Highly Regio-Selective Synthesis of $\beta$-Amino Alcohol by Reaction with Aniline and Propylene Carbonate in Self Solvent System over Large Pore Zeolite Catalyst

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Abstract

The nucleophilic ring opening of epoxides with amines is a well known route for the synthesis of $\beta$-amino alcohols. The use of carbonates offers significant advantages over epoxides as they are far less hazardous materials, safe for handling, do not require high-pressure equipment and most notably the possibility of solvent less reactions. In this work, utilization of zeolite as host catalyst in the reaction media for synthesis of $\beta$-amino alcohols without using solvent is reported.

Keywords: $\beta$-Amino Alcohol, Propylene Carbonate, Zeolite, Catalyst, Chemo-Selectivity

1. Introduction

$\beta$-Amino alcohols are used as intermediates in the synthesis of wide range of biologically active natural and synthetic products [1-3] unnatural amino acids [4,5] and asymmetric synthesis as chiral auxiliaries [6]. The nucleophilic ring opening of epoxides with amines group is a well known route for the synthesis of $\beta$-amino alcohols [7-11]. Generally, in the classical method the cleavage of epoxides is achieved by treatment with excess of amine at suitable temperature. However, the lack of appreciable selectivity, the requirements of high temperature and the need of excess amine in the classical methods have led to the necessity for activation of the epoxides so as to increase their susceptibility to nucleophilic attack by amine.

The various methodologies developed for this include the use of alumina [12], metal amides [13-15], metal alkoxides [16], metal triflates [17-19], transition metal halides [20,21], alkali metal perchlorates [22], rare earth metal halides [23-25] and silica under high pressure [26]. However, these methodologies suffer from one or more disadvantages such as long reaction times, elevated temperatures, moderate yield, use of air and/or moisture sensitive catalysts, requirement of stoichiometric amounts of expensive reagents/catalysts, potential rearrangement of allylic alcohols [27] potential hazards in handling pyrophoric/moisture sensitive reagents in the preparation of the catalyst, and in most cases applicable to aromatic amine only as aliphatic amines often deactivate the organometallic catalyst.

Considering the widespread applications of the resultant $\beta$-amino alcohols, we felt that not only a catalyst choice should be one that is easily available and less expensive, less toxic and operatable under environmentally friendly conditions so as to fulfill the triple bottom line philosophy of green chemistry, but also to develop a more efficient approach for the synthesis of $\beta$-amino alcohols that will lead a new technology. The applications of cyclic carbonates in synthesis of fine chemicals are reviewed recently [28]. The use of carbonates offers significant advantages over epoxides as they are far less hazardous materials, safe for handling, do not require high-pressure equipment and most notable thing is the possibility of solvent less reactions with carbonates as it itself act as solvent in reaction medium [28].

Zeolites can be used for fine chemical synthesis, which is highlighted in a review [29,30]. A choice of solvent is necessary for complete conversion and high yield as the products can be desorbed easily from zeolitic pore and solvent should not compete with the reactant in adsorption process [31]. The feasibility of using Na-Y as catalyst for synthesis of amino alcohols from aniline (AA) and propylene carbonate (PC) has been shown earlier; however, detail study on effect of catalysts and reaction
parameters on the regio and stereo selectivity of product was not reported [32]. We describe here a practical approach that is highly regio selective for synthesis of 1-(phenylamino) propan-2-ol (3) from propylene carbonate and aniline using a solvent less system, and zeolite as recyclable heterogeneous catalyst (Scheme 1). The route is thus aligning with the green chemistry approach.

2. Results and Discussion

When aniline and propylene carbonate in a molar ratio of 1:1 was stirred along with Na-Y zeolite (100 mg) in a round bottom flask at 150°C, mono substituted N-alkylated product, 1-(phenylamino) propan-2-ol (Scheme 1) was formed with > 95% selectivity (analyzed by gas chromatographic). The concentration-time plots for aniline (AN) and β-amino alcohol (AA) at 150°C and 180°C respectively is shown in Figure 1. Solvent less system often results in poor yields due to deactivation of active site by reactant and product molecules. The high concentration of reactants and products many times results in slow reaction due to diffusional resistance, however, in our system even 1:1 moles of reactants results in complete conversion with high regio-selectivity of desired product (3) (See Figure 1). Continuous increase in the formation rate of product makes our system highly precious. These observations support that the propylene carbonate can be best used as self solvent in the reaction medium. The Na-Y zeolite as catalyst of the reaction can be reused with fresh aniline and propylene carbonate with a high conversion of aniline ~ 80% and yields of 1-(phenylamino) propan-2-ol ~ 75% even after 5 recycles.

