1. Introduction

Zeolites, a widely utilized material for various catalytic applications, finds its major application in petroleum refining. Commercially, zeolites are readily available in the micro particle form, whereas its nano particle counterpart is reported to show superior performance owing to a greater surface area and faster diffusion of the reactants and products inside the catalyst. However, there are certain limitations associated with using zeolite nano particles. Firstly, because of their high surface energy, the nano particles tend to agglomerate and hence the accessibility to the active sites is hindered. Secondly, nano particles are difficult to recover after the catalytic test. Owing to the small size of the particles, sometimes, extra effort may be required involving filtration and centrifugation to separate the catalyst from the reaction mixture. To overcome these limitations, there has been growing research on shaping the zeolite particles into fibrous form. The fibers provide an effective way to take advantage of the nanoscale, plus providing weight and volume to the catalyst for easier recovery. For example, Pham-Huu et al. reported superior performance of beta zeolite nanofibers compared to commercial beta zeolite particles for benzoylation of anisole. Liu et al. compared the catalytic activity of commercial micro ZSM-5 particles to ZSM-5 nanoparticles and ZSM-5 fibers for iso-butane cracking. They reported superior conversion, selectivity and better catalyst lifetime of the ZSM-5 fibers compared to the particulate form. The superior activities in both the latter cases were attributed to higher accessibility of the active sites owing less agglomeration. However, only a few reports have been reported till now which provide a comparison of zeolite nano particles to zeolite fibers for catalytic applications. This in itself opens new doors for research into this field related to zeolite fiber fabrication, its characterization and consequent catalytic testing.

Several methods may be adopted to produce zeolite fibers including hydrothermal synthesis, self-assembly techniques and electrospinning. Among these, electrospinning is reported to be a versatile, flexible, scalable yet economical method for fiber production. The method contains a syringe which is filled with an electrospinning solution. The solution may be a combination of zeolite and a polymer. The polymer is essential to provide viscosity for electrospinning. The syringe needle is fixed at a certain electrospinning distance from the collector. Both are connected together through a high voltage supply. The power drives the solution in the syringe to the tip of the needle where is emerges as fine jet from the liquid droplet, which consequently decreases in diameter on reaching the
The flexibility of this method allows us to incorporate certain elements in the electrospinning solution, such as metallic salts. This in turn seems promising in fabricating catalytic fibers which incorporate different elements on zeolite as a support, such as necessary in hydrocracking. In hydrocracking, heavier hydrocarbon molecules are broken into lighter fractions in the presence of hydrogen and a catalyst. However, fabrication and testing of these fibers is still a new research field with only recently the prospect of fabricating and testing the hydrocracking fibers utilizing micro zeolite particles was adopted by our group.

Apart from agglomeration and recovery issues, another major challenge is the zeolite nanoparticle availability. Hydrothermal synthesis of these nanoparticles is usually challenging with rather high centrifugation speeds and times required to recover the particles from the source mixture. Moreover, direct synthesis with a high Si/Al ratio for certain zeolite types such as zeolite-Y is practically impossible and dealumination of the nanoparticles become impractical due to destruction of the crystal structure. A viable solution to this is reducing the particle size of high Si/Al zeolites through ball milling. Recently, Zhuman et al. reported ball milling of high Si/Al ratio zeolite-Y, a viable candidate material for cracking and hydrocracking applications. They used carbon nanostructures (CNS) as a damping material while ball milling zeolite-Y to retain its crystallinity and structure. The CNS was removed after the ball milling process to obtain pure zeolite-Y nano particles with high crystallinity index.

