RSC Advances



PAPER

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Cite this: RSC Adv., 2021, 11, 20836

Received 8th February 2021 Accepted 28th May 2021

DOI: 10.1039/d1ra01062j

rsc.li/rsc-advances

1. Introduction

4-Chlorophenol and 4-chlorophenol derivatives have many applications in the production and application of the chemical industry, especially the alkylation species of 4-chlorophenol. The derivatives of 4-chlorophenol are a kind of important organic chemical intermediate, which are widely used in pharmaceuticals, pesticides and dye intermediates, such as the brain stimulant chlorzoxazone,¹ and the cure of coronary heart disease drug clofibrate; pesticides fenamidone and fenpropion; dye intermediates 5-chloro-2-hydroxyaniline and 1,4-dihydroxyanthraquinone,²⁻⁴ etc. With the development of 4-chlorophenol into a variety of pharmaceutical and new dye intermediates in recent years, chemical industry and commercial demands for 4-

Application of two morphologies of Mn₂O₃ for efficient catalytic *ortho*-methylation of 4-chlorophenol[†]

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Vapor phase ortho-methylation of 4-chlorophenol with methanol was studied over Mn₂O₃ catalyst with two kinds of morphologies. Here, Mn_2O_3 was prepared by a precipitation and hydrothermal method, and showed the morphology of nanoparticles and nanowires, respectively. XRD characterization and BET results showed that, with the increase of calcination temperature, Mn_2O_3 had a higher crystallinity and a smaller specific surface area. N2 adsorption/desorption and TPD measurements indicated that Mn2O3 nanowires possessed larger external surface areas and more abundant acid and base sites. Simultaneously, in the fixed bed reactor, methanol was used as the methylation reagent for the orthomethylation reaction of 4-chlorophenol. XRD, XPS, TG-MS and other characterizations made it clear that methanol reduced 4-chlorophenol and its methide, which were the main side-reactions. And Mn^{3+} was reduced to Mn^{2+} under the reaction conditions. Changing the carrier gas N₂ to a H₂/Ar mixture further verified that the hydrogen generated by the decomposition of methanol was not the reason for dechlorination of 4-chlorophenol compounds. Here we summarized the progress of 4-chlorophenol methylation based on the methylation of phenol. Also, we proposed a mechanism of the 4-chlorophenol dechlorination effect which was similar to the Meerwein-Ponndorf-Verley-type (MPV) reaction. The crystal phase and carbon deposition were investigated in different reaction periods by XRD and TG-DTA. The reaction conditions for the two kinds of morphologies of the Mn_2O_3 catalyst such as calcination temperature, reaction temperature, phenol-methanol ratio and reaction space velocity were optimized.

> chlorophenol and its derivatives are increasing with passing year. Therefore, it is crucial to develop an appropriate catalyst with high efficiency of conversion, high selectivity, long operation lifetime and environmentally friendly, energy-saving reaction modes.

> The traditional way to synthesize alkylated 4-chlorophenol mainly depends on the chlorination of alkylated phenol. A large number of researchers have reported on the chlorination of phenol and its alkylated derivatives. For example, Gusevskaya's group used CuCl₂ as the catalyst and chlorinating reagent (Cu²⁺ is the catalyst and Cl^- are used as halogenating agents), and O_2 is the oxidant. The result of chlorination and monohalogenation of aromatic phenols was satisfactory.5,6 Kee's group used SO₂Cl₂ as the chlorinating agent, and the ratio of *p*chlorine to o-chlorine in the chlorination products exceeded 50.7 Bovonsombat's used *p*-toluenesulfonic acid as the catalyst and N-chlorosuccinimide as the source of chlorine, which can react with para-chlorinated phenol analogs at room temperature.⁸ Sheldon's group applied montmorillonite clay and part of the cation-exchanged L-type molecular sieve as catalyst, SO₂Cl₂ as the chlorinating agent, and 2,2,4-trimethylpentane as the solvent. The conversion of phenol reached up to 96% at 25 °C. The selectivity of 4-chlorophenol achieved 89%, and the ratio of

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Paper

para-chlorinated substitution to ortho-chlorinated substitution reached up to 8.0.2 Xiong et al. adopted microwave-assisted heating with CuCl₂ as the catalyst and hydrochloric acid as the chlorine source to accomplish the complete chlorination of phenol within 60 min.9 Although the conversion rate and selectivity of the above methods are very considerable, it is inevitable to use strong acid and highly toxic SO₂Cl₂ as chlorination reagent. These problems will arise in the practical application of industrialization, whether in terms of the maintenance of industrial equipment or the safety of experimenters and environmental friendly. Accordingly, we converted train of thought and shifted to the research about the methylation of 4chlorophenol. Moreover, the gas-phase methylation of phenol with methanol is one of the first cases of a "green chemistry" applied in industry, in which an electrophilic substitution on an aromatic ring was carried out by applying an alcohol.

As far as we know, the research on the ortho-methylation of 4chlorophenol has not been reported, but there is a lot of works on the ortho-methylation of phenol. Among them, metal oxide catalysts play a vital role, such as α-Fe₂O₃,¹⁰ TiO₂,¹¹ MgO,¹² Mg-Al-hydrotalcite,¹³ Fe-doped ZrO₂ composites,¹⁴ Si-Mn composite oxides,¹⁵ CoFe₂O₄ spinel,¹⁶ V₂O₅-ZrO₂,¹⁷ Mg/Me/O spinel (Me = Al, Cr, Fe),¹⁸ $Cu_xMn_{3-x}O_4$ spinel,¹⁹ Mg-Fe-O composite metal oxide.²⁰ The reason why metal oxides show such high activity and selectivity for ortho-methylation of phenol was mainly based on the vertical adsorption of phenolate species on the surface of the catalyst and the formation of intermediate methyl species. Therefore, in chemical kinetics, methyl species are more likely to attack the ortho position of the phenol ring, forming ortho-substituted phenol (o-cresol) or diortho-substituted phenol (2,6-xylenol).

Herein, two kinds of morphologies Mn_2O_3 catalysts were synthetized by two different approaches, and compared them through a series of characterization analysis, and also evaluated the activity of two kinds of Mn_2O_3 samples in *ortho*-methylation of 4-chlorophenol. The results of productivity were satisfied. At the same time, we also discussed the reaction mechanism of the *ortho*-methylation of 4-chlorophenol and the main side-reaction of this reaction: the dechlorination mechanism of 4-chlorophenol and methylated 4-chlorophenol, and provided a good reference and pavement to develop a series of the highperformance catalysts for *ortho*-methylation of 4-chlorophenol.

2. Experimental

2.1 Chemicals and materials

Manganous acetate (Mn(Ac)₂·4H₂O), 25% ammonia solution (NH₃·H₂O), 4-chlorophenol and hexamethylenetetramine (C₆H₁₂N₄) were purchased from Sinopharm Chemical Reagent Co., Ltd. Methanol, ethanol and acetone were purchased from Beijing Chemical Works. All chemicals and reagents were of analytical grade and were not purified before used. Water used in the experiments was ultrapure deionized water (resistance 18 M Ω cm). Commercial silicon carbide was washed by water and ethanol for 72 h, respectively. After desiccation, silicon carbide was calcined under 600 °C for 6 h.