The effect of temperature on the activity and selectivity of the reaction was studied and results are shown in Figure 2. The results show that the 150°C reaction temperature is suitable to achieve high region-selective product. Further increasing reaction temperature increases conversion but the regio-selectivity of product decreases (See also Figure 1). Therefore, for all further studies we used 150°C as reaction temperature and 48 h as reaction period.

A reaction pathway is shown in Scheme 1 (Route I and II), wherein CO₂ is evolved as a by product. Since, the reaction of CO₂ with propylene glycol is known to give propylene carbonate [33,34], the proposed route becomes even more valuable. The transformation of aniline to regio-selective 1-(phenylamino) propan-2-ol is achieved through zeolitic framework. A mechanism involving nucleophilic attack of aniline on sterically less hindered carbon from propylene carbonate by the loss of CO₂ to form active carbonium ion intermediate.

Zeolitic framework plays vital role in formation of chemo-selective 1-(phenylamino) propan-2-ol product through route-I as zeolitic framework has characteristic
feature of shape selectivity, which leads to a selective product based on molecular shape. The zeolite Y has a framework of basic and cationic sites which act as Lewis base and conjugated Lewis acid pairs [35]. It is likely that benzene ring in aniline is adsorbed on the 12R window of Na-Y framework, which is slightly tilted out from the oxygen plane of the 12R window as its Van der Waals radius (Max. ~ 7.6 Å) is too large to fit perfectly into the 12R window [37]. Therefore, NH₂ group in aniline will be above the ring plane formed by the oxygen and may protrude in the super cage where it interacts strongly with the large electric field associated with the carbon-framework of oxygen. The carbonium ion (active intermediate of the reaction formed by route I and II, Scheme 1) formed on basic sites of Na-Y from propylene carbonate attacks the active hydrogen from NH₂ group leading the formation of N-(phenylamino) propan-2-ol. The example of this is shown in Scheme 1.

Increase of Brønsted acidity of the Na-Y zeolite (increase of H⁺ form) decrease the catalytic activity of reaction as shown in Table 1 (Sr.no. 1 - 5), which further supports our view that Lewis acid-base pairs are responsible for a good activity of catalyst. Thus the Lewis acid and Lewis base in Na-Y zeolite together play an important role in achieving high conversion and high regio-selective ring opening reaction between propylene carbonate and aniline. To understand the role of different alkali metal ions, we exchanged the Na-Y with K and Cs ions and the results are shown in Table 1 (Sr.no. 6 - 7). The conversion and selectivity of product decreased by increase in size of alkali cation exchange (Na < K < Cs), this may be because of hindrance for the benzene ring adsorption on 12R window of Y zeolite. By exchanging Cs the basicity of catalyst increases and highly basic character of the catalyst does not seem to favor the reaction. For understanding the role of basicity, few more experiments were conducted using Lewis base and non-zeolite catalysts such as TEA-Br, Na-Aluminates, Mg-Al (3:1) hydrotalcite, MgO, NaCl. Few more experiments were carried out separately using non zeolite catalysts. With these catalysts lower conversion and lower regio-selectivity for the alkylation reaction is observed and results are shown in Table 1, Sr.no. 8 - 12. In contrast to the result with zeolite Na-Y (see Table 1, Sr.no. 1) in which both basic and acidic sites is available. This aspect was further explored in order to quantify and identify the nature of acid and base sites in NaY. The nature of acid and base sites of NaY zeolite catalyst used in the present work was characterized by some of us earlier [38] by using NH₃ and CO₂ TPD as well as by pyridine adsorption followed by FT-IR analysis. The characterization data showed a marked difference in the nature of acidic and basic active sites of NaY zeolite which is revealed by NH₃ and CO₂ TPD. It was observed from TPD analysis that there exist smaller surface density of acidic sites compared to basic sites in NaY (base to acid ratio 6.25 [38]) indicating that Lewis basicity due to framework oxygen atoms seems to be playing a main role in substrate activation, while on the other hand Lewis acidity is essential but seems to be less important in this case. Nevertheless combination of Lewis acid-base properties is preferred for an efficient catalysis. In this context, it may be noted that in aminolysis of epoxides over solid acid catalysts, catalytic activity decreased with increasing basicity of the amines, presumably due to deactivation of acidic sites of catalyst by amine [39].