In this study, we have utilized zeolite-Y nanoparticles to form hydrocracking fibers through the electrospinning method. Apart from the acidic zeolite support, hydrocracking essentially involves hydrogenation/dehydrogenation components such as nickel oxide and tungsten oxide. The incorporation of the hydrogenation component necessitates an optimum combination of solution and process parameters for the electrospinning of smooth and continuous composite fibers. A detailed study on the fabrication and hydrocracking testing of zeolite-Y micro hydrocracking fibers and particles has already been reported by our group. To the best of our knowledge, no study has been reported till now on the fabrication of hydrocracking fibers made from nano-zeolites. Moreover, it becomes imperative to study and compare the catalytic activities of hydrocracking fibers made from nano zeolite with the hydrocracking fibers made from zeolite-Y micro particles. Further, the fibrous catalysts have been compared with nano zeolite-Y hydrocracking particles for n-heptane hydrocracking. This study will further contribute to the field of catalysis, making significant usage of zeolite in the fibrous form, specifically utilizing nano zeolites for electrospinning zeolite fibers, a field which is still in its infant stages.

2. Experimental

2.1 Materials

Polyvinylpyrrolidone (PVP, $M_w = 1300000$), nickel(II) acetate tetra hydrate (NiAc), ammonium metatungstate hydrate (AMT) and ethanol were purchased from Sigma-Aldrich (St. Louis, MO). Zeolite-Y (CBV 720, BET surface area of 780 m$^2$ g$^{-1}$) was purchased from Zeolyst.

2.2 Catalyst preparation

Nano zeolite Y was produced through the ball milling method as reported in, with a BET surface area of 522 m$^2$ g$^{-1}$. Three catalysts were prepared namely, hydrocracking fibers made from nano zeolite Y, hydrocracking fibers made from as received micro zeolite Y and hydrocracking particles made from nano zeolite Y$^{24}$ particles. Electrospinning was used for fabricating the fibers. An electrospinning solution incorporating zeolite, AMT and NiAc was prepared as follows: NiAc was first dissolved in DI water, after which tungsten salt was allowed to dissolve in the NiAc-water solution. After the salt was completely dissolved, zeolite was added and sonicated (using a BRANSON 1510 model) for about 8 hours to homogeneously disperse the zeolite in the metal salt solution. The ratio of zeolite : NiAc : AMT was kept as 10 : 1 : 15. This ratio was chosen in accordance to literature studies where the Ni : W atomic ratio of 1 : 3 in the overall catalyst showed the highest hydrocracking activity. Zeolite content in the overall catalyst was 87 wt%. After sonication, the suspension was stirred overnight. PVP was then added in a zeolite : PVP ratio of 1 : 1 by weight, and the electrospinning solution was further stirred for 5 hours or more, until a homogeneous suspension composed of zeolite particles- nickel salt- tungsten salt-PVP in water was obtained. The viscous solution was transferred to a syringe. A 22 gauge syringe needle and a Nanon-01A electrospinning setup (MECC, Japan) were used. Solution flow rate, electrospinning voltage and electrospinning distance of 0.5 ml h$^{-1}$, 28 kV and 15 cm were used respectively. The as spun NiAc-AMT-zeolite-PVP composite fibers were collected and dried in a furnace at 80 °C for 6 hours to remove any moisture, following which the fibers were calcined at 550 °C at a heating rate of 1 °C min$^{-1}$ for 2 hours under ambient conditions. This step removes the polymer and decomposes the salt for pure zeolite Y-NiO-WO$_3$ fibers. The calcination temperature of 550 °C was chosen as per previous literature studies which confirmed the removal of polymer and decomposition of the salts at this temperature.

Hydrocracking catalytic particles were also prepared from nano zeolite-Y by the wet impregnation method using similar hydrogenation metal and zeolite compositions for comparison with its fibrous counterpart. Because the particles were not to be electrospun, no polymer was added to it and the impregnated nano zeolite Y was post treated through a similar procedure as adopted for the as-spun fibers: they were first dried at 80 °C, followed by calcination at 550 °C. For convenience, we will be referring to the hydrocracking catalysts as follows: HDC NP for hydrocracking particles prepared from nano zeolite Y and HDC NF and HDC MF for hydrocracking fibers prepared from nano zeolite Y and as-received micro zeolite-Y particles respectively.