2.2 Catalyst preparation

The nanoparticle morphology Mn_2O_3 catalyst (Mn_2O_{3NP}) prepared by the precipitation method was according to previously reported methodology with some minor improvements.¹⁵ In a typical procedure, 14.7 g $Mn(Ac)_2 \cdot 4H_2O$ was dissolved in 100 mL deionized water. Then the above solution was dropwise added into 200 mL 6.5 wt% $NH_3 \cdot H_2O$ diluent at a rate of three drops per second with constant stirring. After the $Mn(Ac)_2$ solution dripped out, aging of the mixture was carried out for 3 h at room temperature. Later, the above liquid was transferred to an evaporating dish, and heated in a water bath at 90 °C for 12 h. Then it was dried in oven at 110 °C for 15 h, and the dried precursor was calcined for 5 h with a 5 °C min⁻¹ ramp rate. The available powders were ready for the *o*-methylation activity measurements after being crushed and sieved (40–60 mesh).

The synthesis of nanowire Mn_2O_3 catalyst (Mn_2O_{3NW}) is as follows. Briefly, 2.45 g $Mn(Ac)_2 \cdot 4H_2O$ and 0.70 g $C_6H_{12}N_4$ were dissolved in 40 mL deionized water. Finally, the mixture was transferred into a Teflon-lined stainless steel autoclave and statically crystallized at 180 °C for 15 h. The solid–liquid mixture was separated by vacuum filtration and washed with deionized water three times. The filter cake was dried at 80 °C for 12 h, and the dried precursor was calcined for 6 h.

2.3 Catalyst characterizations

The crystalline structure was characterized by X-ray diffraction analysis (XRD) with an Empyrean diffractometer from Rigaku equipped with a 9 kW rotating anode Cu source at 40 kV and 40 mA, from 10° to 80° with a scan rate of 10° min⁻¹. Nitrogen sorption experiment for the measurement of surface area and pore volume was carried out at 77 K on a Micromeritics ASAP 2010N analyzer. The surface chemical composition analyzed by X-ray photoelectron spectrum (XPS) was conducted on a Thermo ESCA LAB 250 instrument equipped with Al Ka radiation (1486.6 eV). The 284.55 eV value of polluted carbon C 1s was used as the nuclear effect of the internal standard calibration sample. Morphologies were tested on a field-emission scanning electron microscope (SEM) instrument (HITACHI SU-8020). Simultaneous thermal analysis (STA) mass spectrometry (MS) analysis were performed on Germany NETZSCH STA499F3 QMS403D\Bruker V70 type simultaneous thermal analysis-mass spectrometry-infrared combined instrument at a heating rate of 10 °C min⁻¹ from 40 °C to 900 °C in air.

Temperature programmed desorption (TPD) and temperature programmed reduction (TPR) were recorded on a Chem-BET Pulsar TPR/TPD instrument. The profile is obtained by deducting the blank signal of the sample. The acid and base sites were analyzed by using NH₃ and CO₂ as atmosphere respectively. Typically, 50 mg of catalysts were pretreated at 300 °C in nitrogen flow for 30 min, and then cooled to 50 °C under helium (He) gas. NH₃ or CO₂ gas. After samples were sweeped under He gas for 60 min, the temperature was heated to 600 °C (10 °C min⁻¹). The desorbed gas was determined by a GOW-MAC thermal conductivity detector (TCD). As for H₂-TPR, 15 mg samples were treated in an nitrogen flow at 350 °C



Fig. 1 Vapor phase fixed bed catalytic reactor setup for *ortho*-methylation of 4-chlorophenol.

for 30 min. After the sample was cooled down to 50 °C, 5 vol% H₂/Ar mixture was introduced to quartz tube for the catalysts reduction. The temperature was heated to 800 °C with a ramp rate of 10 °C min⁻¹.

2.4 Catalytic testing

The continuous gas phase reaction of 4-chlorophenol with methanol was carried out in a fixed-bed reactor. It was provided

with temperature sensor at the top and bottom to read and control the reaction temperature. The reactor had a tube length of 400 mm and an inner diameter of 8 mm, which was vertically fixed on the heating bed. In a typical run, the lower part of the reaction tube was filled with rolled wire meshes and quartz wool. The mixture of 500 mg catalyst and silicon carbide were packed onto the quartz wool, and the upper portion of the reactor was filled with silicon carbide, which served as preheating devices. A mass flow controller was provided for continuous and constant flow of carrier gas (N2) to influx the reactor. A HPLC pump was also equipped with reaction system for constant volume flow of reaction mixture. The effluents was collected by a glass bottle containing 1 mL of acetone and cooled with ice water bath. The exhaust was accessed to atmosphere for the pressure balance in the reactor. The reaction system is shown in the Fig. 1.

The composition and content distribution of the products were analyzed by a gas chromatography (Shimadzu GC-14B) equipped with a flame ionization detector (FID) and a capillary column (PV-101; 60 m \times 0.25 mm \times 0.25 µm). The product identification was analyzed by GC-MS (Shimadzu QP-2010 Plus). The major by products were phenol and *o*-cresol while others



Fig. 2 XRD patterns of (A) fresh Mn_2O_{3NP} catalysts calcined at different temperatures; (B) 3 h and 8 h spent Mn_2O_{3NP} catalysts; (C) fresh Mn_2O_{3NW} catalysts calcined at different temperatures; (D) 3 h and 8 h spent Mn_2O_{3NW} catalysts.

3. Results and discussion

3.1 Characterization of the catalyst

Fig. 2A depicts the XRD patterns of Mn_2O_{3NP} at different calcination temperatures. It can be seen that when the calcination temperature was 400 °C, the main diffraction peaks can be assigned as the phase of Mn_3O_4 . The diffraction peaks with 2θ values around 18.0° , 28.9° , 32.3° , 36.1° , 38.0° , 44.4° , 50.7° , 58.5° , 59.6° , and 64.7° were attributed to the characteristic diffraction peaks of Mn_3O_4 . Meanwhile, it can also be observed that the characteristic diffraction peaks of Mn_2O_3 appeared around 2θ values of 32.9° and 55.1° , indicating that a certain amount of Mn_2O_3 would be accompanied by calcination at this temperature. When the calcination temperature went up to 500 °C, the diffraction peak of Mn_3O_4 disappeared and was replaced by Mn_2O_3 , implied that the crystalline phase was transformed from Mn_3O_4 to Mn_2O_3 . With the calcination temperature

increased ulteriorly, the intensity of the diffraction peak gradually heightened. Meanwhile, the widths of the diffraction peaks became narrower, indicating that a clear increase in the crystallite size.¹⁵ Fig. 2C shows the XRD diffraction peaks of Mn_2O_{3NW} at different calcination temperatures. It can be seen that with the calcination temperature increased, the diffraction peak intensity of Mn_2O_{3NW} gradually increased, which was consistent with the tendency of Mn_2O_{3NP} . Under the calcination condition of 400 °C, the XRD patterns of Mn_2O_{3NW} did not show the characteristic diffraction peak of Mn_3O_4 , which proved that there was no Mn_3O_4 crystal phase in Mn_2O_{3NW} , there were all Mn_2O_3 phase.

