Hydro-treating and Hydro-isomerisation of Sunflower Oil using Pt/SAPO-11: Influence of Templates in Ultrasonic Assisted with Hydrothermal Synthesis

Shanmugam Palanisamy¹,*), Durona Palanisamy¹, Mugaishudeen Gul¹, Kannan Kandasamy¹, Borje Sten Gevert²

¹Kongu Engineering College, Erode-638 060, India.
²Kempross AB, Larseredslyckor 14, 42539 Hisingskärre, Sweden.

Abstract

Pt/SAPO-11 mesopores type materials has successfully synthesized using different templates, such as: diethylamine (DEA), dimethylamine (DMA), and n-propylamine (n-PA), under ultrasonication coupled with hydrothermal treatment or independently with hydrothermal treatment. The influences of structure directing agent (SDA) and synthesis method are investigated by different characterization techniques and the role of the material as catalyst in hydrotreating of sunflower oil has examined. The synthesized materials have been characterized by X-ray Diffraction (XRD), Scanning Electron Microscope (SEM), and Fourier Transform Infra Red (FT-IR) techniques. It is found that SAPO-11 material which has synthesized with n-PA as a template has the characteristics of high silicon incorporation. Hydrotreating of sunflower oil is carried out in a fixed bed reactor with Pt impregnated SAPO-11 catalyst and a detailed study on the isomerization is performed by varying the operating parameters like temperature and space velocity. The high selectivity of Pt/SAPO-11 catalyst is achieved by uniform pore size and acidity. Also the pore opening of the catalyst has a major effect in the selectivity of the catalyst. Further, it represents a higher ratio of isomers compared to other synthesized catalysts on hydro-treating of sunflower oil.

Keywords: SAPO-11; Zeolite; Mesoporous; Hydrothermal; Ultrasonication; Hydrotreating; Hydroisomerisation; Hydrodeoxygenation


1. Introduction

The growing demand of fuel is observed and use of renewable feedstock in replacing the conventional fuel in transportation can be the alter solution for the fossil fuel dependency. One such renewable source can be the direct blend of vegetable oil and gas oil in diesel engine [1–5]. But the study proved that the presence of carboxylic group in vegetable oil has increase the NOx emission and corrosiveness in the fuel tank. So, in order to reduce this default, refineries have been upgrading this blend in the hydroprocessing technology by hydrodeoxygenation (HDO) mechanism to remove carboxylic group [1–7]. In other way, direct HDO of tall or vegetable oil may convert into higher straight chain hydrocarbon composition through fatty acids as intermediates. The straight chain hydrocarbon...
from vegetable oil can produce the composition of larger-chain hydrocarbon (n-heptadecane and n-octadecane) [3–7]. This paraffinic gas oil or diesel-like fuel composition may have a problem by increasing the cloud point of the middle distillates. The diesel-like fuel having high content of paraffinic hydrocarbon from different sources is difficult to use under cold weather conditions. To deal the cold properties of upgraded gas oil, it is necessary to reduce the freezing point of higher straight-chain hydrocarbons by increasing the branched carbon compounds in the long chain which can be attained through isomerization mechanism [6–8]. So, the isomeric compounds can reduce the cloud point, however, the properties of cold flow plugging point can minimize by increasing the number of methyl or ethyl branch in the hydrocarbon. Figure 1 shows the change in the melting point of isomeric compounds with methyl branch in hydrocarbon chain, respectively.

In 1962, zeolites were introduced by the Mobil Oil Company to be used as a cracking catalyst in oil refinery technology. It has several industrial applications that provide many advantages to the refinery industry, including ion exchange, shape selectivity, selective absorbent, high activity and selectivity in a wide range of products and high thermal stability. However, the cations of several zeolites reside inside the pore channel for the capability in branching hydrocarbons. The shape selectivity of zeolites is the ability to selectively catalyze reagents or reactants. This ability is achieved through their pore structure and the thermal stability, at which this can be thermally stable up to 600 °C. Several studies have reported minor destruction of the structure caused by dealumination, which modify the pore volume [9,10]. Except the aluminiation on material due to calcined temperature, this may not affect pore volume drastically. The thermal stability of zeolites increases when the SiO$_2$/Al$_2$O$_3$ ratio increases and the decomposition of zeolites occur near a temperature of about 700 °C. The acidic property of zeolites vary by changing the SiO$_2$/Al$_2$O$_3$ ratio [11–14]. In zeolites, acid sites are classified according to the classical Bronsted and Lewis models of acidity [11–17]. A high SiO$_2$/Al$_2$O$_3$ ratio is stable in concentrated mineral acid, whereas a low SiO$_2$/Al$_2$O$_3$ ratio is rather fragile in the presence of acid. In contrast, the high SiO$_2$/Al$_2$O$_3$ ratio is unstable in the basic solution but the low SiO$_2$/Al$_2$O$_3$ ratio can enable increase in stability. Consequently, the acidic property of zeolites is mainly dependent on the SiO$_2$/Al$_2$O$_3$ ratio as well as the operating temperature in the reactor.

