



Immobilized laccase on bentonite-derived mesoporous materials for removal of tetracycline

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HIGHLIGHTS

- BDMMs were expediently obtained from alkali & acid treatment of original bentonite.
- Surface areas and average pore sizes of BDMMs were improved nearly by 74 and 2 times.
- The thermal stability of BDMMs-Lac was highly improved compared to free laccase.
- In the presence of HBT, BDMMs-Lac could remove 60% of TC within 180 min.

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ABSTRACT

Bentonite is a natural and environmentally clay mineral, and bentonite-derived mesoporous materials (BDMMs) were obtained conveniently from the alkali and acid treatment of bentonite. In the present study, BDMMs were explored for immobilization of laccase obtained from *Trametes versicolor*. As a result, bentonite-derived mesoporous materials-Laccase (BDMMs-Lac) was developed for the removal of tetracycline (TC). The enzyme immobilization process was carried out through physical adsorption contact (ion exchange adsorption, hydrogen bond adsorption, and Van der Waals adsorption) between the BDMMs and laccase. The process of immobilization remarkably increased its operating temperature. The BDMMs-Lac exhibited over 60% removal efficiency for TC within 3 h in the presence of 1-hydroxybenzotriazole (HBT). In conclusion, BDMMs-Lac showed more promising potential than free laccase for practical continuous applications.

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1. Introduction

Laccase (EC 1.10.3.2) is an oxidoreductase that belongs to the multicopper oxidase protein family (Huang et al., 2017a,b; Madhavi and Lele, 2009; Zhang et al., 2014). Laccase has the ability to

catalyze some substrates to water (Spina et al., 2015; Huang et al., 2016a,b). In the presence of small molecular weight mediators, laccase has more extensive substrate range and thus exhibits wider applicability in polluted water (Cheng et al., 2016a,b; Chen et al., 2016; Rodriguez and Toca, 2006). The use of laccases also offers a method that is free from secondary pollution during actual wastewater treatment (Lai et al., 2016; Liu et al., 2013; Monje et al., 2010). However, the low stability and high production costs of laccase limit its applicability (Ashe et al., 2016; Li et al., 2018).

Immobilization can overcome the limits of laccase application by enhancing the enzyme properties (Mohamad et al., 2015; Cheng

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et al., 2016a,b). The immobilization methods of laccase have been explored for years (Deng et al., 2013; Guzik et al., 2014; Zhou et al., 2018a,b). Immobilization can increase the stability of enzymes and thus improve the operability of laccase in practice (Lai et al., 2019; Sheldon and van Pelt, 2013). Multifarious carriers have been studied for the successful immobilization of laccase (Zhou et al., 2013; Liu et al., 2012). Clays are low-cost, eco-friendly, recyclable, have low mass transfer, and demonstrate microbial corrosion resistance capacity (An et al., 2015; Li et al., 2015; Liang et al., 2017; Wu et al., 2017). Through activation or etching, they can attain highly specific surface areas and numerous functional groups (Gong et al., 2009; Osuna et al., 2018; Shu et al., 2016; Zeng et al., 2017).

Bentonite, which has layered structure with cations such as Na^+ or Ca^{2+} , shows promising and highly suitable application for the loading of an extensive range of biomolecules (Liang et al., 2017; Ghiaci et al., 2009; Ma et al., 2018). After etching, bentonite exhibited highly improved characteristics, including those relation to cation exchange capacity and surface area (Liang et al., 2017; Bajpai and Sachdeva, 2002; Shu et al., 2014). Furthermore, bentonite, as a natural mineral, is eco-friendly, inexpensive, and accessible (Long et al., 2011; Issaabadi et al., 2017). The application of bentonite for enzyme immobilization has been studied by several research groups (Salem and Salem, 2017; Andjelkovi et al., 2015). Conversely, the utilization of mesoporous and high surface area bentonite for the immobilization of laccase and other different biocatalysts remains to be explored (Xu et al., 2012a,b; Andjelkovi et al., 2015; Zhou et al., 2018a,b).

