La-Faujasite zeolite activated with boron trifluoride: synthesis and application as solid acid catalyst for isobutane–isobutene alkylation

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Abstract
The sodium form of Faujasite Y (Na-FAU) zeolite has been synthesized by the hydrothermal method, and it has been exchanged with ammonium sulphate and later with lanthanum (III) chloride solutions to obtain the La-FAU catalyst. The three zeolites Na-FAU, NH4+-FAU and La-FAU have been characterized by microcrystalline X-ray diffraction, X-ray fluorescence, surface area, pore volume and Brönsted acid sites. The La-FAU catalyst has been successfully activated with boron trifluoride etherate, and it has been tested in the alkylation reaction of isobutane with isobutene up to 112 h of time on stream, since the raw La-FAU catalyst showed a rapid deactivation.

Keywords Faujasite · Lanthanum (III) · Alkylation · Isooctane

Introduction
The alkylation reaction of isobutane with the C4 cut from fluid catalytic cracking (FCC) or steam cracking (SC) allows to increase the amount and quality of the gasoline cut in oil refineries. This reaction is carried out in liquid phase at low temperature and pressure, catalyzed by mineral acids like hydrofluoric acid and sulfuric acid. This fact represents a serious drawback, not only due to the corrosive character of these acids, but also due to the high sulfuric acid consumption and the tropism to form aerosols of hydrofluoric acid. Since many years ago, scientists have tried to replace these liquid acids with solid acids, typically zeolites, but the actual fact is that a solid acid catalyst with the performance of the liquid acids remains still to be found [1–4].

The Faujasite-type zeolites all have the same framework structure, and they crystallize with cubic symmetry. The general composition of the unit cell of Faujasite is (Na2, Ca, Mg)29[Al58Si134O384]·240 H2O. The unit cell contains eight cavities, each of diameter ≈ 1.3 nm. The three-dimensional channels, which run parallel to [110], have 12-ring windows with free apertures of about 0.74 nm. The difference between zeolites X and Y is in their Si/Al ratios which are 1–1.5 and 1.5–3, respectively. Faujasite-type zeolites are one of the most important classes of zeolithic materials and are largely used in industrial processes such as FCC, isomerisation, alkylation and organic and air separations. Y-zeolite is important as catalyst for the industrial reactions in the petroleum refining and petrochemistry such as cracking, hydrocracking, and isomerization. The Si/Al ratio is enhanced by dealumination of NH4-Y for their practical application at high-temperature reactions. The resulting materials are hydrothermally more stable (the so-called ultrastable Y zeolite, USY). The main component of fluid catalytic cracking (FCC) catalysts is rare earth-containing USY zeolites.

In this work, a catalyst based on zeolite Na-Faujasite has been prepared by hydrothermal synthesis and ionic exchange with a La3+ salt. This catalyst has been tested in the alkylation reaction of isobutane (iC4) with isobutene (iC4=), which can be considered as a model of the industrial process, since the main alkylation product, 2,2,4-trimethylpentane (2,2,4-TMP, isoctane, iC8) is the reference compound for the octane rating of gasolines. The sodium form of the zeolite Faujasite has been converted to the acid form by ionic
exchange with lanthanum (III) chloride, since exchanging with rare earth cations greatly improve the hydrothermal stability of zeolites [5]. The hydrolysis of the La\(^{3+}\) cations produced Brönsted acid sites, after Eqs. 1 and 2, where the negative charges of the zeolite framework are neutralized by hydroxylated lanthanum cations and protons:

\[3Y^-\text{La} (\text{H}_2\text{O})^3+ \rightarrow 2Y^-\text{La} (\text{OH})^2+ + Y^-\text{H}^+ , \quad (1)\]

\[3Y^-\text{La} (\text{H}_2\text{O}_2)^3+ \rightarrow Y^-\text{La} (\text{OH})^2+ + 2Y^-\text{H}^+ . \quad (2)\]

X-zeolite [6, 7], Y-zeolite [8–10] and HZSM-5 [11] zeolite have been used previously for this alkylation reaction. Zeolite catalysts have been also tested for the etherification reaction of methanol or ethanol and isobutene, to produce Methyl tert-Butyl Ether (MTBE) and Ethyl tert-Butyl Ether (ETBE), respectively, well used octane boosters for gasoline [12, 13]. Apart from its antiknock characteristics, alkylate oil is also known for its clean burning characteristics due to the absence of sulphur, olefins, and aromatics.

With the use of catalyst based on Faujasite zeolite for alkylation, the refinery could avoid the dangerous use of corrosive liquid acids, that could also leach some sulphur to the fuel streams. The novelty of this work is the activation of a rapidly deactivated La-Faujasite catalyst with boron trifluoride etherate, that allow maintain the catalyst activity for 112 h on stream.

**Experimental section**

**Equipment**

The zeolite Na-Faujasite syntheses were carried out in a 1L autoclave Büchiglasuster. The ionic exchange of the zeolite catalysts was carried out in a 1L stirred glass tank reactor with a heating jacket, controlled with a type-K thermocouple and a proportional integrative differential (PID) controller from PID Eng model TC-10.