2.3 Catalyst characterization

Phase identification and structural characteristics of the prepared catalysts were studied through powder X-ray diffraction (PXRD) and a simultaneous thermogravimetric-Fourier
transform infrared spectroscopy (TGA-FTIR) respectively. PXRD patterns were acquired at room temperature using an X-ray diffractometer (PANalytical, Emperean) operating at 45 kV and 40 mA with Ni-filtered CuKα (λ = 1.5056 Å) radiations in 5–50° half angle range. TGA-FTIR was carried out using a NETZSCH TG 449 F3 Jupiter-Thermogravimetric Analyzer, connected to a Bruker FTIR for gas phase combustion products analysis. The sample was heated from room temperature to 1000 °C using a heating rate of 30°C min⁻¹ under a continuous flow of 20 ml min⁻¹ nitrogen. The infrared spectra were acquired in the range of 4500–700 cm⁻¹. The catalysts were characterized for their morphology through high resolution scanning electron microscopy (HRSEM) using Nova nano field-enhanced SEM operating at 5 kV. SEM samples were gold coated (±5 nm thickness) using a precision etching coating system (Gatan Model 682, Germany). A high resolution transmission emission electron microscope (HRTEM) (Tecnai F20, FEI), operating at 200 kV was also used together with STEM-EDS mode for chemical analysis and EDS mapping. For TEM sample preparation, the catalysts were first sonicated in ethanol for 15 minutes before depositing them onto a carbon coated copper grid. Zeolite acidity zeolite were measured by temperature-programmed desorption of ammonia (NH3-TPD) method. 100 mg of catalyst sample was pretreated in nitrogen at 500 °C for 1 h, cooled to 100 °C and adsorbed NH3 for 30 min. The excess ammonia was flushed by pure nitrogen gas at 100 °C for 45 min, after which temperature-programmed desorption started at a rate of 10°C min⁻¹ from 100 °C to 500 °C. Nitrogen adsorption/desorption experiments were conducted using Brunauer, Emmett and Teller (BET), NOVA®-e Series Model 25 Quantachrome Instruments. The catalyst was first degassed at 300 °C for 4 hours in order to desorb the impurities on the surface. The surface area of the catalysts was calculated through multipoint BET method. Data in the relative pressure range of 0.05–0.30 was used for this purpose.

2.4 Catalytic test

The feed molecule used in the catalytic test is n-heptane (Aldrich, >99 mol%). Catalytic tests were carried out in a continuous fixed-bed down-flow reactor with an inner diameter of 9 mm, operating at a total hydrogen pressure of 5 bars. The catalysts were used as prepared, without forming any extrudate. 500 mg of catalyst was loaded into the reactor sandwiched between quartz wool. For each new test, a fresh catalyst was used, where it was first kept in hydrogen stream for 3 h at 550 °C in order to pre-reduce active metal present in the fiber or particles. Feed was introduced thereafter, and two testing temperatures were chosen for the study, 350 °C and 400 °C. Weight hourly space velocity (WHSV) equal to 4 h⁻¹ with a hydrogen to hydrocarbon molar ratio of 3 : 1 was used. The first sample obtained after 1 hour was flushed out and discarded as drain, after which liquid and gaseous product samples were collected for 3 hours at each 1 hour interval. Liquid products were analyzed by an off line gas chromatography mass spectrum (GC MS), GC 7890B-MS5977A equipped with a 5 column nonpolar column. Nitrogen was used as a carrier gas. Gaseous products were analyzed offline using a 7890A GC system, 8-column RGA with a FID detector for hydrocarbon analysis.

The spent catalyst was treated to remove the absorbed hydrocarbons through toluene extraction after which thermogravimetric analysis (TGA) was performed in order to get the %coke formed. PerkinElmer TGA 4000 was used for this purpose where spent catalyst samples (20–30 mg) were placed in a sample pan and heated from room temperature to 1000 °C at a rate of 10 °C min⁻¹ with an oxygen flow of 20 ml min⁻¹.