Antibiotic pollution has become of increasing environmental concern (Manaia et al., 2016). Antibiotics are widely utilized to treat diseases caused by various bacterial or pathogenic microbes, however, their misuse and over accumulation threaten the environment (Liu et al., 2016; Polese et al., 2016). Tetracycline (TC) is one of the most widely used antibiotics (Nasseh et al., 2018; Gothwal and Shashidhar, 2015). The poor degradation of TC from traditional municipal wastewater treatment plants has led to a latent negative impact on aquatic organisms, thus necessitating the exploration of treatment technologies (Halling-Sørensen, 2002; Huang et al., 2017a,b; Tan et al., 2015). Among the numerous treatment methods, the biodegradation of TC by laccase or immobilized laccase is effective (Islas-Espinoza et al., 2018; Xu et al., 2012a,b).

Although modified bentonite materials have been frequently applied to immobilize enzymes, the use of mesoporous and high surface area bentonite for laccase immobilization has not been explored (Andjelkovi et al., 2015; Ghiaci et al., 2009; Liu et al., 2012). Bentonite can be modified to be mesoporous and to possess a high surface area (Toor et al., 2015; Önal and Sarıkaya, 2007). NaOH-HCl etching modification is an alkali/acid activation composite modification process (Önal and Sarıkaya, 2007). This method has been utilized for the etching of clay materials such as Halloysite, Kaolinite, from which mesoporous materials have successfully obtained (Li et al., 2015; Zhou et al., 2014). However, the use of alkali/acid activation composite modification for bentonite has not been explored. Thus, in this study, bentonite-derived mesoporous materials (BDMMs) were constructed by NaOH-HCl etching. The BDMMs were utilized for laccase immobilization to obtain bentonite-derived mesoporous materials-Laccase (BDMMs-Lac), and the characteristics of BDMMs and the treatment capacity of BDMMs-Lac were explored. BDMMs-Lac was applied for TC antibiotic removal in the presence of the redox mediator 1-hydroxybenzotriazole (HBT). This study is aimed at establishing new eco-friendly, low-cost, and re-usable carriers for immobilizing laccase and for exploring the treatment capacity and removal ability of immobilized laccase for emerging antibiotic pollutants.

2. Material and methods

2.1. Materials

Laccase ($\geq 0.5 \text{ U mg}^{-1}$) from *Trametes versicolor*, HBT, TC, and 2,2-azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) were obtained from Sigma-Aldrich (St. Louis, MO, USA). Bentonite was provided by Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China). All of the other chemicals were of analytical grade.

2.2. Etching of the bentonite

Pristine bentonite was added to NaOH (6 M) and stirred. The bentonite was then washed five times with ultrapure water, dried at 383 K for 12 h, and then added to HCl (5 M) at 353 K with constant stirring for 6 h. The above material was then washed and dried to obtain BDMMs.

2.3. Laccase activity assays

Laccase activity was tested using ABTS as a substrate (Zhang et al., 2014). Briefly, the assay compound consisted of 0.1 M citrate buffer (pH = 5), 1 mM ABTS and free laccase or BDMMs-Lac samples. The activity of BDMMs-Lac and free laccase was detected at an absorbance of 420 nm (UV-2250, Shimadzu Corp., Japan). One unit of laccase activity was defined as the amount of BDMMs-Lac or free laccase required to oxidize 1 μM of substrate per minute.

2.4. Laccase immobilization

The BDMMs were suspended in citrate phosphate buffer (0.1 M, pH = 3–8) containing laccase (0.5–4 mg/mL). The mixtures were then incubated. Later, the sample was centrifuged and the bottom solid was collected and washed several times with citrate buffer (0.1 M, pH = 5). The final solid BDMMs-Lac was obtained after freeze drying at 173 K for 12 h. Fig. 1 depicts the typical process for the stepwise etching of pristine bentonite, and the adsorption loading of laccase.

2.5. Stability assessment

2.5.1. Thermal stability

For temperature stability, free laccase and immobilized laccase were added to centrifuge tubes containing citrate buffer (pH = 5)

