Research Article

A Green Route for Sustainable Nanoporous Solid Acid Catalyst Synthesis Using Bio Template and Analysis of Its Progressive Transformation of CO₂

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Abstract: Green chemistry approach is a most important area in the modern chemical world. The nano (meso) porous materials have the specific application in separation, adsorption and catalysis. A green template (Egg white) is used as a bio template for the synthesis of nanoporous material to replace the hazardous templates. It is a bio-degradable template; it does not create any environmental issues. The synthesized material is characterized by various spectroscopic techniques to confirm its structural formation. Based on pore size, the catalyst named as AlSiO₄-14. The synthesized catalyst (AlSiO₄-14) is active at 175°C. This low temperature activity will not produce coke formation at the same time it does not require any regeneration and the catalyst is continuously active for catalytic reactions. The catalytic activity of carbon dioxide decomposition is achieved at lower temperature. The complete carbon dioxide decomposition of AlSiO₄-14 is 43 % is a great impact of the material.

Keywords: Porous material, Green template, AlSiO₄-14, Complete decomposition, Eco – friendly.

1. Introduction

Nanoporous solid acid catalysts are the most interesting materials for the enormous uses in various fields such as adsorption, petro chemical industries, catalysis, catalytic supports, sensors and fuel cells.[1-7] The mesoporous framework containing materials has the pore size between 2 nm to 50 nm.[2,4] The porous framework materials are used to separate different types of molecules from a mixture of molecules. The majority of the porous solid acid materials had the greatest impact on green chemistry applications, however the molecular sieves synthesize method is not under the green chemistry. Most of the conventional templates are used for the synthesis is not eco friendly. In this present work, we have synthesized the nanoporous aluminosilicate molecular sieves using bio template it does not create any corrosion and pollution to the environment. It is an inexpensive template compared to the conventional templates. Conventional templates like tetraethylammonium hydroxide (TEAOH),[9] organic amines,[10,14] hexamethylenimine,[16,17]alkyltrimethylammonium halides(CnTMAX; X ) Cl or
Br),\[^{[10,13]}\] geminisurfactants type (\([C_nH_{2n+1}-N(CH_3)_2-(CH_2)_sN-(CH_2)_2C_{m}H_{2m+1}]Br_2\) (18-12-18)),\[^{[11]}\] Myristyltrimethylammonium bromide (MTMAB),\[^{[19]}\] dodecyltrimethylammonium bromide,\[^{[13]}\] diblock copolymer, triblock copolymer P123 (nonionic surfactant),\[^{[4,18,24]}\] oligomeric surfactants,\[^{[6]}\] Rhodamine B (dye),\[^{[3]}\] fatty alcohol polyoxyethylene ether (PEO surfactants)\[^{[23]}\] are used as a template. In a few synthesis methods NaOH, butanol and carbon black microspheres is used as a co-surfactant. Cetyltrimethyl ammonium bromide (CTABr)\[^{[18,20,21]}\] is a most common structure directing agent for synthesizing a mesoporous molecular sieves. These structure directing agents are chemically harmful. The synthesized nanoporous materials are mainly used for green chemistry applications, but the synthesizing method is not a green process. To overcome these obstacles, egg white is used as a template (green template).

2. Experimental section

2.1 Green-template

Egg white is used as a bio-template for the synthesis of AlSiO₄ material. Egg white is mainly made-up by water and protein. 55% of the egg white is made up of Albumin and remaining proteins are transferring, mucoid, globulin G2, globulin G3, mucin and lysozyme. To avoid hazardous templates, egg white is used as a green bio-template. AlSiO₄ material is synthesized from inexpensive source material & simple method using green template.

2.2 Materials and methods

\(\text{AlCl}_3\) (98% Merck), \(\text{Na}_2\text{SiO}_3\) (meta) (Loba Chemie) and Egg white is used as a template.

In a distinctive synthesis, 13.5 g (0.0135 mmole) of aluminium chloride is suspended in aqueous medium with constant stirring. After that, 28.5 g (0.0285 mmole) of sodium silicate fine particles is included and 25 ml of egg white is mixed with it. Further the reactants is stirred continuously for 30 minutes and kept aside for 24 hours to achieve the product. The obtained material is continuously cleaned with distilled water as well as dehydrated at 120 °C for 2 hours.

2.3 Calcination

To remove the unwanted volatile compounds and templates from the aperture of the as-prepared material, it is calcinated at 600°C with the help of air. During the calcinations, the white material turns into brown and this is due to the fragmentation of template. This indicated that some organic volatile fragments are present inside the pores of the catalyst. After the complete removal of template, the material became white powder.