Zeolites are crystalline aluminosilicates molecular sieve containing uniform pores and acidic sites, which is widely used in water treatment, fine chemical synthesis, upgrading fuels, gas separation, emission control and adsorbent. Zeolites are synthesized and its properties have been mainly studied with different sources, methods and application by various researches [9–17]. From this series, in the last few decades, synthesis of phosphate incorporated silica-aluminates materials are popularized because of their material as better catalyst performance and stability in hydroprocessing of bio-based fuel synthesis from fat-rich material. Silicoaluminophosphate (SAPO) zeolite is used as a catalyst support in various processes such as upgrading of fuel, chain growth of lighter hydrocarbon, hydrocracking of long chain hydrocarbon and conversion of vegetable oil into bio-diesel. This material has been used for isomerisation, aromatization, hydrogenation, alkylation and dehydrogenation processes in refinery processes. The isomerisation is the branching of hydrocarbons where transformation of molecules which has same number of atoms takes place irrespective of their atomic arrangement on both hydrogenation and dehydrogenation in duel mechanism. Both mechanism such as branching of hydrocarbon and hydrogenolysis of the fatty acids into paraffinic or olefinic hydrocarbons are widely investigated using SAPO as a support material in solid catalyst.

Solid acid catalysts (such as zeolites loaded with transition metal (Pt)) on activation are used in the industrial isomerization process. The catalysts are often called bi-functional as they contain both acid and metal functions.

**Figure 1.** Melting point for different hydrocarbons with position of isomeric compounds. [Source: www.nist.gov]
Here, the functional activity of catalyst can act as acidic solid material and hydrogenation-dehydrogenation process [2,11–12]. Generally, zeolites with alkaline metal cations are colorless powders. It has a crystalline porous aluminium silicate intra-bonds with a vast three dimension network structure of tetrahedral coordinated SiO₄ and Al₂O₃, producing a low-density micro-porous structure. Color presenting on catalysts may occur because they contain transition metals or modifying zeolites by an ion exchange method. Most zeolites can be dehydrated without the collapse of the framework by calcination on temperatures between 350 and 500 °C [12–16].

The research study aimed to hydrotreating the fatty acids in the presence of Pt metal impregnated over the surface of alumina-SAPO-11 support material. The synthesized SAPO-11 is used as a base support for the catalyst for the conversion of vegetable oil into diesel-like hydrocarbon. Though, the diesel-like hydrocarbon is produced in long straight-chain hydrocarbon such as heptadecane (C17) or octadecane (C18), this hydrocarbon can involve the cold plugging problem in fuel injection pipe due to solidification at room temperature. To reduce the plugging problem, it is important to branch the long straight-chain hydrocarbon. So, the importance of synthesis of Pt/SAPO-11 has focused to involve in both hydrotreatment and branching of the fatty acids within the moderate temperature range (290 to 380 °C). Here, the study involves hydrogenation and dehydrogenation over the sunflower oil as reference vegetable oil (refined form of triglyceride) at different operating parameters of temperature and pressure. The synthesis of material can carried out by different templates, such as: diethylamine (DEA), dimethylamine (DMA), and n-propylamine (n-PA), using SiO₂ and Al₂O₃ under ultrasonication coupled with hydrothermal treatment or independently with hydrothermal treatment. The main purpose of the investigation is to analyze the activity and performance of the synthesis of Pt/SAPO-11 using various templates in supported zeolite catalyst for the dual purpose on both hydrotreating of vegetable oil and upgrading into diesel-grade fuel within the optimized condition.

2. Methods and Materials

2.1 Materials

The raw materials used for producing Pt/SAPO-11 and support metal oxide contain colloidal silica (40 wt% suspension in water from Sigma-Aldrich), aluminium hydroxide (Universal Chemicals, India), orthophosphoric acid (85 % Merck), Tetra amine platinum(II) chloride hydrate (99.99% Sigma-Aldrich) dimethylamine (80% Loba chemie), dimethylamine (99% Nice Chemicals) and n-propylamine (99 % Merck).

2.2 Catalyst Preparation

Initially, 10.33 g of aluminium oxide (Al₂O₃) which acts as the alumina source is added to 16.5 mL of distilled water (deionized) and stirred by magnetic stirrer for 50 minutes at room temperature. The gel formation is achieved better in Al₂O₃ than compared to other alumina sources like aluminium chloride (AlCl₃). While stirring, 3 mL of orthophosphoric acid (H₃PO₄) is added in drop wise to the alumina water mixture for 30 minutes. After addition of H₃PO₄ to the mixture, 0.764 g of silica sol is added. Further, 20 mL of ditetylamine (DEA) (S5, S6) is added to the mixture which acts as a structure directing agent (template) and the stirring is allowed to continue for 3 hours. By substituting various templates such as dimethylamine (DMA) (S3, S4) and n-propylamine (n-PA) (S1, S2) SAPO-11 has prepared. The obtained solution is stirred until a gel is formed and it is transferred into an ultrasonic bath in a shielded container at 120 °C for 6 hours maintained at 60 (reported as an ultrasonic – S2, S4, and S6). The ultrasonic method has carried on 39 kHz and 60 W conditions.