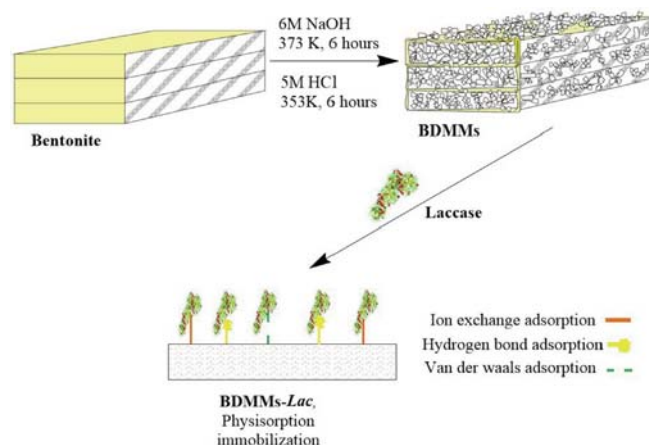


Fig. 1. Schematic of BDMMs preparation and succeeding laccase physisorption immobilization on BDMMs.

and were maintained at 303 K–353 K for 120 min. They reacted with the ABTS and were centrifuged and then measured at 420 nm (UV-2250, Shimadzu Corp.).

2.5.2. Reusability of immobilized laccase

The BDMMs-Lac was dispersed in citrate-phosphate buffer (pH 5) containing 1 mM ABTS and then incubated at 303 K. The sample was centrifuged (6570×g) and the concentration of the transformed ABTS was measured. The BDMMs-Lac was washed with citrate-phosphate buffer. The above procedure was repeated for 10 cycles.

2.6. Immobilized laccase system for the removal of TC

The effect of parameters such as BDMMs-Lac dosage (0.5–4 mg/mL) and reaction time (10–180 min) were studied. The reaction mixture containing BDMMs-Lac and 10 mg/L of TC solution was placed at 303 K for 120 min. TC was tested at the absorbance of 360 nm (UV-2250, Shimadzu Corp.). All of the experiments were examined in triplicate. To determine the possible removal of TC due to adsorption onto the BDMMs, heated-devitalized BDMMs-Lac was used to remove the TC.

3. Results and discussion

3.1. Structural characterization

The morphologies of the bentonite, BDMMs, and BDMMs-Lac samples are presented on Scanning Electron Microscopy (SEM) images (Fig. 2). Fig. 2 (a) illustrates the unbroken structure of the crude bentonite, which consisted of homogeneous particles. Fig. 2 (b) indicates the etching appearance of BDMMs whereby the integrated particles were visually damaged and the interlamellar spacing was enlarged. Relevant Energy dispersive spectroscopic (EDS) analysis confirmed that no obvious elemental change occurred after etching (Fig. 2 (b)). Fig. 2 (c) and Fig. 2 (d) showed no alteration in the structure of BDMMs-Lac before or after degradation in comparison with BDMMs.

The N₂ adsorption-desorption curves of the samples are presented in Fig. 3A. The values of BDMMs were highly elevated in contrast to that of original bentonite. The plot style also changed from III style (H3 hysteresis loop) to V style (H4 hysteresis loop) (Zhang et al., 2016; Yu and Zhang, 2010). The hysteresis loop showed that both bentonite and BDMMs consisted of slit holes, which were formed by the accumulation of flaky particles or layered structures (Yang et al., 2010; Chen et al., 2017). The BET results indicated that the pristine bentonite had a surface area equal to 3.30 m²/g, a pore size equal to 2.73 nm and a pore volume equal to 22.46 mm³/g. Meanwhile, the surface area of BDMMs was 244.62 m²/g, the pore size was 5.53 nm, and the pore volume was 338.8 mm³/g. The specific surface areas were higher than that detected in previous researches (Bajpai and Sachdeva, 2002; Ghiaci et al., 2009).

Fig. 3B shows the FTIR spectra of bentonite, BDMMs, BDMMs-Lac, and BDMMs-Lac after degradation. The broad adsorption band around 3438 cm⁻¹ among all of the samples could be attributed to the stretching vibration of O-H caused by water molecules that are present in the hydrogen bonded interlayer (Jiang et al., 2018; Ztrk et al., 2008). The adsorption band at 1637 cm⁻¹ in all of the samples indicates the stretching vibration of crystal water molecules in the lattice (Ztrk et al., 2008). The band at 1429 cm⁻¹ was presumed to represent the symmetric stretching vibration absorption peak of -COOH (Wen et al., 2019; Chen et al., 2017; Tang et al., 2014). The absorption bands around 1027 and 696 cm⁻¹ of spectrum a, b, c and d were caused by the bending vibration of Si-O-Si and Si-O,