2.4 Catalytic reactor set up

In CO₂ decomposition, curved catalytic reactor is developed. Transformation of carbon dioxide continuously analyzed by the reactor (Figure 1). The reactor is packed with calcinated material and it is kept under engine oil. The reactor is heated in a hot plate and the temperature is monitored by thermometer. 99.6% of unpolluted carbon dioxide is passed through the inlet and the outlet is sealed by stopcocks.
2.5 Regeneration & reusability of the catalyst

For regenerating the catalyst, the catalysts are taken in a U-shape tube with inlet and outlet containing tube, this tube is fitted in a sigma electric Bunsen and it is heated for a high temperature (above 400 °C). Air is generated from aerator and it is passed though KOH solution for removing the CO₂ and it is connected with silica gel (blue) for removing moisture. The CO₂ and H₂O free air is passed to the catalytic bed for regeneration.

2.6 Characterization techniques

The structural formation and crystallinity of the material is analyzed on Bruker D8 advance Low – angle X – Ray Diffraction (XRD) (Cu – α radiation, λ = 1.5418Å). Symmetric and asymmetric stretching is recorded by JASCO – 410 FT – IR (KBr pellet method). Chemisoft TPx V1.02 is used to record the TPD (NH₃ desorption). Removal of template from the material is recorded from TGA Q500 V20.10. Micrometrics, ASAP 2020 V3.00H is used to evidence the porous nature of the material. The morphology and elements of the material is analyzed with a help of Scanning Electron Micrograph (SEM) of Carl Zeiss EVO 18. Carbon dioxide decomposition products are analyzed with Gas Chromatography (Thermo Fischer / chemito GC 1000) using TCD.

3. Results and Discussion

3.1 Small Angle X-Ray Diffraction

The synthesized material is calcinated at 450 °C for the complete elimination of all the templates from the cavities of the material. The XRD pattern (Figure 2) shown that, the material is thermally stable and crystalline. The formation of nanoporous (meso) \(^{[5,4]}\) is confirmed by the intense small angle peak at 1.1 degree and it is in 100 and 110 plane. This XRD analysis proved that the egg white acting as a template for the formation of nanopores in this material.
3.2 FT – IR Analysis

The broad peaks from 3700 to 3000 cm\(^{-1}\) in the as-prepared material is owing to the occurrence of water molecule and the C-H stretching vibrations between 2000 to 1600 cm\(^{-1}\) is caused by the incidence of template in the cavities after calcination these peaks get disappeared (Figure 3)[25]. The material shows the fundamental vibrations of symmetric stretching and bending vibrations at 790, 430 cm\(^{-1}\) respectively. The distinctive peak at 1086 cm\(^{-1}\) proves the formation of tetrahedral frame work [26-29]. This spectrum confirms that the tetrahedral frame work is stable after calcination at higher temperature.

3.3 Thermal Analysis
From thermogravimetry (Figure 4), the weight change is observed from 50 °C to 950 °C. The first weight loss observed from 100 °C to 220 °C, it is due to the elimination of reactive and non-reactive water molecules from the outer layer and pores of the catalyst evidenced by endothermic peaks of DTA curves [27-28]. The second weight loss arises at 500 °C due to entire elimination of bio-template from the cavities with evidence of exothermic peak [30-31]. The final weight loss at 910 °C may be due to the adjacent OH group condensation [30]. TGA curve of AlSiO₄ material shows, it is highly thermally stable up to 950 °C.

![Figure 4: TGA curve for as-synthesized AlSiO₄ (bio-templated)](image)

3.4TPD Analysis

Acidic character of the calcinated material is analyzed from NH₃ Temperature Programmed Desorption (Figure 5). The material has 0.2091 mmole / g of acidity. The catalyst is active at 175 °C and above. The synthesized material has a series of acid sites from weak, moderate and strong [31-32].
From Figure 6, the surface area of the calcinated substance is 42.3077 m²/g and its pore size is 14 nm. Type IV isotherm hysteresis loop from 0.1–0.9 ($p/p_0$) belongs to the adsorption of N₂ in nanoporous walls. From the above result, it is confirmed the formation of mesoporous material [33-34]. The pore size of the material is 14 nm. Hence it is named as AlSiO₄-14. A Physico-chemical property of the AlSiO₄-14 material is tabulated in Table 1.
Table 1: Physico-chemical properties of AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Total acid (mmol / g)</th>
<th>$S_{BET}$ (m$^2$ / g)</th>
<th>Pore size (nm)</th>
<th>$V_{meso}$ (cm$^3$ / g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AlSiO$_4$-14</td>
<td>0.20</td>
<td>42.3</td>
<td>14.1</td>
<td>0.17</td>
</tr>
</tbody>
</table>

3.6 SEM and EDX Analysis

Photographical views of the thermally treated AlSiO$_4$-14 are verified from SEM analysis (Figure 7). SEM analyses give the feature about the particle dimension and contour of the outer face of the material [35]. Type of porosity, pore annulled and particle agglomerations are studied. Moreover the pores are observable and pores are ordered particle sizes are disordered.