The same preparation technique has followed to prepare the gel solution and after sonication the samples are subjected to be kept in a hydrothermal reactor at a temperature of 180 °C for 24 hours (reported as an ultrasonic and hydrothermal process – S1, S3 and S5). The same procedure is repeated for the samples of different templates. The samples are dried at 110 °C for 24 hours and calcined at a high temperature of 600 °C for 52 hours. (The sample was calcined for 52 h in order to burn out the template present over the surface)

Prepared SAPO-11 and γ-Al₂O₃ (Disperal and Locron) is blended in a ratio of 40:60 wt%, and the hydrated blend has extruded in 1/32” material size. The support material was dried at 110 °C for 5 hours and calcined at 350 °C for 6 hours. After drying and calcination, platinum was loaded using Pt(NH₃)₂Cl₂ solutions by impregnation to get the content of 0.5 wt% Pt. The catalysts are dried at 110 °C for 12 hours and calcined at the temperature of 400 °C for 4 hours in air. Finally, H₂ in-situ activation is performed at 400 °C for 4 hours.
2.3 Experimental Set-up

The experimental set-up consists of a feed tank, fixed-bed reactor, product tank, gas collector and dossier pump [18,19]. The fixed-bed reactor is fixed with an electric furnace and connected with an instrumental controller to regulate temperature and pressure conditions. The catalyst is placed at 62.6 mm (31 mL) in height from 619.4 mm as total vertical height of the reactor. The distance of the catalyst from the top or bottom line is 278.4 mm while the remaining part of reactor space can consists of borosilicate glass pellets. The outlet goes to a separator and the reactor pressure is controlled on the gas exit of the separator with manual collection in liquid and gas samples. The gas outlet and liquid products are analyzed with gas chromatography as represented in below section. Hydrogen was supplied at volumetric rate of 1200 mL/h in 3:1 hydrogen to hydrocarbon ratio.

2.4 Analysis Method

The liquid sample analysis is performed using a gas chromatograph (GC) technique (Varian 3400) equipped with a packed column (Elite-5MS column, 30 m x 0.3 mm. I.D., 1.0 m film thickness and dimethyl polysiloxane) and flame ionization detector (FID). The Varian 4270 is the integrator to data computation. Samples (5 µL) were injected into an injector through a micro-syringe. The outlet gas sample from the reactor has analyzed using the Clarus 500 GC online. This GC was connected with 600 link switch controllers, with the signal that has integrated to receive data. The gas analysis has an inlet and outlet sampling value for FID and TCD detectors and has four valves actuated by Nitrogen gas at 0.40 MPa. Gas analyses have consisted of a thermal conductivity detector (TCD) for analyzing CO, CO₂, CH₄ and H₂, as well as the FID for analyzing hydrocarbons. Helium was used as carrier gas for the TCD, which was maintained at 200 °C with oven heat-up of 40 to 60 °C at 2 °C/min and nitrogen as carrier gas for the FID at 60 °C as oven temperature. GCMS-QP 2010 Ultra- Shimadzu GC-2010 Plus consist of capillary column made up of Frontier Labs UA+65 (30 m×0.25 mm I.D. df = 0.1 µm), which was maintained at temperature: 150° C (for 1 min) and ramped at 5 °C/min upon 320 °C (for 20 min). The injection and FID temperature was 300 and 320 °C, the split ratio was 1/100 and m/z intensity range was 20.00–800.00.

The constructive interference of crystal or phase structure can be identified by monochromatic X-ray beam scattering. The uniform or skeletal spacing in crystals are identified by creating the patterns of waves in incident X-rays diffraction (XRD) scattering. XRD patterns of the prepared dried solid catalyst are measured using Cu Kα (λ = 1.54 Å) and Mo-Kα (λ = 0.71 Å) radiation in miniflex and smart-lab Rigaku diffractometer where the sample crystalline structure have analysed at the range of 5 to 70° of 2-theta with a scan step of 0.05° and this performing a counting time of 4 s for each step.

The Scanning Electron Microscope (SEM) had operated at an acceleration voltage of 30 kV. Usually, the samples are used at room temperature after vacuum and dried at 250 °C for 2.5 hours. Before sent to analysis, sample materials are grinded into powder in ethanol solution and involved at vacuum-dried condition on 120 °C for 3–4 hours. After drying, the material has to be placed for identification of grain orientation and texture analysis on an electron backscattered diffraction detector.

The Brunauer, Emmet, and Teller (BET) surface area has been investigated by Quanta Chrome Nova-1000 surface analyzer instrument under liquid nitrogen temperature. During each measurement, the samples are degassed at 300 °C for 24 hours to remove moisture and unwanted absorbed gases in the surrounding.

Fourier-Transform Infrared Spectroscopy (FTIR) with specification of Bruker FTIR A. Model Tensor-27 was used for solid catalyst acidity measurement. Here, 5 gm of solid sample, which is dried in vacuum chamber at 350 °C for 4 hours, has been made as thin wafers by pressed in coin shape by placing in the IR cell. Then, pyridine vapour at 200 °C has introduced to the cell for 60 min and then, the cell was placed at hot inert gas (N₂) which has passed to record the spectrum. Mode of interpretation is transmission and wavenumber (cm⁻¹) in the acidity validation graph.

3. Result and Discussion

The composition, temperature, holding time and nature of the reactants are the major parameters which have to investigate the solid catalyst performance over hydrotreating and branching of the feed. Here, the study mainly focused on the performance of the Pt/SAPO-11 by varying the temperature and pressure of the reactor with constant flow of reactant and its composition of the sunflower oil.

The synthesised catalyst, Pt/SAPO-11, was analysed and tested for uniform pore distribu-
tion, acidic site and particle size using various testing methods such as BET, SEM analysis, FTIR and XRD. To identify and study the crystallinity of the SAPO-11 produced, the support material are examined in SEM and XRD analysis for the understanding. Also, the acidity on the surface metals of support is investigated by Pyridine-FTIR analysis.