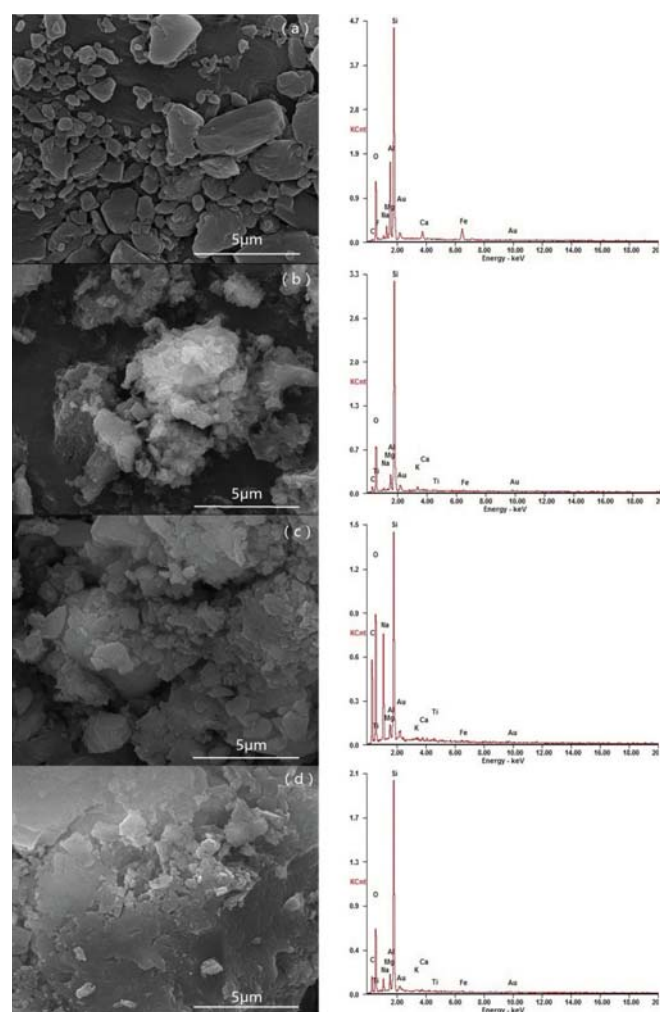


Fig. 2. SEM images and related EDS of a) bentonite, b) BDMMs, c) BDMMs-Lac, and d) BDMMs-Lac after TC degradation.

respectively (Huang et al., 2016a,b; Ztrk et al., 2008). However, the band at 3627 cm⁻¹ was interpreted as the stretching vibration of O-H due to the existence of interlayered adsorption water molecules that disappeared after etching (Huang et al., 2015). The same phenomenon was also observed in the peaks of 2352, 829, and 462 cm⁻¹. The presence of narrow bands at 2352 cm⁻¹ might correspond to the impurities mixed in the bentonite. The other bands in the range of 500–800 cm⁻¹ were the lattice vibration of M-O, M-O-M, and O-M-O (Andjelkovi et al., 2015). Their changes among the different curves may be attributed to the ion exchange and reagent reaction during the etching process (Li et al., 2015).

The X-ray diffraction (XRD) patterns of the bentonite and BDMMs are displayed in Fig. 3C. The characteristic reflection of bentonite at 5.8° belonged to montmorillonite (Chen et al., 2017). It was disappeared after etching. The reductions in BDMMs may be due to the activation of etching reagents. The basal space reflections presented a sharp peak at 2θ = 26.64° in the XRD spectrum of the bentonite and BDMMs samples and indicated a (101) basal spacing of 1.54 nm (JCPDS Card No. 46–1045) (Toor et al., 2015). The characteristic XRD peaks for quartz (2θ = 26.64°, 42.45°, 68.32°), marked by their indices (101), (200), (301), were almost identical between the bentonite and BDMMs. No obvious shifts in the characteristic peaks of the bentonite and BDMMs were observed, demonstrating that there was no expansion in interlamellar

