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Highly Efficient and Selective *N*-alkylation of Amines with Alcohols Catalyzed by *in-situ* Rehydrated Titanium Hydroxide

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Abstract

Catalytic *N*-alkylation of amines by alcohols to produce desired amines is an important catalytic reaction in industry. Various noble metal-based homogeneous and heterogeneous catalysts have been reported for this process. Development of cheap non-noble metal heterogeneous catalysts for *N*-alkylation reaction would be highly desirable. Hereby, we propose the *N*-alkylation of amines by alcohols over a cheap and efficient heterogeneous catalyst - titanium hydroxide. The catalyst provides the selectivity higher than 90% to secondary amines for functionalized aromatic, aliphatic alcohols and amines at high catalytic activity and stability. Mild Brönsted acidity formed by continuous rehydration of Lewis acidity excludes side reactions and deactivation by adsorbed species. The mechanism of the reaction involves dehydration of alcohols to ethers with subsequent C-O bond cleavage by amine with formation of secondary amine and recovery of alcohol.

Keywords: *N*-alkylation; Amines; Alcohols; Heterogeneous; Titanium hydroxide; Alkoxides

Introduction

Complex amines are very important intermediates in chemical industry and life science. They are widely used in the production of agrochemicals, pharmaceuticals, organic dyes, detergents, fabric softeners, surfactants, corrosion inhibitors, lubricants, polymers and so on [1,2,3]. Due to the widespread use of these products, a number of synthesis routes have been developed based on classic nucleophilic substitution, like Buchwald-Hartwig [4], Ullmann reactions [5], and hydroamination [6,7,8]. Conventionally, *N*-alkyl amines are synthesized using alkylating agents, such as alkyl halides, which is environmentally unfriendly due to the toxic nature of halides and production of large amount of inorganic salts as waste. The selectivity to desired amines is generally low due to further over-alkylation [9,10,11].

In the last few decades, three main strategies have been applied for amination of alcohols: (i) "Borrowing Hydrogen" methodology using either noble metal based Ru-[12], Pd- [13], Ir- [14], Rh- [15], and Pt- [16] complexes, and more recently non-noble metal Mn- [17,18], Co- [19,20,21], Ni- [22,23,24,25] and Fe- [26] complexes. Also various of heterogeneous catalysts [27,28,29,30,31,32,33,34] have been used. (ii) Tsuji-Trost type reactions for allylic alcohols using Pd [35] or Ni [36] complexes. (iii) Lewis-acid catalyzed reactions using a variety of salts and ligands [37,38]. Among the best performing catalysts, different metal triflates and triflimides, including the parent triflic acid (HOTf) were found active for synthesis of secondary amines through *N*-alkylation of amines with alcohols based on S_N1 or S_N2 mechanisms [39]. However, these processes have several disadvantages due to use of expensive non-recoverable catalysts, difficulties in catalyst-product separation and indispensable use of additives or co-catalysts such as inorganic or organic bases and stabilizing ligands [40]. Thus, development of easily recoverable and recyclable

non-noble metal based heterogeneous catalysts for *N*-alkylation system is still a challenging topic.

Herein, we firstly report the selective *N*-alkylation of amines by alcohols in the absence of hydrogen, bases and organic ligands under ambient pressure of N_2 by an easily prepared transition metal- and solid acid-based heterogeneous catalyst such as titanium hydroxide. For comparison, a number of metal oxides and hydroxides have been tested in the *N*-alkylation process. Interestingly, titanium hydroxide exhibits the highest activity and selectivity towards the desired amine. In addition, the titanium hydroxide catalyst provides high catalyst stability and good substrate tolerance for amination of different alcohols and substituted amines under optimized conditions. To the best of our knowledge, this new efficient heterogeneous catalyst provides a green route for selective synthesis of amines.

In terms of catalytic mechanism, it is generally accepted that substitution of OH-groups with N nucleophiles proceeds via a S_N1 or S_N2 type mechanism, in the presence of hard Lewis or Brönsted acids [41]. In this work, we show that N-alkylation process of alcohols involves intermediate formation of ethers, which could be a kinetic relevant step of the whole reaction. The high yield of secondary amine can be attributed to the easier C-O bond cleavage in the ether by amines based on effect of mild Brönsted acidity formed by adsorbed water over Lewis acid sites.

