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Magnetic-responsive solid acid catalysts for esterification†

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Two kinds of magnetic-responsive solid acid catalysts were designed and prepared *via* an *in situ* polymerization of poly(ionic liquid)s (PILs) on the surface of Fe₃O₄@SiO₂ NPs for the catalyzed esterification of palmitic acid and methanol. They were characterized using XRD, TGA, VSM, NMR spectroscopy, FTIR spectroscopy, XPS, SEM, and GC techniques. The results confirmed the preparation of solid acid catalysts. Meanwhile, they possessed excellent catalytic activity and recyclability. The effect of the reaction conditions on the esterification was investigated through single-factor analyses, and the proposed catalytic mechanism of the prepared solid acid catalysts in the esterification are also discussed. Under the optimal reaction conditions (10 wt% catalyst, 6 h, 70 °C, and molar ratio (MR) of methanol to palmitic acid of 12 : 1), the conversion rate of palmitic acid could reach 94% and 79% with Fe₃O₄@SiO₂-poly(1-vinyl-3-ethylimidazolium phosphotungstate) (Fe₃O₄@SiO₂-P([VLIM]PW)) and Fe₃O₄@SiO₂-poly(1-vinylimidazole-3-propyl sulfonate) (Fe₃O₄@SiO₂-P([VLIM]SO₃)) NPs serving as catalysts, respectively. Furthermore, the Fe₃O₄@SiO₂-P([VLIM]PW) NPs could still maintain a high catalytic activity even after being reused 5 times without significant deactivation.

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1. Introduction

Esterification is one of the most important reactions in the pharmaceutical and chemical industries. However, the esterification of organic acids to esters is much slower than catalyzed esterification kinetically.¹ Thus, it is critical to develop an appropriate catalyst for esterification. The catalyst could be an acid or base.²⁻⁴ Normally, a base catalyst is more effective than an acid catalyst in the esterification reaction due to its excellent catalytic activities.⁵ However, a base catalyst could directly react with an organic acid and the corresponding salt has to be removed from the product for the purification of the product.⁶ Meanwhile, homogeneous acid catalysts, such as sulfuric acid, hydrochloric acid, and nitric acid, suffer from problems related to equipment corrosion, environmental pollution, and non-recyclability.⁷⁻⁹ Based on the above analysis, solid acid catalysts could avoid the above problems. Meanwhile, the purification processes for the product could be greatly reduced.

Biodiesel is a mixture of different fatty acid esters, which could be prepared by the esterification of fatty acids and short chain alcohols. Compared with traditional fossil fuels, biodiesel

not only is renewable and biodegradable but also possesses similar functions in terms of performance and engine durability.¹⁰⁻¹² Furthermore, it emits less CO and particles in combustion. Therefore, the employment of renewable biodiesel is a feasible solution to help alleviate the current energy and environmental crisis.¹³⁻¹⁵ In the recent years, researchers have paid more attention to developing solid acid catalysts for the preparation of biodiesel. Several kinds of solid acid catalysts, such as polymers,^{16,17} ionic liquids (ILs),¹⁸⁻²⁰ poly(ionic liquids) (PILs),²¹⁻²³ and carbon-derived solid acids and their composites,²⁴⁻²⁶ have been prepared and applied in various fields.²⁷ Among the above-mentioned materials, acidic PILs are the most prominent catalysts owing to their remarkable properties of designability, environmental friendliness, strong acidity, high stability and so on.²⁸⁻³⁰ However, they are easily soluble in polar solvents, which brings difficulties in the recovery of the catalyst and purification of the product. Thus, it would be necessary to immobilize the acid PILs on the surface of recyclable solid materials.

A number of studies have reported that heteropoly acids with the Keggin structure, sulfonic acid, and their salts, including phosphotungstic acid (PTA), vanadium-substituted phosphomolybdic acid, phosphomolybdic acid, heterogeneous cesium hydrogen phosphotungstate, and sulfonated imidazolium ionic liquid, could serve as effective catalysts owing to their strong acid properties.^{9,31-34} Thus, the combination of organic cation and anion groups with strong acid properties is interesting as they would form a kind of novel ionic liquid (IL). These could

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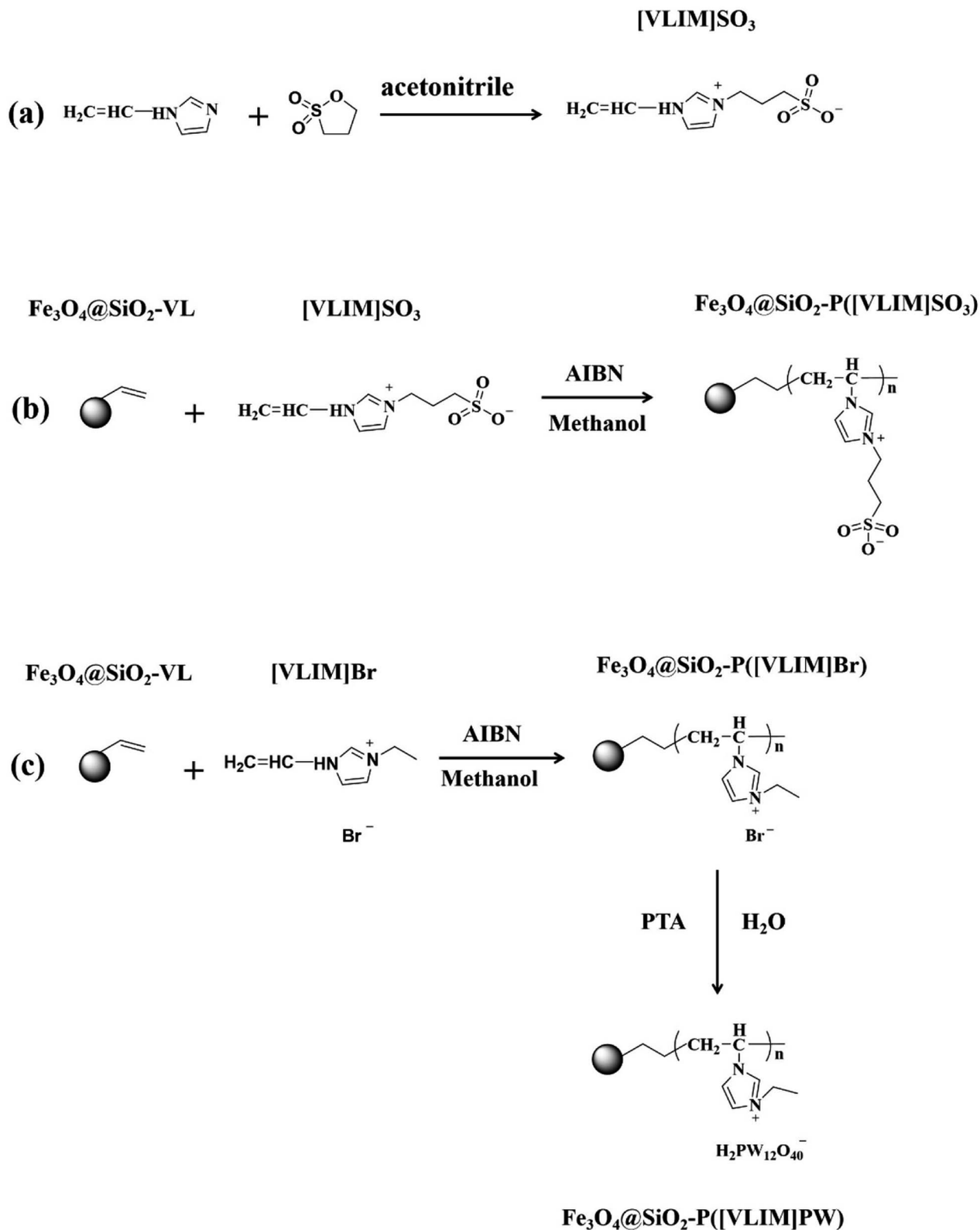


