# RSC Advances

# PAPER

Cite this: RSC Adv., 2017, 7, 48275

### Received 31st July 2017 Accepted 9th October 2017

DOI: 10.1039/c7ra08442k

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# Introduction

Quinolines (quinoline and substituted quinolines) are a type of important heterocyclic compound, which can be widely used as pharmaceuticals, fungicides, herbicides, corrosion inhibitors and functional chemicals.<sup>1-8</sup> Various traditional routes involving liquid phase reaction, such as, Skraup,<sup>9,10</sup> Doebner-Miller,<sup>11,12</sup> Friedländer,<sup>13,14</sup> Combes,<sup>15,16</sup> and Pfitzinger methods,<sup>17</sup> have been developed and utilized in the synthesis of quinolines. However, these routes all suffered from a few drawbacks, e.g., anhydrous reaction condition, volatile organic solvent, expensive or toxic feed, corrosive and/or costly homogeneous catalyst, prolonged reaction time, tedious work-up procedure and difficult recycling of the catalyst.<sup>9-13,15,18,19</sup> In contrast, gas phase reaction route for the synthesis of quinolines employs an appropriate heterogeneous catalyst,<sup>20-25</sup> which can solve effectively the above issues associated with liquid phase reaction route, and thus, it is receiving more and more attentions.

In the gas phase reaction route, up till now, small molecular aldehydes or carbonyl compounds containing active hydrogen on a-carbon, such as, acetaldehyde, propionaldehyde and butyraldehyde, were most frequently employed as the raw materials, due to their high activity in the reaction with aniline

# Heterogeneous catalytic synthesis of quinoline compounds from aniline and  $C_1-C_4$  alcohols over zeolite-based catalysts†

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The synthesis of quinolines from aniline and a  $C_1-C_4$  alcohol was conducted under gas-phase reaction conditions over a series of zeolite-based catalysts. The texture and acid properties of catalysts were characterized by XRD, FT-IR, BET and NH<sub>3</sub>-TPD techniques. It was found that the total yield of quinolines was positively related to the relative content of Lewis acid sites of the catalyst. Among others, the ZnCl<sub>2</sub>/Ni-USY-acid catalyst possessed the best performance. Over this catalyst, the reactions of aniline and most of the alcohols provided a 42.3–79.7% total yield of quinolones under mild conditions, however, those of aniline and methanol, ethanol and iso-propanol predominantly led to N-alkylanilines. Furthermore, the reaction pathways for synthesizing quinolines via aniline reacting with polyhydric alcohols or monohydric alcohols was proposed in our work. **PAPER**<br> **EXERCTS SUBLIC SYnthesis of quinoline**<br> **CREATE ARTICLE STILL SYNTHESIS OF QUINCIINE<br>
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to synthesize quinolines. However, these raw materials are usually toxic, expensive, and particularly, prone to polymerization during reaction.<sup>22,24-26</sup> Relative to carbonyl compounds, small molecular alcohols are hard to polymerize, much more eco-friendly and cheaper, since they can be alternatively manufactured from the fermentation of biomass materials besides the chemical synthesis from fossil-based raw materials. This enables small molecular alcohols to be potential substitutes of carbonyl compounds in the synthesis of quinolines. In fact, few papers dealt with the application of alcohol as feedstock to synthesizing quinolines *via* gas phase reaction route.<sup>20,21,23</sup> For examples, Campanati et al.<sup>21</sup> obtained a 41% yield of 2-methyl-8ethylquinoline from the reaction of ethylene glycol and 2-ethylaniline at 330 °C over K10 montmorillonite catalyst, while Reddy et al.<sup>23</sup> reported a 65% yield of quinoline via the reaction of glycerol and aniline at 425  $\degree$ C over mixed metal oxides catalyst in the presence of air. However, such kind of attempts has been very limited, excepting the above-mentioned reports in the literature. Therefore, an exploration on the utilization of various types of alcohols as feedstock in reacting with aniline to generate quinolines is of signicant importance.

The catalysts for the synthesis of quinolines via gas phase reaction route reported in the literature included mainly metal halide clusters, K10, amorphous silica–alumina and mixed metal oxides.20,22–<sup>25</sup> These catalysts usually presented low to middle activities for the generation of quinolines but low catalytic stabilities, due to their poor acid performance and textural properties. Zeolites represent an important class of porous materials that have found numerous applications in catalysis, especially in the pyrolysis of petroleum to gasoline, because of their unique and tunable acid performance, large stability of framework and well-defined microporous structure.



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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/c7ra08442k

Y-Type zeolite was employed as catalyst in the synthesis of substituted quinolines, showing an appreciable catalytic activity.<sup>24</sup> However, zeolite catalysts usually had to face an issue of transport limitation caused by the slow diffusion rates of reactant and bulky product in microporous.27,28 In this respect, the issue is more serious, since quinolines are relatively bulky, relative to gasoline molecules. To reduce the diffusion limitation, mesostructured ultra-stable Y zeolite (USY) was more frequently employed as catalyst in a series of reactions,<sup>29-31</sup> due to their modified acid performance and large average pore size and therefore high stability and catalytic activity. Mesoporous USY zeolite could be prepared by acid or alkali treatment,<sup>29,31-34</sup> steam dealumination, $30,32,35,36$  or in situ synthesis with mesoporous structure-directing agent.<sup>37</sup> Among others, the acidtreated method of USY zeolite was considered as the most simple and efficient one, and thus, accepted by many researchers. However, to the best of our knowledge, no report on the employment of acid-treated USY zeolite as catalyst for the synthesis of substituted quinolines had been documented in the literature.

In this paper, we reported for the first time the systematical studies on the synthesis of quinolines from aniline and a  $C_1-C_4$ alcohols over modified USY catalysts via gas phase route. It was clarified that the transition metal-modified mesoporous USY could be used as an efficient catalyst for the reaction of various alcohols and aniline to quinolines, exhibiting not only a high catalytic activity but also a good catalyst stability. Furthermore, the related reaction mechanisms, dependent on the type of alcohols, were also proposed in this paper.