The experimentation of the catalytic hydro-treating of the vegetable oil (Sunflower oil) using the synthesized support impregnated with Pt metal. 31 mL (27.83 gm) of Pt/SAPO-11 material (with exact weight of 27.59 g catalyst) is loaded in the reactor, activated using fresh H$_2$ at 350 °C for 4 hours and the investigation on the catalyst performance at space velocity of unity with respect to 31 ml of sunflower oil pumped into the reactor has performed. The space velocities are calculated at liquid hourly space velocity (LHSV = mL of liquid/mL of cat.*h) or (also represented as h$^{-1}$) at preferred reaction pressure (1 to 3 MPa) and reaction

![Figure 2. SEM images of SAPO-11 prepared by using (a) n-propylamine U + H, (b) n-propylamine U, (c) dimethylamine U + H, (d) dimethylamine U, (e) diethylamine U + H, and (f) diethylamine U.](image-url)
temperature (290 to 360 °C). So, the experiment started with 31 mL of catalyst with corresponding 1 h⁻¹ LHSV is performed in a continuous fixed-bed reactor. Also, the tolerance error for mass balances of lighter hydrocarbons and some middle distillate fractions are considered within approximated range of 0.5 to 1 wt%. In addition, traces at the samples of heavy hydrocarbons beyond C19 are considered to be negligible. So, the main components in the residual products are hydrocarbons from dodecane to heavy hydrocarbons (C18≤) and also included with traces of lighter hydrocarbons.

3.1 Microscopic Analysis

The SEM analysis has performed for the synthesized SAPO-11 materials to understand the structure of the sample. It is observed that the sample synthesised with n-propylamine (n-PA) as template under ultrasonic with hydrothermal method (UH) combined together as named as S-1, the particles are observed with a uniform globules having an average particle size of 100.12 nm, as represented in Figure 2 (a). The structure has identified to be irregular which is related to its heterogeneous morphology. Further, while the sample with same template (n-PA) synthesized under ultrasonication process (S-2), exhibit sharp edges of non-uniform cuboids (Figure 2 (b)). The crystal structure in this sample is non-uniform in size and shape. This non-uniform structures show an average particle size of 80.29 nm. Whereas in the sample produced with dimethylamine (DMA) as a template under UH combined together (S-3) have exhibited irregular surface morphology. Thus, it has large body like structures with irregular hollow pits spread along the surface of the body, as shown in Figure 2 (c). The hollow pits seen on the surface indicates that incomplete crystal growth has occurred. The average particle size of the produced sample is 187.96 nm. But the particle size is larger and if there is incomplete formation of structure, the sample produced will have a negative effect on lifetime. In the sample of dimethylamine as template under ultrasonic process with hydrothermal treatment (S-4), Figure 2 (d) observed that there is a formation of few non-uniform clumps of small sized bodies. The small sized bodies formed are irregular in shape and size. These irregular bodies are formed over the surface of support base of S-4 with the average particle size of 167.55 nm.

In the sample synthesized with diethylamine (DEA) as a template under ultrasonication and hydrothermal treatment (UH) coupled together (S-5), Figure 2 (e) observed that there is no proper crystal growth within tested condition. The pores are non-uniform and the morphology is irregular to the subjected synthesis process. The SEM images represent that there a few clusters of irregular shaped bodies with incomplete crystal growth. The average particle size represent in this sample is around 100.69 nm. Whereas, sample of diethylamine (DEA) as a template synthesized under ultrasonic process without hydrothermal treatment (S-6), the Figure 2 (f) shows that there is formation of small sized spherical bodies which has heterogeneous morphology. It is observed that the spherical structures in this sample have just initiated to form flakes. However, the corners of the flakes have figured to catch the pictures of original shapes rather than the entire flake. The average particle size of this sample is found to be 79.99 nm. Table 1 enlists the detailed information of the particle size of the material synthesized from different combinations and template in the method of material synthesis. The smaller particle size in the ultrasonically prepared samples can be accounted to the reduction in agglomeration of the cell crystals rather giving a rise in nucleation and

### Table 1. Average Particle size of the synthesized SAPO materials and the data is extracted from Figure 2.

<table>
<thead>
<tr>
<th>Sample with sample no.</th>
<th>Template</th>
<th>Synthesis</th>
<th>Pa 1 (nm)</th>
<th>Pa 2 (nm)</th>
<th>Average Size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-PA (UH) (S-1)</td>
<td>n-PA</td>
<td>U+H</td>
<td>87.02</td>
<td>113.2</td>
<td>100.12</td>
</tr>
<tr>
<td>n-PA (U) (S-2)</td>
<td>n-PA</td>
<td>U</td>
<td>70.45</td>
<td>90.12</td>
<td>80.29</td>
</tr>
<tr>
<td>DMA (UH) (S-3)</td>
<td>DMA</td>
<td>U+H</td>
<td>116.1</td>
<td>259.8</td>
<td>187.96</td>
</tr>
<tr>
<td>DMA (U) (S-4)</td>
<td>DMA</td>
<td>U</td>
<td>61.12</td>
<td>274</td>
<td>167.55</td>
</tr>
<tr>
<td>DEA (UH) (S-5)</td>
<td>DEA</td>
<td>U+H</td>
<td>56.88</td>
<td>144.5</td>
<td>100.69</td>
</tr>
<tr>
<td>DEA (U) (S-6)</td>
<td>DEA</td>
<td>U</td>
<td>58.07</td>
<td>101.9</td>
<td>79.99</td>
</tr>
</tbody>
</table>
crystal growth. In turn this accounts for the rise in crystallinity [19].

3.2 XRD Analysis

The phase purity and crystallinity of the samples are determined by using Powder X-ray diffraction (XRD) patterns. Crystallinity characterization of the samples has calculated by the equation [20].

\[
\text{Relative intensity} = \frac{\text{Intensity}}{\text{Reference Intensity}}
\]

The line intensities of the XRD pattern at 2\(\theta\) equal to 21.5, 45.7, 59.9 and 67.2° are employed for these calculations.