Fig. 1 Schematic of the preparation of the magnetic-responsive solid acid catalysts.

act as effective solid acid catalysts. The immobilization of ILs through chemical bonds should be considered to avoid the leaching or deactivation. As is well known, magnetic materials

can be easily separated and recycled by an external magnet. Some magnetic-responsive solid acid catalysts have been successfully prepared and applied in esterification.³⁵⁻³⁸



However, there are scant reports on the immobilization of PILs on the surface of magnetic-responsive materials through chemical bonding *via* a simple method. Therefore, in this work, we designed and prepared two kinds of recyclable and efficient solid acid catalysts *via* a series of simple reactions, such as solvothermal, sol-gel, polymerization, and ion exchange: phosphotungstic (PW)- and sulfonic (SO₃)-based PILs-modified Fe₃O₄@SiO₂ NPs, respectively (Fig. 1). These solid acid catalysts were applied in the preparation of biodiesel, which was achieved by the esterification of palmitic acid (model compound) and methanol to evaluate their catalytic activity. The prepared solid acid catalysts were characterized by XRD, VSM, TGA, NMR, XPS, FTIR, SEM, and GC. Meanwhile, the effects of the reaction conditions on the esterification reaction and conversion rate of palmitic acid were investigated through single-factor analysis. Furthermore, the proposed catalytic mechanism of the prepared solid acid in the esterification process and the recyclability of the magnetic-responsive solid acid catalysts were investigated and are discussed as well.

2. Experimental

2.1. Materials

Ethylene glycol, iron (III) chloride hexahydrate, sodium acetate trihydrate, tetraethyl orthosilicate (TEOS), ammonium hydroxide (NH₃·H₂O, 25 wt%), and potassium hydroxide (KOH) were purchased from Sinopharm Chemical Reagent Co., Ltd. 2,2-Azobis-(2-methylpropionitrile) (AIBN), vinyltrimethoxysilane (VL-TMC), 1-vinyl-3-ethylimidazolium bromide ([VLIM]Br), PTA, 1-vinylimidazole (VLIM), 1,3-propanesultone, ethanol, acetonitrile, methanol, and palmitic acid were purchased from Adamas Reagent Co., Ltd. All the reagents were analytical grade and used as received.

2.2. Preparation of Fe₃O₄@SiO₂-VL NPs

First, 0.1 g of Fe₃O₄@SiO₂, which was first prepared as per our previous work,^{39,40} was first dispersed in a solution of 60 mL of ethanol under ultrasonication. Then, 200 μL of VL-TMC was added and stirred for 24 h at room temperature. Afterward, the mixture was heated to 80 °C for 5 h under continuous stirring. After the modification, the obtained black precipitate was collected with a magnet and washed with deionized water and ethanol three times to remove residual reactants. Finally, the product, named as Fe₃O₄@SiO₂-VL NPs, was obtained after drying at 50 °C in a vacuum oven for 24 h.

2.3. Preparation of [VLIM]SO₃

First, 0.94 g of 1-vinylimidazole and 1.12 g of 1,3-propanesultone were slowly added into 60 mL of acetonitrile, placed in an ice bath, and continuously stirred for 0.5 h. After that, the mixture was heated to 60 °C and stirred for 24 h under a nitrogen atmosphere. The product was washed with acetonitrile to remove any unreacted reagents and dried in a vacuum oven at 50 °C for 24 h. The achieved white solid powder was named as [VLIM]SO₃ (Fig. 1a).

2.4. Preparation of Fe₃O₄@SiO₂-P([VLIM]SO₃) NPs

First, 0.1 g of Fe₃O₄@SiO₂-VL NPs, 1 g of [VLIM]SO₃, and 0.01 g of AIBN were dispersed in 60 mL of methanol under ultrasonication for 3 min to form a uniform suspension at room temperature. Then, a nitrogen atmosphere was flowed into the mixture for 0.5 h to remove dissolved oxygen. Afterward, the mixture was heated to 80 °C and continuously stirred for 24 h under a nitrogen atmosphere. The obtained black precipitate was collected with a magnet and washed with methanol to remove any residual reactants. Then it was dried in a vacuum oven at 50 °C for 24 h. The achieved black solid powder was named as Fe₃O₄@SiO₂-P([VLIM]SO₃) NPs (Fig. 1b).

2.5. Preparation of Fe₃O₄@SiO₂-P([VLIM]PW) NPs

Fe₃O₄@SiO₂-P([VLIM]PW) NPs were prepared in a two-step method. The first step was the preparation of Fe₃O₄@SiO₂-P([VLIM]Br) NPs, which followed a similar method to that used for Fe₃O₄@SiO₂-P([VLIM]SO₃), except that the monomer used for the polymerization was [VLIM]Br. In the second step, 0.1 g of Fe₃O₄@SiO₂-P([VLIM]Br) NPs and 1.0 g of PTA were dispersed in 30 mL of deionized water, and then stirred at 80 °C for 24 h. The obtained black precipitate was collected with a magnet and washed with deionized water and ethanol several times to remove any residual PTA, and then dried at 50 °C in a vacuum oven for 24 h. Thus, Fe₃O₄@SiO₂-P([VLIM]PW) NPs were obtained (Fig. 1c).

2.6. Preparation of biodiesel

Fe₃O₄@SiO₂-P([VLIM]SO₃) and Fe₃O₄@SiO₂-P([VLIM]PW) NPs were, respectively, used as the catalysts in the preparation of biodiesel. A stoichiometric amount of catalyst, methanol, and palmitic acid were successively added in to a round-bottom flask equipped with a reflux condenser and mechanical stirrer. The mixture was stirred at 70 °C for a certain time to obtain the biodiesel.

For the recycling studies, the catalyst was washed and recovered with hot water after it was collected by a magnet. Afterward, the catalyst was further dried in a vacuum oven at 50 °C for 24 h for it to be reused. The recycled catalyst was reused under the same conditions.