Experimental Section

Catalyst preparation

Titanium hydroxide was prepared through the hydrolysis of titanium isopropoxide (99%, supplied by J&K) under neutral condition. 200 ml of distilled water was heated up to 60 °C and 10 mL titanium isopropoxide was added dropwise using a peristaltic pump (0.05 mL/min) under continuous stirring (500 rpm) for almost 10 h. Then the

obtained white suspension was centrifuged and washed with ethanol and distilled water for several times followed by drying in vacuum at 80 °C overnight. The obtained sample was denoted as TiOH. The hydrolysis of isopropoxide in hot water is an easy and efficient method to produce amorphous titanium hydroxide with high purity (no NH₃ or Cl⁻) and surface area. This sample has been further calcined at 180 °C and 400 °C for 2 h to produce dehydrated samples denoted as TiOH-180 and TiOH-400, respectively.

The hydroxides of Al (Al-OH) and Zr (Zr-OH) have been prepared in the similar way using alkoxides of aluminium t-butoxide, aluminium isopropoxide, zirconium butoxide and zirconium ethoxide. For comparison, different metal oxides and commercial zeolite ZSM-5 were also used for this reaction as references. The hydrolyzed AlOH sample was calcined under 180 °C, which was denoted as AlOH-180.

Catalytic test

Typical procedure for *N*-alkylation of amines with alcohols was carried out in a 30 mL stainless steel autoclave geared with a pressure gauge and a safety rupture disk. In a typical experiment, the reactor was charged with 4.62 mmol benzyl alcohol, 21.5 mmol aniline and 100 mg of catalyst. 2 g xylene, THF or water was used as a solvent. Then, the reactor was purged with N₂ several times to remove the air. Finally, the reactor was placed on a hot plate equipped with a magnetic stirrer (500 rpm) at 50 °C-180 °C for 0.5-48 h. After the reaction, the reactor was cooled down to room temperature, and the mixture was filtrated and analyzed using an Agilent 7890 GC equipped with a HP-5 capillary column using biphenyl as the internal standard for calculation of alcohol conversion and product yields. After each amination test, the catalyst was washed with ethanol and separated by centrifugation several times and

dried at 80 °C for 10 h for further test of the recyclability. The used catalyst after reaction was denoted as TiOH-80-AR. For the rehydration of the dehydrated catalysts TiOH-180, water was added into the reactor before catalytic test. For characterization, the catalyst was suspended in distilled water overnight followed by drying under vacuum at 80 °C before test. The corresponding rehydrated samples were denoted as TiOH-180-Re, TiOH-400-Re, TiO₂-Re and AlOH-180-Re.

Characterization methods

The powder X-ray diffraction (XRD) patterns were obtained using an X-ray diffractometer (D5000, Siemens) using Cu K α radiation ($\lambda = 0.15418$ nm). Scans were recorded in the 2 θ range of 10-80° using a step size of 0.02° and a step time of 5 s.

The BET specific surface areas and pore size were measured from the N_2 adsorption-desorption isotherms at 77 K on an automated gas sorption analyzer Micromeritics ASAP 2010.

Thermogravimetric analysis (TGA) was carried out in air flow at the temperature range of 20-900 °C with a heating rate of 5 °C ·min⁻¹ on a Mettler Toledo SMP/PF7458/MET/600W instrument.

The FTIR (Perkin-Elmer 1720) spectra of the titanium hydroxide catalysts were measured in the range from 4000 to 1200 cm⁻¹ to trace the variation of surface hydroxyl group of the catalysts. To investigate the nature of acid sites in the catalyst, the pyridine adsorption FTIR technique was employed and spectra were recorded in the range of 1700-1400 cm⁻¹. The self-supported wafers of all the catalysts were prepared by a pellet press instrument and the samples were degassed at 100 °C for 30 minutes followed by saturation with pyridine as reported elsewhere [42]. To explain the mechanism of alcohol dehydration over titanium hydroxide, *in-situ* FTIR was

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performed through heating the pellet from 25 to 180 °C (step 25 °C) in benzyl alcohol vapors (20 Torr).

The NH₃ temperature programmed desorption (NH₃-TPD) was carried out on a Micromeritics AutoChem 2920 with a thermal conductivity detector (TCD). In all the experiments, ~50 mg of sample was calcined at 100 °C for 1 h in helium gas flow (20 mL (STP)/min) and then cooled down to 40 °C followed by saturation with 10% ammonia-helium for 1 h. Then the sample was flushed with helium for 1 h to remove the surface physisorbed ammonia. Finally, the desorption profiles were recorded in the temperature range of 40 °C-800 °C with a heating rate of 5 °C min⁻¹.

