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Biodiesel from batch and continuous oleic acid esterification using zeolite catalysts

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Abstract

The esterification of oleic acid with ethanol was studied in both batch and continuous conditions in a fixed-bed reactor over FAU-type zeolites prepared from shale rock. The addition of Co-Ni-Pt to the zeolite increased the catalyst activity over the entire reaction temperature range. Experiments confirmed that esterification follows pseudo first-order kinetics. Thermodynamic analysis and Thiele modulus calculations show that the reaction is kinetically controlled in batch conditions, while diffusional limitations occur at higher flow rates when conducted in continuous mode. The maximum oleic acid conversions were recorded as 93% for batch and 89% for continuous, which exceed those of any analogous studies.
Keywords: Biodiesel, oleic acid esterification, zeolite, continuous fixed-bed reactor

1. Introduction

Biodiesel is a carbon neutral fuel source that has the potential to provide almost unlimited renewable energy. Untreated biodiesel (natural oils) e.g. sunflower oil, typically consists of fatty acids that require (trans)esterification to produce methyl or ethyl esters that have appropriate viscosity and burning rate (cetane number) for use in existing diesel compression engines thereby making them suitable as fuels. Biodiesel is traditionally prepared using homogeneous catalysts e.g. sulfuric acid. Heterogeneous catalysts have the advantage of being easily removed from the biodiesel produced so it is not surprising that more recent research efforts have focussed on the development and testing of such heterogeneous systems.

Oleic acid is present in natural oils and its acid catalysed esterification has become a model reaction in the study of biodiesel formation. Most of the literature around oleic acid esterification reports its reaction with methanol. Homogenous systems using mineral acid/base catalysts have been well studied so recent innovations in this area tend to focus on increasing the product yield by additional reaction treatment. Examples include the use of a water adsorption apparatus, which increases the reaction yield via water removal to enhance the oleic acid:methanol equilibrium towards the ester [1], and the application of ultrasound during reaction [2]. Heterogeneously catalysed oleic acid:methanol reactions have been reported over the following catalysts; Amberlyst-46 [3]; sulfonated carbon [4]; acid ion-exchange polymer resin [5]; titanium dioxide supported on natural phosphate [6]; 12-tungstophosphoric acid supported on mesoporous silica [7]; and solid acid catalysts prepared from Amazon flint kaolin [8]. Ethyl esters have superior properties to their methyl counterparts, evidenced by higher oil solubility, calorific value, cetane number, cold filtering plugging point, cloud point and pour
point. The catalysis of the oleic acid:ethanol reaction has been examined using: sulfonated cation exchange resin [9]; organophosphonic acid-functionalised silica [10]; sulfonated carbonised bamboo ash [11]; Amberlyst-15 [12] niobic acid [13]; calcium ferrite [14]; and zirconia supported 12-tungstophosphoric acid [15].

Zeolites are high surface area solid acids that have been widely used for water treatment and purification, humidity control, and heterogeneous catalysis [16]. Zeolite acidity originates from the charge imbalance between silica and alumina units bound within the stable crystalline aluminosilicate framework; these properties makes zeolites suitable to the acid-catalysed esterification of free fatty acids in biodiesel production, and a number of reviews have been published [17-22]. In addition to their preparation from laboratory based silicate/aluminosilicates reagents, zeolites may also be prepared from a variety of natural/waste materials; the compositions of silicon and aluminium based minerals in these materials is similar to those of aluminosilicate zeolites, and the literature describes examples of zeolites prepared from fly ash [23-33], kaolin [34-45], fly ash-kaolinite [46] and shale [47-51]. There are few reports in the literature on zeolite catalysed oleic acid esterification reactions, and the majority use methanol; zeolite beta [52,53]; mordenite, ZSM-5 and FAU-type zeolite [54]; and ZSM-5 modified with citric acid [55]. Oleic acid esterification with ethanol was studied in batch conditions over zeolite Y [56] and zeolite beta [57], and via pervaporation-assisted continuous esterification over Amberlyst-15 [58] and NaA zeolite membrane [59].

We recently reported the preparation of pure FAU- zeolite from shale rock, and demonstrated that its catalytic activity was comparable to that of commercial Y zeolite in the esterification of oleic acid with ethanol [51]. Here, we expand on our previous experiments using FAU-type zeolite as a catalyst in both batch and continuous conditions, in which the maximum conversion
is enhanced by the straightforward addition of Co-Ni-Pt metals. Kinetic and thermodynamic parameters are also presented.