As the amount of methanol was excessive and sufficient to drive the reaction forward toward the conversion of palmitic acid to palmitic methyl ester in the esterification, thus the conversion rate of palmitic acid was used to evaluate the catalytic activity, and it was calculated using the expression below.

$$\text{Conversion rate} = \frac{A_1 - A_2}{A_1} \times 100\%$$

where, A₁ and A₂ are the amounts of the initial and final palmitic acid, respectively.

3. Results and discussion

3.1. Characterization of [VLIM]SO₃

The molecular structure of [VLIM]SO₃ was characterized by ¹H NMR and ¹³C NMR spectra, and their data are listed as follows.



These spectra proved that $[\text{VLIM}]\text{SO}_3$ was successfully synthesized.

^1H NMR (500 MHz, CD_3OD) δ : 2.39 (m, 2H), 2.88 (t, $J = 10$ Hz, 2H), 4.51 (t, $J = 10$ Hz, 2H), 5.46 (dd, $J = 10, 5$ Hz, 1H), 5.96 (dd, $J = 20, 10$ Hz, 1H), 7.30 (dd, $J = 20, 10, 1$ Hz), 7.84 (s, 1H), 8.03 (s, 1H), 9.36 (s, 1H) (Fig. S1†).

^{13}C NMR (500 MHz, CD_3OD) δ : 27.03, 109.94, 120.84, 124.60, 129.99, 136.94 (Fig. S2†).

3.2. Characterization of the solid acid catalysts

3.2.1 XRD. The crystal structure of the samples was characterized by XRD (Fig. 2a). In the XRD pattern, the peaks at 2 theta were obtained at 30.1° , 35.4° , 43.0° , 53.4° , 56.9° , 62.5° , and 73.9° , which were respectively assigned to the crystal planes [220], [311], [400], [422], [511], [440], and [533] of Fe_3O_4 (JCPDS: 75-1609).⁹ While, the broad peak at 2 theta = 23.5° was indexed to the amorphous structure of SiO_2 .⁴¹ All these peaks confirmed that Fe_3O_4 and $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs were obtained. It was noteworthy that the peaks in the spectra of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were the same with those of $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs. These results showed that the crystal structure of the samples was not changed, even after being covered with poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs, respectively.

3.2.2 TGA. TGA analysis was conducted to investigate the thermal stability and weight percentage of poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) covered on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs (Fig. 2b). For the prepared solid acid catalysts, there were two weight loss areas that could be observed in each curve. The first area at about $100\text{--}150^\circ\text{C}$ was caused by the removal of absorbed water in the samples,⁴² while the second area above 300°C was associated with the poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) covering on the surface of $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs, which was relatively stable in terms of the thermal properties.⁴³ With the increase in temperature, poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) gradually decomposed and eventually disappeared. Furthermore, the weight percentages of poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) could be calculated through the TGA curves because the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs could not be decomposed under 700°C . They were 40% and 87%, respectively. This meant that there was a large amount of acidic PILs covering the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs.

3.2.3 VSM. To investigate the recyclable properties of the samples, their magnetic properties were measured by VSM in an external magnetic field from $-10\,000$ to $+10\,000$ Oe. In order to calculate the saturation magnetization (M_s), the magnetization (M) was plotted vs. $1/H$. From the extrapolation of the curves, the M_s values of the samples were found to decrease from 71, 57,

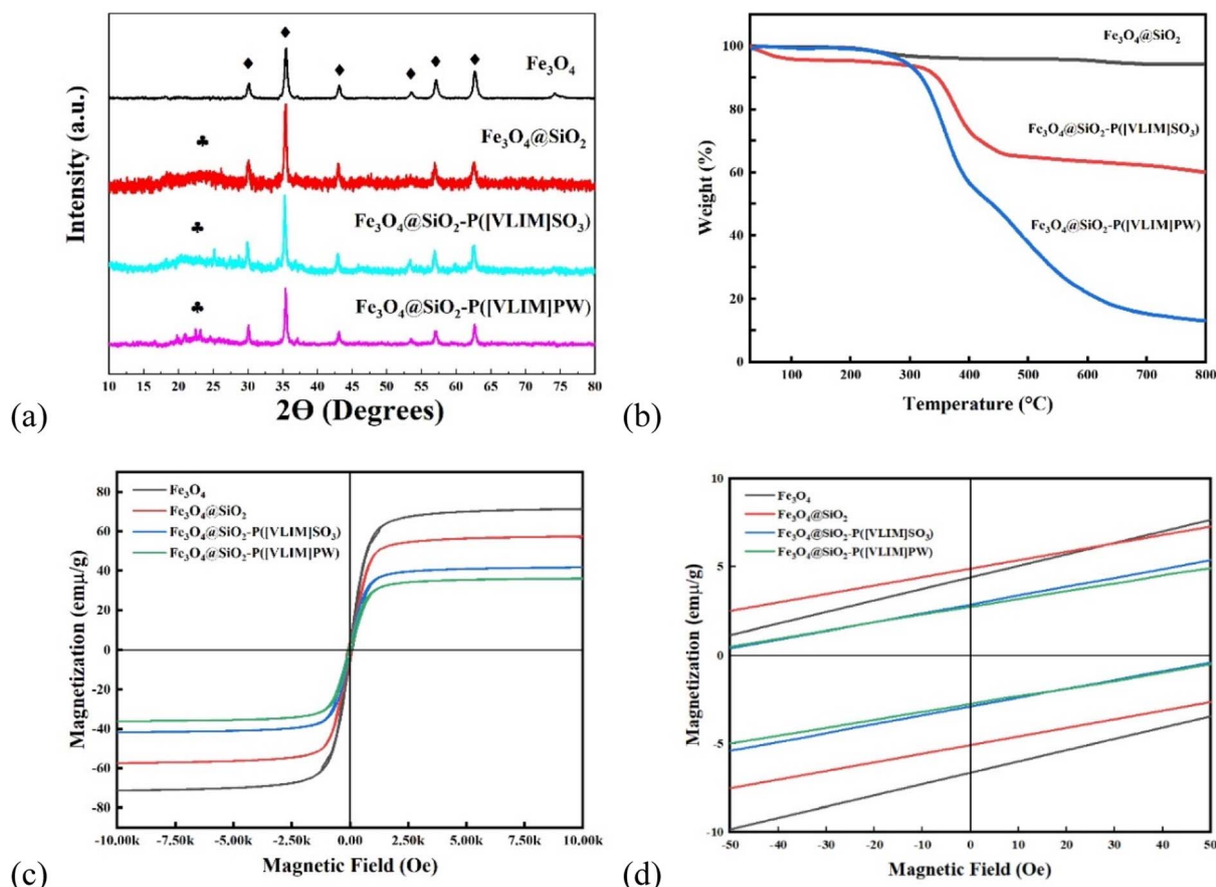


Fig. 2 Characterization of Fe_3O_4 , $\text{Fe}_3\text{O}_4@\text{SiO}_2$, $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$, and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs: (a) XRD, (b) TGA, (c) VSM, and (d) enlarged VSM image.