Results and Discussion

Catalysis over oxides and hydroxides

The additive-free *N*-alkylation reaction of benzyl alcohol (1) with aniline (2) to give corresponding secondary amine *N*-phenylbenzylamine (3) has been used as a model reaction over different non-noble metal hydroxide, oxide and zeolite catalysts (**Table 1**). The product has been identified by GC-MS, ¹H-NMR and ¹³C-NMR analysis (**Figure S1, S2, S3, SI**). The blank test without catalyst did not show formation of desired amine (entry 1). The titanium hydroxide obtained by hydrolysis of titanium isopropoxide showed the highest catalytic activity and gave the desired secondary amine *N*-phenylbenzylamine (3) with the 99.6% yield (entry 2) without formation of secondary imine. Only traces of dibenzyl ether have been detected as byproduct. The product yield was slightly lower for the reaction performed without solvent due to easier formation of tertiary amine through condensation reaction (entry 3). It is interesting to note that the reaction can be performed in aqueous phase (entry **5**), however, organic solvents provide the highest yield of the product (entry **4**). The reaction in air atmosphere provides a bit lower yield than that under N₂ mainly due to

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the partial oxidation of benzyl alcohol to benzoic acid (entry 6). Hydroxides of Al and Zr prepared by hydrolysis of different types of alkoxides could afford the maximum yield of *N*-phenylbenzylamine of only 16.9% (entry 7-10). Additionally, different metal oxides were also tested for *N*-alkylation reaction. The yield was lower than 2% for TiO₂, CeO₂, ZrO₂, MgO, SiO₂ and Al₂O₃ (entry 11-16). The alkylation yield for commercial zeolite ZSM-5 was only 25% (entry 17).

Titanium hydroxide (TiOH-80) was the most active catalyst. The reaction temperature dependence of the secondary amine yield in the *N*-alkylation of benzyl alcohol with aniline on this catalyst was further investigated (**Figure S4, SI**). Below 100 °C, there was nearly no activity in this reaction. The conversion gradually increases as the temperature increases from 100 to 180 °C reaching full conversion. The apparent activation energy of the reaction corresponds to 66.4 kJ/mol (**Figure 1a**). The time course of the secondary amine synthesis over titanium hydroxide (TiOH-80) at 180 °C was investigated (**Figure 2**). The reaction proceeded smoothly to afford the corresponding secondary amine with 99% yield in 15 h. Dibenzyl ether was observed at the initial stage and its yield slowly decreased with time, suggesting the dibenzyl ether could be an intermediate. The reaction seems to proceed in two consecutive steps: 1) transformation of benzyl alcohol to dibenzyl ether and 2) secondary amine formation from dibenzyl ether with aniline.

Characterization

To identify, whether the high performance of secondary amine synthesis comes from the suitable acidic sites formed from surface hydroxyl group of the catalyst, titanium hydroxide obtained by hydrolysis was calcined at different temperatures. The TGA analysis (**Figure 3a**) shows that titanium hydroxide exhibited weight loss in two main steps. The initial weight loss below 250 °C can be assigned to desorption of Page 9 of 42

ACS Catalysis

surface water. The weight loss between 250 °C and 450 °C could be due to the removal of surface hydroxyl groups through dehydroxylation process [43]. No obvious weight loss was detected for TiOH-400 sample. This indicates that high crystalline TiO₂ phase was formed after calcination at 400 °C. The catalyst TiOH-80-AR after reaction showed similar weight loss compared to the fresh catalyst. This indicates similar content of hydroxyl groups and rehydration of the catalyst during reaction. The state of the material during calcination at different temperatures has been studied by XRD. The parent sample and sample after calcination at 180 °C are amorphous, the presence of anatase was detected after calcination at 400 °C [44]. XRD showed that the catalyst was still amorphous after reaction (**Figure 3b**).

FTIR has provided further information about the concentration and acidity of surface hydroxyl groups in titanium hydroxide. As shown in **Figure 4a**, the titanium hydroxide without calcination exhibited a broad peak at 3600 cm⁻¹ and a peak at 1626 cm⁻¹ due to the stretching vibration of hydroxyl groups interacting with adsorbed water and bending vibration of hydroxyl groups of adsorbed water, respectively [45]. As the calcination temperature increased, the intensity of the peak at 1626 cm⁻¹ decreased and finally disappeared at 400 °C with significant narrowing of the peak at 3600 cm⁻¹. It indicates desorption of water and significant decrease in the amount of hydroxyl groups during dehydration of hydroxide and its transformation to anatase phase.