XRD characterization on the crystal phase composition of the two spent Mn_2O_3 catalysts was also performed. As shown in Fig. 2B and D, both Mn_2O_{3NP} and Mn_2O_{3NW} showed the same diffraction peak pattern after an 8 h reaction. The 2θ value at 15.1° was attributed to the characteristic diffraction peak of $MnCl_2$, it was due to the dehalogenation effect of 4-chlor-ophenol. In addition, the 2θ values of 34.9° , 40.5° and 58.6° were the characteristic diffraction peaks of MnO, and the 2θ values of



Fig. 3 SEM images of (A) fresh Mn_2O_{3NP} catalyst; (B) fresh Mn_2O_{3NW} catalyst; (C) 3 h spent Mn_2O_{3NP} catalyst; (D) 3 h spent Mn_2O_{3NW} catalyst; (E) 8 h spent Mn_2O_{3NP} catalyst; (F) 8 h spent Mn_2O_{3NW} catalyst.

70.2° and 74.0° were the characteristic diffraction peaks of Mn_3O_4 . This was indicated that with the reaction proceeding, Mn^{3+} was reduced to Mn^{2+} . One part of the reason was that during the methylation reduced Mn^{3+} to Mn^{2+} , which further formed a more stable $M^{2+}-M^{3+}-O$ phase.²¹ Another part was that formaldehyde was an active specie formed by the dehydrogenation of methanol. During the formaldehyde formation process, part of the O^{2-} from the surface of the catalyst was taken away.¹² Significantly, 3 h of reaction time, Mn_2O_{3NP} did not exhibit a $MnCl_2$ diffraction peak but Mn_2O_{3NW} emerged. The main reason was that Mn_2O_{3NW} possessed more active sites in dehalogenation was a cumulative process on the Mn_2O_3 surface.

The SEM images in Fig. 3 displayed the morphology of these fresh and spent samples. Fresh Mn_2O_{3NP} catalyst was sintered from nanoparticles, and its abundant porosities originated from the nanoparticles sintertion and stacking. After 3 h reaction, owing to coke deposition and the transformation of crystalline phase, it can be seen in Fig. 3C that the surface of catalyst became rough and the number of porosities decreased significantly. With reaction to 8 h, in Fig. 3E, the porosity of the catalyst was further lost. The enlarged SEM image in Fig. 3E showed that there were some microcrystals neighboring the porosities. Integrating the results of XRD characterization, this may be the new generation of $MnCl_2$. In Fig. 3B, Mn_2O_{3NW} exhibited a nanowire structure instead of nanorod, different synthesis methods made Mn_2O_{3NW} show different morphologies from Mn_2O_{3NP} . Mn_2O_{3NW} spent samples in 3 h and 8 h were displayed in Fig. 3D and F, respectively. The enlarged SEM image in Fig. 3D revealed that microcrystal had been formed on 3 h spent Mn_2O_{3NW} , that was different from Mn_2O_{3NP} but in agreement with the XRD results.

In order to investigate the surface chemical composition and valence state of fresh and 8 h spent catalyst, XPS survey scan of fresh Mn_2O_{3NP} catalyst indicated the presence of Mn, O in Fig. 4A while after the reaction mainly contained four elements: Mn, O, C and Cl (see Fig. 5A). The above results indicated that the physical and chemical state on the catalyst surface has changed significantly after the *ortho*-methylation reaction. Obviously the C (1s) peak appeared after 8 h reaction, which can be attributed to carbon deposition. The emergence of Cl (2p) peak implied that the dehalogenation of 4-chlorophenol in pace with the reaction process. For fresh catalyst in Fig. 4B, the Mn 3s spectrogram contains two sets of multiplet splitting peak. The *D*-value of the binding energy (ΔE) was 5.15 eV, it was close to the $\Delta E = 5.4$ eV of Mn³⁺ reported in the literature,²² indicating



Fig. 4 XPS spectrum of fresh Mn₂O_{3NP} samples: (A) survey scan XPS spectra; (B) Mn 3s detailed XPS spectra; (C) Mn 2p detailed XPS spectra; (D) O 1s detailed and fitted XPS spectra.



Fig. 5 XPS spectrum of 8 h spent Mn₂O_{3NP} samples: (A) survey scan XPS spectra; (B) Mn 3s detailed XPS spectra; (C) Mn 2p detailed XPS spectra; (D) O 1s detailed and fitted XPS spectra.

that the main valence of Mn in fresh Mn_2O_{3NP} catalyst was +3. As for the 8 h spent catalyst (Fig. 5B), the ΔE was 6.25 eV which closed to the reported 6.1 eV of Mn^{2+} .²² It was proved that after the *ortho*-methylation reaction of 4-chlorophenol, MnO and Mn_3O_4 were the major components of the catalyst. In addition, the Mn spectrogram embodies two peaks of Mn $2p_{3/2}$ and Mn $2p_{1/2}$ (Fig. 4C). Around Mn 2p doublet spectra of Mn $2p_{3/2}$ and Mn $2p_{1/2}$, the characteristic satellite peak of MnO appeared in the 8 h spent catalyst. It was indicated that the Mn in the Mn_2O_{3NP} catalyst was reduced from +3 valence to +2, which was consistent with the binding energy results of Mn 3s and the XRD patterns. Therefore, with the reaction proceeds, the Mn_2O_3 phase was gradually reduced to MnO and Mn_3O_4 .

The O 1s XPS spectra are presented in Fig. 4D and 5D. The XPS spectra show the broad doublet peaks in fresh and 8 h spent catalysts. For Fig. 4D, the first peak at about 529.6–529.8 eV was attributed to oxygen ions (O^{2-}) in the crystal lattice oxygen²³⁻²⁵ while the second peak near 533 eV could be distinguished into three sets of peaks, which were referred to adsorbed oxygen species (~531.4 eV),²⁵ carbonate species and hydroxyl (~532.4 eV), as well as adsorbed water (~534.0 eV).²⁶ The area integration of the above-mentioned fitted peaks showed that in fresh catalyst, the lattice oxygen species (O^{2-}) accounted for 66.8% of the oxygen elements, and the adsorbed

oxygen species (O^{-}/O_{2}^{2-}) accounted for 20.0%, the proportion of hydroxyl and carbonate species (OH⁻/CO₃²⁻) was 11.0%, and the ratio of adsorbed water was 2.2%. The lattice oxygen occupied a large number of all oxygen elements. It was obviously that the intensity of the O 1s high binding energy region was higher than the low part, which indicated that the lattice oxygen species on the catalyst surface gradually transformed into other forms of oxygen species with the reaction proceeding. Specifically, the lattice oxygen decreased from 66.8% to 13.7%, the adsorbed oxygen species, hydroxyl and carbonate species increased to 52.5% and 31.2%, respectively. Almost, the content of adsorbed water was unchanged. This illustrated that the reactants could be destroy the lattice structure on the catalyst surface, causing the lattice oxygen change to other forms of oxygen species. It is worth noting that the increase in the content of carbonate could be due to the formation of carbon dioxide which produced by the methanol decomposed. Also, the hydroxyl may be formed by the dehalogenation process simultaneously. Similarly, Fig. 6 showed the binding energy of Cl 2p orbital electrons on the surface of the catalyst after 8 h reaction. By fitting the peaks of the Cl 2p XPS spectrum, in the low binding energy region, it can be obtained that the binding energy of the Cl 2p_{3/2} around 199.0 eV was the metal chloride, and the signal near 200.6 eV was metal chloride (Cl $2p_{1/2}$). The



Fig. 6 Cl 2p detailed and fitted XPS spectra of 8 h spent catalyst: (A) Mn₂O_{3NP}; (B) Mn₂O_{3NW}.

spin orbit splitting energy of Cl 2p in metal chloride state was 1.6 eV, which was consistent with previous study.²⁷ In addition, it can be obtained by XPS peak differentiating analysis that the main form of Cl element on the surface of a 8 h spent catalyst was $MnCl_x$, which accounted for 90.9% of all chlorine elements. It was illustrated that the source of Cl was from the reduction of 4-chlorophenol or its methylated derivatives (can be regarded as HCl), instead of 4-chlorophenol or its methides adsorbed on the surface of the catalyst. In the same way, Mn_2O_{3NW} showed a similar distribution of chlorine elements (Fig. 6B).