## Results and discussion

Fig. 1 shows the XRD patterns for various catalysts. One can see that all the catalysts exhibit the characteristic diffraction peaks for the USY zeolite, and the Ni-USY-acid and ZnCl<sub>2</sub>/Ni-USY-acid catalysts do not shown the diffraction peaks for Ni and  $ZnCl<sub>2</sub>$ species. The intensities of diffraction peaks for the USY-acid, Ni-USY-acid and  $ZnCl<sub>2</sub>/Ni-USY-acid$  catalysts are all lower than for the parent USY. It shows that the original structure of USY zeolite is retained, though suffering from a destroy to some extent, after acid treatment, and Ni and  $ZnCl<sub>2</sub>$  species possess very small dimension and have been highly dispersed on USY zeolite. From the enlargement of diffraction peaks, e.g., those ranging from  $9.5^\circ$  to  $12.5^\circ$ , one can find that the diffraction peaks shift slightly towards higher angel over the USY-acid catalyst but obviously towards lower angel over both the Ni-USY-acid and  $ZnCl<sub>2</sub>/Ni-USY-acid}$  catalysts, relative to the parent USY zeolite. As is well known, the difference in the diameters of framework T atoms causes theoretically the expanse or contraction of zeolite framework and thus the whole shift of characteristic diffraction peaks of zeolite,<sup>38</sup> while the diameters of the 4-coordinated ions  $\text{Zn}^{2+}$  (74 pm), Ni<sup>2+</sup> (69 pm),  $Al^{3+}$  (53 pm) and  $Si^{4+}$  (40 pm) decrease sequentially. It is therefore deduced that the partial dealumination occurs during acid treatment of USY zeolite<sup>29</sup> and a proportion of Ni and Zn ions are inserted into the framework during the Ni ion exchange and ZnCl<sub>2</sub> loading over USY-acid zeolite.



Fig. 1 XRD patterns for the unmodified and modified USY zeolite catalysts: (a) USY; (b) USY-acid; (c) Ni-USY-acid; (d) ZnCl<sub>2</sub>/Ni-USY-acid (the inset is the enlargement of the patterns for  $2\theta = 9.5-12.5^{\circ}$ ).

Fig. 2 shows FT-IR the spectra of various catalysts. The bands at ca. 1188, 1064 and 824  $cm^{-1}$  over the USY catalyst are characteristic of TO<sub>4</sub> tetrahedron units, which can be assigned to the external asymmetric, internal asymmetric and external symmetric stretching vibrations of Si-O–T linkages for  $TO_4$  (T = Si and/or Al) tetrahedral. $39-41$  These bands are known to shift



Fig. 2 FT-IR spectra for the unmodified and modified USY zeolite catalysts: (a) USY; (b) USY-acid; (c) Ni-USY-acid; (d) ZnCl2/Ni-USYacid.



Fig. 3  $N_2$ -adsorption/desorption isotherms and BJH pore-size distribution curves for the unmodified and modified USY zeolite catalysts: (a) USY; (b) USY-acid; (c) Ni-USY-acid; (d) ZnCl<sub>2</sub>/Ni-USYacid.

towards higher wavenumbers with increasing the Si/Al molar ratio of zeolite. Besides, the band at 455  $cm^{-1}$  over the USY catalyst is attributed to the internal T–O bending vibration of  $TO<sub>4</sub> tetrahedron<sup>42</sup>$ , and this band is nearly unaffected by the Si/Al molar ratio of zeolite. Compared to the USY catalyst, the Si–O–T stretching vibrations at 1188, 1064 and 824  $\text{cm}^{-1}$  shift respectively to 1192, 1068 and 825  $cm^{-1}$  over the USY-acid catalyst, indicating that the partial dealumination of USY zeolite occurs during the its acid treatment. However, the above Si-O-T stretching vibrations shift respectively to 1178, 1059 and 820  $\mathrm{cm}^{-1}$ , over the Ni-USY-acid and ZnCl<sub>2</sub>/Ni-USY-acid catalysts. Szostak et  $al$ .<sup>43</sup> reported that the incorporation of gallium replaced the framework aluminum of zeolite and in turn led to the significant shift of Si-O-T stretching vibrations towards lower wavenumber, due to the change in the reduced mass of the Si–O–T harmonic oscillator. Accordingly, the above variations in the Si–O–T stretching vibrations can be due to the incorporations of a proportion of Ni and/or Zn ions into the framework of USY-acid zeolite. The above results are consistent with those derived from XRD characterization.

Fig. 3 shows the  $N_2$  adsorption-desorption isotherms for various catalysts, and the corresponding textural properties are summarized in Table 1. One can see from Fig. 3 that, for all the catalysts, the isotherms rise steeply at very low relative pressure, showing the presence of micropores. In addition, the hysteresis loop at  $P/P_0 > 0.4$  is obviously present for the USY-acid, Ni-USYacid and ZnCl<sub>2</sub>/USY-acid catalysts but very unclear for the USY catalyst. It shows that mesopores are generated via the acid treatment of USY zeolite, of which the pore sizes range from 1.5 to 6.0 nm and possess a most possible value at ca. 2.0 nm (see the inset in Fig. 3). From Table 1, one can find that, compared to the USY catalyst, the USY-acid catalyst possesses a larger mesopore volume  $(V_{\text{meso}})$ , total volume  $(V_{\text{total}})$  and external surface area ( $S_{ext}$ ) but smaller micropore volume ( $V_{mic}$ ), micropore surface area ( $S<sub>mic</sub>$ ), micropore size ( $D<sub>mic</sub>$ ) and specific surface area ( $S<sub>BET</sub>$ ). Besides, mesopores with an average size of 2.0 nm are determined for the USY-acid catalyst but not for the USY catalyst. It indicates that the generation of mesopores is a result of the partial dealumination of zeolite by acid treatment,<sup>44,45</sup> and also, a proportion of extracted Al species may have occupied the micropores of zeolite. Compared to the USY-acid, the pore volumes ( $V_{\text{meso}}$ ,  $V_{\text{mic}}$  and  $V_{\text{total}}$ ), surface areas ( $S_{\text{ext}}$ ,  $S_{\text{mic}}$  and  $S<sub>BET</sub>$ ) and pore size ( $D<sub>mic</sub>$  and  $D<sub>mes</sub>$ ) all decrease sequentially over the Ni-USY-acid and ZnCl<sub>2</sub>/Ni-USY-acid. This can be due to the fact that, during the Ni ion exchange and  $ZnCl<sub>2</sub>$  loading, a proportion of  $Ni^{2+}$  and  $Zn^{2+}$  ions enter into the pores and another proportion of  $Ni^{2+}$  and  $Zn^{2+}$  ions are incorporated into the framework of zeolite. Puper  $\frac{1}{2}$ <br>
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Fig. 4 shows the  $NH<sub>3</sub>-TPD$  profiles for various catalysts. The temperatures at the maximum  $(T_i)$  and integral area  $(A_i)$  of the peaks, which correspond respectively to the strength and concentration of acid sites, are summarized in Table 2. The  $T_1$ ,  $T_3$  and  $T_4$  peaks are present over all the catalysts, and they have a maximum at ca. 141-147 °C, 265-280 °C and 330 °C, respectively. The  $T_1$  peak is attributed to the weak adsorption of ammonia molecule over the surface terminal silanol (Si–OH) group,<sup>46</sup> while the  $T_4$  peak can be assigned to the surface Brönsted acid site, i.e., bridged hydroxyl group (Si-OH-Al), in Y-type zeolite.<sup>47</sup> It is known that the extra-framework Al species behave as Lewis acid site and possess a weaker strength than the bridged hydroxyl group in Y-type zeolite.<sup>47</sup> Accordingly, the  $T_3$  peak can be attributed to the Lewis acid site associated with the extra-framework Al specie. Over various catalysts, the area of  $T_4$  peak exhibits an order USY > USY-acid > Ni-USY-acid  $\approx$  $ZnCl<sub>2</sub>/Ni-USY-acid$  and that of  $T<sub>3</sub>$  displays an inverse order,