To identify the acidic properties of the surface hydroxyl groups of the catalyst, FTIR spectra of pyridine adsorbed on titanium hydroxide calcined at different temperatures were recorded (**Figure 4b**). The interaction of pyridine with Lewis acid sites gives rise to the specific bending vibration bands of pyridine ring at 1448 cm⁻¹, 1575 cm⁻¹ and 1605 cm⁻¹ [42]. The peak located at 1540 cm⁻¹ can be assigned to the

ACS Catalysis

 interaction of pyridine with Brönsted acid sites. It can be observed that an increase in the calcination temperature of titanium hydroxide to 180 °C leads to decrease in the intensity of all the peaks related to Lewis acid sites and nearly disappearance of Brönsted acid sites [43]. Further increase in the calcination temperature to 400 °C results in disappearance of all adsorbed pyridine species over the catalyst, which means nearly no acidic sites left on the catalyst. It is interesting to note that the process of dehydration is reversible and treatment of TiOH-180 in water leads to recovery of adsorbed pyridine species corresponding to Brönsted and Lewis acid sites (Figure 4b). However, rehydration process did not lead to acidity generation over the crystalline sample TiOH-400, which could be explained by high stability of anatase structure. Thus, defected labile nature of titanium hydroxide provides both Brönsted and Lewis acid sites, which disappear during dehydration procedure due to the process of healing of defected sites (Figure 5). Indeed, the formation of Brönsted and Lewis acid sites can be proposed as reaction of protonation of hydroxyl groups with subsequent dehydration. The easy rehydration process provide constant presence of both types of acid sites in the catalyst during reaction (Figure 5).

To further identify the amount and strength of the acid sites, temperature programmed desorption of ammonia (NH₃-TPD) has been measured as shown in **Figure 6**. The highest amount of acid sites is observed over pristine titanium hydroxide TiOH-80 with the broad desorption peak at 168 °C and narrow intensive peak at 312 °C corresponding to weak and intermediate acid sites, respectively [46]. Calcination of titanium hydroxide at 180 °C leads to nearly disappearance of intermediate acidity with the weak acidity peak shift to lower temperatures (100 °C). Combined with the results of Py-FTIR, it indicates that high temperature peak (intermediate acidity) can be assigned to desorption of pyridine from Brönsted acidic

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sites and low temperature peak (weak acidity) to desorption of pyridine from Lewis acidic sites. The sample further calcined at 400 °C did not show any obvious NH₃ desorption.

The NH₃-TPD results for titanium hydroxide were compared with those obtained for aluminum hydroxide and zeolites. Aluminum hydroxide also shows the presence of weak and intermediate acid sites. The broad desorption peaks at 220 °C and 420 °C for ZSM-5 can be respectively assigned to intermediate and strong acidities. The amount of acid sites for all the samples calculated through integration and calibration is listed in **Table S1**, **SI**.

The samples pre-treated at different temperatures have been additionally tested in the *N*-alkylation reaction. The initial reaction rate for different samples is shown in **Figure 7**. The initial reaction rate for the sample TiOH-80 (**Figure 1b**, **Figure 7**) was 12.8 mmol/g·h, which was nearly twice higher than that of dehydrated sample TiOH-180 (6.84 mmol/g·h). Calcination at 400 °C almost totally deactivates the TiOH catalyst in *N*-alkylation reaction. It means that anatase structure of TiO₂ without any significant amount of acid sites is not active in *N*-alkylation of benzyl alcohol with aniline. The catalytic test over commercial TiO₂ catalyst further confirms this conclusion (**Table 1**). It is interesting to note that the sample calcined at 180 °C with low content of acid sites (**Table S1, SI**) still provides significant activity in amination reaction. This result can be explained by rehydration of amorphous titanium oxide by water, which produces new acid sites for the reaction. Indeed, the structure of amorphous oxide is very labile and interaction of Brönsted acidity

Crystalline anatase TiO_2 cannot be rehydrated to form acidity during reaction. This suggestion has been further proven by addition of extra water before amination

test with the dehydrated catalysts (Figure 7). The reaction rate (12.55 mmol/g·h) for amorphous titanium hydroxide (TiOH-180) was almost recovered to the initial state after addition of water. We could propose reversible process of dehydration and rehydration for titanium hydroxide during reaction, which provides permanent acidity for amination reaction [47,48] (Figure 5).

To investigate the structure-performance correlation, BET analysis of all catalysts has been performed. There is no obvious relationship between reaction activity and surface area or pore structure of the catalysts (**Table S2, SI**). Indeed, high surface area of hydroxides AlOH and ZrOH or oxides Al₂O₃ and SiO₂ does not provide high activity of these catalysts in amination.