The amount of Al-O-Si in the synthesized material is analyzed by elemental analysis. The selected area of EDX investigation proves the occurrence of distinctive peak of AlSiO$_4$ in Figure 8 and the molar composition is 1:4:1 (Al: 4O: Si). From this outcome, it is confirmed that the material is in tetrahedral framework.

Figure 7: SEM images of calcinated AlSiO$_4$-14

Figure 8: EDX spectrum of calcinated AlSiO$_4$-14
4. Catalytic study of carbon dioxide decomposition

The catalyst is applied for carbon dioxide decomposition reaction to evaluate the catalytic activity and the catalytic reactor setup is shown in Figure 1. Bio-templated AlSiO$_4$-14 decomposes the CO$_2$ molecules by the following reaction mechanism and it is given below.

During the reaction the 2CO$_2$ dissociates as 2CO and O$_2$. The acidic sites of the catalyst further decompose the carbon monoxide into carbon and oxygen.

\[
2\text{CO}_2 \xrightarrow{\text{catalytic cracking}} 2\text{CO} + \text{O}_2 \quad \text{(1)}
\]

Further the catalytic cracking of CO gives carbon and oxygen.

\[
2\text{CO} \rightarrow 2\text{C} + \text{O}_2 \quad \text{(2)}
\]

The reaction mechanism is mainly developed by the presence of active acid sites of the material. In acidic material, the CO$_2$ molecule is decomposed and forms as stable carbon monoxide and release oxygen.
4.1 Effect of temperature

Decomposition of carbon dioxide has been studied from 60 to 200°C for AlSiO$_4$-14 and the conversion percentage is tabulated in Table 2 (Figure 9).

Conversion is more at 70°C and the oxygen product selectivity is poor compared to carbon monoxide. Further increase of temperature, oxygen selectivity increased and carbon monoxide selectivity is decreased. However the carbon dioxide transformation reduces with increase of heat. It is revealed that partial decomposition decreased with increase of temperature or complete decomposition increases with increase of temperature.

Table 2: Result of heat on transformation of CO$_2$ over AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Temperature (°C)</th>
<th>Carbon dioxide transformation (%)</th>
<th>Product conversion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Complete transformation</td>
<td>Partial transformation</td>
</tr>
<tr>
<td>AlSiO$_4$-14</td>
<td>60</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>70</td>
<td>5</td>
<td>15</td>
<td>20</td>
</tr>
<tr>
<td>80</td>
<td>7</td>
<td>4</td>
<td>11</td>
</tr>
<tr>
<td>100</td>
<td>10</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>150</td>
<td>7</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td>200</td>
<td>6</td>
<td>1</td>
<td>7</td>
</tr>
</tbody>
</table>

Figure 9: Result of heat on transformation of CO$_2$ over AlSiO$_4$-14

Catalyst dose: 0.5 g; flow rate: 0.5 ml/min; Time: 1 h

4.2 Effect of flow speed

Effect of flow speed started from 0.5 to 2.5 ml per minute over AlSiO$_4$-14 is exposed in Figure 10. AlSiO$_4$-14 show highest change of carbon dioxide at 0.5 ml per minute, further raise of flow rate, selectivity of carbon monoxide is increased and oxygen decreased. Conversion of carbon dioxide (partial decomposition and complete decomposition) also decreased with increase of flow rate. It may be caused
by the diffusion of cavities with carbon dioxide molecules. This study confirms that the most favorable flow speed for AlSiO$_4$-14 is 0.5 ml per minute (Table 3).