<sup>a</sup> S<sub>BET</sub>, S<sub>ext</sub> and S<sub>mic</sub>: specific surface area, external surface area and micropore surface area, respectively, and S<sub>BET</sub> = S<sub>ext</sub> + S<sub>mic</sub>. <sup>b</sup> V<sub>total</sub>, V<sub>mic</sub> and  $V_{\text{meso}}$ : total pore volume, micropore volume and mesopore volume, respectively.  $c_{D_{\text{micro}}}$  and  $D_{\text{meso}}$ : micropore pore size and average mesopore pore size, respectively.



Fig. 4  $NH_{5}$ -TPD profiles for the unmodified and modified USY zeolite catalysts: (a) USY; (b) USY-acid; (c) Ni-USY-acid; (d) ZnCl2/Ni-USYacid.

while the area of  $T_1$  peak has an order USY  $\approx$  USY-acid > Ni-USY-acid >  $ZnCl_2/Ni$ -USY-acid. The result can be due to the fact that, firstly, the acid treatment and Ni ion exchange both result in the partial dealumination of zeolite, and this decreases the concentration of Si–OH–Al groups but increases that of extra-framework Al species over the USY-acid and Ni-USY-acid catalysts, relative to the USY catalyst; secondly, the Ni ion exchange leads to the replacement of hydrogen ion in the surface Si–OH–Al, and also, to some extent, that in the surface Si–OH group, and this decreases both the concentrations of the two kinds of hydroxyl groups over the Ni-USY-acid catalyst, relative to the USY-acid catalyst; thirdly, it was reported that  $ZnCl<sub>2</sub>$  could react with the surface hydroxyl groups of Mont-K 10 clay to form the Lewis acid sites (–O–Zn–Cl) by the thermal activation.<sup>48</sup> Accordingly, the surface hydroxyl groups (Si–OH and Si–OH–Al) over the Ni-USY-acid catalyst can react with ZnCl<sub>2</sub>, and this decreases the concentrations of surface hydroxyl groups, and also, would generates the new Lewis acid site (–O– Zn–Cl) over the  $ZnCl<sub>2</sub>/Ni-USY-acid}$  catalyst, relative to the Ni-USY-acid catalyst. In fact, a new peak  $T_2$  with its maximum at

ca. 185 °C has been identified over the ZnCl<sub>2</sub>/Ni-USY-acid catalyst but not over the other catalysts, and this peak can be assigned to the new Lewis acid site related to the –O–Zn–Cl groups. It is observed that the desorption temperature for the  $T_2$ peak is lower than that for the  $T_3$  peak, and also, the desorption temperature for the  $T_3$  peak is lower over the ZnCl<sub>2</sub>/Ni-USY-acid and Ni-USY-acid catalysts than over the USY-acid and USY catalysts. This can be due to the facts that the Ni ion exchanged over the zeolite can behave as a kind of Lewis acid site, and thus, it contributes to the  $T_3$  peak, besides the extra-framework Al species over the ZnCl<sub>2</sub>/Ni-USY-acid and Ni-USY-acid catalysts. Also, the electronegativity for the  $\text{Zn}^{2+}$ , Ni<sup>2+</sup> and Al<sup>3+</sup> increases sequentially, and accordingly, the strength for the corresponding Lewis acid sites increases in order. Besides the above  $T_1 - T_4$ peaks, a new peak  $T_5$  with its maximum at ca. 508 °C is also present over the ZnCl<sub>2</sub>/Ni-USY-acid, Ni-USY-acid and USY-acid catalysts but almost absent over the USY catalyst. It was reported that the partial removal of the Al atoms in the secondneighbor positions increased signicantly the strengths of Bronsted acid sites in the Y-typed zeolite.<sup>49</sup> Liu et al.<sup>50</sup> found that the strength of acid sites in the micropores or small cages was stronger than that in the mesopores or larger cavities over the dealuminated USY zeolite. Besides, Fritz et al.<sup>51</sup> and Beyerlein  $et al.<sup>52</sup>$  found that the strength of the bridged Brönsted acid sites (Si–OH–Al groups) were increased obviously due to the inductive effect of extra-framework aluminum species. Accordingly, the peak  $T_5$  can be attributed to the strong Bronsted acid sites, which are generated due to the removal of some adjacent framework Al species, and/or the inducement of the normal Bronsted acid sites, particularly those in the micropores, by the extra-framework Al species. Compared that over the USY-acid catalyst, the areas of peak  $T_5$  over the Ni-USY-acid and ZnCl<sub>2</sub>/ Ni-USY-acid catalysts decrease, particularly for the latter one. This may be due to the partial blockage of the micropores in zeolite by a proportion of Ni and Zn species, which can be evidenced by the  $N_2$ -physisorption determination. From Table 2, one can also see that the ratio for the total concentration of Lewis acid sites to the total Bronsted acid sites over various catalysts, i.e.,  $(A_2 + A_3)/(A_4 + A_5)$ , exhibits an order ZnCl<sub>2</sub>/Ni-USYacid > Ni-USY-acid > USY-acid > USY. **PSC Advances**<br>
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> Table 3 shows the results for the reaction of aniline and n-propanol over various catalysts. One can see that 2-ethyl-3 methyl-quinoline (2E-3MQ) over all the catalysts and 2,3 dimethylquinoline (DMQ), 2-ethylquinoline (2-EQ) and