Catalytic activity is expressed in terms of total % conversion, which is defined as follows:

\[
\text{% total conversion} = \frac{X_i - X_f}{X_i} \times 100
\]

where \(X_i\) is the initial heptane/feed mass, \(X_f\) is the final heptane mass. \(X_c\) is mass of total cracked products, \(X_d\) is mass of individual product, \(X_b\) is the mass of heptane isomers and \(X_o\) is the mass of total olefins formed. All the percentages mentioned in the results are wt% until and unless stated otherwise.

3. Results and discussion

3.1 Catalyst characterization

The hydrocracking catalysts were studied for their morphology and structural characteristics through various techniques. Fig. 1a shows the physical appearance of the as spun and calcined HDC NF. The yellowish color is characteristic of the m-WO₃ phase. Similar color was observed for the HDC MF and HDC NP (Fig. S1†). It is imperative that the fabricated fibers retain their structural stability and do not disintegrate into particles thereafter. Therefore, fiber integrity was tested by sonicating the fibers in ethanol for 15 minutes. It can be seen from Fig. 1b that the fibers do not disperse in solution as reported by pure zeolite fibers in. Another interesting feature of the HDC NF is that the fibrous form occupy more volume compared to the HDC NP. This can be observed in Fig. 1c where equal amounts of catalyst, 200 mg were put into a tube. The higher volume of the HDC NF facilitates easier recovery of the catalyst, as nano particle recovery is also one of the major problems reported.

Fig. 2a shows the TGA graph of as spun HDC NF together with the corresponding FTIR spectra shown in Fig. 2b and c.
Decomposition of NiAc, AMT and PVP is taking place at different temperatures. Dehydration of NiAc and hydrolysis of acetate groups takes place at 200–250°C. The side groups of PVP breakdown during the same temperature range. The sharp CO2 peaks at 345–350°C are due to the combustion of the evolved gaseous species. Major decomposition of the tungsten salt, AMT takes place in the temperature range of 350–450°C. The rest of the decomposition is due to evolution of any secondary products or gases from the fiber.

The XRD patterns for ball milled nano zeolite Y and as received zeolite Y particles are shown in Fig. 3a. As reported by Bota et al., the zeolite structure after ball milling is still retained and all major zeolite peaks below 2θ 20° were registered. The XRD patterns for the three hydrocracking catalysts, HDC NF, HDC NP and HDC MF are shown in Fig. 3b with some major WO3 peak indexed. WO3 registers no peaks before 2θ 20°. The fibers registered similar XRD patterns to the HDC NP, confirming that the electrospinning method does not bring about any structural changes to the catalyst. All major zeolite peaks, as well major peaks of m-WO3 (ref. 25) were registered. All the peaks before 2θ 20° are essentially from zeolite Y. NiO peaks were not detected as NiO was present in very small amounts in the catalyst (less than 5 wt%) and therefore it was not practically possible to detect this phase using the current XRD equipment. Never the less, its presence was detected by EDS as discussed later.

Fig. S2a† shows SEM image of as received zeolite Y micro particles while Fig. S2b† shows the TEM image of the reduced size, ball milled nano zeolite Y particles. The micro particles range from about 500 nm up to 1 μm. These micro zeolite particles were used in the preparation of HDC MF. The micro fibers ranged in diameter from about 1.3 μm to 2.3 μm as shown in Fig. S3.† The ball milled nano zeolite Y particles were used in the preparation of HDC NP and HDC NF. Fig. 4 shows HRSEM images of calcined HDC NF and HDC NP. The micro fibers ranged in diameter from about 1.3 μm to 2.3 μm as shown in Fig. 4a reveals uniform, continuous HDC NF with fiber diameters varying from about 600 nm to 700 nm. A single fiber strand in Fig. 4b shows variation in fiber diameter. A magnified image of the HDC NF is shown in Fig. 4c where a combination of small and large crystallites on the fiber surface can be observed. Similar was observed with HDC MF (Fig. S3b†). Because the nano particles possess high surface energy, the HDC NP tend to agglomerate as seen in Fig. 4d. During calcination, the hydrogenation elements come to the surface of the catalyst. This decreases the surface area of the overall catalyst as components such as WO3 have a low BET surface area of around 10 m2 g−1. Fig. 5a shows the N2 adsorption/desorption curves and the BET surface areas for HDC NF, HDC NP and HDC MF. The HDC MF and HDC NP register similar surface areas of 290 and 296 m2 g−1 respectively while HDC NF gave a surface area of 242 m2 g−1. These surface areas are higher than those reported in the literature for various hydrocracking catalysts.
interesting feature is that the mesoporosity\textsuperscript{33} of the catalyst is retained after it has been formed into a hydrocracking catalyst, for both particles and fibers as evident from the hysteresis loop of the graphs. This is an important feature of zeolite catalysts as their mesoporous character improves the mass transfer properties between the acidic and the metal function. Hence, hierarchical zeolites have known to be more promising for catalytic applications because they facilitate reactants and products within the catalyst and allow synergic metal-support interaction for optimal process efficiency.