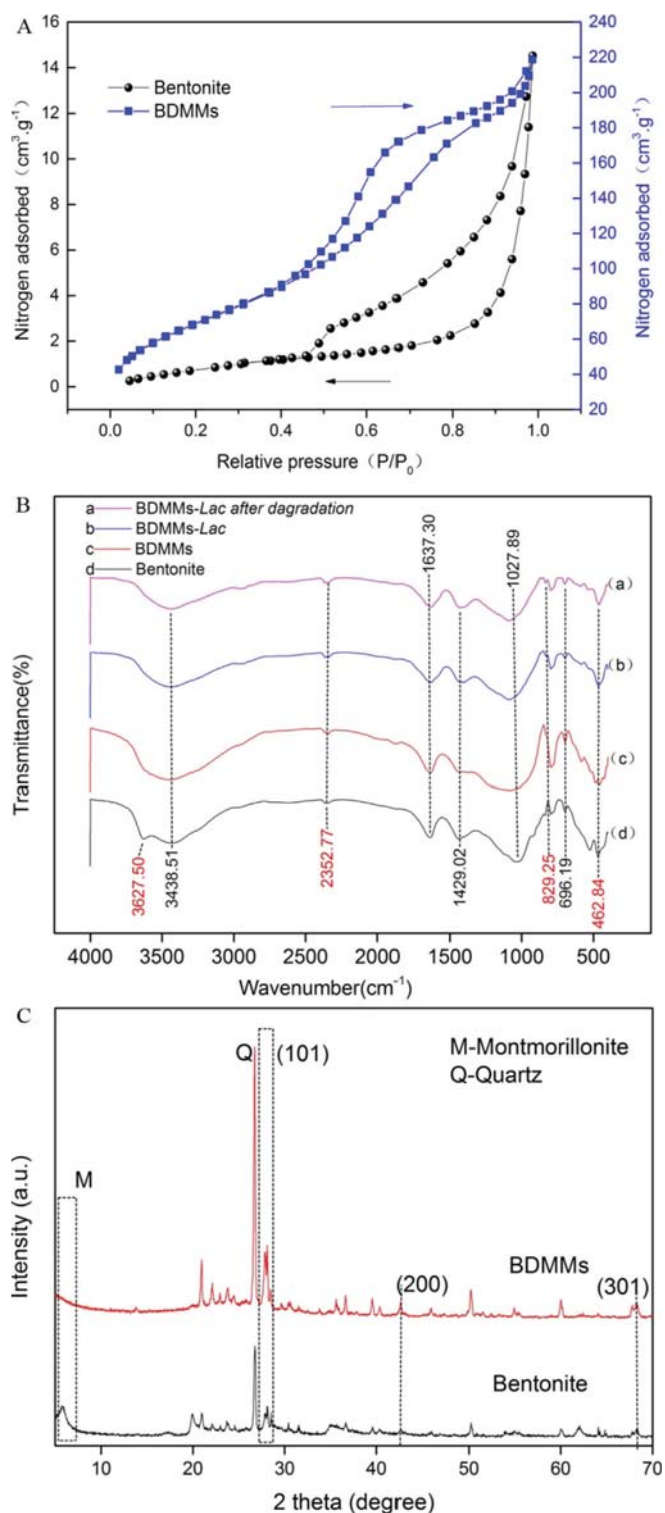


Fig. 3. A) BET nitrogen adsorption/desorption plots of the bentonite and BDMMs. B) FT-IR spectra of bentonite, BDMMs, BDMMs-Lac, and BDMMs-Lac after degradation. C) XRD curves of the bentonite and BDMMs.

spacing. Thus, the same XRD patterns of the bentonite and BDMMs confirmed that they possessed the same crystal structure and interplanar spacing.

3.2. Optimum conditions of laccase immobilization

Immobilization using bentonite as a support material is influenced by many factors (Liu et al., 2012). As shown in Fig. 4A, when the initial laccase concentration increased from 0.5 to 4 mg/mL, the loaded laccase on the bentonite also increased. However, the activity of the immobilized laccase only increased until 2 mg/mL. When the laccase concentration exceeded 2 mg/mL, a decrease in the activity recovery of BDMMs-Lac was observed. Some similar observations have been made in previous studies (Kadam et al., 2017). This phenomenon could be attributed to the overloading of laccase on supports, as the overloading of laccase on the surface of the supports would result in the congestion or crowding of the laccase molecules (Liu et al., 2012). Diffusion-controlled limitations appeared when the laccase loading was high. The agglomeration or crowding of laccase also resulted in the conformational change of the laccase molecules, and a suitable laccase concentration was found to be important for maintaining laccase activity. Thus the optimum laccase concentration was set as 2 mg/mL for the subsequent analyses.