The X-ray diffraction patterns of SAPO-11 samples prepared by two methods are respectively shown in Figure 3. The high intensity of the XRD lines and the low background in the XRD pattern indicate the high crystallinity of the samples and, also the decrease in the intensity of the XRD peaks indicates the probable loss in crystallinity. It can be observed that all these structures are the same and have similar peak positions at Bragg angle 2\(\theta\) = 21.5, 45.7, 59.9 and 67.2°. The characteristic diffraction peak at 2\(\theta\) = 21.5° are in agreement with that of SAPO-11 zeolite framework [21–23]. The intense peaks at a Bragg angle 2\(\theta\) of 21.5° confirm the preferred orientation to the c-axis of hexagonal crystals. The crystallinity of the S-2 catalyst has found to be higher than the other catalysts. So, this has been considered as the reference data while determining the crystallinity of other samples. It is found that the relative crystallinity of all catalysts related to the formation of SAPO-11 phase have decreased in the following order:

n-PA (U) > DMA (U) > DEA (U) > n-PA (UH) > DMA (UH) > DEA (UH).

The diffraction peak intensities or crystallinity of SAPO-11 catalyst synthesized by ultrasonication is generally higher than that of the catalysts synthesized using ultrasonication in hydrothermal condition, which suggests that the ultrasonic coupled with hydrothermal synthesis is more effective for the catalyst synthesis. At low temperature and short period of residence time, ultrasonic transformation is capable of synthesizing catalysts with higher degree of crystallinity. In addition, the reflections of peaks are found to be greater for n-propyl amine than the other two catalysts and diethylamine template can demonstrates the lowest peak intensity in the material validation. The higher crystallinity observed with the ultrasonically synthesized samples is due to the boosted crystal growth and higher external specific surface area.

3.3 FTIR Analysis

The acidity of the zeolites has examined by using Pyridine-FTIR analysis, as represented in Figure 4. The notable bands for the acidic

![Figure 3. XRD images of SAPO-11 prepared by using a) ultrasonic assisted hydrothermal synthesis (UH) method and b) Ultrasonic synthesis (U) method [Note: n-PA= n-propylamine template, DME= dimethylamine and DEA= diethylamine].](image-url)
sites in the pyridine adsorption on the surface of metal site and hydrates present on the surface of metal and oxides are investigated using FTIR. It is observed that the bands occur at wavenumbers around 1400, 1545, 1636, and 3450 cm\(^{-1}\) on both plots of FTIR analysis of samples synthesized by hydrothermal treatment and ultrasonication process are indicated in Figure 4 (a-c). Transmittance percentage exhibited by the sample prepared by ultrasonic with hydrothermal condition is higher than the other samples. Peaks occurring at 1400 cm\(^{-1}\) exhibit Lewis acidity and as the wavenumber increases in 1493 cm\(^{-1}\), Bronsted acidity has initiated to add up with the Lewis acidity. At wavenumber of 1545 cm\(^{-1}\), it is comprised mainly of Bronsted acidity [24–27].

Between 1200 and 1000 cm\(^{-1}\), specific high intensity band at 1068 cm\(^{-1}\) has identified as a distinct feature in the very broad and high intensity band on below 1000 cm\(^{-1}\), the absorption is represented with high intensity that no evaluation of band positions are possible. The acidic peaks are represented in the Figure 4 (a) and (b). The asymmetric stretching vibration at wavenumber of 2080 cm\(^{-1}\) is assigned to the physically adsorbed CO\(_2\). The stretching vibration at wavenumber 1636 cm\(^{-1}\) can be assigned to the H–O–H weakly adsorbed in SAPO-11 [28,29]. The spectrum in the OH region except for the polarization which hasn’t been reported before is well in accordance with literature data obtained from the analysis of powdered polycrystalline samples, recorded in either transmission or diffuse reflectance.

**Figure 4.** FT-IR images of SAPO-11 prepared by using (a) Ultrasonication, (b) Ultrasonication and hydrothermal and (c) hydroxyl-group in catalyst. Note- At (c), Dotted Arrow represents the increase in intensity of the spectrum for –OH group.

**Table 2.** Surface area of the catalyst produced by different methods with various templates (Surface area of the catalyst produced by different methods with various templates).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Template</th>
<th>Mean pore diameter (nm)</th>
<th>Total pore volume (m(^3)/g)</th>
<th>(V_m) (cm(^3)(STP)/g)</th>
<th>Surface area (m(^2)/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-PA (UH) (S-1)</td>
<td>n-PA</td>
<td>10.05</td>
<td>156.63</td>
<td>12.52</td>
<td>474.88</td>
</tr>
<tr>
<td>n-PA (U) (S-2)</td>
<td>n-PA</td>
<td>9.37</td>
<td>72.85</td>
<td>10.75</td>
<td>682.52</td>
</tr>
<tr>
<td>DMA (UH) (S-3)</td>
<td>DMA</td>
<td>11.21</td>
<td>122.53</td>
<td>14.58</td>
<td>493.74</td>
</tr>
<tr>
<td>DMA (U) (S-4)</td>
<td>DMA</td>
<td>9.82</td>
<td>67.13</td>
<td>11.01</td>
<td>731.38</td>
</tr>
<tr>
<td>DEA (UH) (S-5)</td>
<td>DEA</td>
<td>11.88</td>
<td>107.82</td>
<td>13.35</td>
<td>396.51</td>
</tr>
<tr>
<td>DEA (U) (S-6)</td>
<td>DEA</td>
<td>9.85</td>
<td>61.25</td>
<td>11.41</td>
<td>737.80</td>
</tr>
</tbody>
</table>
The spectra of the samples in the OH-stretching vibration domain are shown in Figure 4 (c). The vast band stretching across the range 3000 to 3600 cm\(^{-1}\) can be attributed to the hydroxyl groups from the Si–OH–Al bond [30,31].