2. Experimental

2.1 Materials

The following is a list of the materials’ source/supplier and purity; shale rock was collected from the surface of a tilled field on a working farm in county Wexford, Ireland, washed with water to remove all soil residue and dried at 120 °C for three hours; sodium hydroxide (NaOH) pellets, extra pure, Scharlau; sodium silicate (Na₄SiO₄), 99% purity, BDH Chemicals Ltd.; ammonium chloride (NH₄Cl), Sigma Aldrich; absolute ethanol (C₂H₅OH) Sigma Aldrich; phenolphthalein, 2% in ethanol, Sigma-Aldrich; cobalt (II) nitrate hexahydrate, Co(NO₃)₂.6H₂O, Sigma Aldrich; nickel (II) nitrate hexahydrate, Ni(NO₃)₂.6H₂O, Sigma Aldrich; platinum (IV) oxide hydrate, PtO₂.xH₂O, Sigma Aldrich.

2.2 Catalyst preparation

The zeolite was prepared using a procedure based on our previous paper [51]. The washed shale was crushed in a ball mill apparatus, sieved to <90 µm and calcined in air at 800 °C for 4 h to remove organic matter. 10 g of calcined shale was then refluxed with 40 cm³ of 5 M HCl at 85 °C for 4 h (to remove Fe) and the product recovered by filtration. 1 part (by mass) of calcined shale was mixed with 1.5 parts (by mass) of 40 wt % aqueous NaOH solution, heated at 850 °C in air for 3 h in a furnace to get fused shale, which was crushed to powder form after cooling to ambient temperature. 2 g of fused shale, 1 g sodium silicate and 16 g distilled water were added to a polypropylene bottle, stirred at room temperature for 3 h, then aged under static conditions at room temperature for 18 h, and finally hydrothermally treated at 100 °C for 24 h. The product was recovered by filtration.
To convert the prepared zeolite from Na\(^+\) to NH\(_4^+\) form, 90 g of zeolite were added to 250 cm\(^3\) of 2 M ammonium chloride and stirred in a round bottom flask at room temperature for 2 h. The solid was recovered by filtration, washed with distilled water and the ion-exchange procedure was repeated a further two times using 60 g and 30 g, respectively. The solid was again recovered by filtration, washed with distilled water, dried for 12 h at 120 °C and calcined in air at 500 °C for 4 h to give zeolite in H\(^+\) form, hereafter labelled H-FAU.

Before metal loading, the zeolite was dried in an oven at 110 °C for 2 h to remove any absorbed water. Precious metals were added to the zeolite using incipient wetness; solutions were prepared by dissolving the appropriate masses of Co, Ni, and Pt metal precursors in 0.1 M HCl, which were then added to the zeolite to give loadings of 1 wt.% of each metal. After impregnation, the catalysts were dried overnight at ambient temperature, heated for 24 h at 120 °C, and calcined in air at 500 °C for 4 h to give Co-Ni-Pt-FAU.

2.3 Characterization

Nitrogen adsorption/desorption measurements were carried out using a Micromeritics ASAP 2020 Surface Analyser at -196 °C. Samples were degassed under vacuum (p < 10\(^{-5}\) mbar) for 12 h at 350 °C prior to analysis. BET surface areas of the samples were calculated in the relative pressure range 0.05-0.30. Microscopic images were recorded using a JEOL JSM-5600LV scanning electron microscope (SEM). Semi-quantitative chemical analysis was performed by energy-dispersive X-ray spectroscopy (EDAX) using a detector from Oxford Instruments.

2.4 Catalyst testing

2.4.1 Batch reactor
The esterification reaction of oleic acid with ethanol was performed by reflux in a 500 ml batch reactor placed in a thermostatic oil bath under stirring. The desired amount of catalyst was dried before reaction at 130 °C for 2 h. The reactor was loaded with 50 ml (44.75 g) of oleic acid and the desired amount of pre-heated ethanol was then added to give an ethanol to oleic acid molar ratio of 6. Esterification was carried out at reaction temperatures 40, 50, 60 and 70 °C. 5 ml samples were withdrawn from the reaction mixture at 15 minute intervals, and centrifuged for 10 min at 3000 rpm to separate the solid zeolite from the liquid phase.

