as the content of Co-containing component was lower than the lower detection limitation of XRD. According to a previous report, the existing form of Co was postulated to be CoO2. Spin orbitals of core-level Fe 2p were detected. Fe 2p3/2 and Fe 2p1/2 were located at 711.1 and 724.7 eV, respectively. The spin energy separation between the two spin orbitals was 13.6 eV, indicating that the possible value state for the Fe element was Fe3+. According to the XRD results, the existing form of Fe was postulated to be Fe2O3 and FeMnO3. An obvious spin orbital for O 1s was observed at 531.3 eV, which was attributed to adsorbed oxygen, indicating the oxidation state (O2−) of O element. No obvious spin orbital was detected for Mn 2p, and two spin orbitals were detected for Mn 3s. The distance for spin energy separation of the Mn 3s level was 3 eV, which indicated that the possible value state for Mn was Mn3+. According to the XRD results, the existing forms of Mn were speculated to be Mn2O3 and FeMnO3.

3.6. Radical Quenching Test and ESR Test. The radical quenching tests (Figure 10) were performed to identify the role of ROS in HA removal. "OH, SO4−, and 1O2 were possible ROS for HA removal according to the previous studies. To evaluate the contribution of "OH, SO4−, and 1O2 to HA removal, according to the relative report, EtOH was used as a radical quencher for "OH and SO4− and TBA was used as a radical quencher for SO4−. Tryptophan was used as a radical quencher for 1O2. The results of radical quenching tests indicated that the addition of tryptophan severely inhibited the removal of HA, while EtOH and TBA had only slight inhibition effect for HA removal. According to the effect of the quencher on the HA removal, it was concluded that the order of ROS contribution for HA removal was as follows: 1O2 >> SO4− >> "OH, and 1O2 was the dominant ROS for HA removal.

ESR analysis was employed to determine the ROS produced during PMS activation using OBDCA-sp (Figure 10). DMPO was employed as a radical trapping agent for "OH and SO4− and 1O2 was trapped by TEMPO. For signals of DMPO-adduct, two group signal peaks, DMPO-SO4− and DMPOX, were detected. DMPOX was formed when DMPO was trapped two hydroxyl groups; for signals of TEMPO-adduct, the signal peaks of TEMPO-1O2 were evident. ESR analysis confirmed the presence of "OH, SO4−, and 1O2 in the OBDCA-sp/PMS system. However, due to the complex components of OBDCA-sp, the generation pathway of "OH, SO4−, and 1O2 remains unclear. The possible path generation was postulated as follows (formula 6–11).

\[
\begin{align*}
\text{HSO}_3^- & \rightarrow \text{H}^+ + \text{SO}_4^{2-} \\
\text{SO}_3^- + \text{H}_2\text{O} & \rightarrow \text{O}_2 + \text{SO}_4^{2-} + \text{H}^+ \\
\text{O}_2^- & \rightarrow 2\text{H}_2\text{O} \rightarrow \text{O}_2 + \text{H}_2\text{O}_2 + 2\text{OH}^- \\
\text{"OH} + \text{H}_2\text{O}_2 & \rightarrow \text{HO}_2^- + \text{H}_2\text{O} \\
\text{HO}_2^- & \rightarrow \text{H}^+ + \text{O}_2^- \\
\text{O}_2^- & \rightarrow \text{"OH} \rightarrow \text{O}_2 + \text{OH}^-
\end{align*}
\]

4. CONCLUSIONS

In this work, OBDCA-sp was used to activate PMS for HA removal. 92% of HA removal efficiency was achieved under optimal experimental conditions. The performance of OBDCA-sp for PMS activation was attributed to Fe(III), Mn(III), and Co(III) elements in OBDCA-sp. The ROS scavenger experiments and ESR spectra demonstrated that SO4−, "OH, and 1O2 were responsible for HA removal and that 1O2 was the dominant ROS for HA removal. The study indicated the promising potential for the application of OBDCA-sp in the catalysis field and organic pollution treatment.

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Notes

The authors declare no competing financial interest.
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