Analysis of the structure-catalytic performance relationship shows good correlation between intermediate acidity measured by NH₃ desorption in the range 200-400 °C and reaction rate in amination over different catalysts (Figure S5, SI). It is obvious that the reaction rate increases with increase in the intermediate acidity. However, high content of intermediate acid sites over AI-OH does not lead to high activity in amination. The possible explanation of this effect could be low rehydration of AI-OH at the reaction conditions. Dehydration of AI-OH at 180 °C leads to significantly lower activity in comparison with the parent catalyst AI-OH (Figure S6, SI). Rehydration of AIOH-180 does not show significant increase of the activity in comparison with the rehydration of TiOH-180. This fact could be explained by less labile structure of AI hydroxide in terms of reversible hydration-dehydration process. The activity recovery for the rehydrated sample TiOH-180 was mainly attributed to the continuous regeneration of intermediate Brönsted and Lewis acidic sites, (Figure 5). The lower activity for ZSM-5 was partially because of quasi-irreversible adsorption of aniline on the zeolite strong acid sites under the reaction conditions.

Page 13 of 42

ACS Catalysis

Consequently, we could assume that weak and strong acidities would not play a determining role in amination reaction. Titanium hydroxide with optimum intermediate acidity and able to be rehydrated shows an exceptional catalytic performance for the *N*-alkylation reactions.

Mechanism

Among various strategies for amines synthesis, amination of alcohols using a borrowing hydrogen methodology (or hydrogen auto-transfer process) has been regarded as a highly atom-efficiency choice [49,50,51]. In this approach (**Figure 8 (a)**) alcohol was first dehydrogenated to the corresponding aldehyde with assistance of noble metals like Ru, Pt, Pd, Ir [52,53,54,55]. Then the aldehyde was nucleophilically attacked by an amine to form secondary imine with the eliminating of water as the only byproduct. Finally, imine was hydrogenated to the desired secondary amine. During the whole process, additional hydrogen is usually added for inhibition of the catalyst deactivation and fast hydrogenation step to suppress secondary processes.

In our case, the non-metallic acidic heterogeneous catalyst such as titanium hydroxide without dehydrogenation/ hydrogenation functions was found to be very efficient for the direct *N*-alkylation of benzyl alcohol with aniline to give desired secondary amine. A possible reaction mechanism in this case can be based on S_N2 mechanism of direct nucleophilic attack of α -carbon atom by N of aniline [56,57].The effect of the ratio of aniline to benzyl alcohol from 0 to 10 on the reaction selectivity is displayed in **Figure 9a**. It is interesting and obvious to note the presence of dibenzyl ether, which might be produced by dehydration of benzyl alcohol. The intermolecular dehydration of benzyl alcohol catalyzed by acid sites of metal oxides like alumina has been observed earlier [58,59,60]. Surprisingly, in the presence of

larger amounts of aniline (amine to alcohol ratio >1), the dehydration of benzyl alcohol is completely suppressed. There are two possible explanations of this effect:

- acid sites of titanium hydroxide are responsible for dehydration of alcohol and amination of alcohol. The presence of excess of aniline suppresses the first reaction;
- 2) ether is the primary product of the reaction. The ammonolysis of the ether by aniline proceeds much faster than that of the relevant alcohol.

To provide experimental evidence for the second hypothesis, the catalytic reaction of dibenzyl ether with aniline over titanium hydroxide was investigated (Figure 9 (b)). No product was formed without catalyst. TiOH-80 clearly catalyzed the ammonolysis reaction of dibenzyl ether with high initial reaction rate (80 mmol/g·h), which was about 6 times higher than that of the whole N-alkylation reaction (12.8 mmol/g·h) (Figure 1b, Figure 6). It demonstrates that the ammonolysis of dibenzyl ether through C-O bond cleavage [61,62,63,64] to the desired secondary amine proceeds much faster than the relevant reaction of benzyl alcohol. This indicates that dehydration of alcohol could be the limiting step of the whole *N*-alkylation reaction (Figure 8b). Water was the only byproduct during the whole reaction process. The product yield for the ammonolysis reaction dramatically decreased from 86 % to nearly 0 as the pretreatment temperature increased from 180 to 400 °C (inset of Figure 9b). It indicates that the ammonolysis of dibenzyl ether requires acid sites as well, which is consistent with the previous report [65]. However, the reaction is less sensitive to amount of acid sites in comparison with dehydration of alcohol to ether.