Fig. 6a shows TEM image of a single HDC NF. As these fibers were sonicated for 15 minutes in ethanol before preparing the TEM sample, this further attests to the mechanical integrity of the fibers. Again, the different sizes of crystallites are quite prominent in the TEM image. It is difficult to know the exact location of zeolite and the hydrogenation components as due to the bulk EDS technique, any EDS point gives all the elements in the EDS spectrum. However, we can have an idea about the metal distribution through EDS mapping. Fig. 6b shows STEM-EDS analysis on a small part of the fiber strand as highlighted by the yellow box. All elements were seen to be uniformly distributed within the fiber with the silicon, aluminum and oxygen distribution attesting zeolite uniformity and the nickel, tungsten and oxygen distribution attesting the hydrogenation elements uniformity. Therefore, the electrospinning technique provides the advantage of a uniform elemental distribution.
within a single fiber which is desirable for a hydrocracking catalyst because of close proximity of acid and hydrogenation sites for shorter diffusion lengths of reactant molecules between the zeolite and hydrogenation components.\textsuperscript{15}

### 3.2 Catalyst activity

Table 1 compares the total conversion and major product selectivity for HDC NF, HDC NP and HDC MF at 350 °C and 400 °C. A detailed discussion is provided in Sections 3.2.1 and 3.2.2 where HDC NF conversions and selectivity are compared to HDC MF and HDC NP respectively.

#### 3.2.1 HDC NF compared to HDC MF

Fig. 7a and b compares the total conversion of \textit{n}-heptane for HDC NF and HDC MF at 350 °C and 400 °C respectively on total TOS for 3 hours. At both temperatures, HDC NF shows higher conversion compared to the micro counterpart. This may be attributed to the unique nano-structure and morphology of the catalyst itself,
whereby it can provide more accessibility to the feed molecules to the catalytic active sites with increased surface to volume ratio. The high conversions of the HDC NF till the end of the reaction infer that there are still more active sites for cracking available at the end, while the conversions on total TOS for HDC MF infer that the catalyst is deactivating with time. The HDC NF showed superior catalytic stability where the conversion dropped from 99.3 wt% to only 98.8 wt%, and from 97.3 wt% to

![Fig. 6](a) TEM image of HDC NF (b) STEM-EDS analysis on part of a HDC NF showing uniform distribution of elements within the fiber even in a small fiber portion.