and 42 emu g^{-1} to 36 emu g^{-1} for Fe_3O_4 , $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs, $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$, and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$, respectively (Fig. 2c). We inferred that this was caused by them being covered with a layer of non-magnetic SiO_2 , poly($[\text{VLIM}]\text{SO}_3$), and poly($[\text{VLIM}]\text{PW}$) on the surface of the Fe_3O_4 NPs, respectively. The M_s value of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs was lower than that of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs, which might be caused by the layer of poly($[\text{VLIM}]\text{PW}$) being thicker than that of poly($[\text{VLIM}]\text{SO}_3$) on the surface of the magnetic-responsive $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs. Furthermore, the TGA results showed the weight content of poly($[\text{VLIM}]\text{PW}$) was higher than

that of poly($[\text{VLIM}]\text{SO}_3$) on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs in the as-prepared solid acid catalysts. Thus, we inferred that the thickness of poly($[\text{VLIM}]\text{PW}$) was thicker than that of poly($[\text{VLIM}]\text{SO}_3$) on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs. Also, the coercivity (H_c) of these samples was not zero (Oe), which confirmed that they had the properties for displaying superparamagnetic performances (Fig. 2d). Based on the above analysis, $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs had strong magnetic responsiveness and they could be easily recycled in an applied magnetic field.

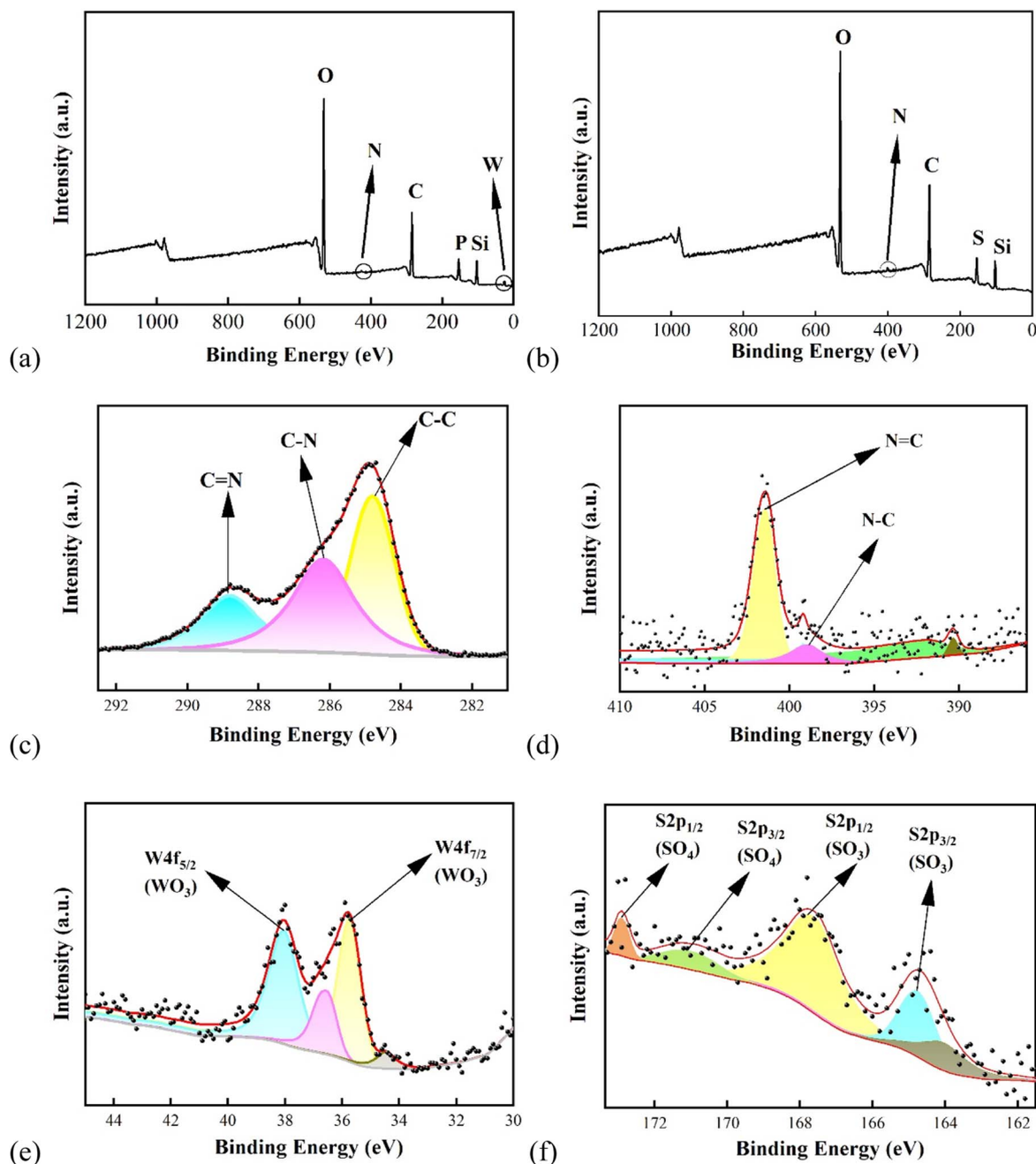


Fig. 3 XPS spectra of the solid acid catalysts. Survey spectra of (a) $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$, (b) $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$; and high-resolution spectra of (c) C 1s, (d) N 1s, (e) W 4f, and (f) S 2p.



3.2.4 XPS. To further confirm that poly([VLIM]SO₃) and poly([VLIM]PW) were covered on the surface of the Fe₃O₄@SiO₂ NPs, the surface composition and chemical state of the elements in Fe₃O₄@SiO₂-P([VLIM]SO₃) and Fe₃O₄@SiO₂-P([VLIM]PW) NPs were characterized by XPS. The survey spectra of Fe₃O₄@SiO₂-P([VLIM]PW) NPs suggested the presence of C, N, P, W, and O elements (Fig. 3a), while C, N, S, and O elements were observed in the spectrum of the Fe₃O₄@SiO₂-P([VLIM]SO₃) NPs (Fig. 3b). The fitting-related data are presented in the ESI (Tables S1 and S2†). Furthermore, the high-resolution spectra of C 1s, N 1s, W 4f, and S 2p were also analyzed. The resulting peaks at 284.8, 286.2, and 288.8 eV in the C 1s spectrum were ascribed to C–C, C–N, and C=N bonds (Fig. 3c). Meanwhile, the peaks at 400.2 and 398.3 eV in the N 1s spectrum were assigned to C=N and C–N bonds⁴⁴ (Fig. 3d).