In Fig. S1 and S2,[†] Mn_2O_{3NW} showed the same tendency as Mn_2O_{3NP} in the XPS total spectrum and the 3s binding energy, 2p orbital binding energy and O 1s binding energy of Mn. The reason was completely consistent with the above explanation.

Nitrogen adsorption/desorption measurements were performed to investigate the porosity of the obtained samples. As represented in Fig. 7A, all the isotherms were type IV as defined by IUPAC, characteristic of mesoporous structure. An obvious H3 hysteresis loop in the region of $0.8 < P/P_0 < 1.0$ was observed. The adsorption/desorption isotherms show a sharp increase at P/P_0 from 0.8 to 1.0 due to the capillary condensation of nitrogen within the mesoporous.28 It can be seen that the amount of N₂ adsorbed gradually decreased with increasing the calcination temperature, which was due to the crystal particles sintering and growth accompanied by a significant decrease in surface area from 11.9 to 3.6 $m^2 g^{-1}$ (Table S1†). Assabumrungrat and Laosiripojana found that the specific surface area decreases with increasing calcination temperature.29 The textural parameters are shown in Tables S1 and S2.[†] The low pore volume and pore diameter (4 V/S) of Mn₂O₃ samples were due to the thermal sintering during the calcination procedure, which led to the blockage of pores. Notably, Mn₂O_{3NW} samples possessed larger external surface areas $(5.6-27.9 \text{ m}^2 \text{ g}^{-1})$ and significantly increased pore volume $(0.010-0.109 \text{ cm}^3 \text{ g}^{-1})$ compared with those of Mn₂O_{3NP} samples. That was due to the hexamethylenetetramine and Mn²⁺ build a sort of topological junction abundant supramolecular aggregates which proceed to 1-dimensional structural precursor.³⁰⁻³² Thus, Mn₂O_{3NW}



Fig. 7 (A) N₂ adsorption/desorption isotherms of Mn₂O_{3NP} catalysts; (B) BJH pore distribution curves of Mn₂O_{3NP} catalysts.

possessed more channels and textural information than the $Mn_2O_{3NP}\!.$

It has been reported that both the acid and base sites are important for the ortho-methylation of phenol.¹⁴ Hence, the strength and qualitative analysis of the acid and base sites on the surface of the Mn₂O₃ catalysts were determined using stepwise TPD of NH₃ and CO₂. The desorption temperature revealed the acid and base strengths of the catalysts. The higher desorption temperature, the stronger acid or basic strength of the catalyst. The desorbed NH₃ appearing at \leq 150 °C should be corresponded to the hydrogen-bonded and physically adsorbed NH3.33 The ammonia molecules desorbed at higher temperature were attributed to acid-bonded NH₃, and the acid sites could be defined as weak (≤150 °C), medium (150–300 °C), strong (300– 450 °C), and very strong (\geq 450 °C).³⁴ From the NH₃-TPD profiles of the Mn₂O_{3NP} under different calcination temperatures in Fig. 8A, it can be seen that all the catalysts present a desorption peak at a range of 150-350 °C. With the calcination temperature increased from 500 to 800 °C, the TPD profiles became inapparently. Thus, the amount of medium acid sites showed a decreasing trend. This explained that the particle size of the catalyst became larger and the specific surface area became smaller with the calcination temperature increased. This was consistent with the characterization results of XRD and N₂ adsorption-desorption experiments. When baked at 400 °C, the

desorption peak of NH₃ appeared in the temperature more than 500 °C, indicating that there were very strong acid sites on the surface of the catalyst. This was due to the calcination of the catalyst at 400 °C, the ambient temperature was not enough for the oxygen to completely oxidize the precursors to +3 valence, but a mixed valence state of +2 and +3. In principle, Mn²⁺ bound less O²⁻ than Mn³⁺, and the Mn on the catalyst surface was in an oxygen-depleted state. Therefore, compared with Mn³⁺, Mn²⁺ exposed more positions to bind with NH3 adsorption, and showed a very strong acid desorption peak on NH3-TPD. NH3-TPD curve of Mn₂O_{3NW} showed a completely different trend from that of Mn₂O_{3NP}. As shown in Fig. 8C, under the calcination conditions of 400-500 °C, the number of acid sites of Mn₂O_{3NW} was numerous, and when calcination exceeded 600 °C, the acid sites were significantly reduced. Since Mn₂O_{3NW} catalyst formed a stable Mn₂O₃ phase at a calcination temperature of 400 °C, the change in the number of acid sites can be attributed to the reduction of the catalyst surface area which caused by thermal sintering.

From the CO₂-TPD profiles of the Mn_2O_3 with different calcination temperatures shown in Fig. 8B and D, it can be seen that all the catalysts present a desorption peak at a range of 100–300 °C which corresponded to the weak base sites.³⁵ From the desorption profiles of CO₂ (Fig. 8B), the amount of weak base sites decreased with the increase of calcination temperatures



Fig. 8 NH₃-TPD and CO₂-TPD profiles of fresh Mn_2O_3 catalysts at different calcination temperatures: (A) NH₃-TPD profiles of Mn_2O_{3NP} samples; (B) CO₂-TPD profiles of Mn_2O_{3NP} samples; (C) NH₃-TPD profiles of Mn_2O_{3NW} samples; (D) CO₂-TPD profiles of Mn_2O_{3NW} samples.



Fig. 9 H_2 -TPR profiles of Mn_2O_3 catalysts: (A) fresh samples; (B) 8 h spent samples.

(500–800 °C). When the calcination temperature increased from 400 to 500 °C, the number of weakly basic sites increased significantly, indicating that the basicity of the catalyst was increased. This was because the Mn_2O_{3NP} calcined at 400 °C was a mixed phase of Mn_2O_3 and Mn_3O_4 . Since the average combined O^{2-} of Mn^{2+} in Mn_3O_4 was less than the number of Mn^{3+} , therefore, less O^{2-} was exposed on the entire catalyst surface, so the alkalinity was less than that calcined at 500 °C.

When calcined at a temperature above 500 $^{\circ}$ C, a stable Mn₂O₃ crystal phase has been formed. With the temperature increased, the thermal sintering of the crystal particles intensified, the specific surface area of the catalyst decreased and the corresponding basic sites on the surface also decreased. The trends of the CO₂-TPD profiles of Mn₂O_{3NW} were similar to its NH₃-TPD, which can also be attributed to the thermal sintering at high temperature. That was the reason why the number of basic



Fig. 10 (A) TG curves of Mn₂O_{3NP} catalyst; (B) DTA profiles of Mn₂O_{3NP} catalyst; (C) TG curves of Mn₂O_{3NW} catalyst; (D) DTA profiles of Mn₂O_{3NW} catalyst.

centers on the catalyst surface to decrease with the calcination temperature increased.