The supernatant layer was purified by rotary evaporator to remove excess ethanol and water generated during reaction and analysed by titration with 0.1 M KOH, using phenolphthalein indicator, to calculate the acid value (AV) as shown in the following equation;

\[
AV = \frac{\text{ml of KOH} \times N \times 56}{\text{Weight of Sample}}
\]  

(1)

From the acid value, the conversion of oleic acid was calculated for each amount of catalyst as shown in the following equation;

\[
\text{conversion\%} = \frac{AV_{t=0} - AV_t}{AV_{t=0}} \times 100\%
\]

(2)

where:

AV\(_{t=0}\) (acid value of the reaction product at time 0)

AV\(_t\) (acid value of the reaction product at time t)

2.4.2 Continuous fixed-bed reactor

Oleic acid esterification was conducted in continuous mode using a vertical fixed-bed quartz reactor (Fig. 1) with internal diameter 10 mm and height 650 mm. 6.35 g of catalyst pellets
were prepared by compressing the catalyst powder at 2 tonnes for 1 min in a hydraulic press, followed by crushing and sieving to the size range 355 to 500 µm. Ethanol and oleic acid were mixed and preheated (using the same quantities as in the batch reaction), and delivered upwards to the reactor using a Perkin Elmer LC pump. The reaction temperature was controlled by circulating water from a bath. Different flow rates were employed to obtain sufficient residence times between the reactants and catalyst inside the reactor column (30, 60, 90, 120, 150 and 180 mins). The product stream was collected in an accumulation tank, during which 10 ml samples were taken for analysis at each flow rate and temperature. The product was purified and analysed in the same manner as that used in the batch reaction.

3. Results and Discussion

3.1 Characterisation
The XRD patterns for H-FAU prepared from shale rock confirm the faujasite structure; these were discussed in detail by the authors in reference 51 so will not be repeated here. The BET surface area of H-FAU decreased from 571 m²g⁻¹ to 490 m²g⁻¹ after the addition of Co, Ni and Pt metal salts by incipient wetness, and subsequent heating and calcination in air. A decrease in surface area is expected as the Co-Ni-Pt-FAU has undergone twice the period of calcination as H-FAU, thereby causing some loss of the metastable zeolite porosity but, nonetheless, retains a high surface area. SEM images, Fig. 2, show the characteristic micron sized particles, approximately 2 µm, of both H-FAU and Co-Ni-Pt-FAU, while EDAX confirms the presence of cobalt, nickel and platinum in addition to silicon and aluminium in the underlying zeolite.

3.2 Esterification reactions
The esterification of oleic acid with ethanol is used as a test reaction to assess the catalytic activity of the shale zeolite. This is a reversible reaction so an excess quantity of ethanol, 6:1
ethanol/oleic acid molar ratio, is used to enhance conversion. The fractional conversions of oleic acid over H-FAU, Fig 3, increase with reaction time at all temperatures where the maximum conversion increases from 45% at 40 °C to 78% at 70 °C. The addition of Co-Ni-Pt increases the catalyst activity over the entire temperature range such that the maximum conversion increases from 58% at 40 °C to 93% at 70 °C. The activity of the Co-Ni-Pt-FAU catalyst exceeds those of the zeolite Y prepared from kaolin in our previous study, and also that of a commercially sourced Y zeolite in H- form (85% at 70 °C) [45].

As previously noted, most of the literature documenting zeolite catalysed oleic acid esterification uses methanol as the alcohol reagent; maximum conversion rates (of oleic acid) were reported as follows; 86% for zeolite beta [52,53]; 81% for mordenite, 80% for ZSM-5, and 78% for FAU-type zeolite [54]; 80% for ZSM-5 [55]; and 83% for citric acid modified ZSM-5 [56]. For zeolite catalysed oleic acid and ethanol batch esterification. Marchetti et al. reported a maximum conversion of 27% after two hours over Na-Y zeolite at 55 °C using an ethanol:oleic acid ratio of 6.3:1; the low value of 27% is likely to be due to using the Na+ form of zeolite, which is much less acidic than the H+ form [56]. Sun et al. reported a conversion rate of 73.6% over zeolite beta after 10 hours at 78 °C using an ethanol:oleic acid ratio of 20:1 [57]. The zeolite prepared here from shale is more active than either of these catalysts, evidenced by conversions of 59% at 50 °C and 71% at 60 °C, versus 27% at 55 °C over Na-Y [56], and 78% at 70 °C versus 73.6% at 78 °C over zeolite beta [57].