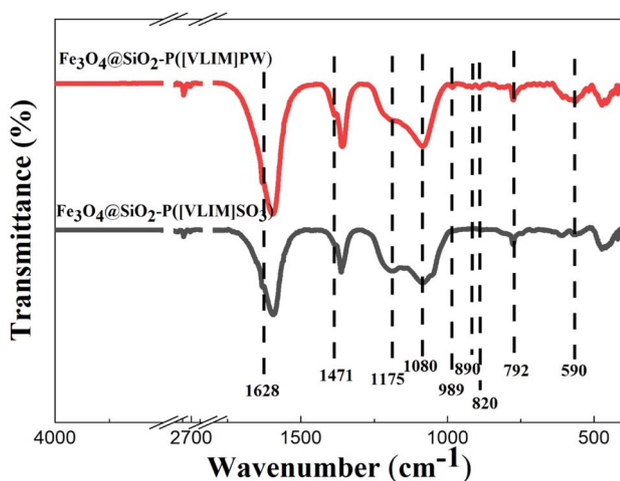


Fig. 4 FTIR spectra of the solid acid catalysts.

These peaks were also both observed in the XPS spectra of Fe₃O₄@SiO₂-P([VLIM]SO₃) and Fe₃O₄@SiO₂-P([VLIM]PW) NPs, suggesting that the poly(vinyl imidazole) group had grafted on to the surface of the Fe₃O₄@SiO₂ NPs.⁴⁵ In addition, the characterized peaks at 36.4 and 38.1 eV in the W 4f spectrum were assigned to W 4f_{7/2} (WO₃) and W 4f_{5/2} (WO₃) of the PW group⁴⁶ (Fig. 3e). The peaks at 164.8 and 167.6 eV in the S 2p spectrum were ascribed to S 2p_{3/2} and S 2p_{1/2} of SO₃, while the peaks at 170.9 and 172.9 eV were attributed to S 2p_{3/2} and S 2p_{1/2} of SO₄ of poly([VLIM]SO₃)⁴⁷ (Fig. 3f). All the spectral results confirmed that poly([VLIM]SO₃) and poly([VLIM]PW) had respectively grafted on to the surface of the Fe₃O₄@SiO₂ NPs, thus confirming Fe₃O₄@SiO₂-P([VLIM]SO₃) and Fe₃O₄@SiO₂-P([VLIM]PW) NPs were achieved.

3.2.5 FTIR. The characteristic groups of the prepared solid acid catalysts were characterized by FTIR (Fig. 4). In the spectra, the peaks at 590 and 1080 cm⁻¹ indicated Fe–O and Si–O–Si bonds, respectively.⁷ Meanwhile, the peaks at 1628, 1471, and 792 cm⁻¹ were associated to the C=C, C=N, and C–H bonds of the imidazole group, respectively.⁴⁸ These above peaks could be observed in both curves. Furthermore, the characteristic peaks at 820, 890, 989, and 1080 cm⁻¹, respectively, indicating O–b1–W, O–b2–W, W=O, and P–O of the Keggin structure, were observed with the Fe₃O₄@SiO₂-P([VLIM]PW) NPs;⁸ while the characterized peak of S=O at 1175 cm⁻¹ was observed with the Fe₃O₄@SiO₂-P([VLIM]SO₃) NPs.^{34,48} All the peaks indicated the as-designed solid acid catalysts were achieved.

3.2.6 SEM. The morphologies of the samples were characterized by SEM. For the Fe₃O₄ NPs, their surface was rough and their mean diameter was about 250 nm (Fig. 5a); while the surface became smooth and the diameter increased to 300 nm after being covered with a SiO₂ layer (Fig. 5b). This meant that the thickness of the SiO₂ layer covered on the surface of Fe₃O₄

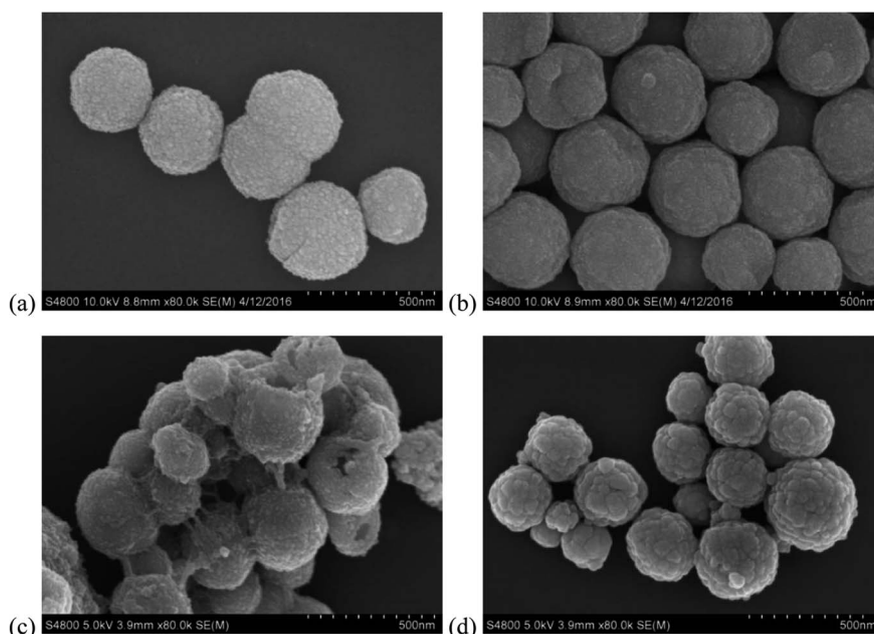


Fig. 5 SEM images of the samples: (a) Fe₃O₄, (b) Fe₃O₄@SiO₂, (c) Fe₃O₄@SiO₂-P([VLIM]SO₃), and (d) Fe₃O₄@SiO₂-P([VLIM]PW) NPs.



NPs was about 25 nm. The diameters of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were both about 350 nm, which was slightly larger than that of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs. Meanwhile, the degree of roughness was increased with poly($[\text{VLIM}]\text{SO}_3$) and poly($[\text{VLIM}]\text{PW}$) covering the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs, respectively. It was noteworthy that the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs were entangled together through the grafted poly($[\text{VLIM}]\text{SO}_3$) on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs (Fig. 5c), and we inferred that this was caused by hydrogen bonds between the SO_3 groups of poly($[\text{VLIM}]\text{SO}_3$) and deionized water in the mixture. However, the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were monodispersed (Fig. 5d), which was probably due to the effect of steric hindrance of the PW groups covered on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs.

3.2.7 Acidity of the solid acid catalysts. The acidity, which was caused by the sulfonate group and phosphotungstic acid group, could be used to determine the catalytic activity of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs when serving as catalysts, respectively. The acidity was thus measured, here by KOH-ethanol solution (5 mg L^{-1}) *via* acid-base titration with phenolphthalein as the indicator.⁴⁹ The acidity was calculated according to the following equation:

$$C_{\text{H}^+} = \frac{C_{\text{KOH}}}{m_{\text{cat}}}$$

where C_{H^+} is the acid concentration of the catalyst (mmol g^{-1}), and C_{KOH} and m_{cat} are the molar concentration of the KOH-ethanol solution and the weight of the catalyst, respectively. The acidity values of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were 2.3 and 3.8 mmol g^{-1} , respectively. It was obvious that the acidity of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs was higher than that of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs, suggesting that the catalytic activity of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs would be better than that of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs.