The influence of different morphologies was also reflected on reducibility as depicted by Fig. 9A. The peak appearing at temperature below 550 °C were assigned to hydrogen consumption by surficial oxygenic species that includes chemically adsorbed oxygen (O_{Chem}) at 372-439 °C and surficial lattice oxygen (O_{Latt S}) at 454-507 °C.³⁶ Obviously, the reduction temperature of Mn₂O_{3NP} was lower than that Mn₂O_{3NW}, which indicated that the oxygen species on the surface of Mn₂O_{3NP} were more active. As shown in Fig. 9B, it is the H₂-TPR spectra of the two catalysts after 8 h reaction. Maybe the coke deposited on the surface of the catalysts, the $O_{\rm Chem}$ and $O_{\rm Latt,S}$ were covered. Hence, the reduction temperatures for O_{Chem} and O_{Latt,S} increased after 8 h reaction. Also, there were two peaks of H₂ consumption in the range (672-719 °C) which was attributed to the reduction peak of the lattice oxygen in the bulk (O_{Latt.B}).³⁷ It is not difficult to find that the reduction temperature of the O_{Latt.B} of Mn₂O_{3NW} was lower than the Mn₂O_{3NP}. That indicates that the nanowire structure inhibited reactivity of oxygenic species on surface of Mn₂O₃, but the strong interaction between $Mn^{3\scriptscriptstyle +}$ and $O_{{\rm Latt},B}$ accelerated migration of $O_{{\rm Latt},B}$ to the surface, which can promote the methanol catalytic oxidation to formaldehyde.

In the methylation of 4-chlorophenol with methanol, with the reaction progressed, Mn³⁺ will be reduced to Mn²⁺ and the oxygen species will be consumed. In addition, the amount of coke deposited on the catalyst surface at different times would be significantly different. In order to analyze the oxygen uptake behavior and coke deposition at different periods of the reaction, TG and DTA analysis were carried out in air. In Fig. 10A and C, it can be seen that the thermal stability of the two fresh Mn_2O_3 catalysts was very good. With the reaction progressed, it was found that the spent catalyst had two curves with weight decrease in the range of 50-200 °C and 350-500 °C. The former was attributed to the thermal weight loss of the catalyst adsorbing water, while the latter was due to the removing of the coke deposited on the catalyst surface. The longer the reaction carried out, the greater the thermal weight loss of the catalyst at 350-500 °C, indicating that the amount of coke deposited increased with the progress of the reaction time. It can be seen that the spent catalyst has a weight increase at 200-350 °C. It was reasonable that the weight increase caused by oxygen regeneration from air.38 Combined with the TG-MS chart of Mn_2O_{3NP} , it can be seen that in the main weight loss part of the catalyst (350–500 °C), peaks of CO₂ and HCl appeared (Fig. 11), which proved that CO₂ and HCl were formed from the catalyst within this temperature range. There is no doubt that CO₂ was generated by the complete combustion of coke deposited in an air atmosphere. Combined with the analysis of XPS and XRD, it can be concluded that the HCl was produced by MnCl₂ in spent catalyst. When the temperature exceeded to 500 °C, the catalyst showed a curve of weight increase. This was the fact that the oxygen in the air supplemented the anion vacancies after the exit of the crystal lattice Cl⁻ under high temperature conditions, and was embedded in the crystal lattice of the catalyst in the



Fig. 11 Quadruple mass spectrometry coupled with thermo gravimetric analysis profiles.

form of O^{2-} , which was also another type of oxygen weight increase.

Fig. 10B and D displayed the DTA spectrum of the catalyst. It can be seen that the spent catalyst had an obvious exothermic peak in the temperature range of 300-400 °C. This was because the catalyst emitted heat when the coke burned in an air atmosphere. The amount of coke deposited on the catalyst surface gradually increased with the progress of the reaction, and the intensity of the exothermic peak of the catalyst for 8 h should be stronger than the 3 h. However, the obtained thermal curves were the opposite conclusion. The reason may be that with the reaction progressed, the crystal lattice Cl on the catalyst surface gradually increased which made the crystal lattice structure more perfect, which was not conducive to the formation of coke deposited though the mass of coke increased. In other words, the increase in the crystal lattice Cl reduced the unsaturation sites exposed by the Mn element, that is, the acid sites decreased, so the formation of coke was hindered. In another aspect, the endothermic effect of decomposing MnCl₂ was greater than the exothermic effect of coke combustion.

3.2 Reaction mechanism of *o*-methylation of 4-chlorophenol and the dechlorination mechanism of chlorophenol

Regarding the research on the reaction mechanism of the *ortho*methylation of phenolic compounds catalyzed by metal oxides, Tanabe initiated pioneering work.³⁹ By comparing the infrared spectra of phenol in acidic oxides and basic oxides, it was found that phenol dissociated and adsorbed on the surfaces of the two types of catalysts. The difference was that for acidic oxides, the aromatic ring of the phenate was adsorbed parallelly on the surface of the catalyst, so every position of the aromatic ring was easily attacked by the methylating agent. As for the basic oxides, due to the mutual repulsion between the large π -bonded electron cloud of the aromatic ring and the electron-rich Lewis base center (O²⁻) on the surface of the catalyst, the aromatic ring was kept away from the surface of the catalyst and adsorbed vertically on the surface of the catalyst. Since the *ortho* position of the hydroxyl O was closer to the surface of the catalyst, it was more likely to be attacked by the methylating agent, thus ensuring extremely high *ortho* selectivity. Regarding the methyl active species in the methylation reaction of methanol and phenolic compounds, Mathew believed that the H⁺ generated by the dissociation and adsorption of phenol and protonated methanol, and then a molecule of H₂O was left to form $CH_3^{\delta+}$, which then attacked the aromatic ring.⁴⁰ Cavani brought forward a completely different view, methanol was first converted to formaldehyde on the surface of the catalyst, and formaldehyde as the actual methylated species attacked the *ortho* position of the phenolic hydroxyl group.¹² Our work proved the correctness of Cavani's view through experiments and referred other similar works.

The first step was the dehydrogenation of methanol. The active site for methanol's dehydrogenation reaction was the terminal Mn=O site, which generated hydroxyl and methoxy species. The terminal oxygen was considered to be the most stable site for hydrogen adsorption⁴¹ (g = gas phase, a = adsorbed species).

$$CH_3OH_g + O_t \rightarrow CH_3O_a + OH_a$$

Next, attacking of the bridging oxygen on the adsorbed methoxy species yielded a hydroxyl and adsorbed formaldehyde. The crack of the C–H bond of the methoxy group was deemed as the rate-determining step in formaldehyde formation, and required a higher temperature for activation than the first process.

$$CH_3O_a + O_b \rightarrow H_2CO_g + OH_a$$

Two hydroxyl groups recombination occurred to yield H₂O from the surface of the catalyst.⁴² The remaining oxygen ion vacancies were filled by the migration of the bulk lattice oxygen inside the catalyst to ensure that the catalyst maintained regular activity and selectivity.

