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Lihui Gao  
China University of Mining and Technology

Lizhang Wang  
China University of Mining and Technology

Shulei Li  
China University of Mining and Technology

Yijun Cao  
Zhengzhou University

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Highly active Fenton-like catalyst derived from solid waste-iron ore tailings using wheat straw pyrolysis

Lihui Gao¹; Lizhang Wang¹; Shulei Li²; Yijun Cao²,³*

1. School of Environment and Spatial Informatics, China University of Mining and Technology, Xuzhou 221116, China
2. National Engineering Research Center of Coal Preparation and Purification, China University of Mining and Technology, Xuzhou 221116, China
3. School of Chemical Engineering and Technology, Zhengzhou University, Zhengzhou 450001, China

Abstract: The pollutants degradation rate of iron ore tailings-based heterogeneous catalysts is the main factor limiting its application. Herein, an iron ore tailings-based Fenton-like catalyst(I/W(3:1)-900-60) with relative fast catalysis rate was constructed by co-pyrolysis(900 °C, 60min holding time) of iron ore tailings and wheat straw with mass ratio of 3:1. With wheat straw blending, the generated I/W(3:1)-900-60 presented a larger surface area(24.3 m²/g), smaller pore size(3.707 nm), reduced iron species (Fe²⁺ from magnetic) and a higher catalytic activity(0.0229 min⁻¹) than I-900-60 (1.196 m²/g, 12.935 nm, 0.012 min⁻¹) pyrolyzed using a single iron ore tailings under the same pyrolysis conditions. In addition, biochar and iron ore tailings in I/W(3:1)-900-60 were tightly combined through chemical bonding. The optimal catalyst remains active after three cycles, indicating its catalytic stability and recyclability. The good Fenton-like MB degradation efficiency of I/W(3:1)-900-60 was ascribed to the sacrificial role of biochar, as well as the electron transfer between biochar and iron active sites or the
redox cycles of \( \equiv \text{Fe}^{3+}/\text{Fe}^{2+} \). This finding provides a facile construction strategy for a highly active iron ore tailings-based Fenton-like catalyst, and thereby had a great potential application in wastewater treatment.

**Key words:** Iron ore tailings; Wheat straw; Heterogeneous catalyst; Biochar; Decomposition rate
1. Introduction

Mining industry is an important activity to extract mineral products around the world. Nevertheless, the mining process always brings a certain degree of pollution. Tailings are solid waste remaining after mining valuable minerals, which are usually disposed at waste dams or landfills that present potential environmental damage (Rico et al. 2008; Kossoff et al. 2014). In March 2020, a miserable accident involving a waste dam occurred when the Yichun waste dam (Harbin, China) for the storage of iron ore tailings collapsed. This accident caused 60,000 m$^3$ of mining tailings disclosing and 3 million m$^3$ of wastewater releasing, leading to 70 kilometers of river pollution and serious economic loss. A number of other tailings dam accidents have occurred in different countries (Batista et al. 2020). According to statistics (Yi et al. 2020), the accumulated of tailings were about 207 billion tons in China, of which the total amount of tailings produced in 2018 was about 12.11 billion tons. Among all types of tailings, iron ore tailings have the largest amount of production, about 4.76 billion tons, accounting for about 39.31% of the total tailings production (Huang et al. 2020).

Therefore, it is imperative to explore new technologies for reducing iron ore tailings and their reuse as raw materials to produce value-added products. Recently, a promising application of iron ore tailings is its utilize as adsorbents or catalysts to remove dye compounds (Silva et al. 2011; Augusto et al. 2018), e.g. as a raw/regenerated efficient Fenton and Fenton-like catalysts for wastewater treatment. Given the high iron contents observed in iron ore tailing, it is reasonable to assume that these wastes are good catalysts for Fenton or Fenton-like processes (Augusto et al. 2018).
2018). However, Fe$^{2+}$ is the effective state for activating the oxidant, while the most common state in iron waste is Fe$^{3+}$ (dos Santos et al. 2016), and the presence of Fe$^{3+}$ modifies the degradation rate (reaction time as long as 24 h) of substance. Some researchers got significant progress in trying to enhance the pollutant oxidation rate through reducing Fe$^{3+}$ to Fe$^{2+}$ (Augusto et al. 2018; de Freitas et al. 2019). Freitas et al achieved good results (80% dye decomposed with 3 h reaction) for the oxidation of methylene blue when iron ore tailing pretreated in a CH$_4$ atmosphere at 550 °C for 2 h.