3.4 Particle Size Analysis

Table 2 shows the surface area of the synthesized SAPO-11 catalysts under ultrasonic method and ultrasonic assisted hydrothermal methods together. The surface area is obtained from the Brunauer–Emmett–Teller (BET) analysis. The results are attained for the samples with the use of different types of templates such as n-propylamine (n-PA), dimethylamine (DMA) and diethylamine (DEA).

From the data shown in the Table 2, it is observed that the samples synthesized using n-propylamine as template has a medium surface area when compared to the other SAPO-11 materials obtained. The SAPO-11 based-catalyst under ultrasonic and hydrothermal methods together with n-PA as template (S-1) has a pore volume of 156.63 m\(^3\).g\(^{-1}\). From the Table 2, it is clearly visible that this sample has the highest pore volume when compared to the other synthesized samples. The Figure 5 indicates the sampling adsorption isotherm on n-PA of both S-1 and S-2 for the indication. The sample produced under ultrasonic method with n-PA as template (S-2) has a pore volume of 72.85 m\(^3\).g\(^{-1}\).

From the results obtained from the BET analysis, it is seen that the sample produced with dimethylamine (DMA) as template under ultrasonic and hydrothermal methods combined together (S-3) has a pore volume of 122.53 m\(^3\).g\(^{-1}\). It is also observed that the sample produced under the same DMA template on ultrasonic condition (S-4) has a pore volume of 67.13 m\(^3\).g\(^{-1}\).

Further, it is analysed from the BET results obtained that the sample produced with the help of diethylamine (DEA) as template under both ultrasonic and hydrothermal conditions coupled together (S-5) has a pore volume of 107.82 m\(^3\).g\(^{-1}\). Similarly the sample obtained with the help of the same template (DEA) under ultrasonic method (S-6), has pore volume of 61.25 m\(^3\).g\(^{-1}\). It is noticed from the results obtained from the BET data that the sample produced with the diethylamine (DEA) as template under ultrasonic and both hydrothermal and ultrasonic conditions combined together has the lowest surface area when compared to other samples produced with n-propylamine (n-PA) and dimethylamine (DMA) as templates. The pore volume of the samples produced with the above mention templates decreased in the order of n-PA > DMA > DEA.

3.5 Hydrodeoxygenation of Sunflower Oil

The synthesized catalyst is investigated in hydrotreating of triglyceride compounds in trickle-bed reactor with LHSV = 1 h\(^{-1}\). The carbonic compound in oxygenated group of triglycerides at sunflower oil is removed as CO\(_2\), CO, and H\(_2\)O within represented mechanism such as Decarboxylation (DCO), Decarbonylation.

Figure 5. a) Hydrothermal synthesis of SAPO with n-Propylamine templates and b) Ultrasonic synthesis of SAPO for 6 hour at 60 °C at 20 kHz.
DCA), and Hydrodeoxygenation (HDO), respectively with paraffins as their main products with a wide range of carbon numbers [32–35]. Table 3 enlists the data on operating conditions for the hydrodeoxygenation of sunflower oil along with the conversion, selectivity, and product composition for the synthesized SAPO-11 with Pt, the GC-MS results of the hydrodeoxygenation is represented in Figure 6. Here, the results obtained using the synthesized Pt/SAPO-11 catalyst samples are also compared with the obtained commercial zeolites which is named as H-β catalyst with Pt on surface. Though H-β material is popular in high thermal stability material and high cracking of hydrocarbon, it has wide application in the fuel upgrading at the refinery process.

From this investigation, it is found that the catalysts prepared under ultrasonic (U) method are superior to that synthesized by the other method (UH) in terms of crystallinity and acid sites distribution. Hence detailed studies on the ultrasonically synthesized catalysts samples are carried by changing the operating temperature and pressure of hydrotreating process with existing catalyst on appropriate conditions. By all comparison as shown in the Table 3, the results obtained using synthesized Pt/SAPO-11 samples exceeded and represent with high conversion rate while compared to the commercial catalyst (H-β). Also, the role of Pt on the catalyst is assumed to have same characteristic and function for this investigation. For n-PA (UH) samples, the conversion efficiency increased with the increase in operating temperature, while maintaining the operating pressure constant, characterized C17 and C18 compositions are identified by GC-MS in Figure 6 for corresponding operating temperature. The lighter hydrocarbon and middle distillate formation is noticed in the Figure 6. However, there is a drop in the yield of mono-branched products when the investigation has repeated with maintained constant operating temperature and while varying the pressure, the conversion efficiency increased with an increase in the yield percentage of mono-branched products.

