IMPROVE THE CATALYTIC ACTIVITY OF FeOOH/BENTONITE MATERIAL BY MECHANICAL ACTIVATION

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ABSTRACT

To improve the catalytic activity of FeOOH/bentonite material used in Fenton-like process, the activation of FeOOH/bentonite by mechanical activation was studied. The optimum conditions for activation of FeOOH/bentonite were as follows: filling ratio of grinding medium 30%, "15 mL of D06 balls plus 15 mL of D10 balls" as combination mode of grinding medium, rotation speed of planet carrier 600 rpm, milling time 40 min, and powder-to-ball ratio 0.5:30 (g·mL-1). The mechanical activation was an effective method to improve the catalytic activity of FeOOH/bentonite. Both the lattice distortion and crystal size decrease happened in α-FeOOH and the change of layer structure of bentonite were increased to the contribute of activation of FeOOH/bentonite.

Keyword: Mechanical activation, Bentonite, α-FeOOH, Fenton-like, Catalytic activity

INTRODUCTION

Even through Fenton oxidation is an effective technology to treat the wastewater containing recalcitrant organic pollutants, some drawbacks existing in conventional homogeneous Fenton system have prevented the wide industrialization of Fenton oxidation in wastewater treatment1-2. In order to overcome the drawbacks, heterogeneous catalysts used in Fenton system have aroused attention of the researchers. The Fenton reaction catalyzed by heterogeneous catalyst usually belongs to the Fenton-like process. Heterogeneous catalysts, which commonly are environmentally benign and easily separated, have been reported to overcome the drawbacks of conventional homogeneous Fenton3-5. Recently, many solid materials used as the heterogeneous catalysts have been prepared to catalyze the Fenton reaction5-6. Bentonite, which is a kind of clay, has particular structures and good properties as well as low cost and abundance in world7-9, so it has been used to prepare bentonite-based heterogeneous catalysts which were successfully used in Fenton-like processes10-11. It has been proved that FeOOH/bentonite, i.e. α-FeOOH (goethite) pillared bentonite, was a high-efficiency catalytic material for the Fenton-like reaction12. However, even high activity of FeOOH/bentonite, the catalytic property of FeOOH/bentonite might be further improved by the activated method. Mechanical activation, a green technique used for synthesizing or modifying materials in high-energy ball mill, in essence is a method to change the physical and/or chemical property of materials via mechanical energy13-16. When the materials was activated in the high-energy ball mill, through the repeated welding and fracturing which happened in the powder particles of material, mechanical activation could increase the specific surface and form the fresh surface areas of the materials, and even destroy the materials' stable crystalline structure17-21.

The aim of this study is to explore the mechanical activation of FeOOH/bentonite material. In our work, the activation of FeOOH/bentonite by mechanical activation was firstly investigated in detail, and then based on the XRD characterization of material, the mechanism for improvement of catalytic activity by mechanical activation was also discussed preliminarily.

MATERIALS AND METHODS

Materials and chemical reagents

Bentonite was purchased from Shanghai No.4 Reagent and H.V. Chemical Limited Company, China. All chemicals and reagents used in the study, including Fe(NO₃)₃, Na₂CO₃, HNO₃, H₂O₂ (30%) and orange II, were of analytical reagent grade.

Preparation and mechanical activation of FeOOH/Bentonite catalyst

Na₂CO₃ powder was slowly added into the solution of 0.2 mol·L⁻¹ Fe(NO₃)₃ under room temperature, and the molar ratio of added Na₂CO₃ to Fe(NO₃)₃ was about 0.325:1. The gained solution was stirred for one day and then aged for one day at room temperature. After that, the intercalation solution was added slowly into bentonite dispersion (5%) under vigorous stirring until the dosage of Fe³⁺ to bentonite amount in the dispersion was about 10 mmol/g. After the mixture was stirred for 8 h and then aged for one day at 60 °C, the mixture was centrifuged, washed several times with deionized water, dried at 105 °C overnight, and then ground to less than 0.074 mm. After that, 15 mL of solution of H₂O₂ (30%, 19.6 mmol·L⁻¹) and 0.2 g·L⁻¹ catalyst in the suspension at 3500 rpm for 10 minutes. The remaining orange II in sample solution was determined using a VIS-722 spectrophotometer (Shanghai Yoke Instrument Co., Ltd., China) at 485 nm which is the maximum absorbance wavelength of orange II solution. The catalytic property of catalyst was evaluated by the removal ratio of orange II which was calculated with the following formula:

\[ \eta = \frac{C_0 - C_1}{C_0} \times 100\% \]

where \( C_0 \) and \( C_1 \) represent the time-dependent concentration and the initial concentration, respectively. After the degradation of orange II, the content of total iron ions (C_total, mg·L⁻¹) and the content of ferrous iron ions (C_ferrous, mg·L⁻¹) in the solution were measured by the method of o-phenanthroline spectrophotometry.

XRD patterns of catalyst were measured with a D8 ADVANCE X-ray diffractometer (Bruker, Germany) equipped with Cu and Ka radiation at 40 kV and 40 mA, and the recorded 20 was in the range of 5°-80°.

RESULTS AND DISCUSSION

Effect of filling ratio of grinding medium on FeOOH/bentonite catalytic activity

0.5 g of FeOOH/bentonite was placed into the milling chamber, and the following conditions were used for mechanical activation: stainless steel milling balls with a diameter of 10 mm as grinding medium, rotation speed of planet carrier 600 rpm, and milling time 60 min. The effect of filling ratio of grinding medium on the catalytic activity of activated FeOOH/bentonite is shown in Fig. 1.

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3407
As shown in Fig. 1, when the filling ratio of grinding medium was less than 30%, the catalytic activity increased with the filling ratio increase, and the catalytic activity reached its maximum at the filling ratio of 30%, with the corresponding removal ratio of orange II of 86.1%. When the filling ratio was over 30%, the catalytic activity declined with the further increase of filling ratio. Therefore, the filling ratio of 30% was used for the mechanical activation in all subsequent experiments. When the filling ratio was too low, the contact between stainless steel milling balls and FeOOH/bentonite in the milling chamber was not enough, so the collision and friction between stainless steel milling balls and FeOOH/bentonite was not sufficient. With the increase of filling ratio, the collision frequency between stainless steel milling balls and FeOOH/bentonite was increased, which could make the mechanical activation effect on the FeOOH/bentonite increase quickly. However, when the filling ratio of grinding medium was too high, the void in the milling chamber became small, which formed a “dead zone” for the mechanical activation. In the “dead zone”, the grinding medium and the material only could move slowly. And, furthermore, the free path of stainless steel milling balls in the “dead zone” was shortened. Hence, the mechanical activation effect was greatly weakened as a result of the high filling ratio of grinding medium.

Effect of combination mode of grinding medium on FeOOH/bentonite catalytic activity

Two kinds of stainless steel milling balls with diameters of 6 mm and 10 mm were used to study the effect of combination mode of milling balls on the catalytic activity of activated FeOOH/bentonite. The stainless steel milling balls with diameter of 6 mm and 10 mm were designed as D06 balls, and D10 balls, respectively. The conditions of mechanical activation were as follows: mass of FeOOH/bentonite 0.5 g, filling ratio of grinding medium 30%, rotation speed of planet carrier 600 rpm, and milling time 60 min. The removal ratios of orange II catalyzed by the FeOOH/bentonite catalysts which were activated under different combination modes of milling balls are listed in Fig. 2.

As shown in Fig. 2, the catalytic activity of FeOOH/bentonite catalyst activated under the combination mode of “D06,15%+D10,15%”, i.e., 15 mL of D06 balls plus 15 mL of D10 balls, was the highest. During the activation of FeOOH/bentonite, the voids between stainless steel milling balls were inevitably generated in the cylindrical milling chamber. However, when the stainless steel milling balls with different diameters were selected as the grinding medium, the bigger balls could increase the impact force and the shear force, both of which came from the grinding between balls. Furthermore, the smaller balls could easily fill the void between the bigger balls, so that the contact surface between stainless steel milling balls and FeOOH/bentonite material increased during the process of mechanical activation, and the friction between stainless steel milling balls and FeOOH/bentonite material also increased. The combination mode of “D06,15%+D10,15%” was chosen as the optimum condition in all subsequent mechanical activation experiments.