3.3. Catalytic properties of the solid acid catalysts

The catalytic properties of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were investigated by the catalyzed esterification of palmitic acid and methanol. The esterification reaction was carried out at 70 °C for 6 h while the MR of methanol to palmitic acid was 18 : 1. As shown in Fig. 6, $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs both presented strong catalytic activity in the esterification. The conversion rate was significantly improved with the increase in the dosage of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs in the mixture, reaching up to 75% and 92%, respectively. Meanwhile, the results indicated that the catalytic activity of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ was higher than that of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs in the esterification. We inferred that this could be attributed to the high loading content and monodispersion of poly($[\text{VLIM}]\text{PW}$) covered on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs. These results were consistent with those of the TGA and SEM images. When the dosage of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs was 10 wt% (weight percentage based on palmitic acid), the conversion rate could reach up to 92%.

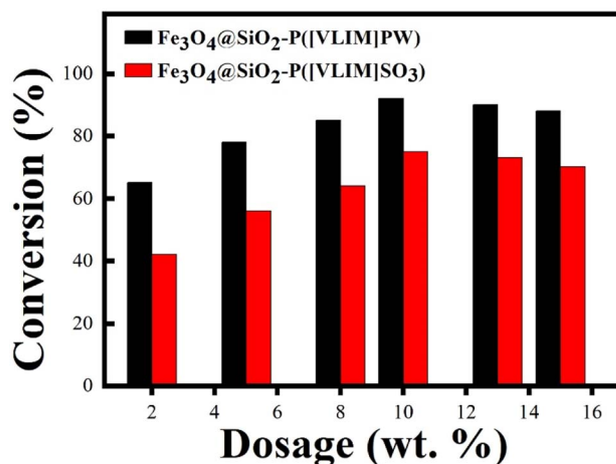


Fig. 6 Conversion rate of palmitic acid in the esterification catalyzed by $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs with different dosages (70 °C, 6 h, MR of methanol to palmitic acid of 18 : 1).

However, the conversion rate gradually decreased when the dosage of solid acid catalysts was more than 10 wt% in the mixture, which was probably caused by aggregation of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs in the mixture, and thus their catalytic activity decreased.

A series of experiments were carried out according to single-factor analyses to determine the optimal reaction conditions for the preparation of biodiesel by the esterification of palmitic acid and methanol. According to the results obtained from Fig. 6, the dosage of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs was selected as 10 wt%. The effects of changing the reaction factors, such as reaction time, reaction temperature, and MR of methanol to palmitic acid, were investigated, and the corresponding results are shown in Fig. 7. The results indicated that the optimal reaction conditions were as following: reaction time, temperature, and MR of methanol to palmitic acid of 6 h, 70 °C, and 12 : 1, respectively. Under the above reaction conditions, the conversion rate could reach up to 94%.

3.4. Catalytic stability

From the viewpoints of economical consideration and environmental concerns, the stability properties of catalysts are greatly important. Thus, experiments to investigate the catalyst reuse potential were conducted in the esterification of palmitic acid and methanol with $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs as the catalyst under the optimal reaction conditions mentioned above. The results are shown in Fig. 8, where it can be seen that even after being reused for 5 times, the conversion rate was still as high as 84%. These results confirmed that the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs possessed excellent reusability potential, and thus they could be suitable to serve as efficient solid acid catalysts in the esterification.

The acidity of the reused $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs was also investigated, and the value was 3.2 mmol g^{-1} after recycling the catalyst 5 times. The $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs showed a slight decrease in acidity, whereas their SEM image and VSM



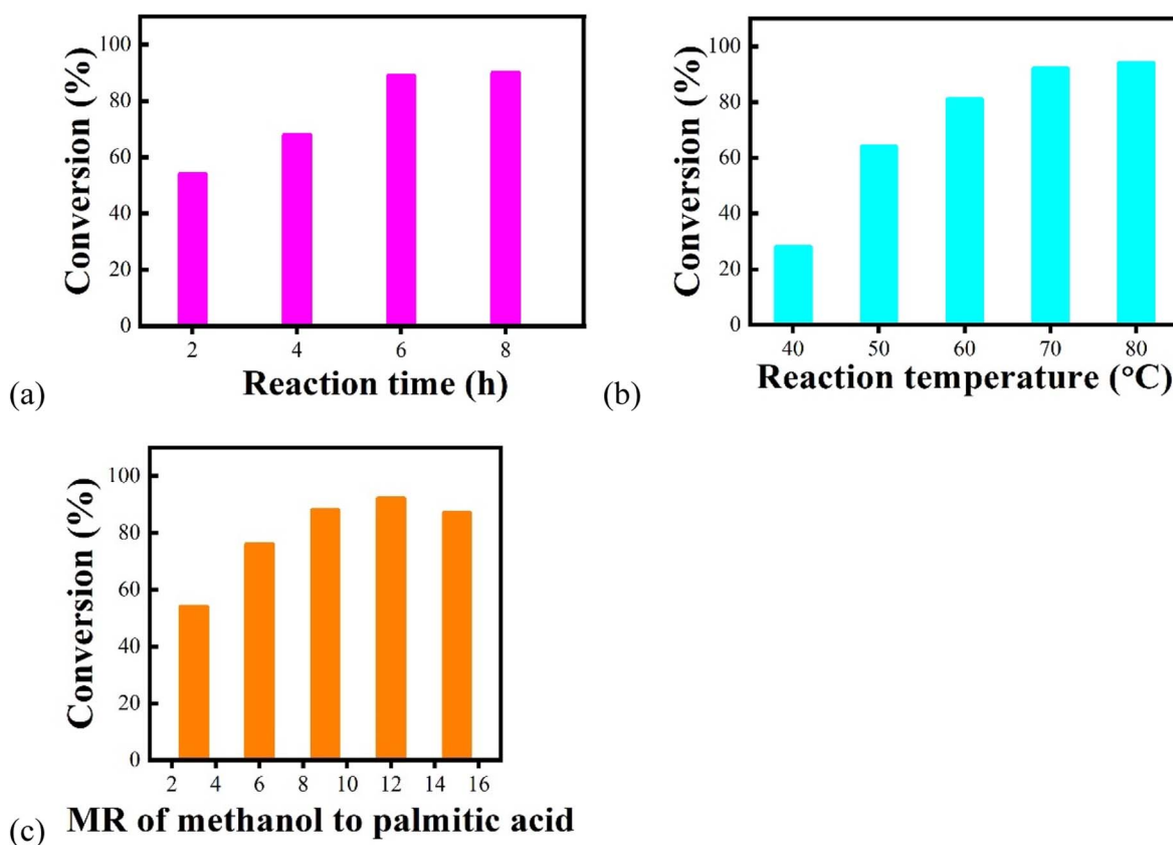


Fig. 7 Effects of changing different reaction factors on the conversion rate with 10 wt% $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs as the catalyst: (a) reaction time (70 °C, MR of methanol to palmitic acid of 18 : 1), (b) reaction temperature (6 h, MR of methanol to palmitic acid of 18 : 1), and (c) MR of methanol to palmitic acid (70 °C, 6 h).

value showed no change. However, in the FTIR spectra of the used $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs, there was a decrease in the intensity of the characteristic peak for phosphotungstate (Fig. S3†), indicating a slight deactivation of the reused catalyst.