Table 3. HDO of sunflower oil in reactor at LHSV = 1 h⁻¹.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Template</th>
<th>Oper. Temp. (°C)</th>
<th>Oper. Press. (MPa)</th>
<th>Conversion (%)</th>
<th>Selectivity HDO/(DCO+DCA)</th>
<th>Mono-branched (%)</th>
<th>Di-branched (%)</th>
<th>Cracked (&lt;C₈) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H-β</td>
<td></td>
<td>320</td>
<td>2</td>
<td>81</td>
<td>1.1</td>
<td>46</td>
<td>14</td>
<td>40</td>
</tr>
<tr>
<td>S-1 n-PA (UH)</td>
<td>320</td>
<td>2</td>
<td>91</td>
<td>1.5</td>
<td>55</td>
<td>15</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>300</td>
<td>2</td>
<td>84</td>
<td>1.3</td>
<td>67</td>
<td>13</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>2</td>
<td>92</td>
<td>1.2</td>
<td>63</td>
<td>22</td>
<td>15</td>
</tr>
<tr>
<td>S-2 n-PA (U)</td>
<td>340</td>
<td>2</td>
<td>98</td>
<td>1.5</td>
<td>57</td>
<td>21</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>1.5</td>
<td>92</td>
<td>1.1</td>
<td>55</td>
<td>18</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>2.5</td>
<td>94</td>
<td>1.5</td>
<td>61</td>
<td>22</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>3</td>
<td>95</td>
<td>1.6</td>
<td>66</td>
<td>22</td>
<td>21</td>
</tr>
<tr>
<td>S-3 DMA (UH)</td>
<td>320</td>
<td>2</td>
<td>84</td>
<td>1.4</td>
<td>52</td>
<td>13</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>300</td>
<td>2</td>
<td>71</td>
<td>1.3</td>
<td>51</td>
<td>9</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>2</td>
<td>82</td>
<td>1.5</td>
<td>52</td>
<td>12</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td></td>
<td>340</td>
<td>2</td>
<td>89</td>
<td>1.5</td>
<td>55</td>
<td>14</td>
<td>31</td>
</tr>
<tr>
<td>S-4 DMA (U)</td>
<td>320</td>
<td>1.5</td>
<td>84</td>
<td>1.5</td>
<td>52</td>
<td>15</td>
<td>33</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>2.5</td>
<td>91</td>
<td>1.6</td>
<td>56</td>
<td>17</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>320</td>
<td>3</td>
<td>93</td>
<td>1.6</td>
<td>59</td>
<td>18</td>
<td>23</td>
</tr>
<tr>
<td>S-5 DEA (UH)</td>
<td>320</td>
<td>2</td>
<td>88</td>
<td>1.4</td>
<td>53</td>
<td>12</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>300</td>
<td>2</td>
<td>78</td>
<td>1.1</td>
<td>72</td>
<td>11</td>
<td>17</td>
</tr>
<tr>
<td>S-6 DEA (U)</td>
<td>320</td>
<td>2</td>
<td>88</td>
<td>1.3</td>
<td>62</td>
<td>17</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>340</td>
<td>2</td>
<td>96</td>
<td>1.5</td>
<td>56</td>
<td>19</td>
<td>25</td>
</tr>
</tbody>
</table>
and it is observed that HDO/(DCO+DCA) ratio also increases constantly and linearly concurs with the literature fact that DCA/HDO ratio decreases as temperature increases [36–39]. The DCO and DCA mechanism forms C16 and C17 hydrocarbons, whereas C18 can be formed by HDO mechanism. The conversion efficiency peaked for both n-PA (UH) and DEA (UH) on the operating condition of T = 340 °C and P = 2 MPa. Using DMA (UH) sample, the conversion peaked at operating conditions of T = 320 °C and P = 3 MPa. Unlike the other side processes, selectivity for hydrodeoxygenation is observed to be expressed as a drastic rise of C17 with increase in the operating pressure for the investigated experiments as indicated in between 10 to 11 mins of retention time at GC-MS.

3.6 Hydroisomerisation of n-Octadecene

The HDO of sunflower oil contains unsaturated double bonds of triolein, trilinolein, tripalmitin, and tristearin triglycerides in the sunflower oil. These higher level unsaturated triglycerides demand higher consumption of hydrogen on hydrotreatment i.e., HDO pathway [35–40].

In this study, we used a gas chromatography-mass spectrometry (GC-MS) analysis (Figure 6) to determine the presence of the paraffin in the end product. Figure 7(a) displays the variation in concentration in products (wt %) over varying operating temperature at constant LHSV = 1 h⁻¹ and 5.0 MPa. The results from both these tests are found to be in consensus with each other and the interpretations are given as following:

Peaks beyond the retention time 12 min in the Figure 6 and higher depiction of hydrocarbons with boiling point above 300 °C in Figure 7 (a) are appeared due to the feed subjected by octadecene (n-C18=) paraffin. The fairly elevated presence of these traits at the operating temperature 270 °C establishes that the hydro-

![Figure 6. GC analysis of hydroisomerisation of n-C18= using Pt/SAPO and 5 MPa.](image)

![Figure 7. Hydro-isomerisation of n-C18= of a) different operating temperature and 5 MPa and b) different space velocity of feed at T = 310 °C and 5 MPa.](image)
Figure 8. Proposed mechanism of a) Hydrodeoxygenation of triglycerides into hydrocarbons and b) Hydro-isomerisation of n-C18= into branched hydrocarbons.
genolysis is initiated at its preliminary heating stage. The distortion in between retention times 10 and 12 seconds in the following processes with operating temperature above 270 °C are attributed to the hydro-isomerized products of the n-C18= hydrocarbon whose boiling points falls in the range of 275–300 °C. An abrupt increase in the peaks below the retention time 2 seconds when processed at temperatures above 310 °C corresponds to the low boiling pointed alkanes ranging from C8 to C16. This is attributed on the account of undesirable hydrocracking taking place at higher operating temperatures. Detailed hydroprocessing and hydroisomerisation of this mechanism is represented in Figure 8 (a) and (b).

