ORIGINAL RESEARCH



Biodiesel from rapeseed oil (*Brassica napus*) by supported Li₂O and MgO

Jerry L. Solis^{1,2,3} · Albin L. Berkemar³ · Lucio Alejo¹ · Yohannes Kiros³

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Abstract Vegetable oils are a vast triglyceride source for biodiesel production; i.e. fatty acid methyl esters (FAME), with methanol and a catalyst via transesterification reaction. The aim of this work was to study heterogeneously catalysed biodiesel production with solid oxides such as mayenite (Ca₁₂Al₁₄O₃₃) and alumina (Al₂O₃) as catalyst carriers using edible rapeseed oil as feedstock. These oxides were impregnated to have Li₂O and MgO concentrations of 5-10 and 5-30 wt% on each carrier, respectively. The catalysts were characterized using N₂-physisorption (BET/BJH), scanning electron microscopy (SEM), and X-ray diffraction (XRD) analyses. The synthesized catalysts were mesoporous ranging from 119 to 401 Å and their chemical phase composition was confirmed by the XRD. The catalyst coating (MgO/Li₂O) was studied, along with the catalyst amount in the reactor and the assessment of the transesterification reaction kinetics. The reaction was studied at 60 °C, atmospheric pressure, agitation rate of 180 rpm, and a reaction time of 2 h in a 6:1 molar ratio of methanol to oil. For each catalyst, loadings of 2.5, 5, and 10 wt% relative to the oil weight were evaluated. The highest biodiesel yield was obtained by 5 wt% (relative to oil weight) impregnated mayenite catalyst coated with 10 wt% of Li₂O. The kinetic data fits to a

Keywords Biodiesel · Transesterification · Heterogeneous catalyst · Mayenite · Alumina, MgO, Li₂O

Introduction

Vegetable oils (triglycerides) feedstock and market are affected by several factors; e.g. population growth expressed through supply and demand, biodiesel access rate and production, and fossil oil price fluctuations. Among the adversities of biodiesel production are the lower prices of petroleum barrel, which reduces the biodiesel production cost advantage (no competitive cost) and the food security policies that disregard integrated food-biofuel policies to boost biofuel production [1, 2]. Despite such adversities, worldwide biodiesel production from vegetable oils is constantly growing [3, 4]. Broad availability of vegetable oils has made biodiesel (BD) production from these feedstocks (first generation) to be well-known and optimized processes, which are commercialized in many parts of the world [5–7].

Even though vegetable oil production is increasing [8], the driving force towards eco-friendly biofuels has changed the attention towards second and third generation feed-stocks; i.e. non-edible oils, waste cooking oil, animal fats and algae. Algae was first regarded as a second generation feedstock but as it showed higher oil production yields with lower resource inputs than other feedstock, it was placed as a third generation feedstock. Oil yield of algae is higher than soybean oil and rapeseed oil by 130 times and 50 times, respectively [9–11].



pseudo-first-order model having a reaction rate constant equal to $0.045~\mathrm{min}^{-1}$ under these mild reaction conditions.

[☑] Jerry L. Solis jlsv2@kth.se

Centro de Tecnología Agroindustrial, FCyT, UMSS Universidad Mayor de San Simón, Cochabamba, Bolivia

² Centro de Biotecnología, FCyT Universidad Mayor de San Simón, Cochabamba, Bolivia

Department of Chemical Engineering and Technology, KTH Royal Institute of Technology, 100 44 Stockholm, Sweden

Micro-emulsions, thermal cracking and transesterification have been proven to reduce the use of fossil fuel and achieve a vegetable oil-based biofuel [12]. Renewability, high cetane number, lower emissions, and higher combustion efficiency are the advantages of transesterification over the other BD production methods [13]. In transesterification, triglycerides are separated to glycerol and fatty acid-alkyl esters (FAAE) with alcohol as a reactant and a catalyst. Under homogeneous reaction conditions, the reaction mixtures (containing BD, alcohol, catalyst, waste water, and glycerol) have to undergo a series of purification, separation, and final disposal processes, which are the drawbacks in high-scale BD production. Even though homogeneous transesterification has high reaction rates, non-reusability of the catalyst and further downstream process hinders the economic competitiveness with respect to fossil diesel [14].

Different catalysts and feedstocks were used to overcome BD production restraints, along with the reaction conditions such as temperature, catalyst loading and alcohol to oil molar ratio [12, 15, 16]. Heterogeneous, enzymatic and even with supercritical reactant (alcohol) transesterifications have been studied as alternatives to improve BD production. Heterogeneously produced BD studies have increased as the preferred transesterification method, since it is environmentally benign, needs no waterwashing and product separation is much easier [17, 18]. Different feedstocks, from non-edible oils [19, 20] to algae [11, 21], have also been tested for BD production. However, BD production with such feedstocks is challenging due to high free fatty acid (FFA) contents and water presence [15]. FFAs and water have a negative effect over alkali catalysed transesterification. Soap formation has been reported among the negative effects, followed by deactivation of the catalyst [10, 22, 23]. As a result, developing a high activity heterogeneous catalyst despite the FFA content and water content has become a challenge [24, 25].

The catalysts for transesterification must have a support that can hold the reactive oxides, being stable and compatible with the active material to be adsorbed. Some of the most used supports found in literature include aluminium, zirconium and titanium oxides, ferromagnetic nanoparticles, mesoporous silica, ion-exchange resins, and zeolites [26]. These supports might have some active sites which increase the yields of the BD production. Under the support requirements listed above, alumina and mayenite are good candidates for catalyst support. Alumina (Al₂O₃) is a widely used mineral, as a catalyst and as a catalyst support, due to its favourable properties; e.g. structural and chemical properties [27, 28]. Alumina is a strong Lewis acid, specifically the tetrahedral aluminium sites. Among the applications of alumina are Fischer–Tropsch synthesis

(FTS), dimethyl ether catalysed production [29], and partial hydrogenation of hydrocarbon based fuels [30]. Alumina has been studied as heterogeneous catalyst for BD production by several authors [31–36], using mild transesterification conditions such as 60 °C, atmospheric pressure, and mixing rates of 250 rpm. Mayenite (Ca₁₂Al₁₄O₃₃) is a mineral common in high-temperature, thermally metamorphosed, impure limestone included in volcanic rocks [37]. Applications of mayenite as a material include its use as a transparent conductive oxide (TCO), as a catalyst for the combustion of volatile organic compounds (VOC) or as an ionic conductor [38]. Catalytic hydrogenation, steam reforming, and catalysed transesterification using mayenite as the catalyst support have also been reported [39–42] showing the suitability of such material as a desired support in the formulation of a novel catalyst for transesterification reaction promotion.