As depicted in Fig. 4B, the activity and the relative activity of BDMMs-Lac changed with the increase in immobilization time from 15 to 180 min. The relative activity of BDMMs-Lac increased remarkably until 30 min, following which the relative activity remained the same from 30 to 120 min. The activity of BDMMs-Lac almost reached 800 U/g, following which the activity and relative activity began to decline. The activity of the immobilized enzymes depends on the nature of the enzyme protein (Liu et al., 2012). As time progressed, the possible amounts of inactivated laccase increased during immobilization, and the laccase flexibility declined. With the increase in physical adsorption immobilization time, the adsorption site on BDMMs was eliminated. The relevant steric hindrance and diffusion limitations might have also resulted in the decrease in laccase activity (Liu et al., 2012).

The effect of solution pH on the activity of free and BDMMs-Lac was explored at different pH values ranging from 3.0 to 8.0 (Fig. 4C). The free and immobilized laccase typically demonstrated maximal activity at pH 4.0 and pH 5.0. The variation in optimum pH was also previously surveyed in immobilized laccase on magnetic bimodal mesoporous carbon (Liu et al., 2012). It may be attributed to the electrostatic interaction affected by the support microenvironment around the laccase. Different pH values resulted in different microenvironments. The isoionic point influenced the net charge of the laccase and carrier such that the laccase activity could be hindered or invoked (Chen et al., 2015; Liu et al., 2012; Zhang et al., 2015). BDMMs-Lac showed better adaptability when the pH value was above 5. As the pH increased to 6, the free laccase and BDMMs-Lac maintained 37% and 48% of their relative activity, respectively. To a certain extent, this result indicated that immobilization could retain laccase activity.

3.3. Properties of BDMMs-Lac

Operational stability is important for determining processing costs (Liu et al., 2012). The results presented in Fig. 5A showed that BDMMs-Lac lost 37% and 64% of its original activity after three and five cycles, respectively. The physical adsorption immobilization exhibited weak binding forces between enzyme and carrier. Thus, the activity loss may have resulted from the laccase leaching during the washing stages (Skoronski et al., 2017).

The thermostability of free laccase and BDMMs-Lac was explored over a temperature range of 303 K–353 K. As indicated in Fig. 5B, BDMMs-Lac was more stable than the free laccase, and both free laccase and BDMMs-Lac presented their highest stability at 313 K. Furthermore, between 323 K and 353 K, the immobilized