Better degradation results could be achieved in case of iron waste pretreatment under reducing atmosphere (H$_2$, CO, CH$_4$). Biomass could produce reducing gases and compounds during pyrolysis (Williams and Besler 1996; Wang et al. 2010). In our previous paper, we found that a unite mass wheat straw could produce about 5 mg/g H$_2$ and 18 mg/g CH$_4$ (Gao and Goldfarb 2019). Therefore, if iron tailings and wheat straw blends were pretreated through pyrolysis to realize the conversion of ferric iron to a low-valent state, it would not only improve the catalytic oxidation efficiency of organic matter, but also realize the resource utilization of wheat straw, which is of great significance.

Therefore, this paper applies methylene blue as a model pollutant to investigate the influence of pyrolysis conditions on the catalyzed degradation rate of pollutants. The pyrolyzed products’ morphology, physical-chemical properties, stability and the catalytic mechanism were discussed. The research in this article aims to provides a theoretical basic for iron ore tailing catalyst with high degradation rate.

2 Materials and methods
2.1 Materials

Iron ore tailings were kindly provided by “Hainan mining” industry, located in Hainan, China. The sample was hematite tailings (T), and ground to 74 μm prior to use. Wheat straw selected as biomass because of the high production in China, which was ground in a coffee mill and sieved to a particle size between 100 and 300 μm (Gao and Goldfarb 2019). Table S1 showed the characteristics of wheat straw and hematite tailings. Methylene blue was purchased from Aladdin Chemical Reagent Co.Ltd. China.

2.2 Synthesis of the catalysts

Hematite tailings and wheat straw mixed with mass ratio 1:1, 2:1, 3:1, 4:1 and 5:1. Then, these mixtures were treated in tube furnace using a heating rate of 10 °C/min (Wang et al. found feedstock pyrolyzed at this heating rate could produce more reducing gas, such as CO and H2) (Wang et al. 2018) until different temperatures (500, 600, 700, 800 and 900 °C) for different holding times (30, 45, 60, 90 and 120 min). The obtained catalyst was denoted as I/W(a:b)-T-t, where I iron ore tailings, W wheat straw, a:b the mass ratio (w/w), T finally temperature, t holding time.

2.3 MB decomposition experiments

To evaluate the decomposition rate of catalyst produced from iron ore tailings, the bath catalytic experiments were carried out using 0.3 g of pyrolyzed product in contact with 100 mL dye solution with initial concentration of 60 mg/L and 16 mmol H2O2 under a shaker. 1.5 mL of solution was pipetted at given time intervals, centrifuged for 1 min and the dye concentration was detected by a UV-vis spectrophotometer (Unico
To investigate the stability and reusability of catalyst, the optimal pyrolyzed product was reused/recycled several times for the MB decomposition under the same Fenton-like condition.

2.4 Analytical method

The phase transformation of ferrochemical group in pyrolyzed solid was analyzed by XRD (Bruker D8 Advance, Germany). The morphology and composition were characterized by SEM-EDX (FEI QuantaTM 250, USA). The textural properties were test by BET (BELSORP-max, Japan). The functional groups of catalysts were investigated by FTIR (Bruker Vertex 80v, Germany). Electron paramagnetic resonance (EPR, Bruker 300E spectrometer, Germany) was applied to test the main reactive oxidative species (ROS) generated in the system. Fe valence and carbon functional group variation of catalysts were characterized by XPS (Thermo Fisher Escalab 250 Xi, USA), and the C1s1/2(284.6 eV) was used as the binding energy calibration standard.