<sup>a</sup>  $T_i$  refers to the temperature (°C) at the maximum of desorption peak *i*.  $\frac{b}{A_i}$  refers to the integral area of desorption peak *i*, and it also means the concentration (mmol  $g^{-1}$ ) of acid sites corresponding to the desorption peak i.





<sup>a</sup> Reaction condition: LHSV (aniline) = 0.4 h<sup>-1</sup>; molar ratio of propanol/aniline = 3; reaction temperature = 410 °C. <sup>b</sup>  $\chi$ : conversion of aniline. <sup>c</sup>  $S_i$ : selectivity to component *i*. <sup>d</sup>  $S_{QS}$ : total selectivi dimethylquinoline. <sup>h</sup> 2-EQ: 2-ethylquinoline. <sup>i</sup> 2-MQ: 2-methyl quinoline. <sup>j</sup> NPA: N-propylaniline. <sup>k</sup> DNPA: N,N-dipropylaniline. <sup>i</sup> Others: 2propylaniline, 4-propylaniline and 3-methylindole.

3-methylquinoline (3-MQ) over some catalysts are generated. These products have an order  $2E-3MO \gg DMO \gg 2-EO$  $\sim$  3-MQ. Besides, the byproducts, including predominantly N-propylaniline (NPA) and small amounts of 4-propylaniline, 2-propylaniline, N,N-dipropylaniline, and 3-methyl indole, are also identified in the product mixtures. Over the unmodified catalysts (entries 1, 3, 6, 9 and 11), the conversion of aniline exhibits an order USY-acid > USY >  $SiO_2/Al_2O_3$  > HZSM-5 > HY. This order is found to be approximately reverse to that for the concentrations of Brönsted acid sites over  $SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub>$ , HZSM-5 and HY is usually in the range of *ca.* 0.01-0.03 mmol  $g^{-1}$ , 0.1–0.3 mmol  $g^{-1}$  and 1–3 mmol  $g^{-1}$ , respectively; because of dealumination, the concentration of Brönsted acid sites over HY, USY and USY-acid decreases sequentially. Therefore, the lower concentration of Brönsted acid sites is favorable to the conversion of aniline, probably due to the smaller tendency for the deactivation of basic aniline by the protons of Brönsted acid sites. However, USY catalyst is also known to possess a higher concentration of Brönsted acid sites than  $SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub>$  catalyst, while the former catalyst exhibits a larger conversion of aniline for the synthesis of quinolines than the later one. This is most probably because USY catalyst usually possesses a larger specific surface area (>700  $m^2$   $g^{-1}$ ) than SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> (<350  $\mathrm{m^2~g^{-1}}$ ), enabling more active sites, *e.g.*, Lewis acid sites, to be exposed for the activation and conversion of aniline. Both the total selectivity to quinolines  $(S_{OS})$  and total yield of quinolines  $(Y_{QS})$  have an order USY-acid > USY > HY > HZSM-5 >  $SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub>$ . This can be due to the fact that the pores size of the zeolites employed in this work have an order USY-acid > USY > HY > HZSM-5, while quinolines possess a larger molecule size than the byproducts as mentioned above and therefore their

formation is favorable in the larger pores than in the smaller pores. Moreover, zeolites are known to possess a narrow distribution of pore size and thus favor the selective formation of quinolines through cyclisation–aromatization, relative to the amorphous  $SiO_2-Al_2O_3$  that has a broad pore size distribution. One may have noticed that the selectivity to NPA as the predominant byproduct over the unmodified catalyst has an order USY-acid < USY < HY < HZSM-5 <  $SiO_2$ - $Al_2O_3$ , being just reverse to that for the total selectivity to quinolines. It hints that the generation of quinolines can be a result of the further conversion of NPA. Over the mono-metal-modified catalysts (entries  $2, 4, 5, 7, 8, 10$  and  $12-16$ ), particularly those modified by Ni, the conversion of aniline as well as the total selectivity to quinolines and total yield of quinolines are all increased, relative to the corresponding unmodified catalysts (entries 1, 3, 6, 9) and 11). Moreover, the total yield of quinolines over the modi fied catalysts exhibits the same order as that for the corresponding unmodified catalysts. It indicates that the modification of metal has presented a positive effect on the catalytic performance, and the promotion effect of metal display an order Ni > Fe > Co > Cu > Zn. The Ni-USY-acid catalyst (entry 16) provides a 38.5% total yield of quinolines with 68.6% conversion of aniline and 56.1% total selectivity to quinolines. Over the  $ZnCl<sub>2</sub>$  further modified Ni-USY-acid catalyst, i.e.,  $ZnCl<sub>2</sub>/$ Ni-USY-acid (entry 17), the total yield of quinolines is found to increase significantly, providing an as high as 97.6% conversion of aniline with 79.7% total selectivity to quinolines. Besides, compared to that (40.1%) over the Ni-USY-acid catalyst, the selectivity to NPA (9.3%) decreases obviously over the  $ZnCl<sub>2</sub>/Ni-$ USY-acid catalyst. As is well known, transition metal ions behave as Lewis acids, while  $ZnCl<sub>2</sub>$  as a Lewis acid catalyst has exhibited good performances in many reactions like Friedel– Crafts alkylation.<sup>50,51,53</sup> Therefore, the larger Lewis acidity is highly desired for the catalyst in the synthesis of quinolines from aniline and propanol, and  $ZnCl<sub>2</sub>$  promotes not only the  $N$ -alkylation of aniline and  $n$ -propanol to generate NPA but also the further conversion of NPA to quinolines. After the carrier gas  $H_2$  is replaced with air, the total yield of quinolines is decreased obviously over the  $ZnCl<sub>2</sub>/Ni-USY-acid$  catalyst (cf. entries 17 and 18). It is known that the formation of quinolines from aniline and a carbonyl compound involves the cyclization-dehydrogenation step.<sup>22</sup> Accordingly, the presence of low valent metal species, acting as both Lewis acid sites and dehydrogenating active centers, is vital for a catalyst to provide high performance in the synthesis of quinolines from aniline and alcohol. The above results indicate that the  $ZnCl<sub>2</sub>/Ni-USY$ acid exhibits the highest catalytic performance among all the examined catalysts.