3.5. Mechanism of the solid acid-catalyzed esterification

Based on the results from recent reports about catalyzed esterification kinetics,^{34,50,51} we think the catalyzed esterification reaction in this work was accomplished *via* the chemisorption of palmitic acid on the surface of the magnetic-responsive solid acid catalysts. The esterification was a nucleophilic reaction. The electron cloud density of poly([VLIM]PW) and poly([VLIM]SO₃) was high, thus $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]SO_3)$ and $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs could provide protons to palmitic acid. This resulted in palmitic acid being easily absorbed on the surface of $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]SO_3)$ and $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs. The $C=OH^+$ was generated from the reaction of the active sites on the magnetic-responsive solid acid with the $C=O$ of palmitic acid. Then, the $-COOH$ group of palmitic acid was protonated and a carbocation was formed. After that, nucleophilic attack from methanol to the above carbocation was performed, and then the tetrahedral intermediate was achieved. Finally, the tetrahedral intermediate was respectively moved away from the surface of $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]SO_3)$ and $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs *via* proton migration, and thus palmitic acid methyl ester was obtained. Meanwhile, the catalytic activities of $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]SO_3)$ and $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs were regenerated after the esterification, respectively. Furthermore, we think the enhanced catalytic performance of the prepared solid

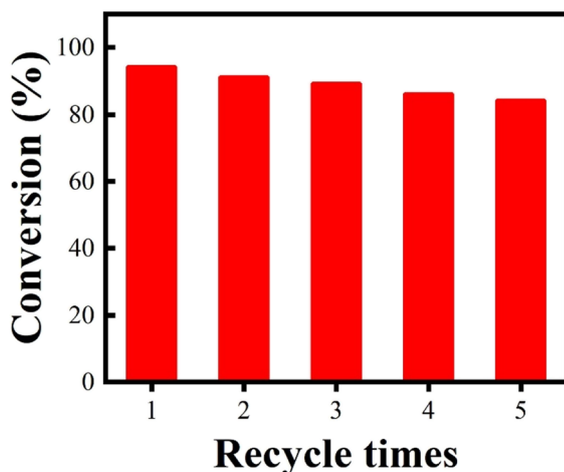


Fig. 8 Reusability of $\text{Fe}_3\text{O}_4@SiO_2-P([VLIM]PW)$ NPs as catalysts for the esterification of palmitic acid and methanol. Reaction conditions: RM of methanol to palmitic acid of 15 : 1, 10 wt% catalyst, 70 °C, 6 h.



acid catalysts in this work was caused by the PILs being uniformly dispersed on the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs with a large specific surface area. Thus, a greater number of activated sites on the magnetic-responsive solid acid catalysts could be applied in the esterification and the catalytic performance was improved.

4. Conclusions

In summary, two kinds of magnetic-responsive solid acid catalysts were designed and prepared by an *in situ* polymerization on the surface of magnetic nanoparticles. They were characterized by XRD, VSM, NMR, FTIR, TGA, XPS, SEM, and GC. The results showed that the solid acid catalysts with magnetic responsiveness were about 350 nm in diameter, and their surface was rough due to the coverage of poly([VLIM]PW) and poly([VLIM]SO₃) on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs, respectively. The weight contents of the poly([VLIM]PW) and poly([VLIM]SO₃) coverings were about 87% and 40%, respectively. The catalytic activity of the solid acid catalysts was evaluated by the catalyzed esterification of palmitic acid and methanol. By comparison, the catalytic properties of $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs were better than that of the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{SO}_3)$ NPs, which was due to the greater number of activated sites and high loading content of poly([VLIM]PW) covered on the surface of the $\text{Fe}_3\text{O}_4@\text{SiO}_2$ NPs as presented in the esterification. Under the optimal reaction conditions (10 wt% catalyst, 6 h, 70 °C, and MR of methanol to palmitic acid of 12 : 1), the conversion rate of palmitic acid in the esterification could reach up to 94% when the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs served as catalysts. Furthermore, the $\text{Fe}_3\text{O}_4@\text{SiO}_2\text{-P}([\text{VLIM}]\text{PW})$ NPs could still maintain a high catalytic activity even after being reused 5 times without significant deactivation. More importantly, this work proposes a simple method to cover acidic PILs on the surface of magnetic particles. This method has great potential for the preparation of solid acid catalysts with magnetic responsiveness, which could be used in many domains, such as catalysis and waste treatment.

Author contributions

Dan Xue: conceptualization, writing – original draft & review & editing, investigation, visualization, methodology, validation, formal analysis, data curation, supervision, resources. Yun Jiang: investigation. Fang Xia Zheng: investigation.

Conflicts of interest

The authors state that there are no conflicts to declare.