Alkali and alkaline earth metal oxides have been extensively tested as catalytically active phases for heterogeneous transesterification of triglycerides [43–46]. Lithium and magnesium oxides—from the alkali metal and the alkaline earth metal groups, respectively—have been proven to be dominant catalysts due to their physical and chemical properties. Lithium oxide surpasses the other oxides of its group due its high catalytic activity, crystallite ordering effect, and high basic strength, which are advantageous properties for vegetable oil heterogeneous transesterification [47]. Biodiesel production was achieved with this element in mixed oxides [48, 49] and over different supports [34, 50]. Magnesium oxide catalysts have also been studied (as the active phase and as a catalyst support) and it is reported to be vastly active towards transesterification reactions. Depending on the support and the magnesium species involved some variations of the catalytic capability have been reported [46, 51, 52]. Such differences might happen due to several variables such as synthesizing conditions in the calcination procedure and the precursors used. To prepare the catalyst for the transesterification reaction, lithium and magnesium oxides were impregnated over the supports, as they were reported to be catalytically active and suitable to be adsorbed on supports to be used in transesterification reactions [18].

Several reaction conditions are considered in heterogeneous catalysed BD production, among them are reaction temperature, agitation rate, methanol to oil molar ratio, catalyst loading, etc. [13, 26, 53]. Different authors presented high BD yields with high temperatures [54] and long reaction times [55], but the present study focused in BD production at rather mild conditions (low temperature and short reaction times). Temperature is an important factor as its increase or decrease affects the reaction yield [56–58] and will also affect the reaction time and agitation rate [59]. If a catalyst is weakly promoting the





transesterification reaction, higher reaction temperatures will be needed to carry the reactants through the transesterification reaction. In low activity catalyst conditions, agitation rate or reaction times are affected negatively, resulting in the need of higher mixing agitation rates or longer reaction times [59]. In contrast, transesterification studies reported by Boz and Kara [55] and Endalew et al. [25], among others, show high BD yields working at 60 °C with improved solid catalysts. The reaction time was set to reach the maximum possible conversion in 2 h as reported in other studies for heterogeneously catalysed systems [60]. Agitation rate reported range is quite wide, depending on the transesterification study for homogeneous or heterogeneous catalysts, the agitation may vary from 1200 rpm [61] to 100 rpm [62], respectively. Finally, the methanol to oil molar ratios determines the yield of the transesterification reaction. Reported data shows a variation ranging from 6:1 to even 30:1 [15, 54], meaning a fivefold requirement of methanol compared to other reaction setups. The molar ratio of methanol used has to be more than the stoichiometrically needed and it has been proved to be useful at least at a 6:1 molar ratio related to oil to achieve biodiesel production [12, 15]. In this study, we kept that ratio to decrease the effects of over using such reactant.

The present work deals with the preparation and application of catalysts, which are found to be effective for the transesterification of rapeseed oil by solid oxides. The catalysts made of a stable support (synthesized mayenite or commercial alumina) and a highly active phase (Li₂O or MgO) were used to produce biodiesel at low energy consumption and reactant use as possible compared to literature. Translated to reaction conditions, the reaction temperature was 60 °C with an agitation rate of 180 rpm. In terms of reactant use, the methanol to oil molar ratio was kept low at 6:1 for all the experiments. The best catalysts were chosen afterwards to analyse the kinetics of the BD production and their reusability potential.

Materials and methods

Materials

Commercial refined rapeseed oil from the brand Zeta was used, along with methanol puriss p.a. (\geq 99.8% GC, Sigma-Aldrich), aluminium nitrate nonahydrate ACS (\geq 98.0%, Sigma-Aldrich), lithium hydroxide monohydrate ACS (\geq 98.0%, Sigma-Aldrich), isopropyl alcohol (\geq 99.7% FCC, Sigma-Aldrich), calcium hydroxide ACS (\geq 95.0%, Sigma-Aldrich), magnesium acetate tetrahydrate p.a. (\geq 99.0% GC, Fluka), propyl acetate (\geq 96.0% GC, Fluka), alumina pellets (Norton 1/16"), and *n*-heptane (\geq 99.0% HPLC, Carlo Erba reactifs) were used for the present study.

Synthesis of the support and catalyst

Alumina based catalysts

To prepare alumina-supported catalysts (Li₂O/Al₂O₃ and MgO/Al₂O₃), alumina (Al₂O₃) was dried at 100 °C until the weight of the sample remained constant, then it was cooled to room temperature. Lithium hydroxide and magnesium acetate were then separately dissolved in methanol. The dissolved salts were then soaked in Al₂O₃ in weight proportions to achieve 5–10 wt% Li₂O (A₁ and A₂) and 5–30 wt% MgO (A₃, A₄, and A₅) on the support. The mixtures were dried at 100 °C overnight, calcined at 650 °C for 2 h, cooled down and stored for further use.

Mayenite based catalysts

Mayenite support was produced by mixing stoichiometric amounts of calcium hydroxide and aluminium nitrate. Isopropanol was added when slurry was formed and it was heated up to 80 °C until the mixture was clear, then it was taken to the oven to be dried at 100 °C. After 72 h drying process, the mixture was milled in a mortar and calcinated at 650 °C for 2 h. At the end of calcination the produced mayenite was cooled down and mixed with the catalytic material as in the alumina impregnation described above to produce mayenite-Li₂O and mayenite-MgO catalysts, with 5-10 and 5-30 wt% of impregnated oxides, respectively. Mayenite-supported catalysts were also named and numbered after the support in increasing order from M₀ (bare mayenite), M₁-M₂ (lithium impregnated and M_3-M_4 (magnesium impregnated mayenite), mayenite).