3 Results and discussion

3.1 Determination of optimal pyrolysis based on degradation rate

The pyrolysis preparation mass ratio (iron ore tailings to wheat straw), reaction time and heating rate were fixed at 3:1, 60 min and 10 °C/min, respectively, and catalysts were obtained under different pyrolysis temperature. Fig.1(a) showed the comparison of MB removal efficiency versus time and pseudo-second-order kinetic constant \(k_2, R^2>0.994\) under different catalysts, which -5 min means the \(\text{H}_2\text{O}_2\) was added after 5 min adsorption. Only approximately 20% of MB was removed after 1 h
with a kinetic rate of 0.0139 min\(^{-1}\) when the product prepared at 500 °C used. However, the removal rate and efficiency were substantially improved as the pyrolyzed temperature increased. The best decomposition rate (0.0182 min\(^{-1}\)) and efficiency (74%) were reached using a catalyst prepared at 900 °C, which was much higher than the generated iron ore tailings catalyst reported by Augusto et al (7.4 × 10\(^{-4}\) min\(^{-1}\), 63%) (Augusto et al. 2018). Considering the dye decomposition rate and efficiency, the optimal pyrolysis temperature for synthesizing catalyst was determined to be 900 °C.

Setting the pyrolysis temperature as 900 °C and final holding time as 60 min and heating rate as 10 °C/min, the removal efficiency and rate of MB were investigated by pyrolysis products at different mass ratio. As shown in Fig.1(b), only 10% MB was removed with a removal rate \(k_2\) of 0.0120 min\(^{-1}\) when the product prepared by hematite tailings only, indicating that the pyrolyzed product synthesized from iron ore tailings might not be active for dye degradation. While the decomposition efficiency of MB (\(k_2\) increased from 0.0120 min\(^{-1}\) to 0.0182/0.0183 min\(^{-1}\)) gradually increased when the products made from hematite tailings and wheat straw blends were used as catalyst. This may be due to biomass blends helped to reduce the \(≡\text{Fe}^{3+}\) to \(≡\text{Fe}^{2+}\) or low-valent iron (Ellison and Boldor 2021). The mechanism of \(\text{H}_2\text{O}_2\) activation by iron ore tailings based catalyst with \(≡\text{Fe}^{2+}\) may involve the following reaction processes (Luo et al. 2010).

Firstly, a complex assigned as \(≡\text{Fe}^{2+} \cdot \text{H}_2\text{O}_2\) may form between the hydrous surface of \(≡\text{Fe}^{2+} \cdot \text{H}_2\text{O}\) and \(\text{H}_2\text{O}_2\) (Eq.1), where \(≡\text{Fe}^{2+} \cdot \text{H}_2\text{O}\) represents the reduced sites on the iron ore tailings catalyst surface. The formed \(≡\text{Fe}^{2+} \cdot \text{H}_2\text{O}_2\) can produce \(\cdot \text{OH}\) by \(\text{H}_2\text{O}_2\) activation, which is ready to decompose and oxidize MB (Eq.2 and 3)).
\[
\equiv \text{Fe}^{2+} \cdot H_2O + H_2O_2 \rightarrow \equiv \text{Fe}^{2+} \cdot H_2O_2 \\
\equiv \text{Fe}^{2+} \cdot H_2O_2 \rightarrow \equiv \text{Fe}^{3+} + \cdot OH + OH^- \\
\cdot OH + \text{MB} \rightarrow \cdots CO_2 + H_2O
\]

(1)

(2)

(3)

However, the radical formation mechanism by \equiv \text{Fe}^{3+} and H_2O_2 is proposed as follows:

\[
\equiv \text{Fe}^{3+} + H_2O_2 \rightarrow \equiv \text{Fe}^{3+} \cdot H_2O_2 \\
\equiv \text{Fe}^{3+} \cdot H_2O_2 \rightarrow \equiv \text{Fe}^{2+} + \cdot OOH + H^+ \\
\equiv \text{Fe}^{3+} + \cdot OOH \rightarrow \equiv \text{Fe}^{2+} + O_2 + H^+
\]

(4)

(5)

(6)