Fig. 5 illustrates the effects of various conditions on the conversion of aniline as well as the total selectivity to quinolines and the total yield of quinolines for the reaction of aniline and propanol over the ZnCl<sub>2</sub>/Ni-USY-acid catalyst, using  $H_2$  as carrier gas. Fig.  $5(A)$  shows the effect of  $ZnCl<sub>2</sub>$  loading in the catalyst. One can see that, with the increase of  $ZnCl<sub>2</sub>$  loading, the conversion, selectivity as well as yield all increase obviously, achieving their maxima at 10 wt%  $ZnCl<sub>2</sub>$  loading, and then decrease. While the loading of  $ZnCl<sub>2</sub>$  introduces more Lewis

active sites for the generation of quinolines, too large amounts of ZnCl<sub>2</sub> may block the channels of zeolite catalyst so as to inhibit the diffusion of reactants and products. Fig. 5(B) shows the effect of temperature, which indicates that, with increasing the temperature, the conversion increases all through, however, both the selectivity and yield exhibit a maximum at 410  $\degree$ C. This occurs owing to the fact that, though the increase in temperature promotes the activation and conversion of aniline and propanol over the catalyst, too higher temperatures may also lead to some side reactions, e.g., the cracking of reactants and products and the formation of carbon deposition. Fig. 5(C) shows the effect of the molar ratio of propanol/aniline, which indicates that, with increasing the molar ratio of propanol/ aniline from 1 to 3, both the conversion and yield increase and the selectivity decreases. However, with further increasing the molar ratio of propanol/aniline, the conversion is almost unchanged, and both the selectivity and yield are decreased. This is due to the fact that, as mentioned from the discussion of Table 1, the generation of quinolines can be a result of the further conversion of NPA. Therefore, the increase in the molar ratio of propanol/aniline may lead to not only the conversion of NPA to N,N-dipropylaniline (DNPA) but also the direct reaction between aniline and propanol to other byproducts such as 2-propylaniline and 4-propylaniline, competing with the formation of quinolines. Fig. 5(D) shows the effect of LHSV (aniline). It can be seen that with increasing the LHSV, the **BSC Advances**<br>
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Fig. 5 Effect of reaction conditions on the synthesis of quinolines from aniline and propanol over the ZnCl<sub>2</sub>/Ni-USY-acid catalyst. (A) ZnCl<sub>2</sub> loading; (B) reaction temperature; (C) molar ratio of propanol/aniline; (D) LHSV of aniline (the reaction is conducted under the conditions of temperature = 410 °C, molar ratio propanol/aniline = 3, LHSV = 0.4 h<sup>-1</sup> and molar ratio of propanol/aniline = 3, one among which is varied for (A)–(D) while the others remaining unchanged).

conversion decreases all through and both the selectivity and yield exhibit a maximum, which appear at 0.6  ${\rm h}^{-1}$  and 0.4  ${\rm h}^{-1},$ respectively. This is because that the increase in LHSV decreases not only the contact time and thus the conversion of aniline over catalyst, but also the possibility of side reactions due to the fast diffusion of products away from the catalyst. However, a too higher LHSV would also lead to an insufficient adsorption and activation of aniline the thus its transformation into quinolines over the catalyst.

From Fig. 5, it is obtained that the optimized reaction conditions for the generation of quinolines from aniline and propanol over the  $ZnCl_2/Ni$ -USY-acid catalyst are:  $ZnCl_2$  loading  $= 10$  wt%; carrier gas  $= H<sub>2</sub>$ ; temperature  $= 410$  °C; molar ratio propanol/aniline = 3; and aniline LHSV =  $0.4 \text{ h}^{-1}$ . Under those optimized reaction conditions, the conversion of aniline and total yield of quinolines are respectively 97.6 mol% and 79.7 mol% at 81.7 mol% total selectivity to quinolines. This provides a 60.6 mol% (54.3 wt%) GC yield of 2-ethyl-3 methylquinoline, according to Table 3.

Fig. 1S† shows the GC-MS chromatogram of the products mixture received from the reaction of aniline and propanol under the above optimized conditions. It is identified that 2-ethyl-3-methylquinoline has been generated as the main product. After separation and purification, the structure of the recovered 2-ethyl-3-methylquinoline is further confirmed by both  ${}^{1}$ H NMR(Fig. 2S†) and  ${}^{13}$ C NMR (Fig. 3S†) characterizations (<sup>1</sup>H NMR:  $\delta$  = 8.03 (d, 1H, J = 8.4 Hz), 7.84 (s, 1H), 7.70  $(d, 1H, J = 8.0 Hz)$ , 7.61 (dd, 1H,  $J = 8.0, 7.2 Hz$ ), 7.44 (dd, 1H,  $J =$ 7.6, 7.2 Hz), 3.02 (q, 2H,  $J = 7.6$  Hz), 2.50 (s, 3H), 1.37 (t, 3H,  $J =$ 7.6 Hz); <sup>13</sup>C NMR:  $\delta = 163.30, 146.67, 135.73, 129.40, 128.52,$ 128.29, 127.33, 126.70, 125.61, 29.51, 19.10, 13.08). Besides, the purity of 2-ethyl-3-methylquinoline is determined as 98.6% from the <sup>1</sup>H NMR characterization and the recovery yield of 2-ethyl-3-methylquinoline after separation and purification is determined as 90.2%, which corresponds to a 44.0 wt% separated yield (see ESI†).