Characterization of the support/catalyst

Characterization of the support/catalyst morphology was performed using Zeiss Ultra 55 scanning electron microscopy (SEM) in the range of 1 µm to 200 nm. Determination of the support/catalyst porosity, pore volume, and surface area were determined using a Micromeritics ASAP 2000 N₂-physisorption apparatus. The catalyst samples were degassed at a temperature of 523 K for approximately 5 h. The adsorption isotherms were carried out at a temperature of 77 K to perform BJH and BET analyses. X-ray diffraction analyses were also performed with a Siemens diffractometer D5000 using Cu-Kα radiation source ($\lambda = 1.5406 \text{ Å}$) in a 2θ degree range from 20° to 80°. The diffractograms were analysed with EVA software and the identification of the chemical species was performed correlating the data to powder diffraction file (PDF) numbers.



Transesterification and product characterization

The transesterification of rapeseed oil was performed in a round-bottom flask reactor that included a thermometer, a magnetic stirrer and a condenser. Methanol and rapeseed oil (6:1 molar ratio) were mixed in the reactor with the catalyst loadings of 5 wt% (relative to oil mass) at 180 rpm, 60 °C, and atmospheric pressure for 2 h. After the end of the reaction the mixture was transferred into a graduated cylinder and left to settle down for the separation of the liquid phases.

Transesterification samples were analysed with gas chromatography (Agilent GC 6890) using a capillary column (HP-FFAP polyethylene glycol 30 m \times 530 μ m \times I.D. 1 μ m) at a temperature of 215 °C with helium at 7.2 mL/min as the carrier phase and a split ratio of 80:1. The GC is equipped with a flame detector ionization detector (FID) working at 300 °C. The samples were prepared by taking 113 μ L from the FAME phase, mixed with 11 μ L of propyl acetate (as the internal standard) and 500 μ L *n*-heptane (as the solvent). The quantification of biodiesel yield was elaborated by fitting the results in a standard curve made with different solutions containing a FAME standard of known concentration.

Impregnation amount and catalyst concentration effects in BD production

The experiments to test the potential use of the produced catalysts were carried with the catalysts with different Li_2O or MgO impregnated amounts over the supports (alumina or mayenite). The transesterifications were performed for each catalyst at the experimental conditions described above and a fixed catalyst concentration of 5 wt% relative to oil mass as the results were recorded and further used in the following steps. Afterwards, all the catalysts were used in different concentrations for the transesterification reaction to observe the effects of the catalyst concentration. The concentration range of the catalysts that were tested included 2.5, 5.0, and 10.0 wt% of catalyst relative to oil mass (excluding A_0 and M_0). All the remaining reaction conditions were kept as indicated above.

Transesterification kinetics

Kinetic studies were carried out on the transesterification reaction using the catalyst with the highest biodiesel yield. The experimental setup was the same as in section 2.4 and 1 mL samples were taken with glass Pasteur pipettes every 20 min from the core of the reactor during 2 h of reaction time. The samples were left to settle down and taken for GC analysis to determine their FAME content.





Catalyst reuse potential

Two catalysts were selected based on their activity inferred from their BD production yields. The reuse capability was studied for a second use of such catalysts after a cycle of reaction; i.e. an experimental run using 5.0 wt% of such catalysts with a 6:1 methanol to oil molar ratio, an agitation rate of 180 rpm, a temperature of 60 °C, and atmospheric pressure for 2 h of reaction time. The mixture was transferred to a graduated cylinder to settle down. After the formation of the different component phases—catalyst, glycerol, biodiesel, and methanol—the catalyst was separated, filtered, and washed with CH₃OH. Once the catalyst was separated from glycerol, the second cycle of transesterification under the same conditions was performed.

Results and discussion

Characterization of the support/catalysts

Alumina and mayenite were characterized for their ability as support materials. Li₂O and MgO were impregnated as active phase on the supports to have a total of nine catalysts, which had BET specific surfaces, pore volumes, and pore diameters, as shown in Table 1.

BET analyses reveal small surface areas on mayenitederived catalysts, as reported by Li et al. [63] and Ruszak et al. [64], having surface areas of 1 and $5 \text{ m}^2\text{g}^{-1}$, respectively. The impregnation of Li₂O and MgO notably show an increased surface area, more than ten times in the best case (M₄). In addition to a larger surface area, Li₂O impregnation increases pore diameter and pore volume of the mayenite-derived catalysts compared to bare mayenite. Such advantageous properties enhance the catalytic activity in two possible ways. First, the large pore volumes provide more room to large triglyceride molecules to reach the active sites and proceed to adsorption (increased pore diffusion). The second improvement is that the catalyst has more adsorption area for the methoxy complex and the triglyceride-catalyst complex to form and react through the transesterification mechanism [65]. Such desired properties are seen and confirmed by the larger biodiesel yield when the reaction is carried with the catalyst M₂. The impregnation of MgO on mayenite increases the surface areas and pore volume (M₃ and M₄) but the increase of the pore diameter is smaller compared to the Li₂O impregnation case, which may alter the accessibility to the active sites.

On the other hand, alumina surface properties, e.g. surface area and pore volume, are larger than mayenite, as shown in Table 1. High surface area of 102 m²g⁻¹ and 650 °C calcination temperature suggests that alumina might be in gamma phase [66]. The effect of Li₂O and

Table 1 Surface properties of the synthesized catalysts

Catalyst type	Support	Catalytic additive ^a	$\begin{array}{c} BET \\ m^2 g^{-1} \end{array}$	Pore D ^b Å	Pore V ^c cm ³ g ⁻¹
$\overline{A_0}$	Alumina	_	102	181	0.464
A_1	Alumina	5% Li ₂ O	79.2	183	0.363
A_2	Alumina	10% Li ₂ O	68.4	176	0.300
A_3	Alumina	5% MgO	86.9	175	0.381
A_4	Alumina	30% MgO	77.3	165	0.320
A_5	Alumina	30% MgO and lumps	76.0	178	0.340
M_0	Mayenite	_	1.85	267	0.012
M_1	Mayenite	5% Li ₂ O	9.46	297	0.078
M_2	Mayenite	10% Li ₂ O	15.2	401	0.170
M_3	Mayenite	5% MgO	7.61	181	0.038
M_4	Mayenite	30% MgO	21.2	119	0.072

^a The additives are Li2O or MgO impregnated on the support in wt%

MgO wet impregnation over alumina reduced to an extent such properties. Despite alumina-supported catalyst showed reduction of surface areas compared to bare alumina, pore volumes are still larger than mayenite-supported catalysts offering more space for reactants to access the active sites.