Further, the reaction of aniline and propylene carbonate carried out over different zeolites (Na-ZSM-5, Na-Mordenite and Na-L) gives different conversion and selectivity of the 1-(phenylamino) propan-2-ol as shown in Table 1, Sr. no. 13 - 15. The aniline in different zeolitic framework adsorb with different molecular orientation. The Na-Y zeolite may be suitable for this reaction as the active NH₂ group of aniline adsorbed in Na-Y is located in super cage and which is easily accessible as an alkylation group formed by propylene carbonate with particular shape leading to regio-selective

Table 1. Activity over zeolites, alkali exchange zeolites and non zeolite catalysts.

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Catalyst</th>
<th>Al/(Al + Si) ratio</th>
<th>Na⁺ form, (%)</th>
<th>Conv., (%)</th>
<th>Yield, (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Na-Y</td>
<td>0.29</td>
<td>100</td>
<td>80.5</td>
<td>79.9</td>
</tr>
<tr>
<td>2</td>
<td>NaH-Y</td>
<td>0.29</td>
<td>82</td>
<td>35.3</td>
<td>33.9</td>
</tr>
<tr>
<td>3</td>
<td>NaH-Y</td>
<td>0.29</td>
<td>45</td>
<td>27.1</td>
<td>25.9</td>
</tr>
<tr>
<td>4</td>
<td>NaH-Y</td>
<td>0.29</td>
<td>10</td>
<td>24.2</td>
<td>21.6</td>
</tr>
<tr>
<td>5</td>
<td>H-Y</td>
<td>0.29</td>
<td>0</td>
<td>23.8</td>
<td>20.1</td>
</tr>
<tr>
<td>6</td>
<td>K-Y</td>
<td>0.29</td>
<td>100</td>
<td>74.4</td>
<td>73.3</td>
</tr>
<tr>
<td>7</td>
<td>Cs-Y</td>
<td>0.29</td>
<td>100</td>
<td>69.2</td>
<td>67.8</td>
</tr>
<tr>
<td>8</td>
<td>Na-Alumin-ate</td>
<td>-</td>
<td>-</td>
<td>42.2</td>
<td>21.4</td>
</tr>
<tr>
<td>9</td>
<td>Mg-Al Hydrotalcite</td>
<td>-</td>
<td>-</td>
<td>41.7</td>
<td>32.8</td>
</tr>
<tr>
<td>10</td>
<td>TEA-Br</td>
<td>-</td>
<td>-</td>
<td>48.3</td>
<td>40.9</td>
</tr>
<tr>
<td>11</td>
<td>MgO</td>
<td>-</td>
<td>-</td>
<td>5.0</td>
<td>4.5</td>
</tr>
<tr>
<td>12</td>
<td>NaCl</td>
<td>-</td>
<td>-</td>
<td>15.1</td>
<td>14.0</td>
</tr>
<tr>
<td>13</td>
<td>Na-MOR</td>
<td>0.15</td>
<td>100</td>
<td>13.4</td>
<td>12.9</td>
</tr>
<tr>
<td>14</td>
<td>Na-L</td>
<td>0.24</td>
<td>100</td>
<td>12.2</td>
<td>11.1</td>
</tr>
<tr>
<td>15</td>
<td>Na-ZSM-5</td>
<td>0.03</td>
<td>100</td>
<td>17.1</td>
<td>16.0</td>
</tr>
</tbody>
</table>

*Temperature: 150°C; Time: 48 h; AN/PC ratio: 1:1 mole: mole.
formation of 1-(phenylamino) propan-2-ol. Consistent with these mechanistic schemes, the overall yield of alkylation is highly dependent on the ring size and substitution pattern of the zeolite framework.