Fig. 6 shows the conversion of aniline as well as the total selectivity to quinolines and the total yield of quinolines as functions of reaction time over the  $ZnCl<sub>2</sub>/Ni$ -USY-acid catalyst under the above optimized conditions. One can see that, within 30 h of reaction, the average level of the conversion of aniline, selectivity to quinolines and yield of quinolines remains at ca. 93%, 82% and 76%, respectively. For the reaction time longer than 30 h, the catalytic performance turns to decrease. This occurs probably due to either the formation of carbon deposition over the catalyst or the gradual loss of active  $ZnCl<sub>2</sub>$  from the catalyst. In any way, the residual conversion of aniline, selectivity to quinolines and yield of quinolines still keep a level of 77%, 65% and 50%, respectively for the reaction time of 65 h. The above result shows that the  $ZnCl<sub>2</sub>/Ni$ -USY-acid possesses an appreciably good stability of catalytic performance.

Table 4 shows the results for the reactions of aniline and  $C_1$ – $C_4$  alcohols (methanol, ethanol, *n*-propanol, iso-propanol, n-butanol, ethylene glycol, 1,2-propanediol and glycerol) over the  $ZnCl<sub>2</sub>/Ni-USY-acid catalyst.$  One can see that, for the reactions of aniline and  $C_1-C_2$  monohydric alcohols, no quinolines but N-alkylaniline (NAA) is generated, and the yield of NAA is as



Fig. 6 Lifespan of  $ZnCl<sub>2</sub>/Ni-USY-acid$  catalyst for the reaction of aniline and propanol to quinolones (reaction condition: carrier gas  $=$  $H_2$ ; molar ratio of *n*-propanol/aniline = 3; LHSV (aniline) = 0.4 h<sup>-1</sup>; temperature  $= 410$  °C).

high as 86.3–88.7%. For the reaction of aniline and polyhydric alcohols or  $C_3 - C_4$  monohydric alcohols, quinolines are generated as the main products, of which the total yield of quinolines is in the range of 42.3–79.7%, while N-alkylaniline is formed as the main byproduct. However, exception to the above regularity occurs for the reaction of aniline and iso-propanol, which leads to the generation of NAA (56.5% yield) as main product and quinolines (1.2% total yield) as minor ones. This probably because the steric effect of isopropyl group is disadvantageous to the dimerization of N-iso-propylaniline, which is the key step for the cyclization. It is identified that the most predominant quinolines generated by employing various alcohols are as follows: 2-ethyl-3-methyl-quinoline (n-propanol), 2-propyl-3 ethyl-quinoline (n-butanol), 2-methyl-quinoline (ethylene glycol), 2-ethyl-3-methyl-quinoline (1,2-propanediol) and quinoline (glycerol). This hints that, while quinoline is formed via the reaction between equimolecular aniline and glycerol, alkylquinolines are formed via the reaction between one molecular aniline and two molecular other proper  $C_2-C_4$  alcohols. The above results also indicate the reaction of aniline and n-propanol, among all other alcohols, has provided the highest total yield of quinolines (79.7%).

From the above results for the catalytic performance evaluation and catalyst characterization, it is obtained that both the total yield of quinolines from the reaction of aniline and n-propanol to quinolines and the relative content of Lewis acid sites exhibit an order  $ZnCl<sub>2</sub>/Ni$ -USY-acid > Ni-USY-acid > USYacid > USY, but the porosity of catalyst displays a different order. Accordingly, the catalytic performance is positively related to the relative content of Lewis acid sites of catalyst.

Besides, for methanol, ethanol or iso-propanol as reactant, NAA is generated as the predominant product, almost without the generation of quinolines. For  $C_2-C_4$  polyhydric alcohols or  $C_3-C_4$  monohydric alcohols as reactant, the quinolines constructed theoretically by one molecular aniline and two





<sup>a</sup> Reaction condition: LHSV (aniline) = 0.4 h<sup>-1</sup>; molar ratio of propanol/aniline = 3; reaction temperature = 410 °C; carrier gas = H<sub>2</sub>. <sup>b</sup> NAA: Nalkylaniline, including NPA and DNPA. <sup>c</sup> Others: other substituted quinolines, such as, 2-ethylquinoline, 2-methyl-3-ethylquinoline.

molecular alcohols are generated as the predominant products, with NAA being generated as the most abundant byproduct. Exception to the above regularity is the case for glycerol as reactant, in which the predominant product quinoline is theoretically constructed by equimolecular aniline glycerol.

It has been reported that the mechanism for the synthesis of quinolines from aniline may involve either unsaturated aldehyde/ketone<sup>20,23,30</sup> or Schiff base/Schiff base dimer<sup>54,55</sup> as the key intermediate. As is well known, a  $C_1 - C_4$  monohydric alcohols very hard to be transformed into the corresponding aldehyde/ ketone but easily dehydrated into the corresponding alkene over solid acid catalyst,<sup>56–58</sup> and the reaction of amine and monohydric alcohol over molecular sieve catalyst has constituted the most important route for the synthesis of N-alkyl compound.<sup>58</sup> Contrastively, it is feasible to convert a  $C_2-C_4$ polyhydric alcohol into the corresponding aldehyde/ketone over solid acid catalyst, while aldehyde/ketone can be further converted into unsaturated aldehyde/ketone via Aldol reaction.<sup>59</sup>

Particularly, glycerol can be dehydrated directly into acrolein at high yield.<sup>23</sup> In addition, the stable Schiff base is usually prepared from the reaction of aniline and aldehyde/ketone.<sup>58</sup>

Basing on the above facts, it is proposed that the reaction of aniline and alcohol to quinolines may follow the mechanisms as described by Scheme 1. For a  $C_2-C_4$  polyhydric alcohol as reactant (Scheme 1A), it is first dehydrated into the corresponding aldehyde, which is further converted into unsaturated aldehyde via Aldol reaction. Then, the unsaturated aldehyde reacts with aniline via Michael addition, followed by cyclization and dehydrogenation-aromatization, and this finally generates quinolines. It is also possible that the aldehyde obtained from a  $C_2-C_4$  polyhydric alcohol first reacts with aniline to generate the corresponding Schiff base. Then, the Schiff base is subjected to dimerization, cyclization and dehydrogenation– aromatization, and this finally generates quinolines. Specially, for glycerol as reactant, it is first directly dehydrated into acrolein, being an unsaturated aldehyde, and then subjected to the