The prepared catalysts and bare supports without impregnation were analysed by powder XRD from 20.0 to $80.0~2\theta$ angles. XRD diffractogram shows peaks at 32.5° , 37.2° , 46.0° , corresponding to alumina (Al₂O₃ PDF-01-074-4629) [67, 68]. Figure 1 shows the complete set of diffractograms for alumina-derived catalysts.

Regarding Li₂O-alumina catalysts (A₀, A₁, and A₂), not only characteristic peaks of Li₂O are present (PDF-00-012-0254 $2\theta = 33.5^{\circ}$, 38.2° , 67.0°) [69] but also peaks that suggest the presence of the specie lithium aluminate (LiAlO₂ PDF-01-073-1338) [70, 71] are found in the A₂ diffractogram. Lithium aluminate has three allotropic forms [72], two of which— γ -LiAlO₂ and β -LiAlO₂—have signals at 22.3°, 23.3°, 28.8°, 34.1°, 50.5°, 58.8° and 60.4° [70]. The preparation method was successful in the adsorption of Li₂O on alumina surface but also in the formation of LiAlO₂ species. The formation of this species stabilizes the material, improving its reuse potential.

In the case of MgO impregnation over alumina (A_3 , A_4 , and A_5), the most intense signal is caused by cubic MgO formation at high concentration (30 wt%), showing high signals at 43.0° and 62.0° (PDF-01-071-1176). Characteristic peaks of orthorhombic magnesium aluminate spinel (MgAl₂O₄ PDF-00-033-0853) are also shown in the diffractogram A_4 and A_5 , which show signals at 28.2°, 32.2°, and 43.0°. Magnesium spinel powders are advantageous due to chemical stability and resistance, and chemical inertness [73, 74].

Mayenite XRD diffractogram signal (M₀) in Figs. 2 and 3 shows peaks at 24.0°, 31.2°, 34.0, 37.8°, and 54.2° (PDF-00-009-0413). Mayenite-supported catalyst XRD diffractograms (M₁ and M₂) show that Li₂O and MgO were not only adsorbed on the mayenite structure but also show that other species have been formed, such as tetragonal LiAlO₂ $2\theta = 35.0^{\circ}$, 45.4° , 61.6°) and (PDF-01-073-1338 orthorhombic Li_5AlO_4 (PDF-00-024-0596 $2\theta = 22.7^{\circ}$, 37.8° , 45.7°). The diffractograms of the catalysts M_0 , M_1 , and M₂ show that the increase of lithium (0, 5, and 10 wt%) induces the formation of the specie Li₅AlO₄ (PDF-01-071-1736 $2\theta = 32.1^{\circ}, 37.7^{\circ}, 45.9^{\circ}, 67.1^{\circ}$) by the increase of the signal at 37.7°. Characteristic peaks of cubic CaO are also present at 32.5, 37.5, 54.8, 64.3, and 67.6° (PDF-01-070-5490). The M₁ catalyst diffractogram also suggests the presence of CaCO₃ by the characteristic signals at 29.8° and 39.6°, which might emanate from CO₂ in the air during calcination.

A cubic MgO phase found over mayenite catalysts was identified by the peaks at 37.2° , 43.3° , and 62.7° (PDF 01-071-1176) in the M_3 and M_4 diffractograms in Fig. 3. The diffractograms M_3 and M_4 show that not only a phase of MgO over mayenite support was successfully achieved but also shows that some magnesium aluminate (MgAl₂O₄ PDF-00-033-0853) has been formed over mayenite. The diffractogram shown in Fig. 3 confirm that the major components of the catalysts M_3 and M_4 are MgO and mayenite. The formation of additional compounds might be a result of contact with the atmosphere, impurities from the initial reagents, and the conditions of the catalyst preparation; e.g. temperature and atmosphere in the oven.

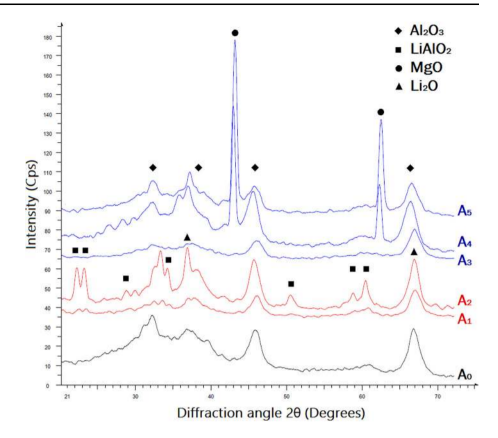
In addition to N₂-physisorption and XRD characterization procedures, SEM micrographs were performed on samples of the catalysts to determine the particle



^b D stands for pore diameter

^c V stands for pore volume

Fig. 1 XRD diffractograms of alumina-supported catalysts



morphology. The SEM micrographs of mayenite show that its particles have a rectangular and intertwined agglomerate with an average size of 266 nm and varying cavities between 20 and 180 nm (Fig. 4a). Mayenite impregnated with 5 wt% of Li₂O presented lotus-like formations (Fig. 4b), crisscrossing the surface with shallow holes.

In contrast, alumina SEM micrographs show a cloudy-like agglomeration of γ -alumina (Fig. 5a). Li₂O seems to give more order to the alumina surface but still shows that no specific shape can be the attribute of its particles (Fig. 5b). Finally, MgO over alumina micrograph (Fig. 5c) suggests amorphous formation on the alumina catalysts that were impregnated with MgO (A₃, A₄, and A₅). Due to the amorphous nature of the alumina-derived catalyst particles, it was not possible to have an average particle size.