To investigate the alcohol dehydration process on the acid sites of titanium hydroxide, in-situ FTIR experiments were performed (Figure 10, Figure S7, SI).

ACS Catalysis

Adsorption of benzyl alcohol leads to appearance of the bands at 1230, 1380, 1360, 1447, 1470 and 1494 cm⁻¹. The bands at 1447 and 1494 cm⁻¹ can be assigned to the in-plane C-O-H bending modes [66]. The bands at 1230 and 1470 cm⁻¹ are associated usually with methylene group [67]. The bands at 1360 and 1380 cm⁻¹ can be assigned to aromatic skeletal vibrations [68].

As the temperature increased from room temperature to N-alkylation temperature, the additional bands appears at 1600 cm⁻¹ and 1575 cm⁻¹ (**Figure 10, Figure S7, SI**) These bands can be assigned to bending vibrations of strongly adsorbed water over Lewis acid sites and C=C stretching vibrations in benzyl ether [69]. Additionally the band at 3510 cm⁻¹ of OH stretching vibration increases at higher temperature (**Figure S7, SI**). This indicates dehydration of alcohol with formation of water and ether.

The FTIR spectrum of the catalyst after reaction and intensive washing in xylene shows the peaks at 1380 and 1360 cm⁻¹ similar to FTIR spectra after benzyl alcohol adsorption (**Figure 11**). In order to verify if these species could be intermediates during catalytic reaction, the catalyst after reaction has been treated at the reaction conditions in pure xylene. The liquid phase after reaction contained 0.33 mmol (3.3 mmol ether/g_p) of dibenzyl ether which was the only product of the reaction (**Figure S8**, **SI**). The amount of the product corresponds well to the amount of acidic active sites determined by TPD of NH₃ (**Table S1**, **SI**). FTIR analysis shows that surface species have disappeared after reaction (**Figure 11**). In order to identify the localization of these species, Py has been adsorbed over the catalyst before and after treatment in solvent (**Figure 11**). It is interesting to note that the peak attributed to adsorption of Py over Lewis acid sites decreased significantly after reaction but has been totally recovered after treatment in xylene. The possible explanation of this effect could be formation of surface alkoxide species by interaction of alcohol with

ACS Catalysis

Lewis acid sites (Figure 11). The presence of intermediate surface alkoxides have been observed earlier over alumina [70]. FTIR analysis of low wavenumber region (Figure 11) of the catalyst after reaction (TiOH-80-AR) shows intensive bands at 1126, 1070 and 1041 cm⁻¹, which has been earlier assigned to vibration modes of Ti-O-C species in alkoxides [71].

The kinetic analysis of the reaction of benzyl alcohol dehydration to benzyl ether shows that it corresponds to the second order similar to the whole N-alkylation reaction (Figure S9, SI).

Based on the catalytic and characterization results discussed above, a possible reaction mechanism of ether formation on acidic titanium hydroxide as key intermediate for further ammonolysis to *N*-alkylation product was proposed in **Figure 12**. The alcohol reacts with Lewis acid sites over the TiOH catalyst with formation of surface alkoxides. Protonation of oxygen of alkoxide species by hydroxyl group with SN2 attack of α -carbon by oxygen of another alkoxide results in formation of ether and partial hydration of the catalyst. Subsequent dehydration leads to regeneration of the active sites. Thus, synergetic effect of mild Brönsted and Lewis acid sites over titanium hydroxide for activation of alcohol able to transform it to ether by dehydration is necessary. These properties make titanium hydroxide unique for this reaction.

Adsorption of pyridine over other catalysts confirms this assumption. Oxides containing mainly Lewis acid sites and zeolite with Brönsted acidity provide low activity in *N*-alkylation reaction (**Table S3, SI**). Presence of both Brönsted and Lewis acid sites can be observed also over other hydroxides like Al-OH, however, catalyst cannot be rehydrated in-situ during reaction loosing activity in time (**Figure S6, SI**).

ACS Catalysis

 Thus, amount, nature and regenerability of acid sites are the key factors affecting activity of the catalyst in N-alkylation reaction.

To investigate the stability and reusability of titanium hydroxide, the used catalyst was washed with ethanol and separated from the liquid phase by centrifugation for several cycles and dried at 80 °C for 10 h. As shown in **Figure 13**, the titanium hydroxide could be used for at least three catalytic cycles without obvious change in reaction rate and amine yield, indicating good stability and recyclability for titanium hydroxide.