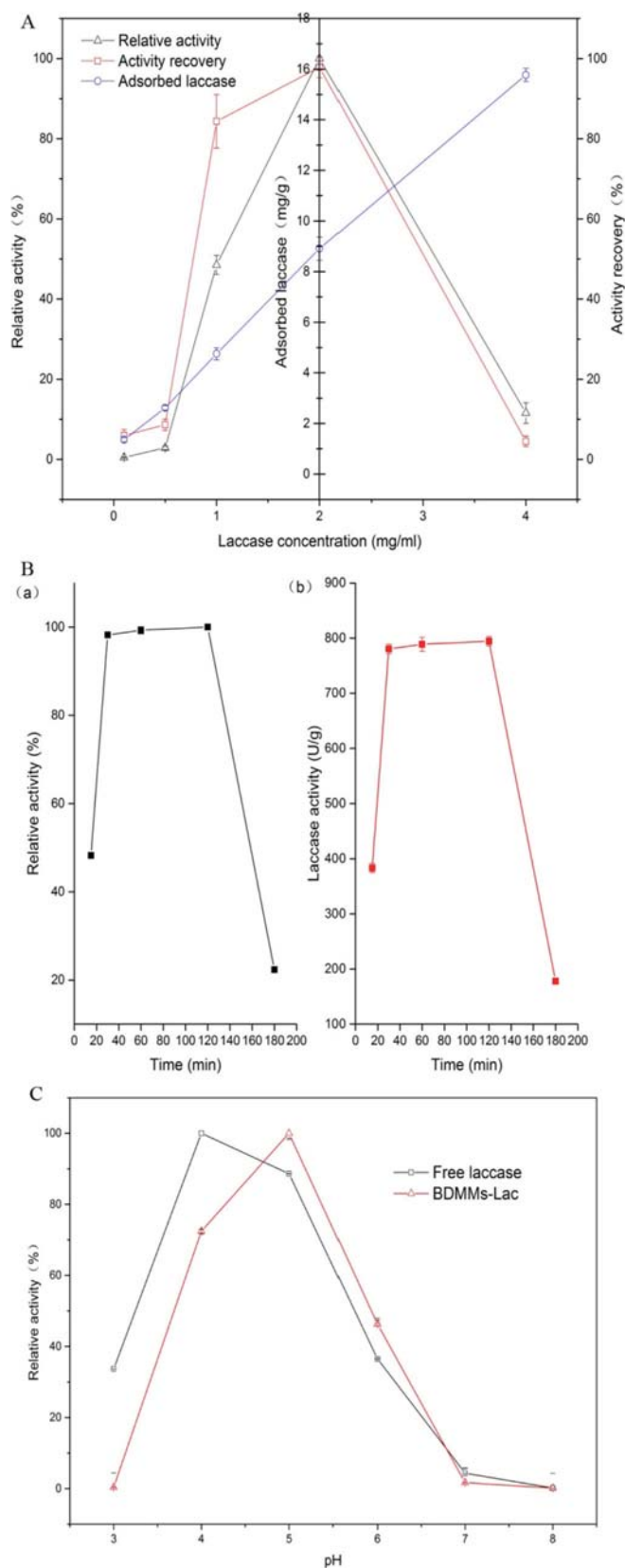


Fig. 4. A) Effect of laccase concentrations from 0.5 mg/mL to 4 mg/mL on the activity of the immobilized laccase. B) Effect of time from 15 min to 180 min on the activity of the immobilized laccase. C) Effect of pH from 3.0 to 8.0 on the activity of the free and immobilized laccase.

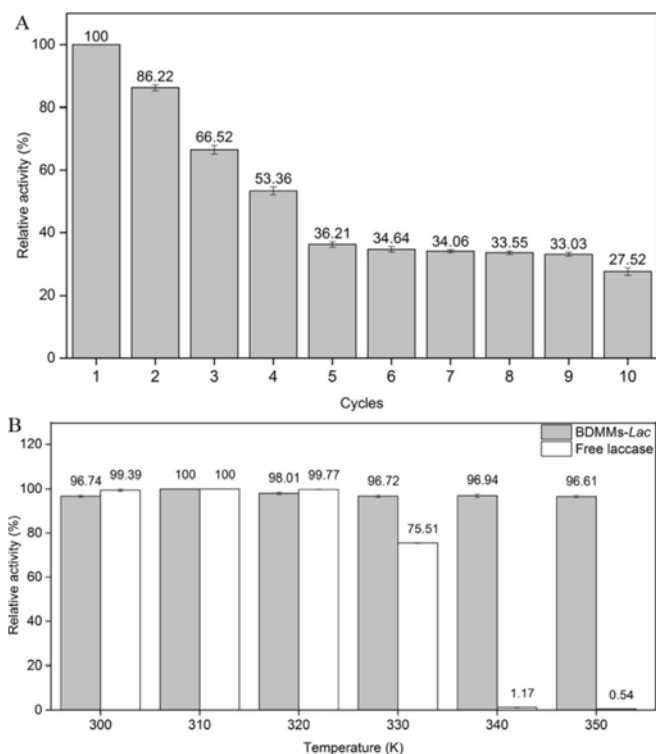


Fig. 5. A) Operational stability of BDMMs-Lac in continuous cycles. B) Thermal stability studies of free laccase and BDMMs-Lac at 303–353 K for up to 120 min.

laccase maintained 96% of its initial activity, while free laccase could only retain 0.54% of its initial activity when the temperature exceeded 343 K. The results were attributed to the high thermostability of BDMMs-Lac towards denaturation. Immobilization increased laccase rigidity and decreased laccase conformational flexibility (Andjelkovi et al., 2015). The highly improved thermal stability of BDMMs-Lac benefits its application in high-temperature industrial processes (Menezes-Blackburn et al., 2011).