The formed \equiv \text{Fe}^{2+} subsequently produce \cdot OH (Eq.(1)-(3)). Although \cdot OH can be generate from H_2O_2 when either \equiv \text{Fe}^{2+} or \equiv \text{Fe}^{3+} is present, the generation rates are much faster between \equiv \text{Fe}^{2+} and oxidant (Kwan and Voelker 2002). Although the degradation efficiency for catalysts produced from mass ratio 1:1, 2:1 and 3:1 was equivalent, degradation rate \(k_2\) values of mass ratio 3:1(0.0182 min\(^{-1}\)) and 2:1 (0.0183 min\(^{-1}\)) were higher than 1:1(0.0158 min\(^{-1}\)). In order to realize the iron ore tailings bulk utilization and construct a catalyst with higher degradation rate, we determined the optimal mass ratio of iron ore tailings to wheat straw was 3:1.

Fixing the pyrolysis temperature as 900 °C, the ratio of iron ore tailings to wheat straw as 3:1 and heating rate as 10 °C/min, the optimum holding time was investigated in Fig.1(c). The dye decomposition efficiency of pyrolyzed product was enhanced when the holding time for the synthesis catalyst stretched from 30 to 60 min. However, the degradation efficiency of MB was not significantly improved when the holding time further extended to 90 and 120 min. Interestingly, \(k_2\) showed a first increasing then
decreasing trend as the holding time extending, and reached the maximum (0.0182 min$^{-1}$) when the holding time was 60 min. Thus, 60 min was the optimal catalyst holding time among those investigated parameters.

To sum up, the relative optimal pyrolysis condition was a blends ratio of 3:1, a pyrolysis temperature of 900 °C and a holding time of 45 min. pH as an important factor for the efficiency of Fenton-like reaction, the effect of pH was investigated in Fig 1(d). We could see that pH could significantly affect the degradation of MB in iron ore tailings catalyzed Fenton-like process. As shown in Fig.1(d), the degradation efficiency was relatively low and the $k_2$ values were 0.0104 min$^{-1}$ and 0.0182 min$^{-1}$ at pH 8.1 and 6.8 (did not adjust). While the decomposition of MB gradually increased as the pH value decreasing and got a highest/fastest degradation efficiency (84% MB removal, 0.0229 min$^{-1}$) at around pH 3, which was consistent with previous studies (Hu et al. 2011). The generation of ·OH from H$_2$O$_2$ is the key step in the entire Fenton-like process, and ·OH catalyzed by iron ore tailings are gradually limited with pH increasing. The higher pH with more OH$^-$ will cause the reaction (Eq. (7)) to shift back and reduce the activity of Fenton reagent (Zheng et al. 2016), which resulted in a slow decomposition rate of I/W(3:1)-900-60.

$$\text{Fe}^{2+} + H_2O_2 \leftrightarrow \text{Fe}^{3+} + OH^- + \cdot OH \quad \text{Eq.(7)}$$
Figure 1 Degradation of MB by pyrolyzed product prepared under different conditions

3.2 Catalyst characterization
Since the catalytic efficiency was apparently different by I-900-60, WS-900-60 and I/W(3:1)-900-60, XRD patterns were collected to indicate the phase information of these samples, shown in Fig.2. The XRD pattern of I/W(3:1)-900-60 showed diffraction peaks at \(2\theta=30.3^\circ, 43.3^\circ, 53.8^\circ, 57.5^\circ, 68.2^\circ\) corresponded to \(\text{Fe}_3\text{O}_4\), which exhibits magnetic characteristics. The peaks of WS-900-60 were noted at \(2\theta=21.8^\circ, 26.5^\circ, 28.0^\circ\), corresponding to \(\text{SiO}_2\) crystallites. The diffraction peaks of raw iron ore tailings (RT) were attributed to \(\alpha\)-\text{FeOOH}, and \(\alpha\)-\text{FeOOH} was converted to \(\text{Fe}_2\text{O}_3\) when heated to 900 °C under \(\text{N}_2\) atmosphere (Zhang et al. 2018). Therefore, only the pyrolysis of iron ore tailings or wheat straw cannot produce a composite with low-valent iron.