Impregnation effect in transesterification

The prepared catalysts were used for the biodiesel production tests, where the supports without impregnation of active phase were used as control samples (M_0 and A_0). The assessment of the catalyst effect per oil mass on FAME yields was performed on several batches using each different catalyst batchwise (Figs. 6, 7).

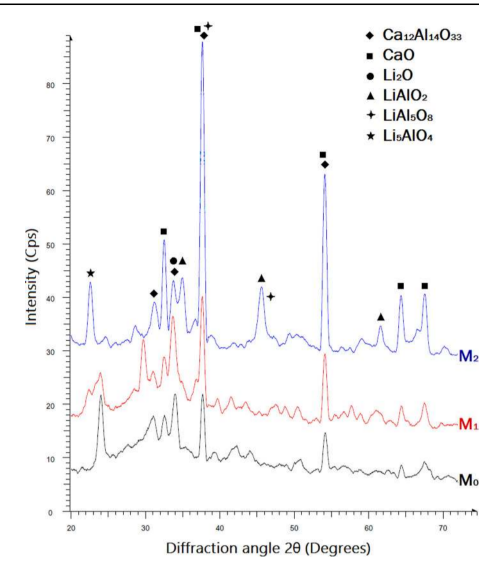
The fact that alumina is slightly catalytic for the transesterification becomes evident when observing Fig. 6 reaching 14.0% of BD yield. In the case of lithium oxide impregnation, the experiments showed that an increase of lithium oxide concentration on the support reduced the BD yield. The composition increase from 5.0 to 10 wt% of Li₂O on the alumina support was followed by a reduction of 13% BD yield (relatively comparing A₁ and A₂ BD yields in Fig. 6). It is apparent that the presence of Li₂O and Al₂O₃ in the surface of catalyst A₁ has higher catalytic activity than the LiAlO₂ specie found in the catalyst A₂, as shown in the XRD analysis in Fig. 2. This further suggests that the basic nature of the Li₂O and the acidic nature of the Al₂O₃ interaction increase the catalysts activity more than the formation of the stabilized structure of LiAlO₂, as well as the reduction of the available surface area and pore volume, which increases the mass transfer limitations towards the formation of the transesterification intermediates [65].

When impregnating the alumina with MgO (A_3 , A_4 and A_5), the BD production reaches a yield of 84.3% (catalyst A_3 shown in Fig. 6). The same hindrance to the reaction yield at the higher impregnation concentration appears again for the case of the MgO impregnated catalysts (from the catalyst A_3 to A_4), at higher MgO concentration (A_4) the BD yield decreases to 83.4%. Considering bare alumina activity, the BD overall yields might be achieved by the simultaneous action of alumina (acidic) and magnesium oxide (basic) but it is clear that the formation of different oxide species, such as MgAl₂O₄ as determined by XRD





Fig. 2 XRD diffractograms of Li_2O -mayenite catalysts M_0 , M_1 , and M_2



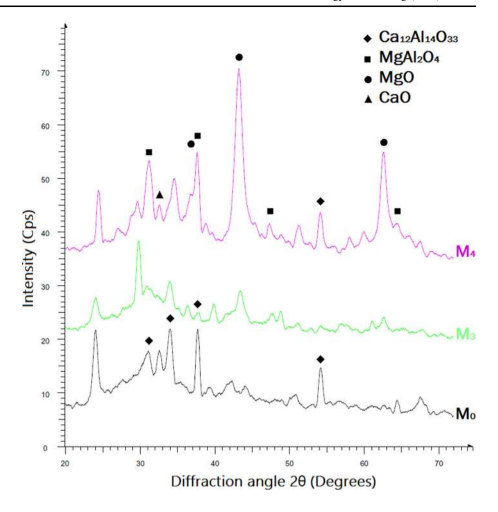
analysis in Fig. 1, decrease the catalytic activity towards transesterification (catalysts A₄ and A₅ shown in Fig. 6). This reasoning is further strengthened when the BD yield slightly increases in the transesterification using the catalyst A₅ (MgO precipitate/lumps with the impregnated alumina as shown in Fig. 6). The catalysts A₃, A₄, and A₅ might also be at their maximum activity limit regarding their capability for biodiesel production in this conditions as reported by Montero et al. on MgO nanocrystals [75], since the yields are similar between the reactions using the catalysts A₃, A₄, and A₅. However, in the alumina-based catalysts approximately 10% of BD production yield difference shows that Li₂O-alumina (A₁) is the best aluminabased catalyst. High impregnation ratios of MgO or Li₂O affects the catalytic activity in a negative way, while low impregnation ratios increase the catalytic potential surpassing by more than sixfold what bare alumina can provide on its own.

The studies with the mayenite-supported catalysts showed better results reaching maximum conversion with one of the catalysts (M₂). The yields for the different Li₂O-mayenite catalysts (M₁ and M₂) are shown in Fig. 7, from which it can be concluded that the activity increase is exponential with the increasing amount of Li₂O and thus is highly active catalysing triglyceride conversion to biodiesel. Such results are in agreement to previous reports on Li₂O activity for transesterification [25, 47] but on other supports.

The yields with the increasing amount of MgO (M₃ and M₄) show an increase of triglyceride conversion (Fig. 7). However, the activity increase from pure mayenite to 30 wt% of MgO impregnation is not high, from 7.6% BD to 23.6% BD yield, respectively. Such behaviour shows that the impregnation of MgO over mayenite is not very active under these conditions and is in accordance with the literature regarding MgO on other supports [18, 76, 77],



Fig. 3 XRD diffractograms of MgO-mayenite catalysts M_0 , M_3 , and M_4



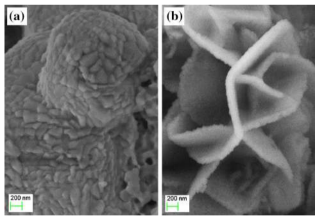


Fig. 4 SEM micrographs of mayenite (a) and Li₂O-mayenite (b)

suggesting that MgO alone has a relative inferior catalytic activity among the alkaline earth metals.