Substrate scope

The scope of the present titanium hydroxide system with regard to alkylation of various amines and alcohols was also examined. **Table 2** shows that the catalyst can be used for selective conversion with aniline of structurally diverse aromatic and biomass based alcohols to the secondary amines (entry 1-5). The *o*-methyl benzyl alcohol (entry 4) was less reactive, which could be due to the steric hindrance effect of the methyl group at the *o*-position [34]. Aniline derivatives bearing an electron-donating or electron-withdrawing group also reacted with benzyl alcohol to afford the corresponding secondary amines with the yields higher than 90 % (entry 6-9).

Aliphatic amines (butylamine and dibutylamine) can be also used for the synthesis of secondary and tertiary amines by reaction with benzyl alcohol, respectively (entry 10 and 11). However, the yield of the product in this case is lower than in the case of aromatic amines (<50 %) with significant contribution of ethers in the products due to the lower reactivity of aliphatic amine. The reaction conditions have been modified: the time of the test, amount of the catalyst were increased, while

the loading of reagents were decreased (entry 12 and 13). Higher yield of the products comparable to the aromatic amines ($\approx 90\%$) was obtained.

Titanium hydroxide provides high selectivity at reasonable conversions for the reaction of octanol, tetrahydrofurfuryl alcohol and isopropanol with aniline (entry 14-16). There is still a significant contribution of ether but the yield of the target product can be easily improved by increase of the contact time as it has been demonstrated earlier (entry 12 and 13).

Titanium hydroxide also provides reasonable yield of secondary aliphatic amine through *N*-alkylation of aliphatic alcohols (primary and secondary) with aliphatic amines (entry 17-20). The main difference in comparison with aromatic alcohols and amines is higher contribution of ether in the case of aliphatic amines and alcohols (**Table 2**). Note that variation of the product yields with time of aliphatic amine remains consistent with the hypothesis about the role of ether as an intermediate of alcohol ammonolysis. Indeed, *N*-alkylation of isopropanol with aniline demonstrate continuous increase of the yield of the product and a volcano curve for the di-isopropyl ether as a function of time (**Figure S10, SI**). Thus, unique structure of titanium hydroxide provides efficient amination of aromatic and aliphatic alcohols by aromatic and aliphatic amines toward secondary and tertiary amines.

Conclusions

In summary, we have demonstrate that a non-noble metal solid titanium hydroxide acts as a stable, reusable and efficient heterogeneous catalyst for *N*-alkylation of amines with alcohols to secondary amines without hydrogen, bases or organic ligands promotion. The conversion and selectivity of titanium hydroxide were much higher than other hydroxides, metal oxides and zeolite due to the availability of mild

 intermediate Brönsted and Lewis acidities formed by rehydration-dehydration processes. The high yield for direct secondary amine synthesis can be attributed to the acidic catalytic intermolecular dehydration of alcohol to form corresponding ether followed by easier ether ammonolysis.

Supporting Information

GC-MS, ¹H NMR, ¹³C NMR, FTIR, reaction activity, catalytic reaction rate, kinetic analysis, ether formation, acidity of the catalysts and BET analysis.

This information is available free of charge on the ACS Publications website.

Data availability

All data are available from the authors upon reasonable request.

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ACS Catalysis

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Table 1. Amine synthesis from benzyl alcohol and aniline over various non-noble metal oxides and hydroxides (benzyl alcohol (4.62 mmol), aniline (21.5 mmol), xylene 2 g, catalyst 0.1 g, reaction temperature $180 \circ C$, 15 h, atmospheric N₂)

| Ċ | ^он + | NH ₂ 180 °C, xyler | 15 h, cat ➤ ne (2 g) | N H | + H ₂ O |
|-------|--------------------|-------------------------------------|----------------------------|------------------|--------------------|
| 1 | Catalant | 2 | F 4 | 3 | |
| Entry | Catalyst | Y leid of 5 (%) | Entry | Catalyst | Y leid of 5 (%) |
| | Hydroxide | 25 | | Oxides | |
| 1 | — | 0 | 11 | TiO ₂ | 1.4 |
| 2 | Ti-OH | 99.6 | 12 | MgO | <1 |
| 3 | <i>a</i> Ti-OH | 92.4 | 13 | SiO ₂ | <1 |
| 4 | ^b Ti−OH | 99.2 | 14 | Al_2O_3 | <1 |
| 5 | ^c Ti-OH | 97 | 15 | CeO ₂ | 0 |
| 6 | ^d Ti-OH | 89.8 | 16 | ZrO ₂ | <1 |
| 7 | eAl-OH | 16.9 | | Zeolites | |
| 8 | fAl-OH | 10 | 17 | ZSM-5 | 25 |
| 9 | <i>g</i> Zr-OH | 8.1 | | | |
| 10 | ^h Zr-OH | 12 | | | |

*a*No solvent used; *b*under solvent THF; *c*under water; *d*under air; *b*hydrolysis from aluminium t-butoxide; *f*hydrolysis from aluminium isopropoxide; *g*hydrolysis from zirconium butoxide; *h*hydrolysis from zirconium ethoxide.