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References

- D. D. Bala, M. Misra and D. Chidambaram, *J. Cleaner Prod.*, 2017, **142**, 4169–4177.
- J. Gaidukevič, J. Barkauskas, A. Malaika, V. Jasulaitienė and M. Kozłowski, *Appl. Surf. Sci.*, 2021, **554**, 149588–149700.
- K.-S. Lin, N. V. Mdlovu, H.-Y. Chan, K. C.-W. Wu, J. C.-S. Wu and Y.-T. Huang, *Catal. Today*, 2022, **397–399**, 145–154.
- M. O. Abdelmigeed, E. G. Al-Sakkari, M. S. Hefney, F. M. Ismail, A. Abdelghany, T. S. Ahmed and I. M. Ismail, *Renewable Energy*, 2021, **165**, 405–419.
- A. Naeem, I. Wali Khan, M. Farooq, T. Mahmood, I. Ud Din, Z. Ali Ghazi and T. Saeed, *Bioresour. Technol.*, 2021, **328**, 124831–124838.
- M. Racar, I. Šoljić Jerbić, Z. Glasovac and A. Jukić, *Renewable Energy*, 2023, **202**, 1046–1053.
- J. Li and X. Liang, *Energy Convers. Manage.*, 2017, **141**, 126–132.
- R. Sree and S. Kuriakose, *J. Energy Chem.*, 2015, **24**, 87–92.
- H. Li, J. Wang, X. Ma, Y. Wang, G. Li, M. Guo, P. Cui, W. Lu, S. Zhou and M. Yu, *Renewable Energy*, 2021, **179**, 1191–1203.
- P. X. Pham, K. T. Nguyen, T. V. Pham and V. H. Nguyen, *ACS Omega*, 2020, **5**, 20842–20853.
- W. Xie, Y. Xiong and H. Wang, *Renewable Energy*, 2021, **174**, 758–768.
- G. Knothe and L. F. Razon, *Prog. Energy Combust. Sci.*, 2017, **58**, 36–59.
- A. K. Sodhi, S. Tripathi and K. Kundu, *Clean Technol. Environ. Policy*, 2017, **19**, 1799–1807.
- M. Hamza, M. Ayoub, R. B. Shamsuddin, A. Mukhtar, S. Saqib, I. Zahid, M. Ameen, S. Ullah, A. G. Al-Sehemi and M. Ibrahim, *Environ. Technol. Innovation*, 2021, **21**, 101200–101220.
- X. Zhang, N. Li, Z. Wei, B. Dai and S. Han, *Renewable Energy*, 2022, **196**, 737–748.
- Y. Wang, G. Yang, J. He, G. Sun, Z. Sun and Y. Sun, *Process Saf. Environ. Prot.*, 2020, **141**, 333–343.
- N. A. Negm, M. A. Betiha, M. S. Alhumaimess, H. M. A. Hassan and A. M. Rabie, *J. Cleaner Prod.*, 2019, **238**, 117854–117869.
- I. Tankov, Z. Mustafa, R. Nikolova, A. Veli and R. Yankova, *Fuel*, 2022, **307**, 121876–121887.
- F. Huang, Y. Su, Y. Tao, W. Sun and W. Wang, *Fuel*, 2018, **226**, 417–422.
- X. Lin, X. Ling, J. Chen, M. Li, T. Xu and T. Qiu, *Green Chem.*, 2019, **21**, 3182–3189.
- X.-X. Han, Y.-F. He, C.-T. Hung, L.-L. Liu, S.-J. Huang and S.-B. Liu, *Chem. Eng. Sci.*, 2013, **104**, 64–72.
- Y. Bian, Q. Shan, C. Guo, C. Liu and J. Zhang, *Catal. Lett.*, 2021, **151**, 3523–3531.
- Y. Leng, J. Wang, D. R. Zhu, X. Q. Ren, H. Q. Ge and L. Shen, *Angew. Chem., Int. Ed.*, 2009, **48**, 168–171.
- Z. Yu, X. Chen, Y. Zhang, H. Tu, P. Pan, S. Li, Y. Han, M. Piao, J. Hu, F. Shi and X. Yang, *Chem. Eng. J.*, 2022, **430**, 133059–133069.
- J. Yu, Y. Wang, L. Sun, Z. Xu, Y. Du, H. Sun, W. Li, S. Luo, C. Ma and S. Liu, *ACS Omega*, 2021, **6**, 7896–7909.



- 26 M. Fan, Z. Si, W. Sun and P. Zhang, *Fuel*, 2019, **252**, 254–261.
- 27 T. Parangi and M. K. Mishra, *Comments Inorg. Chem.*, 2019, **39**, 90–126.
- 28 F. F. Roman, A. E. Ribeiro, A. Queiroz, G. G. Lenzi, E. S. Chaves and P. Brito, *Fuel*, 2019, **239**, 1231–1239.
- 29 M. A. Hanif, S. Nisar and U. Rashid, *Catal. Rev.*, 2017, **59**, 165–188.
- 30 A. S. Amarasekara, *Chem. Rev.*, 2016, **116**, 6133–6183.
- 31 J. Zhang, X. Liu, M. N. Hedhili, Y. Zhu and Y. Han, *ChemCatChem*, 2011, **3**, 1294–1298.
- 32 J. Zhang, X. Liu, M. Sun, X. Ma and Y. Han, *ACS Catal.*, 2012, **2**, 1698–1702.
- 33 J. Zhang, M. Sun, X. Liu and Y. Han, *Catal. Today*, 2014, **233**, 77–82.
- 34 P. Fan, S. Xing, J. Wang, J. Fu, L. Yang, G. Yang, C. Miao and P. Lv, *Fuel*, 2017, **188**, 483–488.
- 35 Y.-T. Wang, X.-X. Yang, J. Xu, H.-L. Wang, Z.-B. Wang, L. Zhang, S.-L. Wang and J.-L. Liang, *Renewable Energy*, 2019, **139**, 688–695.
- 36 P. Ke, D. Zeng, J. Wu, J. Cui, X. Li and G. Wang, *ACS Omega*, 2019, **4**, 22119–22125.
- 37 H. Jing, X. Wang, Y. Liu and A. Wang, *Chin. J. Catal.*, 2015, **36**, 244–251.
- 38 J. Gardy, E. Nourafkan, A. Osatiashtiani, A. F. Lee, K. Wilson, A. Hassanpour and X. Lai, *Appl. Catal., B*, 2019, **259**, 118093–118094.
- 39 D. Xue, Q. B. Meng and X. M. Song, *ACS Appl. Mater. Interfaces*, 2019, **11**, 10967–10974.
- 40 D. Xue, X. Song and F. Liang, *RSC Adv.*, 2017, **7**, 25450–25454.
- 41 C. Sun, D. Xu, C. Hou, H. Zhang, Y. Li, Q. Zhang, H. Wang and M. Zhu, *J. Mech. Behav. Biomed. Mater.*, 2021, **121**, 104593–104601.
- 42 Y. Yin, X. Huang, W. Wang and X. Liu, *Solid State Sci.*, 2023, **136**, 107109–107118.
- 43 S. Y. Lai, K. H. Ng, C. K. Cheng, H. Nur, M. Nurhadi and M. Arumugam, *Chemosphere*, 2021, **263**, 128244–128266.
- 44 D. Yuan, J. Zhao, Q. Zhang, P. Lu, Y. Wang, Y. He, Z. Liu, Y. Liu, X. Zhao and C. Meng, *J. Mater. Chem. A*, 2022, **10**, 12656–12668.
- 45 C. Liu, S. Qiu, P. Du, H. Zhao and L. Wang, *Nanoscale*, 2018, **10**, 8115–8124.
- 46 S. Zou, H. Wang, S. Li, B. Lu, J. Zhao and Q. Cai, *Appl. Surf. Sci.*, 2022, **574**, 11.
- 47 X. Li, Z. Zhou, Y. Zhao, D. Ramella and Y. Luan, *Appl. Organomet. Chem.*, 2020, **34**, 13.
- 48 Y. Wang, D. Zhao, L. Wang, X. Wang, L. Li, Z. Xing, N. Ji, S. Liu and H. Ding, *Fuel*, 2018, **216**, 364–370.
- 49 Y. Wang, D. Wang, M. Tan, B. Jiang, J. Zheng, N. Tsubaki and M. Wu, *ACS Appl. Mater. Interfaces*, 2015, **7**, 26767–26775.
- 50 D. D. Bala, V. Vasquez and D. Chidambaram, *Ind. Eng. Chem. Res.*, 2018, **57**, 14948–14956.
- 51 Y. Wang, D. Zhao, G. Chen, S. Liu, N. Ji, H. Ding and J. Fu, *Renewable Energy*, 2019, **133**, 317–324.