![Figure 2 XRD patterns of the raw and prepared samples](image-url)

Significant transformation of \(\text{Fe}_2\text{O}_3\) to \(\text{Fe}_3\text{O}_4\) was observed at 700-900 °C. When the temperature reached above 700 °C, most of \(\text{Fe}_2\text{O}_3\) peaks disappeared, and the diffraction peaks of magnetite appeared. It can be concluded that \(\text{Fe}_2\text{O}_3\) with trivalent could be reduced to \(\text{Fe}_3\text{O}_4\) during high-temperature pyrolysis. More \(\text{Fe}_2\text{O}_3\) spindles were converted to \(\text{Fe}_3\text{O}_4\) as the temperature increasing. This might due to that there were organic matters such as cellulose, hemicellulose and lignin in wheat straw, and
these organic matters could be cracked and devolatilized into reducing gas or liquid, such as H$_2$, CH$_4$, which lead to the ferric iron reduction (Gong et al. 2012; Sharma et al. 2015; Xun et al. 2019; Yunji et al. 2019). In summary, in the process of co-pyrolysis of wheat straw and iron ore tailings, the reducing substances such as H$_2$ or CH$_4$ produced from wheat straw exhibited strong reducibility to reduce iron ore tailings to magnetite. Furthermore, the reduction degree increased as temperature increasing.

To illustrate the effect of biomass/wheat straw on the high degradation efficiency catalyst formation, the morphologies of I-900-60 and I/W(3:1)-900-60 were further compared. The surface morphologies of I-900-60 and I/W(3:1)-900-60 were shown in Fig.S1. The I-900-60 presented a large flaky structure with a flat and non-porous surface after pyrolysis, mainly because the natural structure of iron ore tailings. In the case of I/W(3:1)-900-60, SEM images showed that most particles had a smaller flake structure, with porous surface. This result indicated that mixing wheat straw promote the formation of a porous and smaller flake structure.

| Samples           | Surface area(m$^2$/g) | Pore volume(cc/g) | Average pore size(nm) |
|-------------------|-----------------------|-------------------|-----------------------|
| I-900-60          | 1.1963                | 0.2749            | 12.935                |
| I/W(3:1)-900-60   | 24.319                | 5.5875            | 3.707                 |

As observed from SEM images, I/W(3:1)-900-60 had a smaller particle than I-900-60. N$_2$ adsorption-desorption isotherm was applied to calculate the surface area, pore volume and pore size distribution. As shown in Table 1 and Fig.S2, the average pore size of T-900-60 and I/W(3:1)-900-60 was 12.95 nm and 3.707 nm. However, the total pore volume was 0.2749 cc/g and 5.5875 cc/g, respectively. Thus, we can conclude that wheat straw addition increased the pore volume, and decreased average pore size.
Additionally, although I/W(3:1)-900-60 had a much larger specific surface area (24.319 cm$^2$/g) than I-900-60 (1.1963 cm$^2$/g), its surface area was smaller than those reported iron-load activated carbon adsorbent (300-600 m$^2$/g) (Park et al. 2015; He et al. 2016; Saleh et al. 2017). This result further verified that prepared catalysts from iron ore tailings had weak adsorption ability. Therefore, we speculated the MB degradation by I/W(3:1)-900-60 was due to catalysis instead of adsorption. I/W(3:1)-900-60, with relatively larger surface area and richer pore volume, could provide greater active catalysis sites and increase catalysis performance (Neamţu et al. 2004; Duarte et al. 2012), agreed with the improved MB degradation efficiency catalyzed by I/W(3:1)-900-60.