<table>
<thead>
<tr>
<th>Catalyst/testing temperature</th>
<th>Conversion of (n)-heptane (wt%)</th>
<th>Selectivity for (n)-C6 (wt%)</th>
<th>Selectivity for i-C6 (wt%)</th>
<th>Selectivity for C3 (wt%)</th>
<th>Selectivity for C4 (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HDC NF [350 °C]</td>
<td>98.8</td>
<td>19.8</td>
<td>39.3</td>
<td>2.3</td>
<td>9.9</td>
</tr>
<tr>
<td>HDC NF [400 °C]</td>
<td>96.8</td>
<td>50.7</td>
<td>18.9</td>
<td>2.3</td>
<td>1.4</td>
</tr>
<tr>
<td>HDC NP [350 °C]</td>
<td>98.3</td>
<td>20.8</td>
<td>33.7</td>
<td>3.7</td>
<td>6.1</td>
</tr>
<tr>
<td>HDC NP [400 °C]</td>
<td>95.7</td>
<td>42.0</td>
<td>24.2</td>
<td>3.1</td>
<td>2.4</td>
</tr>
<tr>
<td>HDC MF [350 °C]</td>
<td>63.1</td>
<td>0.5</td>
<td>0.9</td>
<td>20.9</td>
<td>24.9</td>
</tr>
<tr>
<td>HDC MF [400 °C]</td>
<td>31.8</td>
<td>0.6</td>
<td>0.7</td>
<td>19.0</td>
<td>24.9</td>
</tr>
</tbody>
</table>
96.8 wt% at 350 °C and 400 °C respectively. While for HDC MF, there was a sharp decrease in conversion since the first hour at 350 °C, where the total conversion dropped from 88.1 wt% to 63.1 wt% by the end of total TOS. Similar was observed at 400 °C, where the conversion dropped from 38.1 wt% to 31.8 wt% at the end of the reaction. However, the drop in conversion was less drastic in the latter case. Moreover, higher conversions were registered at a lower temperature. This might be explained in terms of coke formation. Coke formation has a strong impact on the performance of the catalyst. After the production of unsaturated molecules like olefins, the coke formation reactions rather proceeds easily. The percentages of coke calculated from TGA graphs (Fig. S4†) were as follows, 0.40 wt% and 1.05 wt% for HDC NF at 350 °C and 400 °C respectively while 1.46 wt% and 1.10 wt% for HDC MF at 350 °C and 400 °C respectively. It is interesting to see that similar percentages of coke were formed for both HDC NF and HDC MF at 400 °C, but the HDC NF showed far superior conversion than HDC MF at the same temperature. At a higher temperature, it is postulated that coke forms immediately and hence the catalyst deactivates faster, registering lower conversion from the beginning. This might be a reason for low conversion of HDC MF at 400 °C. The deactivation of zeolite-based catalysts by coke is caused either by poisoning of acid sites or by pore blockage. In the former case, one coke molecule usually blocks one or more active sites, affecting the activity linearly. It is quite probable that the coke formed with HDC MF blocked the catalyst pores due to slower diffusion kinetics and hence decreased the catalyst activity gradually. With the HDC NF, due to faster kinetics, the coke formed did not hinder catalyst activity. Also, less coke is formed with HDC NF compared to HDC MF at 350 °C, even though HDC NF gave greater conversions. The greater coke percentage might be the reason for pore blockage and hence faster deactivation in HDC MF.

From Table 1, it can be observed that the major products for HDC MF are butane (C4) and propane (C3). These results are consistent with the literature where n-heptane usually cracks through a bi-functional mechanism with an olefin forming from dehydrogenation on the hydrogenation site, which diffuses to the acidic site and gets cracked through β-scission mechanism. Moreover, from Fig. 8, it might be inferred that the HDC MF is behaving not only as a cracking catalyst, but also

![Graphs showing total conversion for HDC NF and HDC MF at 350 °C and 400 °C.](image1)

![Graphs showing evolution of iC7 with TOS for HDC NF and HDC MF at 350 °C and 400 °C.](image2)
Fig. 9  % olefin formation for HDC NF and HDC MF at (a) 350 °C and (b) 400 °C.

Fig. 10  Comparison of total conversions for HDC NF and HDC NP on total TOS at (a) 350 °C and (b) 400 °C.

Fig. 11  Isomerization to cracking ratios of HDC NF and HDC NP at (a) 350 °C and 400 °C.
as a hydro isomerization catalyst. The amount of heptane isomers (iC7) obtained with the HDC MF is much higher than that registered by HDC NF. This typical product feature of high iC7 has been observed previously when n-alkanes were hydrocracked with low to medium overall conversions. Another important feature is the increase in iC7 with TOS for both HDC NF and HDC MF. A possible reason for this is again deactivation of the catalyst. As the catalyst deactivates, isomerization of the feed increases and cracking decreases.