One important observation from Fig. 7 is that mayenite itself is slightly catalytic for the transesterification reaction. Despite a rather small activity, the BD yield is 8%, such active support characteristics demonstrates a potential of mayenite as a support for the catalysed reaction.

Catalyst concentration in the transesterification

The effect of the catalyst loading was evaluated based on the final BD yields. Batches with varying amounts of catalyst were run using the same reaction conditions. The results from these variations are presented in Figs. 8 and 9.

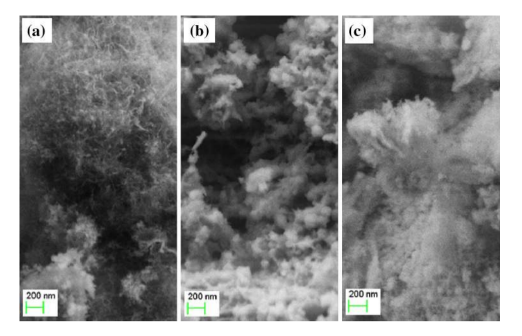
Mayenite-supported catalysts were used in three different loadings of catalysts in relation to oil mass. The trend for the lithium impregnated catalysts (mayenite with lithium oxide impregnation of 5 and 10%, M_1 and M_2) in the three loadings of catalysts into the reactor (2.5, 5.0, and 10.0 wt%) is quite different (Fig. 8). The first catalyst, which had lower Li_2O impregnation (M_1), had almost constant conversion for the batches with 2.5 and 5.0 wt% of catalyst loaded into the reactor. When increasing the amount up to 10 wt% of M_1 catalyst into the reactor, the yield had no significant increase (27.0% of BD).

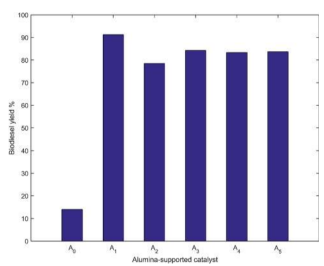
The seemingly constant yield between the 2.5 and 5.0 wt% loading of catalysts might be explained by the lack of more active sites, being at the lower limit of catalytic activity. The transesterification reaction occurs due to the reduction of the activation energy of the three consecutive reversible reactions. It has been determined by





Fig. 5 SEM micrographs of alumina (**a**), Li₂O-alumina (**b**), and MgO-alumina (**c**)





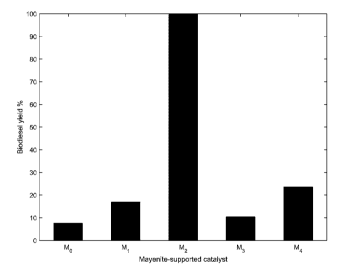


Fig. 6 Biodiesel yields using different alumina-supported catalysts

Fig. 7 Biodiesel yields using different mayenite-supported catalysts

other authors that such reduction of the activation energy might be achieved by high reaction conditions (temperature and pressure [53, 78]) as well as by the activity of the catalyst. The reactions to produce diglycerides and monoglycerides from triglycerides can be achieved with the conditions stated above, but the final glycerol production tends to be kinetically slower in the absence of a catalyst. As the catalytic material is relatively low, in the reactions with the loadings of 2.5 and 5.0 wt% into the reactor, the yield will be low as well. The fact that the reaction yield slightly increases by the highest load of catalyst (M_1 at 10 wt% loading into the reactor) suggest that more catalytic material is needed to increase the catalytic sites for the reaction to proceed at higher rates.

The yields of the reactions by M₂ catalyst (Fig. 8) increase between the two lower loadings (2.5 and 5.0 wt%) and then decreases in the last case (10 wt%). The results in Fig. 8 show that Li₂O is a very active specie on the mayenite. The increased concentration of lithium oxide on the mayenite (from 5 to 10 wt% of impregnation) and the support catalysts larger surface properties, give rise to low mass transfer limitations and high triglyceride conversions. The slight drop in BD yield when 10 wt% of the catalyst M₂ is loaded into the reactor suggests that the steady agitation rate—this variable was kept constant at 180 rpm for all the experiments—might affect the reaction environment reducing the turbulent regime in the reactor and leading to an increase of the mass transfer limitations.



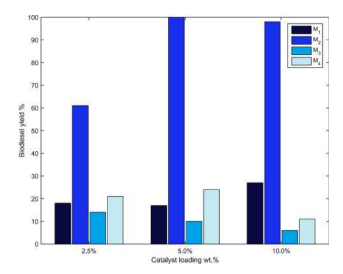


Fig. 8 Loadings of catalysts (M₁ to M₄) for BD production

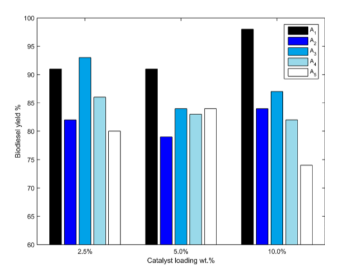


Fig. 9 Loadings of catalysts (A₁ to A₅) for BD production

From Fig. 8 it can be seen that higher yields are generally rendered from the catalysts with a higher level of MgO impregnation $(M_3 \text{ and } M_4)$. However, when increasing the amount loaded into the reactor, the batches with M₃ drop in yield with increased amount of catalyst. This is not the case for M₄, where the yield first goes up slightly and then levels off to a BD yield of 11% when increasing the catalyst amount in the reactor up to 10 wt%. The reason for this may be the agitation rate, enough to keep the medium in a laminar regime, which increases the mass transfer limitations and results in reduced BD yields. The catalytic activity of MgO 5.0 wt% impregnated over mayenite (M_3) is low as demonstrated by the experiments with 2.5, 5, and 10 wt% of such catalyst into the reactor (Fig. 8). When impregnating with higher concentration of MgO (M₄), the catalytic activity of the catalyst is high enough to enhance the yield slightly when adding more catalyst to the reactor. In the case of mayenite impregnated with 30 wt% MgO (catalyst M_4) the surface area and pore volume increase significantly, which in turn reduces the mass transfer limitations. This would then make the reaction more kinetically controlled [79]. When subsequently increasing the catalyst amount to 10 wt% into the reactor, the yield goes down, probably again due to the mass transfer limitation as a result of the reduced turbulence. The M_4 catalyst had the highest surface area of the mayenite catalysts, making it occupy larger volumes. As a consequence, a 10 wt% catalyst loading into the reactor took up more space than the other catalysts (M_1 , M_2 , and M_3), decreasing the turbulence in the reactor and causing the reaction to proceed at slower rate.