![Figure 3](image_url)

**Figure 3** (a) the photos of the products attraction to a magnetic device; (b) FTIR spectra of I-900-60 and I/W(3:1)-900-60

There is an important question to further discuss that whether composites (wheat straw biochar and iron ore tailings) were simply mixed and exist alone or integrated together. Fig.3(a) showed the catalysts were attracted as a whole by a magnetic device, indicating iron ore tailings and biochar combined as a whole. This binding form is
beneficial to remove the catalyst from wastewater after reaction. In addition, FTIR
spectra was shown in Fig.3(b). The peak at 3643/3646 cm\(^{-1}\) was attributed to O-H
stretching and bending vibrations (Zhang et al. 2018). Similar I-900-60 peaks were
observed in the spectra of both I-900-60 and I/W(3:1)-900-60, including Si-O stretching
vibrations of the Si-O-Si, Si-O-Al, Si-O-Fe groups (1074 cm\(^{-1}\), 957/981 cm\(^{-1}\), 472 cm\(^{-1}\)), although their intensities varied (Doelsch et al. 2003). The silicon content in iron ore
tailings was much higher than that in wheat straw (Table S1). Therefore, the intensity
of Si-O stretching vibration in I-900-60 was stronger than in the spectrum of I/W(3:1)-
900-60. Meanwhile, new peaks associated with the -C=O and -C-H stretching vibration
at 1224 cm\(^{-1}\) and 876 cm\(^{-1}\) were observed in the spectrum of I/W(3:1)-900-60; these
peaks were assigned mainly to the formation of small organic molecules (such as,
phenol, furfural, styrene) that are devolatilized, oxidized or creaked from wheat straw
during pyrolysis process. In addition, a new peak at 573 cm\(^{-1}\) attributed to an
asymmetric Fe-O stretching vibration was observed. It might be caused by loading iron
into biochar or Fe\(_3\)O\(_4\) itself (Yuan and Dai 2014). Therefore, the new bond Fe-O on
I/W(3:1)-900-60 may indicate the combination of iron and biochar occurred through
chemical bonds.

3.3 The catalyst’s stability and reusability analysis

It is important to evaluate the stability of a heterogeneous catalyst. As illustrated
in Fig.53, I/W(3:1)-900-60 was stable in first three runs and remained high MB
degradation efficiency. At the 4\(^{th}\) run, the activity of I/W(3:1)-900-60 reduced slightly
but the degradation efficiency still higher than 80%, showing the iron ore tailings based
catalyst can be reused for at least 4 Fenton-like cycles without significant activity loss. This slightly activity loss is probably due to the small molecules produced during MB degradation occupying part of the active sites, leading to a decrease in catalytic efficiency (Zhang et al. 2018). In addition, the concentration of leaching iron ions after first three runs was measured. As shown in Table S2, the concentration of leached iron was 0.089 mg/L, 0.085 mg/L, and 0.093 mg/L, which was only 0.8% of the iron content in I/W(3:1)-900-60. Low leached iron concentration also indicated heterogenous Fenton catalysis was the dominant reaction for MB removal (Gao et al. 2017). Meanwhile, XRD and SEM were applied to examine the structure stability of I/W(3:1)-900-60. As illustrated in Fig.S4, compared with fresh catalyst, the crystalline nature and morphology of used I/W(3:1)-900-60 did not change significantly. These results indicated that the cost-effective I/W(3:1)-900-60 was a promising heterogenous catalyst in Fenton-like catalytic degradation of organic wastewater due to its significant stability and reusability.

3.4 Reactive oxidative species and catalysis mechanism

![Figure 4](image_url)

**Figure 4** (a)DMPO-ROS adducts generated from I/W(3:1)-900-60 Fenton catalysis reactions at 10min (b)influence of radical scavenger (TBA) on the catalytic degradation of MB

ROS produced in I/W(3:1)-900-60 was detected by an EPR spin-trap technique
with DMPO. Fig.4(a) showed the EPR spectrum of the generated adducts during the 10 min reaction. On the EPR spectrum, there was observed a four-fold peak with an intensity of 1:2:2:1, which was labeled to the DMPO-OH (Yang et al. 2013). However, the four-fold peak was not appeared in the presence of 300 mmol/L T-Butyl alcohol (TBA, • OH scavenger). Therefore, •OH was the key ROS produced in I/W(3:1)-900-60 catalyzed Fenton-like reactions. Fig.4(b) showed the effect of •OH on MB degradation. We can see the removal efficiency of MB significantly decreased from 84% to 29% in the presence of 300 mmol/L. This result indicated that •OH played a dominant role in MB degradation in I/W(3:1)-900-60 catalyzed Fenton-like reactions.