Reaction rate constants were calculated from plots of ln oleic acid concentration versus time, Fig. 4; the gradients are the pseudo first-order rate constants (ethanol in excess) which are then converted to the ‘true’ second-order rate constants by accounting for the concentration of ethanol. The best fits were found to occur up to and including 1 hour reaction time; thereafter
the reaction approaches/reaches equilibrium where the extent of the reverse hydrolysis reaction is such that the measurement of oleic acid alone is insufficient for a meaningful kinetic study. All plots showed excellent linearity (regression coefficient 0.99 or greater). The pseudo first-order and second-order rate constants, Table 2, increase with both reaction temperature and the addition of Co-Ni-Pt to the zeolite. The vast majority of papers in the literature do not report the second-order rate constants. This is unfortunate, as their inclusion would allow a direct comparison of catalyst activities between research groups doing similar studies. The pseudo first-order rate constant is sometimes mentioned but, on its own, is not an accurate means of disseminating catalyst activity for the purpose of comparison, due to the numerical value not accounting quantitatively for the excess alcohol concentration.

The Arrhenius parameters of oleic acid esterification were calculated from the second-order rate constants, Fig. 5, giving activation energies of 28.6 kJ mol\(^{-1}\) for H-FAU and 31.9 kJ mol\(^{-1}\) for Co-Ni-Pt-FAU, Table 1. The slightly higher activation energy over the Co-Ni-Pt-FAU catalyst is surprising, given the associated higher rates of reaction relative to H-FAU. However, activation energy is a measure of the temperature response of an activated reaction and the results here may simply indicate a different reaction mechanism over Co-Ni-Pt-FAU (at least for a portion of reacting molecules). This is somewhat confirmed by the higher value of the Arrhenius frequency factor; 16804 dm\(^3\) mol\(^{-1}\) h\(^{-1}\) for Co-Ni-Pt-FAU versus 3886 dm\(^3\) mol\(^{-1}\) h\(^{-1}\) for H-FAU. It is not possible to make a direct comparison of these Arrhenius frequency factors to those from other studies as most papers make no such mention, and for those that do, the numerical value and units are calculated using the pseudo-first order rate constants. The Arrhenius frequency factor is an approximate measure of the collision rate of reacting molecules. Accordingly, we tentatively attribute the higher Arrhenius frequency factor to the greater affinity of oleic acid to the metal surface(s) (as opposed to the underlying zeolite),
which concentrates the oleic acid solution near the zeolite acid sites thereby increasing the 
reaction rate. (It is important to note that the esterification reaction is catalysed by the zeolite 
acid sites, and not necessarily Co, Ni or Pt.) This increased affinity may be due to the direct 
adsorption/interaction between the oleic acid double bond and the metal phase e.g. Pt. The 
resulting increase in oleic acid concentration near the zeolite surface could also increase any 
ionic interaction between the carboxylic acid group of oleic acid and the zeolite surface, 
providing further reactant concentration and an associated increase in reaction rate.

Activation energies >20 kJ mol\(^{-1}\) are consistent with kinetically controlled reactions, so the 
results here confirm that the oleic acid esterification reaction over FAU-zeolite is kinetically 
controlled i.e. not diffusion limited [60]. The extent of diffusion may be evaluated more 
precisely using the Thiele modulus, \(\phi\), which for a spherical particle of radius \(R\) (such as that 
in the present paper of SEM diameter 2 μm) is calculated using the following equation [61]:

\[
\phi = \frac{R}{3} \sqrt{\frac{k'}{D_{\text{eff}}}}
\]  

(3)

This approach thus combines a theoretical calculation of the effective diffusion coefficient, 
\(D_{\text{eff}}\), with the experimental value of the pseudo first-order rate constant \(k'\). A \(\phi\) value < 0.4 
indicates that the reaction concentration profile does not drop significantly within the catalyst 
pores i.e. the reaction is not diffusion limited. The effective diffusion coefficient was calculated 
using the following equation [62]:

\[
D_{\text{eff}} = \frac{D_{AB} \epsilon_p}{\tau_p}
\]  

(4)

where \(D_{AB}\) is the molecular diffusion coefficient of the limiting reactant A in solvent B, \(\epsilon_p\) is 
the catalyst particle porosity and \(\tau_p\) is the catalyst pore tortuosity. For the study presented here,
the molecular diffusion coefficient of oleic acid in ethanol was calculated using the Wilke-Chang method [63] to be $5.27 \times 10^{-11}$ m$^2$ s$^{-1}$ at 40 °C, which is similar to that calculated for triglyceride in methanol, $6.1 \times 10^{-11}$ m$^2$ s$^{-1}$ [64]. The Thiele modulus values were calculated for each batch reaction using an $\varepsilon_p$ value of 0.345, determined from the nitrogen adsorption void fraction, and a $\tau_p$ value of 4 [62]. The calculated values, Table 1, range from $5.68 \times 10^{-4}$ to $8.01 \times 10^{-4}$, which is well below 0.4 thereby providing further evidence that the reaction is kinetically controlled i.e. not diffusion limited.