Effect of rotation speed of planet carrier on FeOOH/bentonite catalytic activity

To analyze the effect of rotation speed of planet carrier on the catalytic activity of FeOOH/bentonite, FeOOH/bentonite was activated at different rotation speed under the condition: mass of FeOOH/bentonite 0.5 g, combination mode of “D06,15%+D10,15%”, filling ratio 30%, and milling time 60 min. The effect of rotation speed of planet carrier on the catalytic activity of FeOOH/bentonite is shown in Fig. 3.

With the increase of rotation speed of planet carrier, the removal ratio of orange II increased until the rotation speed was 600 rpm, and the removal ratio reached its maximum at 600 rpm. The results indicated that the activated FeOOH/bentonite reached its highest activity at the rotation speed of 600 rpm. At low rotation speed, the movement of stainless steel milling balls in the milling chamber was in a motion state of cascading. In this motion state, the stress on the FeOOH/bentonite would be mostly the compression and shear stress between milling balls or milling balls and milling chamber wall. With the increase of rotation speed, even though no detachment of balls from the wall or the other balls occurred, the compression and shear stress on the FeOOH/bentonite increased quickly with the rotation speed increase, which resulted in the increase of catalytic activity of FeOOH/bentonite. However, when the rotation speed was 600 rpm, the movement of milling balls might shift from the cascading motion into the cataracting motion, in which a fraction

![Graph](image1.png)

**Fig 1.** Effect of filling ratio of grinding medium on the removal ratio of orange II.

![Graph](image2.png)

**Fig 2.** Effect of combination mode of different balls on the removal ratio of orange II: (A) D06,30%; (B) D10,30%; (C) D06,15%+D10,15%

![Graph](image3.png)

**Fig 3.** Effect of rotation speed of planet carrier on the removal ratio of orange II.
of the milling balls detached from the grinding chamber wall and followed a parabola trajectory in flight before impacting into the bulk of balls. Besides the compression and shear stress, as existed in the cascading motion, the impact stress on the FeOOH/bentonite from dropping down balls also occurred in the catacracting motion. Hence, the catalytic activity of FeOOH/bentonite reached its maximum at the rotation speed of 600 rpm. A slight drop of catalytic activity with the further increase of rotation speed which was over 600 rpm might be because that part of milling balls were in the centrifugation motion, in which the centrifuged milling balls aligned on the chamber wall and followed its rotation with almost no relative velocity, which made the stress on the FeOOH/bentonite decline slightly.

**Effect of milling time on FeOOH/bentonite catalytic activity**

The effect of milling time on the catalytic activity of FeOOH/bentonite was investigated, and the parameters used for mechanical activation were as follows: mass of FeOOH/bentonite 0.5 g, combination mode of “D06,15%+D10,15%”, filling ratio 30%, and rotation speed of planet carrier 600 rpm. The result is shown in Fig. 4.

![Fig 4. Effect of milling time on the removal ratio of orange II](image)

The catalytic activity of FeOOH/bentonite first increased with the increase of milling time from 20 to 40 min, and then remained almost stable when milling time was over 40 min. The reason for such a result was that with the increase of milling time, the FeOOH/bentonite in the milling chamber was fully activated. The milling time of 40 min was chosen as the optimum condition in all subsequent mechanical activation experiments.

**Effect of powder-to-ball ratio on FeOOH/bentonite catalytic activity**

To investigate the influence of powder-to-ball ratio in the process of mechanical activation on the catalytic activity of FeOOH/bentonite, FeOOH/bentonite material was activated at different powder-to-ball ratio which was defined as the ratio of mass of activated powder to packing volume of milling balls (Rptb, g mL⁻¹). And the other parameters used for mechanical activation were as follows: combination mode of “D06,15%+D10,15%”, filling ratio 30%, rotation speed of planet carrier 600 rpm, and milling time 60 min. Moreover, in order to identify the enhancement effect on catalytic activity by mechanical activation, a comparative experiment of orange II degradation in Fenton-like process catalyzed by FeOOH/bentonite, which was without the treatment of mechanical activation, was also carried out. The results are listed in Table 1.