Analysing the results of the alumina-supported catalysts, it can be seen that for the A₁ catalyst the yields constantly increase with the increasing catalyst amount (Fig. 9). For the higher impregnations of Li₂O (A₂), the yield first drops from the first batch to the second and then finally rises to a yield higher than the initial one (in the range from 2.5 to 10.0 wt% of catalyst load). The trend for A₁ suggests that there is no mass transfer limitation with increasing the amount of catalyst in the reactor, at the same time shows that between 2.5 and 5.0 wt% of catalyst load the yield is not significantly increasing, and at an utterly high amount of catalyst (a catalyst increase from 2.5 to 10.0 wt%) the BD yield has an increase from 91 to 98% BD yield suggesting that the catalyst A₁ is lacking of more active sites to achieve higher BD yields. The trend with catalyst A2 suggests that there is a mass transfer hindrance at lower catalyst amounts, but when increasing the catalyst amount to 10 wt%, the sheer amount of catalyst makes the yield go up since more active catalytic sites are involved in the reaction.

The alumina-supported catalysts impregnated with MgO are shown in Fig. 9. Starting with A₃, the yield at first is very high, then drops to a level of 84% of BD yield, and then again increases up to 87% of BD yield. The reason for this behaviour of the yields might be due to a poor MgO adsorption over the alumina support. As a result, not all of the MgO is impregnated over the alumina support, and is in a "free state" as MgO powder. When the catalyst was later fed into the reactor, the ratio between MgO-alumina catalyst and MgO powder might have induced such differences showing the instability of the catalysts A_3 , A_4 , and A_5 . The case for the A₄ catalyst shows that the yield continuously drops with increasing amounts of loaded catalyst. Since the MgO particles on alumina have a lower pore volume than the bare alumina, it would lead the mass transfer limitations to occur earlier than in the case with free MgO particles.

The trend of A₅ experiments (catalyst with MgO particles) is that the activity first increases from the lower





catalyst amount to the middle one, only to finally decrease below the value of the initial yield. This also makes sense, considering that probably the mass transfer limitations occurs due to the constant agitation rate instead of adjusting the agitation rate to each catalyst loading in the reactor.

Transesterification kinetics

The reaction rate for a solid phase catalysed transesterification, as suggested by other authors, might evolve through three stages [53, 80]. Initially the conversion rate of BD is fairly low. Considering a three-phase reaction system (methanol, oil, and catalyst), such rate might be due to possible mass transfer limitations, and the time needed to form the reactive methoxide phase over the catalyst surface may delay the BD production, all this followed by a possible pseudo second-order reaction rate on the early stages of the reaction [81]. In the second stage, when the BD production increases, the liquid phase of the reactant mixture might become more uniform and the methoxide complex forms faster in a two-phase reaction system (liquid-solid) leading to an increase in the reaction rate and at this stage the reaction rate has been reported to follow a pseudo-first-order reaction rate [53, 80, 82-86]. The third stage is characterized by a decrease in the reaction rate that occurs due to the oil depletion (triglyceride source).

A mass balance over the reactor is carried out defining the design as a stirred tank reactor batch process, leading to the reaction rate shown in Eq. 1.

$$V_{\text{Cat}}r_{\text{A}} = \frac{dC_{\text{A}}}{dt}V_{\text{Tot}} \rightarrow r_{\text{A}} = \frac{dC_{\text{A}}}{dt}\frac{V_{\text{Tot}}}{V_{\text{Cat}}}$$
(1)

The global reaction obeys Eq. 2 in the presence of a catalyst, where A is the triglyceride; B is the methanol; C is the biodiesel; and D is the glycerol.

$$A + 3 \cdot B = 3 \cdot C + D \tag{2}$$

Following Eq. 2, the overall conversion should follow a fourth order reaction rate law [84]. The way to steer the reaction towards the fatty acid methyl ester production is to use the methanol in excess. If an excess of methanol is used in the reaction and considering that the second stage of the reaction is predominant (two phase reaction system), it is possible to approximate its overall kinetics to a pseudo-first-order reaction rate by simplifying it as shown in the Eq. 3:

$$r_A = -kC_A C_B^3 \cong -k^* C_A \tag{3}$$

where k^* is the pseudo-first-order rate constant. Replacing Eq. 1 in Eq. 3 for r_A and rearranging the terms to have the model relating the variables of concentration to density and mass (Eq. 4):

$$\frac{\mathrm{d}C_A}{\mathrm{d}t} = -k^* C_A \to \ln\left(\frac{C_A}{C_{Ao}}\right) = -k^* \left(\frac{V_{\mathrm{Cat}}}{V_{\mathrm{Tot}}}\right) t + C$$

$$= -k^* \frac{\rho_{\mathrm{Tot}}}{\rho_{\mathrm{Cat}}} \left(\frac{m_{\mathrm{Cat}}}{m_{\mathrm{Tot}}}\right) t + C \tag{4}$$

Such equation in terms of conversion and suggesting a constant density ratio between the density of the total volume and the density of the catalyst that can be included in the apparent reaction rate constant as k^{**} , will result in (Eq. 5):

$$-\ln(1 - X_A) = k^{**} \frac{m_{\text{Cat}}}{m_{\text{Tot}}} t + C$$
 (5)

The model in Eq. 5 describes the reaction behaviour, which could be plotted, i.e. $-\ln(1 - X_A)$ vs. t, and is used to determine k^{**} having the units min⁻¹.