Scheme 1 Reaction pathways for the synthesis of quinolines via the reactions of aniline and polyhydric alcohols (A) or monohydric alcohols (B)

cyclization and dehydrogenation–aromatization, generating quinoline. For a  $C_1-C_4$  alcohol as reactant (Scheme 1B), it first reacts with aniline to generate the corresponding N-alkylaniline, which is subsequently dehydrogenated into Schiff base. Then, the Schiff base is subjected to dimerize, cyclization and dehydrogenation-aromatization, and this finally generates quinolines. It should be addressed that the dimerization of Schiff base involves the generation of carbanion  $Ph = N-R$ , which is impossible for  $R =$  methyl and very unstable for  $R =$ ethyl but relatively stable for larger alkyls. Therefore, only N-alkylanilines are generated from the reactions of aniline and methanol and ethanol, while quinolines can be predominantly formed from the reactions of aniline and *n*-propanol or n-butanol. Besides, there is a large steric effect for the dimerization of N-iso-propylaniline, and this accounts for the very low selectivity to quinolines but high selectivity to N-alkylaniline from the reaction between aniline and iso-propanol. In the above mechanism for the reaction of aniline and alcohol, the presence of Lewis acid sites promotes the adsorption activation of aniline and alcohols and the subsequent various steps, while the presence Bronsted acid sites may cause a few negative effects, e.g., the formation of inactive protonated aniline and the inhibition to the formation of carbanion Ph  $= N-R^{-}$ . Therefore, the employment of the catalyst with high concentration ratio of Lewis acid sites to Bronsted acid sites is largely favorable to the synthesis of quinolines from aniline and alcohol, and this has been evidenced by the above results for catalyst characterization and catalytic performance evaluation. Paper<br>
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## **Experimental**

### Chemicals

All organic reagents for the synthesis of quinolines and inorganic salts for the preparation of catalysts were purchased from commercial companies and had purities higher than 99%. These chemicals were employed as received, without further purification.

Zeolites USY (Si/Al = 10) and HY (Si/Al = 6) as well as amorphous  $SiO_2$ -Al<sub>2</sub>O<sub>3</sub> (Si/Al = 6–9) catalysts were all received from SINOPEC Catalyst Changling Division, and HZSM-5 (Si/Al  $=$  50) were purchased from Shanghai Fuxu Zeolite Company. These catalysts were first calcined at 550  $\degree$ C for 5 h, and then, they were either used directly as catalysts or further subjected to the procedures for the preparation of modified catalysts as described in the subsequent section.

#### Preparation of modified catalyst

Acid treatment of USY. The commercial zeolite USY were first dispersed in a 0.5 M hydrochloric acid aqueous solution, according to a weight ratio liquid/solid  $= 10/1$ , and then refluxing treated at 60 $\degree$ C for 30 min under strong stirring. After that, the suspension was filtrated, washed thoroughly with deionized water and dried at 100 °C. The thus-prepared catalyst was denoted as USY-acid.

Ion exchange of USY-acid. The above-prepared USY-acid powder was further modified respectively by the nitrate salts

of various transition metals via ion-exchange method. In the typical procedure, USY-acid was first dispersed into aqueous solution of a transition metal nitrate salt, according to a weight ratio liquid/solid = 10/1, and then refluxing treated at 90 °C for 6 h under strong stirring. After that, the suspension was filtrated, washed thoroughly with deionized water and dried at 100 °C, followed by calcination at 550 °C for 4 h. The resultant powder was repeatedly subjected to the above procedure for three times, with the fresh salt solution being employed for each time. The thus-prepared catalyst was denoted as M-USYacid ( $M = Fe$ , Co, Ni, Cu and Zn, respectively).

Ion exchange of Y, USY and ZSM-5. As comparisons, a series of M-Y, M-USY and M-ZSM-5 catalysts  $(M = Fe \text{ or } Ni)$  were also prepared by the same ion-exchange method as described above for M-USY-acid.

Wet impregnation of USY. The above-prepared Ni-USY-acid was first impregnated with a calculated amount of 0.1 M  $ZnCl<sub>2</sub>$  aqueous solutions at room temperature for 12 h under vigorous stirring. Then the slurry was dried at 120 °C overnight, followed by calcination at 500  $^{\circ}$ C for 4 h in air. The thusprepared catalyst was denoted as  $ZnCl<sub>2</sub>/M-USY-acid$ , and it contained 5-40 wt% of  $ZnCl<sub>2</sub>$ . Unless otherwise mentioned specially, the content of content of  $ZnCl<sub>2</sub>$  in the  $ZnCl<sub>2</sub>/M-USY$ acid catalyst was 10 wt%.

Deposition-precipitation of  $SiO_2-Al_2O_3$ . A calculated amount of  $Ni(NO<sub>3</sub>)<sub>2</sub>$  and an excess amount of urea were first dissolved in distilled water at the room temperature. Then, a certain amount of  $SiO_2-Al_2O_3$  was dispersed into the above solution under stirring. After that, the resultant slurry was heated to 90  $\degree$ C and refluxed for 3 h under strongly stirring. Finally, the slurry was filtrated room temperature, dried at 120 °C overnight, and followed by calcination at 550 °C for 4 h. The thus-prepared catalyst was denoted as  $Ni/SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub>$ , and it contained 10 wt% of Ni in oxide form.

#### Characterization of catalyst

X-ray diffraction spectroscopy (XRD) was performed with a Bruker D8-Advance X-ray diffractometer. The operation conditions were as follows: Cu target K $\alpha$  ray ( $\lambda = 1.54187$  Å); scanning voltage 40 kV, scanning current 40 mA; scanning speed 0.2 s, scanning step  $0.02^{\circ}$ .