The generated formaldehyde had good positive carbon electrophilic properties and the aromatic ring possessed the electron rich delocalized electron cloud.²⁰ The carbon of formaldehyde attacked the 4-chlorophenate species, forming 5-chloro-salicylic alcohol species with hydroxymethyl moieties. In detail, the positively charged carbon attacked the *ortho* position of the aromatic ring O to form a transition state intermediate species. Because of the aromatic ring was in an electron-deficient state, the electron from the C-H covalent bond restores the aromatic ring original appearance. The generated H^+ combined with the O of the aldehyde group to generate a hydroxymethyl moiety (Scheme 1).

The resulting 5-chloro-salicylic alcohol generated 4-chloro-2cresol through a series of complex processes (Scheme 2). Significantly, 5-chloro-salicylic alcohol can generate almost equimolar 5-chloro-salicylic aldehyde and 4-chloro-2-cresol through an intermolecular disproportionation reaction. A



Scheme 1 Dissociation and adsorption of 4-chlorophenol and the attack of formaldehyde to aromatic ring.

slightly higher formation of 5-chloro-salicylic aldehyde can be attributed to the dehydrogenation of the alcohol.²⁰ The same type of reaction has been widely reported, such as the intermolecular disproportionation of ethanol into ethane and acetaldehyde, catalyzed by reduced vanadium oxide.⁴³ Also, benzyl alcohol disproportionated into toluene and benzoic aldehyde.⁴⁴

Unlike the main side reactions of *ortho*-methylation of phenol with methanol (excessive methylation of phenol), the dechlorination of 4-chlorophenol compounds was the main side reaction. The dechlorination reaction of 4-chlorophenol compounds was most directly and easily associated with the hydrogen generated by the decomposition of methanol in high temperature. However, by changing the carrier gas N_2 to H_2/Ar mixture with 10% H_2 content, we found that the selectivity of 4-chlorophenol compounds has not changed, and even the selectivity has increased slightly (Table S11†). Therefore, it showed that the hydrogen generated by the reason for dechlorination of 4-chlorophenol methanol was not the reason for dechlorination of 4-chlorophenol methanol was not the reason for dechlorination of 4-chlorophenol methanol was not the reason for dechlorination of 4-chlorophenol was not the reason for dechlorination for 4-chlorophenol was not the reason for dechlorination of 4-chlorophenol was not the reason for dechlorination for 4-chlorophenol was not the selective was not was not



Scheme 2 Summary of the hypothesis for transformation of the intermediately formed 5-chloro salicylic alcohol into 4-chloro-2-cresol during 4-chlorophenol methylation with Mn_2O_3 catalysts.



Scheme 3 A possible path way of 4-chlorophenol reduction.

chlorophenol compounds. We believed that methanol was used as a nucleophile and reducing reagent to reduce 4-chlorophenol compounds. This was due to a mechanism similar to Meerwein-Ponndorf-Verley-type (MPV) reaction.45 Next, it involved the substitution of chlorine from methanol hydrides (Scheme 3). Specifically, methanol dissociated and adsorbed on the surface of Mn_2O_3 , and the dissociated methoxy ions and H^+ were adsorbed on metal ions and oxygen ions, respectively. Chlorine in 4-chlorophenol compounds (take 4-chlorophenol as an example) was electron-rich, so it was easier to combine with oxygen-coordinated unsaturated Mn. At the same time, the negative hydrogen attacked the C-Cl bond, forming a thermodynamically stable six-membered cyclic transition state. Finally, negative hydrogen replaced the chlorine, and the adsorbed methoxy ions became formaldehyde and removed from the catalyst surface. Even though methanol was used as H transfer reagents, the MPV reaction usually occurred in the reduction of aromatic aldehydes and ketones. However, in the process of 4chlorophenol dechlorination, the manganese oxide catalyst introduced new chloride ions and hydrogen oxygen, this was consistent with the XPS characterization results of the catalyst after the reaction (the increase in the content of hydroxide ions and carbonate ions and the chlorine element was a metal chloride), which further illustrated the rationality of the explanation of the dechlorination mechanism.

3.3 Catalytic activity results

In this reaction, the products were 4-chloro-2-cresol, phenol, 2cresol, 2,6-xylenol, 2,4-xylenol, 2,4,6-trimethylphenol, 4-chloro-2,6-xylenol. Probably due to the catalyst and the reaction temperature range, this reaction did not generate 4-chloroanisole or anisole substances. In many previous work, phenol and methanol produced anisole instead of *o*-cresol under alkaline catalysts (such as MgO) and at low temperature. However, in this work, the formation of anisole or 4-chloroanisole was not detected.

Table S3[†] showed the reaction activity and product selectivity of Mn_2O_{3NP} under different calcination temperatures. It can be seen that with the calcination temperature increased, the activity of the catalyst gradually decreased. The sintering effect was responsible for low catalytic activity. The sintering effect led to the decrease in the number of active centers on the surface of the catalyst, which was also proved by the NH_3/CO_2 -TPD characterization. With the increase of the calcination temperature, the number of medium acid and weak base centers decreased significantly. The catalyst calcined at 400 °C had more acid sites on the surface, so the dechlorination effect was more significant. The selectivity of phenol and phenol methylate was higher than that of other calcination temperatures. In addition, the poly-methyl substituents of phenol (2,6-xylenol, 2,4-xylenol, 2,4,6-trimethylphenol) were also slightly higher than others, this conclusion was consistent with similar work.¹⁴ The insight was the stronger the acidity, the longer the average time the phenoxy anions were adsorbed at the acidic sites, then the easier it was for the resulting mono-substituted product to continue to react with the methyl species to generate di- or trisubstituted products.

The result of Mn₂O_{3NW} was slightly different from Mn₂O_{3NP} (Table S4[†]). This was attribute to Mn₂O_{3NW} was the same Mn₂O₃ crystal phase under investigation. The difference was that the specific surface area (the number of medium acid and weak base centers), despite the specific surface area of Mn₂O_{3NW} was larger than that of Mn₂O_{3NP}, the two samples behaved quite similarly. In a previous paper, researchers had made the hypothesis that active sites in MgO were corner and steps at crystallite surface.¹² Based on this, it might be expected that the high-surface-area Mn₂O_{3NW} possessed both a greater overall number and a greater density of defect sites, because of its wirelike structure. Nevertheless, this was in contrast with the experimental evidence. One possible explanation was that the particle efficiency in the high-surface-area Mn₂O_{3NW} was low, which was due to the internal diffusion limitations. This implied that only active sites located at the external surface of the catalyst particles contributed to reactivity. If this were the case, regardless of the difference in specific surface area, the reactivity of Mn₂O₃ would not differ too much.

The reaction temperature played a crucial role in deciding the reactivity and the selectivity of products in either vapor phase or liquid phase reaction. In temperature studies, the reactions were carried out at temperature from 350 to 450 °C. It was observed as given in Table S5 and S6,† the increased of temperature led to higher conversion of 4-chlorophenol. That was attribute to the process from methanol to formaldehyde was more likely to occur with the increase of reaction temperature. However, at 450 °C, the conversion of 4-chlorophenol was slightly decreased (Table S5[†]). This was due to the coke generated during the reaction were deposited on the surface of the catalyst, which reduced the porosity of the catalyst.⁴⁶ Simultaneously, the dechlorination effect of 4-chlorophenol also intensified with the increase of temperature. Maybe the increase of temperature enhanced the dissociation and adsorption of methanol on the catalyst surface, the hydride transfer reaction made 4-chlorophenol and 4-chlorophenol methyl compounds easier to dechlorine.