57



Figure 1 The Arrhenius plot of secondary amine yield for *N*-alkylation from benzyl alcohol and aniline over titanium hydroxide TiOH-80 (a) and time dependence for *N*-alkylation with catalysts before and after dehydration (b)





Figure 2 Time dependence on amine yield for *N*-alkylation from benzyl alcohol and aniline over titanium hydroxide. (Benzyl alcohol 4.62 mmol, aniline 21.5 mmol, xylene 2 g, catalyst 100 mg, 180 °C, atmospheric N₂.)

а





2 θ (degree)

Figure 3 Characterization of titanium hydroxide catalysts using TG analysis (a) and XRD (b)





Figure 4 FTIR spectra of titanium hydroxide catalysts (a) and Py adsorption over titanium hydroxide catalysts (b)



Figure 5 Proposed process of dehydration and rehydration for titanium hydroxide





Figure 6 NH₃-TPD profiles of different samples



Figure 7 Comparison of the initial reaction rate for titanium hydroxide and calcined samples at different temperature (Benzyl alcohol 4.62 mmol, aniline 21.5 mmol, xylene 2 g, catalyst 100 mg, 15 h, atmospheric N_2).





Figure 8 (a) General hydrogen borrowing mechanism and (b) proposed non metallic mechanism through ether formation.





Figure 9 (a) Products distribution under different amine/alcohol molar ratio. Condition: benzyl alcohol 4.62 mmol, aniline/benzyl alcohol molar ratio 0-9.28, xylene 2 g, catalyst 100 mg, 15 h, atmospheric N_2 . (b) Time dependence for amine formation directly from dibenzyl ether. Condition: dibenzyl ether 5 mmol, aniline 20 mmol, xylene 2 g, catalyst 100 mg, atmospheric N_2 (inset: reaction performed with different catalysts for 1 h).



Figure 10 In-situ FTIR spectra of titanium hydroxide exposed to benzyl alcohol vapor calcined under different temperature.



Figure 11 FTIR spectra of titanium hydroxide (a) and Py adsorption (b) before reaction (TiOH-80), after reaction (TiOH-80-AR) and treatment in xylene (TiOH-80-AR-treated)









Figure 13 Reusability test of *N*-alkylation of benzyl alcohol with aniline using titanium hydroxide. Reaction conditions: alcohols (4.62 mmol), amines (21.5 mmol), xylene 2 g, catalyst 100 mg, reaction temperature 180 °C, 15 h, atmospheric N₂.

| R_1 -OH + R_2 -NH ₂ $\xrightarrow{\text{TiOH-80}}$ R_1 -N- R_2 + H_2 O | | | | | | | | | |
|---|----------|--------------------|----------|------------|--------|---------------|--|--|--|
| Entry | Alcohols | Amines | Products | Conv. % | Selc.% | Ethers (%) | | | |
| 1 | ОН | NH ₂ | N-C | 99.4 | 100 | 0 | | | |
| 2 | СІ | NH ₂ | | 99 | 100 | 0 | | | |
| 3 | ОН | NH ₂ | O-C-N-C | 97 | 98 | 0 | | | |
| ^b 4 | ОН | NH ₂ | N-C | 90 | 91 | 2 | | | |
| 5 | От | NH ₂ | | 100 | 98 | 0 | | | |
| ^b 6 | ОН | NH ₂ | N H | 99 | 95 | 0 | | | |
| 7 | ОН | CI NH ₂ | N CI | 99 | 97 | 0 | | | |
| 8 | ОН | NH ₂ | N-C-O | 100 | 99 | 0 | | | |
| 9 | ОН | NH ₂ | N N N | 100 | 99 | 0 | | | |

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^aReaction conditions: alcohols (4.62 mmol), amines (21.5 mmol), xylene 2 g, catalyst 100 mg, reaction temperature 180 °C, 15 h, atmospheric N₂; ^b24 h; ^c48 h; ^d48 h (alcohols 0.5 mmol, amines 2.3 mmol, xylene 2 g, catalyst 200 mg); ^e72 h; ^f96 h.

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