3.4. Removal of TC

The effect of reaction time on removal of TC is displayed in Fig. 6A. The removal of TC could be attributed to the combined effects of degradation by BDMMs-Lac and the adsorption by the BDMMs support. As shown in Fig. 6A, approximately 60% of the TC was removed in 120 min by BDMMs-Lac. The more important contribution of the laccase catalytic process could thus be confirmed, as the adsorption only contributed approximately 20% of the removal. However, the result also revealed the benefit of employing BDMMs as immobilization support in the removal process. The accumulation of the catabolite might inhibit the removal process, which was reported in a previous study (Yang et al., 2017).

The relationship between immobilized laccase dosage and TC removal is presented in Fig. 6B. The removal efficiency of TC gradually increased with increased in immobilized laccase dosage from 0.4 to 4 mg/mL. When the dosage was 4 mg/mL, the removal efficiency reached 52%. When the dosage was 2 mg/mL, the removal amount reached 1.85 mg/g. The removal efficiency began to decrease when the BDMMs-Lac dosage was higher than 4 mg/mL, and this phenomenon was attributed to excessive dosage, leading to contact site reduction between TC and mediator HBT, as well as contributing to the consumption of laccase activity (Sun et al., 2017).

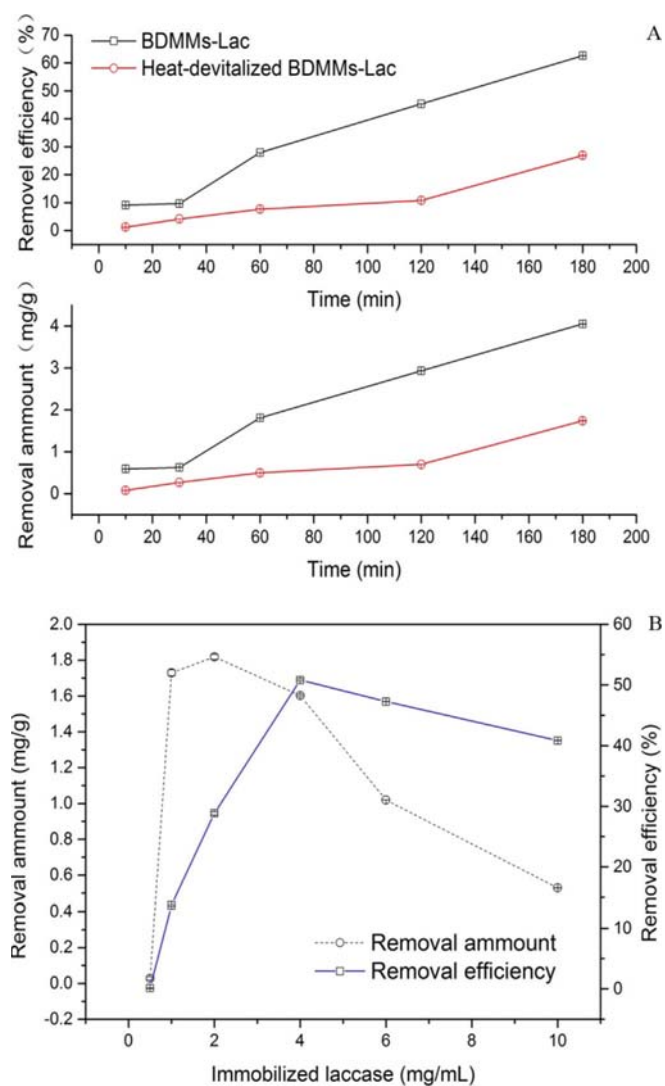


Fig. 6. A) Time-course of the removal and adsorption rates for TC by BDMMs-Lac and the heated-deactivated BDMMs-Lac. B) Effect of immobilized laccase dosage on the removal rates of TC by BDMMs-Lac.

4. Conclusions

The surface area, average pore size, and pore volume of BDMMs obtained from this study were all increased ($3.3 \rightarrow 244.62 \text{ m}^2/\text{g}$, $2.73 \rightarrow 5.53 \text{ nm}$, $22.46 \rightarrow 338.8 \text{ mm}^3/\text{g}$). The stability of BDMMs-Lac was improved compared to free laccase, particularly thermal stability. The biodegradation rate of BDMMs-Lac for TC reached nearly 60%. This study showed that BDMMs could be conveniently and efficiently obtained and has potential applicability in further practical biomacromolecule immobilization. Furthermore, as the obtained BDMMs-Lac is an economical and eco-friendly biocatalyst, it has wide applicability for the elimination of micropollutants from wastewater.

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