The electron exchange between Fe(II)/Fe(III) and H2O2 can induce the formation of • OH in the heterogeneous Fenton-like reaction. XPS was applied to analyze the chemical state of iron species on I/W(3:1)-900-60 before and after catalysis reaction. Fig. 5 shows XPS results of Fe2p in fresh and used I/W(3:1)-900-60. The peaks located at 724.8 eV and 710.9 eV were attributed to Fe 2p1/2 and Fe 2p3/2 states of Fe2p orbits, respectively (Gao et al. 2017; Li et al. 2018). Furthermore, the Gaussian-Lorentzian was applied to decompose these two peaks into 6 different fitting peaks(Ding; et al. 2016). Among them, the fitting peaks located at 719.1 and 729.9 eV attribute to satellite peaks, as well as at 712.0 and 725.3 eV assign to Fe3+, and at 710.6 and 723.8 eV correspond to Fe2+, respectively (Li et al. 2018). A summary of deconvoluted peaks’ area and the Fe2+/Fe3+ ratio was presented in Fig.5(a). Apparently, the ratio value of Fe2+/Fe3+ decreased from 2.18 to 2.17 after reaction, demonstrating only a small amount of ≡Fe2+ lost electrons and oxidized to ≡Fe3+ during the catalysis reaction. These XPS
results are in accordance with the good recyclability and stability of I/W(3:1)-900-60 catalyst, which may attribute to the protective effects of the biochar (Li et al. 2018).

To further verify the protective role of biochar during Fenton-like reaction. XPS analysis was conducted to investigate the functional groups changes of I/W(3:1)-900-60 catalyst. As illustrated in Fig.5(b), the C1s spectra can be decomposed into three fitting peaks with C=C sp²/C-Csp³ (284.7 eV), C-O (286.1 eV) and C=O (288.8 eV) (Li et al. 2017b). Apparently, compared with the fresh I/W(3:1)-900-60, the relative content of C=C sp²/C-Csp³ carbon was reduced by 5.8%, C-O and C=O increased by 4.4% and 1.4% after catalysis reaction. This indicates that biochar was oxidized during the Fenton-like process with turning C=C sp²/C-Csp³ carbon to C-O or C=O. Therefore, the recyclability and stability of I/W(3:1)-900-60 catalyst may owe to the existence of biochar which acted as a sacrificial role and limited the oxidation of iron active sites in catalyst. Actually, the biochar can act as a catalyst with electron donor-accepter for the induction of H₂O₂ into ·OH or ·OOH. The persistent free radicals (PFRs) on the surface of biochar formed by the thermal decomposition of organic compounds can be the reduced and oxidized active sites through electron transfer to form radical species ((Khachatryan and Dellinger 2011; Zhu et al. 2018). Fang et al found that PFRs on the
surface of biochar have an important influence on the production of OH by H$_2$O$_2$
activation (Fang et al. 2014). In addition, the PFRs on the biochar surface lead to the
existence of unpaired electrons, which can exchange electrons directly with organic
matter, accelerating the MB degradation efficiency (Fang et al. 2013; Yang et al. 2016).

According to the above analysis results and discussion, the possible Fenton-like
reaction mechanism was illustrated in Fig. 6. At the beginning, the MB molecules were
adsorbed onto biochar of I/W(3:1)-900-60 from aqueous solution through surface
action and pore diffusion. Then, the iron active sites of ≡ Fe$^{2+}$ and PFRs in biochar
simultaneously transfer electron to H$_2$O$_2$ to generate ·OH for MB degradation.
Moreover, the electron transfer between PFRs and iron active sites or the redox cycles
of ≡Fe$^{3+}$/Fe$^{2+}$ combined results in the enhanced degradation efficiency and rate of
heterogenous Fenton-like reaction. Significantly, the stable performance of I/W(3:1)-
900-60 with good reusability is due to the sacrificial effect of biochar for limiting the
oxidation of iron active sites. In addition, the presence of unpaired electrons in PFRs
contributes to a certain extent to improve the degradation efficiency of pollutants. Table 2 showed the MB decompose rate $k_2$ for various Fenton-like catalysts. We can see that I/W(3:1)-900-60(0.0229 min$^{-1}$) exhibited a higher $k_2$ than or comparable to most reported kinetic rate data, which indicates that the co-pyrolysis of iron ore tailings and biomass waste is an effective way to improve the degradation rate/efficiency of iron ore tailings based heterogeneous catalysts. Meanwhile, it is of great significant to realize the resource utilization of iron ore tailings. However, these kinetic rate dates are far less than the degradation rate of homogeneous Fenton catalysis (Gou et al. 2021). Thus, the performance optimization of iron ore tailings-based heterogeneous catalysts based on porous and low-valent iron still needs further studied.