In order to extend the usefulness of this protocol further, optically pure enantiomer (R or S, 98% ee by GLC, and used as received from Aldrich, USA) of propylene carbonate was allowed to react with aniline and the results are represented in Table 2. Reaction of (R or S) 1,2-propylene carbonate (~98% ee) with aniline gave four possible chiral \( \beta \)-amino alcohols illustrated in Table 2. As expected, an excellent selectivity of (2R)-1-(phenylamino) propan-2-ol or (2S)-1-(phenylamino) propan-2-ol is obtained with yields, 96:4 (3:4) in favor of nucleophilic attack of amine at the sterically less hindered methylene carbon of 1,2 propylene carbonate (Table 2 Sr. no. 1 and 2). Two isomers viz. 1-(phenylamino) propan-2-ol (3) showed the characteristic ion peak at m/z \( M^-45 \) due to loss of CH\(_3\)CHOH whereas, 2-(phenylamino) propan-1-ol (4) showed the ion peak at m/z \( M^-31 \) due to loss of CH\(_2\)OH in mass spectra [40]. Thus it was observed that chiral amino alcohol can be synthesized using stable chiral 1,2-propylene carbonates.

3. Conclusions

High activity and high selectivity for synthesis of 1-(phenylamino) propan-2-ol is achieved over recyclable Na-Y zeolite. It has been shown that Lewis acid-base pairs play an important part in activity of a catalyst. The halide free and solvent free reaction of aniline and propylene carbonate is a clean, safe, and reproducible, method that uses less hazardous materials and does not require high pressure equipments usually required in ring opening reaction of epoxide by aniline. The synthesis method proposed here could impose practical value in the common applications for a wide range of amino alcohol derivatives. No operational problems that are foreseen for the large-scale version of this green approach. Synthesis process and technical refinement should further increase the synthetic regio-selectivity of the mono 1-(phenylamino)propan-2-ol product, which of great interest in medicinal, biological active synthetic and natural product, artificial amino acids, and chiral auxiliaries for asymmetric synthesis. Finally, it was also shown that starting from chiral carbonates; chiral \( \beta \)-amino alcohols can be synthesized retaining high ee value.

4. Experimental

4.1. Materials

Aniline and propylene carbonate were purchased from either S.D. fine chemical Ltd. India or Aldrich, USA. Zeolite Na-Y was provided by Südz-Chemie, India, other zeolites such as, K-Y, Cs-Y, Na-MOR, Na-L and Na-ZSM-5 already synthesized and well characterized in our laboratory were used as such [41,42].

4.2. Typical Reaction Procedure

Typical procedure used for production of 1-(phenylamino)propan-2-ol from propylene carbonate and aniline is as follows. In a 20 ml round-bottomed flask equipped...
with a magnetic stirring and reflux condenser was charged with 4.66 g (50.1 mmol) of aniline, 5.21 g (51.1 mmol) of propylene carbonate and 0.10 g Na-Y zeolite as catalyst. The mixture was vigorously stirred at room temperature for 10 min and then mixture was heated at 150°C. After reaction the viscous colorless liquid products were analyzed by gas chromatograph using Hewlett Packard 6890 Series GC equipped with auto sampler instrument, on a HP-5 capillary column (30 m × 320 μm × 0.25 μm film thickness, on a 5% phenyl methyl siloxane stationary phase).

After completion of reaction, reaction mixture was diluted with acetone and filtered (sartorius-393 grade filter paper) to separate the catalyst. More washings of acetone were given to the catalyst to remove adhered organic impurities. Then the catalyst was dried at 120°C for 24 h in vacuum oven, cooled and reused. The above procedure was repeated. This reusability of catalyst was done further five times and the reactant charges according to the amount of catalysts recovered keeping reactant to catalysts ratio same.