**Table 3:** Result of flow rate on transformation of CO$_2$ over AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Flow rate (mL/min)</th>
<th>Carbon dioxide transformation (%)</th>
<th>Product conversion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Complete transformation</td>
<td>Partial transformation</td>
<td>Total conversion</td>
</tr>
<tr>
<td>AlSiO$_4$-14</td>
<td>0.5</td>
<td>5</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>1.5</td>
<td>3</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>2</td>
<td>14</td>
</tr>
</tbody>
</table>

**Figure 10:** Result of gas flow rate on transformation of CO$_2$ over AlSiO$_4$-14

Catalyst dose: 0.5 g; temperature AlSiO$_4$-14 (70°C); Time: 1 h

**4.3 Result of catalyst dosage**

Loading of catalyst dosage verified from 0.5 to 2.5 g (Table 4 & Figure 11). Utmost transformation of carbon dioxide is found at 2.5 g over AlSiO$_4$-14. Product selectivity of oxygen is also good with increase of catalyst dosage. AlSiO$_4$-14 does not undergo any recombination process when there is boost of catalyst quantity, the conversion & product selectivity (mainly O$_2$) is raised and CO selectivity is decreased.

**Table 4:** Result of catalyst dosage on transformation of CO$_2$ over AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Catalyst dosage (g)</th>
<th>Carbon dioxide transformation (%)</th>
<th>Product conversion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Complete transformation</td>
<td>Partial transformation</td>
<td>Total conversion</td>
</tr>
<tr>
<td>AlSiO$_4$-14</td>
<td>0.5</td>
<td>5</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>1.5</td>
<td>13</td>
<td>24</td>
</tr>
</tbody>
</table>
Figure 11: Result of catalyst dosage on transformation of CO$_2$ over AlSiO$_4$-14

Flow speed: 0.5 ml / min; temperature AlSiO$_4$-14 (70°C); Time: 1 h

4.4 Effect of time on flow

Time on flow has been studied over AlSiO$_4$-14 catalyst continuously up to 5 hrs with optimized conditions (Figure 12). The conversion (complete & partial decomposition) of CO$_2$ and product selectivity of carbon monoxide and oxygen are shown in Table 5. There are no significant changes in CO$_2$ conversion and product selectivity (CO & O$_2$). It indicated that the catalyst is active for long time. The carbon may be deposited in the pores. It may also promote the reaction; hence, activity of the catalyst is maintained during the reaction.

Table 5: Effect of time on flow for the decomposition of CO$_2$ over AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Contact time (hrs)</th>
<th>CO$_2$ transformation (%)</th>
<th>Product conversion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Complete transformation</td>
<td>partial transformation</td>
</tr>
<tr>
<td>AlSiO$_4$-14</td>
<td>1</td>
<td>43</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>51</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>51</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>52</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>48</td>
<td>45</td>
</tr>
</tbody>
</table>
4.5 Complete transformation of carbon dioxide

Complete transformation of carbon dioxide is shown in Table 6. AlSiO$_4$-14 decomposed nearly 43 to 50% of carbon dioxide (average 49%) during the time on stream and its partial decomposition average is 39%. The surface area is low and pore size is large. However, the conversion is achieved 49%. It may be due to the nature of template, acidity and cavity size of the catalyst.

Table 6: Complete transformation of carbon dioxide on AlSiO$_4$-14

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Optimized Temperature (°C)</th>
<th>Total acid (mmol / g)</th>
<th>$S_{BET}$ (m$^2$/g)</th>
<th>Pore size (nm)</th>
<th>Complete CO$_2$ decomposition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AlSiO$_4$-14</td>
<td>70</td>
<td>0.2091</td>
<td>42.3</td>
<td>14.1</td>
<td>43</td>
</tr>
</tbody>
</table>

5. Conclusion

Eco friendly and economically inexpensive bio-material based AlSiO$_4$-14 catalyst is synthesized and characterized. From the analytical studies, the synthesized catalyst has high thermal stability up to 900 °C, surface area is 42.3077 m$^2$ / g and its pore size is 14 nm. From TPD, the synthesized material is active at 175 °C and above, acidity of the material is 0.209 mmol / g. Egg white created the specific features in the material. One of the main advantages of the material is, there is no coke formation during the reaction due to the presence of low temperature active acid sites. Finally its catalytic activity is experienced by carbon dioxide decomposition reaction and various experimental conditions are optimized for maximum conversion and selectivity. Comparing with other material bio-templated AlSiO$_4$-14 produced 75 to 93% of conversion and 70 (CO) & 30 (O$_2$) percentage of product selectivity. The catalyst can decompose the carbon dioxide molecules constantly. The complete decomposition of CO$_2$ molecules are achieved 43% by AlSiO$_4$-14 catalyst.
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Summary of Interest

The authors Dr. M.A. Mary Thangam and Dr. C. Kannan have no conflict of interest.

References