It is interesting to note that the major products obtained with the HDC NF are n-hexane (n-C6) and iso-hexane (i-C6). The % selectivity changes with a change in temperature, where a higher temperature of 400 °C is producing more n-C6 concurrently with a lower amount of i-C6 when compared to 350 °C.

Fig. 9a and b compares the percentage of unsaturated products formed by HDC NF and HDC MF at 350 °C and 400 °C respectively. Less olefin formation is registered with the HDC NF hinting towards either more accessibility and/or less deactivation of the hydrogenation sites in catalytic fiber. Interestingly, the percentage of olefins formed with both the catalyst is quite small ranging from about 0.6 wt% to about 2.5 wt% of the total converted products. The major olefins obtained from GC were benzene derivatives and toluene.
3.2.2 HDC NF compared to HDC NP. Fig. 10a and b compares the total conversion of \( n \)-heptane for HDC NF and HDC NP at 350 °C and 400 °C respectively on total TOS. Both the catalysts registered high conversions reaching more than 95 wt%. Unlike HDCMF, it is difficult to compare and conclude that one is giving superior conversion over the other, because the percentage of total conversion is quite similar for both HDC NF and HDC NP during each hour at both the temperatures. Never the less, it is prominent that between the two catalysts, HDC NP is showing greater catalytic stability with less deactivation. At 350 °C, the conversion for HDC NP sharply drops after the first hour from 99.1 wt% to 98.4 wt%, while for HDC NF, it drops from 99.3 wt% to only about 99.25 wt%. At the end of the reaction, the HDC NF registered a conversion of 98.8 wt% while HDC NP gave a total conversion of 98.3 wt%. At 400 °C, the stability of the HDC NF is much more prominent with a conversion decrease from 97.3 wt% to 96.8 wt% at the end of total TOS. HDC NP showed a sharp decrease in conversion from 98.5 wt% to 95.9 wt% after the first hour. This signifies that the HDC NP catalyst deactivates faster and hence registers a sharp decrease in conversion from the beginning. Because the preparation procedure for the developed hydrocracking catalysts is similar in terms of the way the nickel and tungsten salts were impregnated onto zeolite and the fact that the percentages of the metal oxides was kept similar for all catalysts, the acidity measured for HDC NF can said to be representative for HDC NP as well. Because the preparation procedure for the developed hydrocracking catalysts is similar in terms of the way the nickel and tungsten salts were impregnated onto zeolite and the fact that the percentages of the metal oxides was kept similar for all catalysts, the acidity measured for HDC NF can said to be representative for HDC NP as well. According to NH\(_3\)-TPD results (Table S1, Fig. S5†), the catalyst has total acidity of 598 \( \mu \)mol g\(^{-1}\), and hence there are enough acidic sites for the catalyst to crack \( n \)-heptane. Hence, this deactivation of HDCNP might be related to the percentage of coke formed which in turn might be responsible in blocking of the acidic sites. Higher percentage of coke is registered for HDC NP (Fig. S4†), 1.0 wt% at 350 °C and 2.5 wt% at 400 °C when compared to HDC NF (0.4 wt% and 1.05 wt% at 350 °C and 400 °C respectively). Another approach to study the deactivation might be adopted by calculating the isomerization to cracking ratios of the two catalysts. With subsequent deactivation, the isomerization reaction increases with more isomerized products than the cracked products. From Fig. 11, it can be seen that the isomerization: cracking (iso: cracking) ratio for HDC NP at 400 °C increases rapidly till the end of the reaction while for HDC NF, the increase in ratio is gradual. This steep increase for HDC NP can be attributed to an increase in deactivation of the catalyst, most probably due to high percentage of coke formation as already discussed above. At 350 °C, the iso: cracking ratio for both HDC NF and HDC NP increase after the first hour, after which the ratio almost becomes constant suggesting much lower deactivation of the catalysts. This can be related to the lower amount of coke formed for both HDC NF and HDC NP at 350 °C.