4.3. Analytical Procedures

About 300 mg of the reaction crude diluted with acetone was coated on ~3 g column grade silica gel support (Gold Silica 20-40 μm spherical; Catalog # 60-5394-480; Purchased from RediSep® Media, USA) using rotary evaporator. Acetone is slowly evaporated under vacuum till the support changes to a free flowing powder which is than loaded onto the chromatogram instrument and the products were separated by flash chromatography on a 4 g normal phase silica RediSep® column employing n-hexane-ethyl acetate as the eluent with gradient programming. Liquid chromatography was performed using Combi-Flash Companion, supplied by Teledyne ISCO, USA. 1-(phenylamino)propan-2-ol was thus isolated in pure form. Products were characterized and confirmed by GC-MS, IR and 1H and 13C NMR spectroscopy. Regio-selectivity of isomers was determined by GC and GC-MS analysis.

1-(phenylamino)propan-2-ol: 1H NMR (200 MHz, CDCl3): δ = 1.25 (d, J = 6.3 Hz, 3H; -CH3); 2.64 (brs, 2H; NH & OH); 2.98 (dd, J = 8.4 & 12.8 Hz, 1H; -NCH- ); 3.21 (dd, J = 3.4 & 12.8 Hz, 1H; -NCH- ); 3.95 - 4.09 (m, 1H; -OCH- ); 6.64 - 6.67 (m, 3H; -CH, Ar); 7.15 - 7.25 (m, 2H; -CH, Ar). 13C NMR (50 MHz, CDCl3): δ = 20.74 (-CH3); 51.59 (-NCH2); 66.28 (-OCH2); 113.21 (-CH, Ar); 117.83 (-CH2, Ar); 129.23 (-CH, Ar); 148.12 (-C, Ar). IR (film): νmax = 3394 (OH, NH), 2970 (CH), 1602...
Figure 3. (a) Chiral HPLC Spectrum of racemic products (Major: 3a + 3b; minor: 4a + 4b; peak No. 2, 3, 1 and 4, respectively); (b) Chiral HPLC Spectrum of compound 3a (t, 44.625 min); (c) Chiral HPLC Spectrum of compound 3b (t, 57.483 min).
obtain H+ form. The degree of exchange was determined from RT to 540°C at a rate of 3°C/min. The amount of NH4 form of NaY zeolite. (the reported NH4NaY that is formed by adsorption of ammonia on zeolite is synthesized by solid-liquid ion exchange process.)

NH4 bending modes were observed. The absorption at 1490 cm–1 for ammonia, adsorbed on HNaY zeolite, [43] is at 1680 cm –1, 1500 cm –1, 1450 cm –1 and 1700 cm–1, 1640 cm –1, 1425 cm –1 and 1370 cm–1 due to C=C, the location of NH4 cation in zeolite framework is nitrogen ion location close to Si-OH-Al group. Therefore, the extent of Na + exchange was monitored to differ from when NH4 cation that is formed by NH3 adsorption on HNaY and when by ion exchange of NaY zeolite is synthesized by solid-liquid ion exchange process.

One gram of Na-Y zeolite was exchanged with 25 ml, 1M ammonium acetate solution by stirring the mixture at several times, dried at 120°C overnight and calcined (under the flow of air at 540°C for 12 h by increasing the temperature from RT to 540°C at a rate of 3°C/min) to obtain H+ form. The degree of exchange was determined by recording the absorbance intensity of NH4NaY exchanged samples in the range 2000 - 1000 cm –1. Bands at 1700 cm –1, 1640 cm –1, 1425 cm –1 and 1370 cm –1 due to NH bending modes were observed. The absorption at 1370 cm –1 was monitored for the purpose of calculating the amount of NH4 form of NaY zeolite. (the reported absorbance is at 1680 cm –1, 1500 cm –1, 1450 cm –1 and 1490 cm –1 for ammonia, adsorbed on HNaY zeolite, [43] in the present case, however, we believe that NH4NaY zeolite is synthesized by solid-liquid ion exchange process, the location of NH4 cation in zeolite frame-work is much more similar to Na location in original NaY, while NH4NaY form that is formed by adsorption of ammonia on Si-OH-Al groups of HNaY zeolite could have ammonium ion location close to Si-OH-Al group. Therefore, the interaction of NH4 cation with frame-work Al seems to differ from when NH4 cation that is formed by NH3 adsorption on HNaY and when by ion exchange of NaY zeolite). Three samples were thus prepared with degree of H+ form, 18%, 55% and 90% respectively.

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6. References