When observing the yield vs. time plot (Fig. 10), it is clear that initially the reaction is not limited as a three stage system. Instead, the conversion rate is constant for the first 40 min of the reaction. This suggests that enough agitation was used to skip the first stage of the reaction, where the mass transfer limitations have detrimental effects on the reaction rate. The particle size might have aided as well, as a study by Pugnet et al. [87] shows that for catalyst powders with a particle size smaller than 500 μm such mass transfer limitation are avoided. After the initial 40 min, the conversion rate start decreasing and the process reaches the third stage due to the depletion of reactants in the transesterification reaction.

The kinetics were determined on the transesterification reaction with the catalyst M_2 (mayenite plus Li_2O 10 wt%), using a catalyst load of 5 wt% relative to oil, reaction temperature of 60 °C, agitation rate of 180 rpm, a molar ratio of 6:1 (CH₃OH:oil), and a reaction time of 2 h.

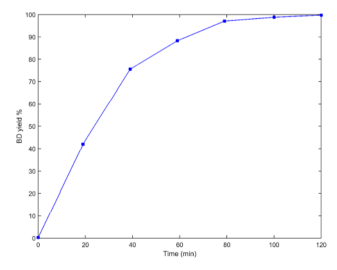


Fig. 10 BD yield plotted against the reaction time in the kinetic study



The reaction rate constant, was obtained by the use of Eq. (5) and plotted as displayed in Fig. 11.

The apparent rate constant (k^{**}) value according to the slope of Fig. 11 was found to be $0.0447 \, \mathrm{min}^{-1}$. The R^2 value is nearly 0.97 indicating that the curve fitting is acceptable. Furthermore, the obtained reaction rate expression can be compared with previously reported transesterification kinetic studies. Deshmane and Adewuyi [53] reported a pseudo-first-order reaction rate for homogeneous transesterification that has a reaction rate constant close to the value found in this study $(k^* \approx 0.036 \, \mathrm{min}^{-1})$. Supported on their study is also possible to confirm that the properties of the catalyst M_2 (mayenite-lithium oxide) enable the possibility to reduce the mass transfer limitation, avoiding the initial delay in the reaction (described as the first stage of the transesterification [53, 80]), and being comparable to a homogeneous transesterification system.

Catalyst reusability

Two different catalysts were selected to study their ability to be reused. The yields of the two different catalysts— M_2 and A_4 —are shown in Fig. 12. To prepare the catalysts for reuse tests, they were filtered after one transesterification, to separate them from the reaction mixture, as the glycerol is the by-product of the reaction to which the catalysts are attached for some time. When all the biodiesel had been filtered off the reaction mixture, the catalyst was cleaned with methanol, rendering it ready to be reused.

The alumina catalyst (A_4) did not perform well in the reusability test (Fig. 12) as its catalytic activity dropped to less than half the original value. The catalyst might have been deactivated due to catalytic material loss during the reaction. This experiment also confirms the weak

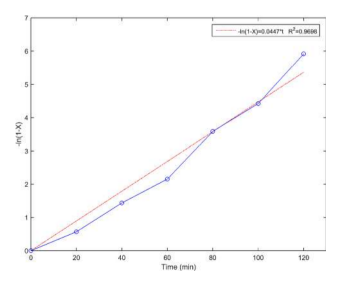


Fig. 11 Reaction rate plot, fit to Eq. (5)





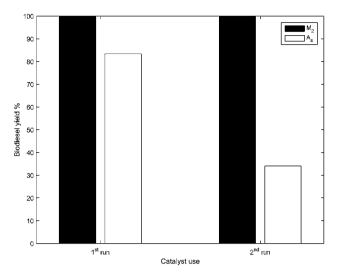


Fig. 12 Catalyst reuse test in terms of BD yield

impregnation of the MgO over alumina and that it was rather a powder surrounding the support, which readily loses the catalytic material in the filtering and washing processes. The filtration of the catalyst M_4 was a slow process due to free MgO particles. This process might also wash away the impregnated MgO making the catalyst lose even more its transesterification activity.

On the other hand, the mayenite-supported catalyst (M_2) showed an outstanding performance for two-cycle reuse transesterification reaction. In both cases the catalyst M_2 was able to achieve a 100% of BD yield. The preparing process for the M_2 catalyst reusability proved to be straightforward since the glycerol and biodiesel were separated from the catalyst with ease. Furthermore, the results of M_2 in this test suggests that Li_2O has been integrated in the mayenite support with the desired material stability. The prospect of reusing the mayenite catalyst (M_2) in more transesterification cycles might hold rather promising results.

Conclusions

Lithium and magnesium oxides were impregnated over alumina and mayenite to be used as catalysts for transesterification reaction of rapeseed oil. The prepared catalyst compositions over such supports were from 5 to 10 wt% and from 5 to 30 wt% of Li₂O and MgO, respectively, producing a total of nine different catalysts. The preparation of the catalysts lead to material with advantageous physical properties of surface area, pore diameter, and pore volume (mesoporous catalysts). The method used to prepare the mayenite-based catalysts in this study was able to increase surface properties when Li₂O was impregnated. The increase of MgO concentration had the same

increasing effect on surface areas while pore diameters were decreased. In the case of alumina-based catalysts, any increase of Li₂O or MgO impregnation concentration led to a decrease of surface area, pore diameter, and pore volume. After the tests of the catalytic activities of the produced catalysts on rapeseed oil transesterification, a series of experiments with different loadings of catalyst in the same reaction conditions were performed. As a result of such experiments, the mayenite-supported catalyst with 5 wt% of impregnated Li₂O (M₂) showed a significantly higher activity compared to other catalysts. The catalyst M₂ showed an increase from 60 to 100% of BD yield, which also showed the advantageous performance over the catalysts supported on alumina (only reaching 96% of BD yield in the best case). Thus, the catalyst M₂ was selected for the reaction kinetics study. The reaction rate experiments showed that the three-phase system (methanol-oil-catalyst) was not controlling the BD production with mayenitesupported catalyst with 5 wt% of impregnated Li₂O. Accordingly, the reaction follows a pseudo-first-order reaction rate with an apparent rate constant of 0.045 min⁻¹. Finally, the reusability tests of the catalysts showed that the catalyst M₂ has potential due to its ability to maintain its complete activity after two cycles.

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