The increased reaction rate using Co-Ni-Pt-FAU is also evident when the reaction is conducted on a continuous basis in the fixed-bed reactor. The maximum conversions of oleic acid, Fig. 6, reach 41% at 40 °C and 75% at 70 °C over H-FAU, while the equivalent figures over Co-Ni-Pt-FAU are 54% and 89%, respectively. Pseudo first-order kinetics were confirmed for both H-FAU and Co-Ni-Pt-FAU catalysts when used in the fixed-bed reactor, Fig. 7, which matches with the rate law found in batch. It is interesting to note that the kinetic treatment showed the greatest linearity of fit at lower LHSVs, particularly so in the case of Co-Ni-Pt-FAU. Further evidence of the superior catalytic activity by the addition of Co-Ni-Pt can be seen in Fig. 7, where the absolute reaction rates (measured as the quantity of oleic acid converted) are plotted. Visual inspection of both H-FAU and Co-Ni-Pt-FAU gradients shows that, overall, the reaction rates for most catalysts increase relatively linearly up to one hour reaction time, and, thereafter, there is a noticeable reduction in the degree of increase. This change in response is due to diffusional limitations where the reactant flow rates are sufficiently high to prevent complete reaction (these effects are currently under investigation and will not be discussed here as they are beyond the scope of this paper). However, it is interesting to examine the dynamics of rate changes at times greater than one hour. In the case of H-FAU, the reaction rates decrease slightly between one and two hours at temperatures 40 °C and 50 °C, but increase at 60 °C and
70 °C. For Co-Ni-Pt-FAU, the rates increase at all temperatures over the same range. This effect is due to the higher reaction rate caused by the Co, Ni and Pt on the zeolite; while the contribution by the metals is insufficient to eliminate the diffusion effects entirely, it does nonetheless partially mitigate the diffusional limitations, and increases the absolute reaction rate overall.

To our knowledge, there are no other reports of zeolite catalysed oleic acid esterification with ethanol under continuous conditions. Zeolites have however been used to remove water by pervaporation, where a non-zeolitic material was the active catalyst. Han et al. reported a maximum conversion of 84.2% over a cation exchange resin catalyst, that increased to 87.1% when a NaY zeolite membrane was added for pervaporation [59]. The maximum conversions of the same reaction under continuous conditions, but which do not incorporate zeolites, are up to 55% from 50-70 °C using Amberlyst-15, and up to 90% at 249 °C using niobic acid [13]. Considering the reactions in this paper used a maximum temperature 70 °C and reaction time 1.5-2 h, our conversion results, 93% batch and 89% continuous, obtained without the need for pervaporation, are particularly promising.

4. Conclusions

Shale rock is found to be an excellent natural material to prepare FAU-type zeolite for the catalysed esterification of oleic acid and ethanol, a model reaction for biodiesel production. The conversion rates increased in both batch and continuous conditions with temperature and the addition of Co, Ni and Pt to the zeolite. Kinetic and thermodynamic studies proved that the reaction is pseudo first-order, with respect to oleic acid, and is kinetically controlled. On a wider level, these findings confirm the suitability of using zeolites as heterogeneous catalysts in biodiesel preparation.
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Fig. 1: Schematic of continuous fixed-bed reactor.
Fig. 2: SEM and EDAX images for H-FAU (top) and Co-Ni-Pt-FAU (bottom).
Fig. 3: Oleic acid conversion vs. reaction temperature in batch reactor: H-FAU (top) and Co-Ni-Pt-FAU (bottom).
Fig. 4: Pseudo first-order plots vs. reaction temperature in batch reactor, FAU (top) and Co-Ni-Pt-FAU (bottom).
Fig. 5: Arrhenius plots for oleic acid esterification in batch reactor.
Fig. 6: Oleic acid conversion vs. LHSV and reaction temperature in continuous fixed-bed reactor: H-FAU (top) and Co-Ni-Pt-FAU (bottom).
Fig. 7: Pseudo first-order plots vs. LHSV and reaction temperature in continuous fixed-bed reactor: H-FAU (top) and Co-Ni-Pt-FAU (bottom).
Fig. 8: Reaction rate (of oleic acid) vs. LHSV and reaction temperature in continuous fixed-bed reactor: H-FAU (top) and Co-Ni-Pt-FAU (bottom).
Table 1: Pseudo first-order rate constants, second order rate constants, Thiele modulus and thermodynamic data for oleic acid esterification using H-FAU and Co-Ni-Pt-FAU in batch conditions.

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