![Table 1. Removal ratios of orange II catalyzed by different catalysts.](image)

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>Rptb (g mL⁻¹)</th>
<th>η(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activated FeOOH/bentonite</td>
<td>0.25:30</td>
<td>75.5</td>
</tr>
<tr>
<td>Activated FeOOH/bentonite</td>
<td>0.5:30</td>
<td>92.3</td>
</tr>
<tr>
<td>Activated FeOOH/bentonite</td>
<td>0.75:30</td>
<td>80.7</td>
</tr>
<tr>
<td>Activated FeOOH/bentonite</td>
<td>1.0:30</td>
<td>78.5</td>
</tr>
<tr>
<td>FeOOH/bentonite</td>
<td>N/A</td>
<td>26.3</td>
</tr>
</tbody>
</table>

As shown in Table 1, with the increase of Rptb, i.e. with the increase of mass of FeOOH/bentonite added into the milling chamber, the catalytic activity of FeOOH/bentonite increased quickly until the Rptb was 0.5:30 (g mL⁻¹), and then the catalytic activity decreased with the further increase of Rptb. With the increase of Rptb, the increase of FeOOH/bentonite mass increased the contact between FeOOH/bentonite and milling balls. The stress effect on FeOOH/bentonite also changed more sufficient for enough collision between powder and milling balls, so that the mechanical energy from the milling balls could easily transform into the internal energy of FeOOH/bentonite powder. However, when the value of Rptb was over 0.5:30 (g mL⁻¹), the dosage of FeOOH/bentonite was too excessive in the milling chamber, and the powder accumulation formed in the gaps between milling balls or milling balls and milling chamber wall. Therefore, partial FeOOH/bentonite could not get enough mechanical stress, which produced a negative effect on the activation of FeOOH/bentonite. Hence, the Rptb of 0.5:30 (g mL⁻¹) was chosen as the optimum condition for the mechanical activation of FeOOH/bentonite. Moreover, as shown in Table 1, compared with the high removal ratios of orange II catalyzed by the activated FeOOH/bentonites, the removal ratio catalyzed by the FeOOH/bentonite was only reached 26.3%, indicating that the mechanical activation was an effective method to improve the catalytic activity of FeOOH/bentonite.

**Catalyst characterization and mechanism of catalytic activity improvement**

The X-ray diffraction patterns of FeOOH/bentonite and activated FeOOH/bentonite are shown in Fig. 3. As shown in Fig. 3, FeOOH/bentonite revealed the presence of signals in 2θ positions corresponding to 53.23°, 31.79°, 34.75°, 39.98°, and 53.23°, which were characteristic peaks of α-FeOOH (JCPDS: 29-0713), indicating that the iron phase immobilized on bentonite was mainly in form of α-FeOOH. Likewise, the pattern of activated FeOOH/bentonite exhibited diffraction peaks at 2θ 21.22°, 33.24°, 39.98°, and 53.23°. However, compared to the characteristic peaks of α-FeOOH on FeOOH/bentonite, the broadening of diffraction peaks of α-FeOOH was observed from the pattern of activated FeOOH/bentonite, indicating that under mechanical activation, lattice distortion and crystal size decrease might happen in α-FeOOH, which was immobilized on bentonite. Moreover, as shown in Fig. 5, compared to FeOOH/bentonite, the layer structure of bentonite in the activated FeOOH/bentonite was destroyed in a certain degree after mechanical activation, which could make the adsorption ability of activated material increase. Hence, under the action of mechanical activation, the microstructural changes in activated FeOOH/bentonite, including the lattice distortion and crystal size decrease happened in α-FeOOH, and the change of layer structure of bentonite, were contributed to the increase of catalytic activity of activated material.

![Fig 5. XRD spectra of samples](image)

**CONCLUSIONS**

The optimum conditions for activation of FeOOH/bentonite by mechanical activation were as follows: filling ratio of grinding medium 30%, “15 mL of D06 balls plus 15 mL of D10 balls” as combination mode of grinding medium,
rotation speed of planet carrier 600 rpm, milling time 40 min, and powder-to-ball ratio 0.5: 30 (g·mL⁻¹). The mechanical activation was an effective method to improve the catalytic activity of FeOOH/bentonite. Both the lattice distortion and crystal size decrease happened in α-FeOOH and the change of layer structure of bentonite were contributed to the increase of catalytic activity of activated FeOOH/bentonite.

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