The influence of the molar ratio of chlorophenol to alcohol on this reaction is shown in Tables S7 and S8.[†] For the two morphologies catalysts, as the molar ratio increased from 1 : 1 to 1 : 20, the conversion of 4-chlorophenol generally increased, the selectivity of phenol poly-methylation species increased, and the dechlorination of chlorophenols has been inhibited. With the chlorophenol-to-alcohol molar ratio increased, the methanol densities increased, more formaldehyde intermediates produced, which increased the conversion of 4-chlorophenol. Meanwhile, the generated mono-substituted phenol or chlorophenol methyl compounds were more likely to react with formaldehyde to generate di- or multi-substituted methvlated phenol or chlorophenol under an abundant formaldehyde condition. The dechlorination effect of chlorophenols was inferior, which was seems to contrary to the mechanism we proposed, because more methanol generated more hydrides to reduce the chlorophenols. Considering that the amount of methanol molecules was larger than the amount of dechlorination sites even in the ratio was 1:1, thus it was believed that increased methanol ratio could not be change dechlorination effect visibly. In other words, the dehydrogenation of methanol formaldehyde and chlorophenol dechlorination was to occurred on different sites, perhaps the number of the former is far greater than that of the latter.

Tables S9 and S10[†] showed the reactivity and selectivity of the two Mn₂O₃ catalysts at different liquid hourly space velocity (LHSV). It can be observed that with the increase of volumetric space velocity, for Mn₂O_{3NP} catalyst, the conversion of 4-chlorophenol gradually decreased, the selectivity of phenol increased, the selectivity of 4-chloro-2-cresol slightly increased, as well the selectivity of o-cresol was basically unchanged, and the selectivity of poly-methyl substituted phenol decreased. The increase in LHSV indicated that the reactants flowing through the catalyst per unit time increased. When it reached to an extreme level, the catalyst did not have the ability to deal with so many reactants, which caused the conversion dropped. Accordingly, the LHSV increased, especially too much methanol, which generated more coke depositions, and blocked the active sites on the catalyst surface, so the conversion decline. The selectivity of 4-chloro-2-cresol was increased due to more 4chlorophenol went through the surface of the catalyst per unit time. The chlorine removed by the dechlorination effect made the dechlorination site filled untimely. Therefore, when the sampling time was 3 h, the selectivity of 4-chloro-2-cresol would naturally increase. The variety in the selectivity of phenol and poly-methyl substituted phenol can be attributed to the increase in LHSV, the less contact time between phenol species and catalyst.⁴⁶ Mn₂O_{3NW} showed a larger LHSV result than Mn₂O_{3NP}. That was because of the Mn₂O_{3NW} had a larger external surface area, which made it possible for more 4-chlorophenol dissociation adsorption.

Without the presence of any catalyst, N_2 and H_2/Ar mixture were used to the carrier gas for this reaction. It was found that the conversion of 4-chlorophenol was very low, and the conversion in both cases were almost no difference (Table S11†). Even in the H_2/Ar mixed gas atmosphere, phenol was formed, but the conversion was too low, indicating that there was no further conversion of 4-chlorophenol. That implied that the H_2 produced by methanol decomposition did not the reason why chlorophenol dechlorination. Repeating the above experiment with the presence of catalyst, it was clear that the selectivity of 4-chloro-2-cresol remained largely the same, and the conversion of 4-chlorophenol was significantly declined. This is because in the H₂/Ar mixed gas condition, the lattice oxygen of the catalyst was easier to take away, and inhibited 4-chlorophenol dissociated and adsorbed on the surface of the catalyst. In addition, due to the removal of lattice oxygen, the cations on the catalyst surface were excessive and the acidity increased, and the adsorption effect of phenol oxygen anions on the catalyst surface became stronger, which aggravated the excessive methylation of phenol and 4-chlorophenol. Hence, the selectivity of phenol poly-methylation species was increased. Under the same conditions, when formalin solution was used as the methylating agent, the dechlorination of 4-chlorophenol was effectively inhibited, indicating that methanol was the cause of dechlorination of 4-chlorophenol and its analogues. In addition, thanks to formalin was directly used as the methylation reagent, the process of methanol to formaldehyde was skipped, so there were more methyl active species (formaldehyde) around the adsorbed phenol oxygen anions though traces of methanol. Therefore, the selectivity of phenol and o-cresol decreased, the poly-methyl substituted phenol increased.

4. Conclusions

Two kinds of morphologies Mn₂O₃ catalysts were prepared by precipitation and hydrothermal methods and calcined under different temperatures. Increasing calcination temperature increased the crystallinity of the catalysts. Mn₂O_{3NP} need higher calcination temperature to form a Mn_2O_3 phase than Mn_2O_{3NW} . As the reaction to, coke deposition on the catalyst and MnCl₂ crystalline grain emerged, and the Mn³⁺ was reduce to Mn²⁺. The specific surface area of the Mn₂O_{3NW} catalyst was larger than that of the Mn₂O_{3NP}. And the specific surface area and pore volume of each catalyst decreased with the increase of the calcination temperature, which leaded to the decline of the number of acid-base centers on the catalyst surface. The catalytic activity and selectivity of the two catalysts had little difference, mainly because the methylation reaction of 4chlorophenol with methanol occurred on the outer surface of the catalyst. Temperature affected the process of methanol dehydrogenation to formaldehyde. High reaction temperature caused excessive coke deposition on the surface of the catalyst due to methanol decomposition, which affected the subsequent catalytic reaction. Increasing the ratio of alcohol to 4-chlorophenol produced more formaldehyde. Overabundance methanol could not promote the dechlorination of 4-chlorophenol and its methyl compounds, therefore, it did not cause excessive reduction of 4-chlorophenol, but enhanced the degree of polymethylation of 4-chlorophenol. The increase in the LHSV of the reactants reduced the conversion of 4-chlorophenol, and the selectivity of 4-chloro-2-cresol increased slightly. At the same time, the Mn₂O_{3NW} catalyst can maintain the reactivity at a higher LHSV. The conversion was corresponding to particle efficiency instead of simply specific surface area. The dechlorination effect of 4-chlorophenol and its analogues was due to the hydride produced by methanol on the surface of the catalyst substitute the chlorine on the phenol ring, rather than the hydrogen produced by methanol thermal decomposition.

Conflicts of interest

The authors declare not competing financial interest.

Acknowledgements

This work was supported by Jilin Province Science and Technology Research Plan (Key Scientific Research Project) (No: 20150204020GX).