From Table 1, it can be seen that the major products for both HDC NP and HDC NF are quite similar with large percentages of \( n \)-C\(_6\) and i-C\(_6\) obtained. Fig. 12 highlights the distribution of liquid products after each hour for 350 °C and 400 °C. It can be seen that the production of heptane isomers is much less compared to \( n \)-C\(_6\) and i-C\(_6\), which implies that the main reaction is cracking rather than isomerization unlike in HDC MF discussed in the previous section. Secondly, higher percentage of iC\(_7\) is obtained for HDC NP compared to HDC NF. There is a slight difference in selectivity for both HDC NP and HDC NF at the two temperatures. At 350 °C, the percentage of i-C\(_6\) obtained is greater than \( n \)-C\(_6\), while at 400 °C, more \( n \)-C\(_6\) is obtained than i-C\(_6\). One possibility is that after the production of \( n \)-C\(_6\), the molecule may isomerize to i-C\(_6\). At a lower temperature, this happens readily. While at 400 °C, most of the \( n \)-C\(_6\) does not isomerize and remains as \( n \)-C\(_6\). Further comparing the selectivity at 400 °C, it can be observed that the HDC NF register more percentage of \( n \)-C\(_6\) while HDC NP gave more percentage of
i-C6. This might be attributed to an increased isomerization of n-C6 with catalyst deactivation.

Fig. 13 compared the amount of olefin formed at the two testing temperatures for HDC NF and HDC NP. Interestingly, higher percentages of olefin are formed with HDC NP, compared to both HDC NF and HDC MF. Thus the fibrous form provides a close proximity of hydrogenation and acidic sites due to the uniform distribution of the elements compared to the particles.

The superior catalytic performance of nano zeolite-Y for the hydrocracking application suggests that the study may be extended for other applications whereby advantage of the nanostructure of the zeolite can be taken together with a uniform elemental distribution obtained through electrospinning. For example, it may be applied in producing hydrocracking catalysts with other hydrogenation components such as Co-Mo/nano Y, Ni-Mo/nano Y and for Mo/nano ZSM-5 for methanol oxidation and many others.

4. Conclusion

Hydrocracking fibrous catalysts utilizing nano zeolite Y were prepared from the electrospinning method whereby a suspension of nano zeolite Y, nickel and tungsten salts and PVP polymer was electrospun. The nano zeolite Y-NiO-WO3 fibers were characterized for their morphology, mechanical strength, structural properties and elemental analysis through various techniques including HR-SEM, HR-TEM, BET, XRD and STEM-EDS. The fibers showed good mechanical stability in ethanol after sonication with a uniform distribution of the hydrogenation and acidic component within the fibers. These fibers were compared for n-heptane hydrocracking activity with hydrocracking fibers made from micro zeolite Y and hydrocracking particles made from nano zeolite Y at 350 °C and 400 °C. The nano zeolite Y hydrocracking fibers showed superior conversions of about 98.8 wt% and 96.8 wt% compared to hydrocracking micro zeolite Y fibers with 63.1 wt% and 31.8 wt% at 350 °C and 400 °C respectively. Similar conversions for nano zeolite fibers and nano zeolite Y particles were registered, however, the former showed better catalytic stability over the total time on stream of 3 hours. The fibrous form gives an added advantage of easier recovery of the catalyst. In addition, less coke was formed with the nano zeolite Y fibers compared to hydrocracking nano particles. This in turn was attributed to much lesser deactivation of the fibrous catalyst. These results for nano zeolite Y-NiO-WO3 composite catalysts prove to be promising for utilizing zeolite in the nano form for hydrocracking application and further, this study can be extended to a wider context for different zeolite types for various catalytic applications where zeolite is often used in conjunction with other metals or metal oxides.

Conflicts of interest

There are no conflicts to declare.

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