The numeric data of the GC-MS Analysis is provided in the Table 4. Conversion as well as the composition of both iso-alkanes and n-alkanes shows a significant increase with increasing operating temperatures. An increase in iso-alkanes is desirable reaction but the undesirable escalation in n-alkanes is due to the cracking which also contributes to the decline in the ratio of isomerization to cracking. Augmentation in this ratio may result in blocking of pores and a prominent acceleration in the degradation of catalyst life. Thus, with increase in temperature, the rate of conversion increased which is also accompanied with cracking at higher temperature and lowered selectivity to iso-paraffins.

Study on the effect of LHSV on the hydroisomerization reaction is also studied altering the LHSV rate at constant temperature 310 °C and pressure 5 MPa and the results are shown in Figure 7(b). At low LHSV 0.5 h⁻¹, presence of n-C18 was found to be a relatively negligible amount when compared to its presence at higher velocities. Adding on with point that the products with boiling point range falling between 288 and 281 °C, higher conversion rate was confirmable. At 0.75 h⁻¹ LHSV, conversion was good but with a small observable amount of C18 hydrocarbons falling in the boiling point range 300 and 315 °C. At 1.0 h⁻¹ LHSV, the conversion is lower that it showed a higher intensity peak at the concentration of C18 hydrocarbons. So, at given temperature and pressure, hydro-isomerization is favoured at a considerably lower LHSV.

### Table 4. Numerical data of the GC-MS Analysis on conversion and presence of straight chained and isomerized alkanes in percentage and ratio by SAPO catalyst.

<table>
<thead>
<tr>
<th>Oper. Temp. (°C)</th>
<th>Conversion (%)</th>
<th>mono branched i-16</th>
<th>mono branched i-C16</th>
<th>iso-alkanes</th>
<th>n-alkanes</th>
<th>Cracking</th>
<th>Total cracking</th>
<th>Isomerization/.cracking</th>
<th>iso-alkanes/n-alkanes</th>
<th>Selectivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>270</td>
<td>43.5</td>
<td>13.7</td>
<td>8.7</td>
<td>28.2</td>
<td>4.8</td>
<td>2.4</td>
<td>7.2</td>
<td>3.9</td>
<td>5.9</td>
<td>64.8</td>
</tr>
<tr>
<td>290</td>
<td>79.1</td>
<td>17.9</td>
<td>9</td>
<td>48.9</td>
<td>16.6</td>
<td>5.3</td>
<td>22</td>
<td>2.2</td>
<td>2.9</td>
<td>61.9</td>
</tr>
<tr>
<td>310</td>
<td>91</td>
<td>23.2</td>
<td>10.4</td>
<td>58</td>
<td>18.6</td>
<td>5.8</td>
<td>24.4</td>
<td>2.4</td>
<td>3.1</td>
<td>63.7</td>
</tr>
<tr>
<td>330</td>
<td>94.3</td>
<td>18.3</td>
<td>8.7</td>
<td>61.5</td>
<td>19.2</td>
<td>6.3</td>
<td>25.4</td>
<td>2.4</td>
<td>3.2</td>
<td>65.2</td>
</tr>
</tbody>
</table>

Copyright © 2021, ISSN 1978-2993

### 4. Conclusion

A unique mesoporous crystalline structure of SAPO under structural representation as SAPO-11 support is obtained. The prepared materials are analysed and tested under the performance in the hydro-treating condition after activation with H₂ atmosphere. The XRD analysis of the synthesized samples indicated that the sample prepared under ultrasonic method had high degree of crystallinity when compared to the other samples prepared with hydrothermal and ultrasonic methods combined together. The loss of crystallinity in the combined methods is mainly due to the release of high amount of heat during the oxidation process. The acid site concentrations of the prepared SAPO-11 catalysts are analysed using FTIR spectroscopy. It is observed that the ratio of medium strength Bronsted acid sites to the strong Lewis acid sites is higher for the samples synthesized by ultrasonic method. The higher Bronsted acid sites also accounted for a higher catalytic activity. The ratio of acid sites has comparatively less in the samples produced under both ultrasonic and hydrothermal conditions combined together. The SEM analysis is carried out in the produced samples to analyse the size, growth and formation of the crystals in the SAPO-11 catalyst. Also, it is observed that the crystal shape and size are relatively crystalline in the samples produced under ultrasonic method. The crystal formation and the growth of the crystal construction are better for the SAPO-11 material which prepared under ultrasonic method. Also, the results revealed that catalyst samples synthesized with n-PA as template exhibited high...
crystallinity and uniform distribution of acid sites over the larger surface area relative to the samples synthesized using other templates. This also attributes to the high conversion efficiency in the hydrodeoxygenation of the sunflower oil into desired large-chain hydrocarbons which are consists of branched hydrocarbons or unsaturated straight-chain hydrocarbons. Thus, the synthesized Pt/SAPO-11 catalysts can be used for various applications such as hydrocarbon processing and conversion of vegetable oil into bio-fuel with enhanced properties of fuel to use as replacement in conventional diesel.

Acknowledgement

The authors thank the SEED Grant Research Scheme (KEC/R&D/SGRS/05/2020), KVIT Trust and FIST-DST, Department of Science and Technology (SR/FST/COLLEGE-096/2017), India for the financial support.

References


Copyright © 2021, ISSN 1978-2993