References

- 1 J. Blanco, P. Avila and F. V. Melo, *Bull. Chem. Soc. Jpn.*, 1983, 56, 909–913.
- 2 J. M. Gnaim and R. A. Sheldon, *Tetrahedron Lett.*, 2004, 45, 9397–9399.
- 3 O. Shoji, T. Kunimatsu, N. Kawakami and Y. Watanabe, *Angew. Chem., Int. Ed.*, 2013, **52**, 6606–6610.
- 4 Y. Li, B. Li, L.-F. Geng, J. Wang, Y. Wang and J. Huang, *Catal. Lett.*, 2015, **145**, 1014–1021.
- 5 L. Menini and E. V. Gusevskaya, *Appl. Catal., A*, 2006, **309**, 122–128.
- 6 L. Menini and E. V. Gusevskaya, *Chem. Commun.*, 2006, 209–211.
- 7 E. A. Bugnet, A. R. Brough, R. Greatrex and T. P. Kee, *Tetrahedron*, 2002, **58**, 8059–8065.
- P. Bovonsombat, R. Ali, C. Khan, J. Leykajarakul, K. Pla-on, S. Aphimanchindakul, N. Pungcharoenpong, N. Timsuea, A. Arunrat and N. Punpongjareorn, *Tetrahedron*, 2010, 66, 6928–6935.
- 9 Y.-W. Xiong, H.-D. Duan, X. Meng, Z.-Y. Ding and W.-C. Feng, *J. Chem.*, 2016, 2016, 1–5.
- 10 H. Huang, J.-Y. Liu, X.-X. Liu, J.-F. Xiao, S. Zhong, X.-C. She, Z.-H. Fu and D.-L. Yin, *Fuel*, 2016, **182**, 373–381.
- 11 A. R. Gandhe, J. B. Fernandes, S. Varma and N. M. Gupta, *J. Mol. Catal. A: Chem.*, 2005, **238**, 63–71.
- 12 F. Cavani, L. Maselli, S. Passeri and J. A. Lercher, *J. Catal.*, 2010, **269**, 340–350.
- 13 S. Velu and C. S. Swamy, Appl. Catal., A, 1994, 119, 241-252.
- 14 C. F. Braganza and A. V. Salker, *Chin. J. Catal.*, 2016, 37, 1991–1996.
- 15 Y.-L. Wang, Y.-Y. Song, W.-T. Huo, M.-J. Jia, X.-Y. Jing, P.-P. Yang, Z. Yang, G. Liu and W.-X. Zhang, *Chem. Eng. J.*, 2012, **181**, 630–635.
- 16 N. Ballarini, F. Cavani, S. Passeri, L. Pesaresi, A. F. Lee and K. Wilson, *Appl. Catal., A*, 2009, **366**, 184–192.
- 17 K. V. R. Chary, K. Ramesh, G. Vidyasagar and V. Venkat Rao, J. Mol. Catal. A: Chem., 2003, 198, 195–204.
- 18 V. Crocellà, G. Cerrato, G. Magnacca, C. Morterra, F. Cavani, L. Maselli and S. Passeri, *Dalton Trans.*, 2010, **39**, 8527–8537.
- 19 A. S. Reddy, C. S. Gopinath and S. Chilukuri, *J. Catal.*, 2006, **243**, 278–291.
- 20 T. Tabanelli, S. Passeri, S. Guidetti, F. Cavani, C. Lucarelli, F. Cargnoni and M. Mella, *J. Catal.*, 2019, **370**, 447–460.

- 21 M. Yadav, K. Y. Rhee, S. J. Park and D. Hui, *Composites, Part B*, 2014, **66**, 89–96.
- 22 E. S. Ilton, J. E. Post, P. J. Heaney, F. T. Ling and S. N. Kerisit, *Appl. Surf. Sci.*, 2016, **366**, 475–485.
- 23 N. A. Merino, B. P. Barbero, P. Eloy and L. E. Cadús, *Appl. Surf. Sci.*, 2006, 253, 1489–1493.
- 24 A. F. Carley, M. W. Roberts and A. K. Santra, *J. Phys. Chem. B*, 1997, **101**, 9978–9983.
- 25 K. Sutthiumporn and S. Kawi, *Int. J. Hydrogen Energy*, 2011, 36, 14435–14446.
- 26 J. L. G. Fierro, Catal. Today, 1990, 8, 153-174.
- 27 H. Raaf and N. Schwentner, Appl. Surf. Sci., 2001, 174, 13-34.
- 28 J. Newnham, K. Mantri, M. H. Amin, J. Tardio and S. K. Bhargava, *Int. J. Hydrogen Energy*, 2012, 37, 1454–1464.
- 29 N. Laosiripojana and S. Assabumrungrat, *Appl. Catal., B*, 2005, **60**, 107–116.
- 30 G. R. Patzke, F. Krumeich and R. Nesper, *Angew. Chem., Int. Ed.*, 2002, **41**, 2446–2461.
- 31 L. Vayssieres, K. Keis, A. Hagfeldt and S. E. Lindquist, *Chem. Mater.*, 2001, **13**, 4395–4398.
- 32 D.-B. Kuang, A.-W. Xu, Y.-P. Fang, H.-Q. Liu, C. Frommen and D. Fenske, *Adv. Mater.*, 2003, 15, 1747–1750.
- 33 V. N. Kalevaru, N. Madaan and A. Martin, *Appl. Catal., A*, 2011, **391**, 52–62.
- 34 H. Zhao, C.-C. Zuo, D. Yang, C.-S. Li and S.-J. Zhang, *Ind. Eng. Chem. Res.*, 2016, **55**, 12693–12702.
- 35 G.-L. Zhang, H.-H. Zhang, D. Yang, C.-S. Li, Z.-J. Peng and S.-J. Zhang, *Catal. Sci. Technol.*, 2016, **6**, 6417–6430.
- 36 Q.-Q. Li, Z. Huang, P.-F. Guan, R. Su, Q. Cao, Y.-M. Chao,
 W. Shen, J.-J. Guo, H.-L. Xu and R.-C. Che, ACS Appl. Mater. Interfaces, 2017, 9, 16243–16251.
- 37 Y.-X. Gao, W.-D. Wang, S.-J. Chang and W.-X. Huang, *ChemCatChem*, 2013, 5, 3610–3620.
- 38 D.-C. Wang, L.-J. Jin, Y. Li and H.-Q. Hu, *Fuel*, 2017, **210**, 803–810.
- 39 K. Tanabe and W. F. Hölderich, *Appl. Catal., A*, 1999, **181**, 399–434.
- 40 T. Mathew, B. B. Tope, N. R. Shiju, S. G. Hegde, B. S. Rao and C. S. Gopinath, *Phys. Chem. Chem. Phys.*, 2002, **4**, 4260–4267.
- 41 C. Brookes, M. Bowker and P. P. Wells, *Catalysts*, 2016, **6**, 92–118.
- 42 J. S. Chung, R. Miranda and C. O. Bennett, *J. Catal.*, 1988, **114**, 398–410.
- 43 Y. Nakamura, T. Murayama and W. Ueda, *ChemCatChem*, 2014, 6, 741–744.
- 44 E. Nowicka, J. P. Hofmann, S. F. Parker, M. Sankar, G. M. Lari, S. A. Kondrat, D. W. Knight, D. Bethell, B. M. Weckhuysen and G. J. Hutchings, *Phys. Chem. Chem. Phys.*, 2013, 15, 12147–12155.
- 45 T. Pasini, A. Lolli, S. Albonetti, F. Cavani and M. Mella, J. Catal., 2014, 317, 206–219.
- 46 K. H. Bhadra and G. D. Yadav, *Appl. Catal., A*, 2018, **562**, 67–78.