| Catalyst                  | MB (mg/L) | Catalyst dosage (g/L) | Time (min) | $k$ (min$^{-1}$) | Reference               |
|---------------------------|-----------|-----------------------|------------|------------------|-------------------------|
| I/W(3:1)-900-60           | 60        | 3                     | 60         | 2.29×10$^{-2}$   | This study              |
| Fe$_3$O$_4$               | 100       | 3                     | 30         | 1×10$^{-3}$      | (Costa et al. 2008)    |
| Fe$_3$O$_4$/H$_2$/300/1h  | 100       | 3                     | 30         | 4×10$^{-3}$      | (Costa et al. 2008)    |
| Fe$_3$O$_4$/H$_2$/400/1h  | 100       | 3                     | 30         | 2×10$^{-2}$      | (Costa et al. 2008)    |
| Ferrocene                 | 10        | 0.372                 | 120        | 6.17×10$^{-3}$   | (Wang et al. 2014)     |
| Fe$_3$O$_4$/rGO           | 10        | 0.3                   | 120        | 2.6×10$^{-3}$    | (Liu et al. 2013)      |
| Fe$_3$O$_4$/SiO$_2$/C     | 50        | 1                     | 140        | 3.6×10$^{-2}$    | (Liu et al. 2013)      |
| Fe$_3$O$_4$/CeO$_2$       | 100       | 1                     | 120        | 2×10$^{-2}$      | (Li et al. 2017a)      |
| Fe$_3$O$_4$/galic acid/GO | 64        | 1                     | 200        | 1.2×10$^{-2}$    | (Hua et al. 2017)      |
| N,C/CuO-Fe$_2$O$_3$       | 75        | 0.1                   | 180        | 1.08×10$^{-2}$   | (Ren et al. 2019)      |
| FeNi/C-300                | 30        | 1                     | 30         | 1.05×10$^{-2}$   | (Li et al. 2020)       |

4 Conclusions

An iron ore tailings-based Fenton-like catalyst(I/W(3:1)-900-60) with relative fast catalysis rate was constructed by co-pyrolysis(900 °C, 60min holding time) of iron ore
tailings and wheat straw with mass ratio of 3:1. Compared with single pyrolyzed iron ore tailing, the catalytic efficiency and rate of I/W(3:1)-900-60 (0.0229 min⁻¹, 84%) were considerably enhanced for the decomposition of MB due to the electron transfer between biochar and iron active sites or the redox cycles of $\equiv \text{Fe}^{3+}/\text{Fe}^{2+}$. As a result of the sacrificial effect of biochar, oxidizing $\text{C}═\text{Csp}^{2}$ bonds and limiting the deactivation of iron active sites($\equiv \text{Fe}^{2+}$), I/W(3:1)-900-60 showed a good reusability and stability. Moreover, the presence of unpaired electrons in persistent free radicals (PFRs) of biochar accelerated the electron exchange and further enhanced the MB decomposition rate. This work opens up a way to synthesize an iron ore tailings-based Fenton-like catalyst with higher degradation rate, as well as realize the utilization of solid wastes.

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**Author contributions**

Lihui Gao: Writing-original draft, methodology, Data curation; Lizhang Wang: Writing-Reviewing and Editing; Shulei Li: Investigation, Sample preparation; Yijun Cao: Supervision and Editing.

**Availability of data and materials**: All data generated or analyzed during this study are included in this published article.

**Declarations**

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