Fourier transform infrared (FT-IR) spectroscopy was recorded on a Varian 3100 spectrometer equipped with a GTGS detector. The catalyst was first mixed with KBr  $(1:100)$  by thoroughly grounding, and then, the mixture was pressed into the sample holder and mounted in the cavity of the spectrometer. Each spectrum was obtained by scanning for 32 times at a resolution of 2  $cm^{-1}$ .

N2-physisorption was performed at liquid nitrogen temperature using a QuantaChrome Autosorb-1 instrument. Before the adsorption measurement, the specimen degassed for 16 h at a temperature of 300 $^{\circ}$ C under a residual pressure lower than 4  $\times$  10<sup>-4</sup> Pa in the degas port of the adsorption apparatus. The specific surface area was calculated by using the multipoint BET equation. The pore volume and area was calculated by using the t-plot micropore analysis method. The pore size and its distribution of microporous were obtained from the  $N_2$ desorption branch employing the SF method, and those of mesoporous were determined from the  $N_2$ -desorption branch by the BJH method.

Ammonia temperature programmed desorption (NH3- TPD) was determined on a Micromeritics AutoChem II 2920 instrument equipped with a thermal conductivity detector (TCD). The catalyst specimen was first degassed in a flow of helium with a flow rate of 50 mL min<sup>-1</sup> at 400 °C for 30 min, followed by cooling to 100 °C. Then,  $NH<sub>3</sub>$  was repeatedly pulse-injected until a saturation adsorption over the specimen had been achieved. After that,  $NH<sub>3</sub>$  was desorbed by heating the specimen from 100 to 800  $\degree$ C at a rate of 10 °C min. During the adsorption and desorption of  $NH<sub>3</sub>$ , the helium flow was retained and its flow rate was maintained constant at 60 mL  $min^{-1}$ .

The metal contents of catalysts were determined over a Varian 240AA atomic absorption spectrometer. The operation conditions were as follows: sample aspiration rate  $=$  $3.0 \text{ mL min}^{-1}$ ; lamp current =  $3.0 \text{ mA}$ ; slit width =  $0.2 \text{ nm}$ ; air flow rate =  $0.8$  mL min<sup>-1</sup> and acetylene flow rate =  $5.0 \text{ mL min}^{-1}$ .

#### Catalytic performance evaluation

Catalytic performance test was carried out in a fixed-bed tubular quartz reactor (i.d. 6 mm, total length 400 mm) at atmospheric pressure. The catalyst (1.0 g) was loaded in the middle of reactor, with the upper space of catalyst bed in the reactor being filled with quartz granulates (20 meshes). The mixture obtained by dissolving aniline into a  $C_1-C_4$  alcohols (methanol, ethanol, n-propanol, iso-propanol, butanol, ethylene glycol, 1,2-propanediol and glycol) at a certain molar ratio (aniline/alcohol  $=$  $1/1-1/5$ ) was employed as feedstock. Firstly, the catalyst was in situ pre-treated at certain reaction temperature ranging from 310 to 430 °C for 1 h in a flow of carrier gas ( $H_2$  or air; 10  $\rm{mL}\,\rm{min}^{-1}$ ). Then, the feedstock was pumped into a preheater to be vaporized at 200 $\degree$ C, at a liquid hourly space velocity (LHSV) ranging from 0.2–0.8  $h^{-1}$ . Then, the feedstock was introduced in to the reactor at a pre-set temperature, with the carrier gas being kept. After the reaction had been run stably for 2 h, the products were collected by a condenser which was placed in an ice-water bath.

The composition of products mixture was analysed by a Varian CP-3800/Saturn 2200 gas chromatography-mass spectrometry (GC-MS). Two CP8944 capillary columns (VF-5, 30 m  $\times$ 0.25 mm  $\times$  0.25 µm) were respectively connected to mass detector and flame ionization detector (FID) for the quantitative and qualitative identifications.

Basing on the converted aniline, the conversion of aniline  $(\chi)$ , selectivity to component *i*  $(S_i)$  and yield of quinolines  $(Y_{OS})$ were respectively calculated as follows:

Conversion of aniline (%):

$$
\chi(\%) = \frac{N_{\text{aniline in feed}} - N_{\text{aniline in products mixture}}}{N_{\text{aniline in feed}}} \times 100
$$

Selectivity to component  $i$  (%):

$$
S_i\ \left(^{0}\!\!/_{\!0}\right)=\frac{N_i}{\sum N_i}\times 100
$$

Yield of component  $i$  (%):

$$
Y_{\rm QS}~(^{0}\!/_{0}) = \chi \times S_{\rm QS} \times 100,
$$

where  $N_{\text{aniline in feed}}$  and  $N_{\text{aniline in products mixture}}$  referred to the moles of aniline in the feed and in the products mixture,  $N_i$  to the moles of component i in the products mixture and  $S_{OS}$  to the total selectivity to quinolines, respectively.

# Conclusions

The novel route for the synthesis of quinolines from the reaction of aniline and a  $C_1-C_4$  alcohol under gas-phase reaction condition over heterogeneous catalyst had been developed in this work. It was identified that, for the reaction of aniline and propanol, the USY zeolite-based catalysts possessed obviously higher performances, relative to other catalysts. Over various modified USY catalysts, both the total yield of quinolines and the concentration ratio for Lewis acid sites to Bronsted acid sites exhibited an order USY < USY-acid < Ni-USY-acid <  $ZnCl<sub>2</sub>/$ Ni-USY-acid. Under the optimized conditions over the  $ZnCl<sub>2</sub>/Ni-$ USY-acid catalyst, the reaction of aniline and most of alcohols provided a 42.3–79.7% total yield of quinolines, however, the reaction of aniline and methanol, ethanol or iso-propanol predominantly generated N-alkylanilines. It was proposed that the mechanism for the formation of quinolines from aniline and alcohol involved mainly Schiff base as the key intermediate, while that involved unsaturated aldehyde also could not be excluded. BSC Advances Articles. The method of micropromas were obtained from the R<sub>1</sub> Selectivity to component  $f(\theta) = \sum_{N} \lambda_N \times 100$ <br>
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# Conflicts of interest

There are no conflicts of interest.

# Acknowledgements

We are grateful to the financial supports from the project #21376068 for National Natural Science Foundation of China (NSFC), Program for Lotus Scholar in Hunan Province, and also, Program for New Century Excellent Talents in University.

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