The reactor used for the reaction experiments was a stainless steel (SS) tubular reactor (47.5 cm length \(\times\) 4.0 cm o.d.), heated by three independent heating jackets, which were controlled by an adaptive predictive control software from SCAP [14]. This reactor was connected: upstream to a stainless steel (SS) vaporizer (10.5 cm length \(\times\) 5.0 cm o.d.) filled with glass Raschig rings, and downstream consecutively to: (i) a SS heat exchanger (tube and shell, 15.5 cm length \(\times\) 5.0 cm o.d.) cooled by an ethylene glycol–water solution using a circulation pump from Electro AD model H-5P3; (ii) a SS gas–liquid separator (15.5 cm length \(\times\) 5.0 cm o.d.), and (iii) a SS liquid storing tank (10.5 cm length \(\times\) 5.0 cm o.d.).

The feed of gases (isobutane, isobutene, synthetic air and nitrogen) was controlled using mass flow controllers from Brook Instrument BV, model 5850 TR, with a range 0–6 NL/h and 1% accuracy. The back pressure regulator was also from Brook Instrument BV, model 5866, with a range 0–2 bar (0–20 hPa) and 0.5% accuracy. The feed of liquids was done with a linear actuator from Sage Instruments, model 341B, with a speed range from 0.022 to 0.321 m/h, and a flow range from 0.67 to 180 mL/h depending on the syringe size. All connections between these items of equipment were done with ¼” SS tube using Hoke fittings and electro-valves from Hirschmann model 12B GDM. Pressure measurements were carried out with a sensor DS-Europe, model LP634, and temperature measurements were done with type-K thermocouples from Sirsa SA. Figure 1 shows a scheme of this pilot plant.

X-ray diffraction analysis (XRD) of microcrystalline solids were carried out in an instrument PANalytical PW1710 with copper Kα radiation (\(\lambda=0.154\) nm) set at 40 kV and 40 mA, scanning from 4° to 60° at 2°/min.

X-ray fluorescence analysis (XRF) were carried out in a wavelength dispersive instrument PANalytical PW1404 with Sc-Mo tube, using pearls of lithium tetraborate. For the analysis of lanthanum, pressed pellets have been used, since the pearls were elaborated with a tiny amount of lanthanum oxide.

Atomic absorption spectrophotometry analyses were carried out in an instrument Philips PU9100X with a N₂O/C₂H₂ flame and selecting for B analysis the line with \(\lambda=249.8\) nm.

Gas chromatography analysis (GC-FID) was carried out in a chromatograph Agilent 5840 with FID detector, in the analytical conditions shown in Table 1.

Thermogravimetric analysis (TGA) was carried out in an instrument TGA2 Mettler heating the sample from ambient temperature to 800 °C at a rate of 10 °C/min under a constant nitrogen flow.

**Materials**

Neutral sodium silicate solution, with 27%w/w of SiO₂ and 8%w/w of Na₂O was bought from Panreac. Sodium aluminate, ammonium sulphate and sodium hydroxide were bought from Probus. Lanthanum (III) chloride was bought from Merck. Boron trifluoride etherate and isooctane were bought from Sigma-Aldrich. High purity isobutane (> 99.9%) and isobutene (> 99.9%) were bought from Air Liquide and they were used without further purification.

Table 2 summarizes the composition of the nucleation and crystallization gels, for a total reaction volume of 500 mL and 20%w/w of nucleation gel, and the amounts of the above commercial reagents necessary to achieve these compositions.
Catalyst preparation

Formation of the nucleation gel

In the autoclave, the sodium hydroxide was dissolved in distilled water at room temperature, and later the sodium aluminate was added, stirring the mixture until complete dissolution of both reagents. Afterwards, the sodium silicate was slowly added, forming a gel which was stirred at 500 rpm during 1 h. This nucleation gel was left standing at room temperature for 48 h without any agitation.
Crystallization procedure

One hour before the end of the nucleation gel aging, a new gel was prepared in exactly the same way as indicated for the nucleation gel, but with the reagent amounts shown in the Difference column of Table 2. This gel was stirred at 500 rpm for 1 h. After aging, the nucleation gel was activated heating it at 60 °C in a water bath, and it was mixed with the recently prepared additional gel at 60 °C under stirring. This full crystallization gel was stirred at 500 rpm in the autoclave until the crystallization temperature of 100 °C was reached. Afterwards, the reaction mixture was kept at 100 °C without stirring during 4 h. Later, the reaction mixture was centrifuged at 1000 rpm, the mother liquid was discarded and the solid was washed with distilled water several times, eliminating the washing water by filtration, until neutral pH was reached in the washing water. The white solid obtained was dried at 120 °C for 12 h. XRD and XRF analyses showed that it consisted of the sodium form of Faujasite (Na-FAU).

Preparation of the active zeolite Faujasite catalyst

The sodium form of zeolite Faujasite (Na-FAU) was converted to the catalytically active form by ionic exchange in solution as follows:

To prepare the zeolite catalyst, the sodium form of the zeolite Na-FAU was stirred at room temperature with a 5% w/v solution of ammonium sulphate during 24 h. The solid was filtered and washed until absence of sulphate in the washing waters (checking by silver sulphate precipitation with a diluted solution of silver nitrate). This ionic exchange procedure was repeated three times [4]. To obtain the zeolite La-FAU catalyst, the NH4-FAU zeolite was exchanged with a 0.5% w/v solution of lanthanum (III) chloride for 24 h at room temperature with stirring. The solid was filtered and washed with distilled water until absence of chlorides in the washing waters (checking by silver chloride precipitation with a diluted solution of silver nitrate). The La-FAU zeolite was dried at 120 °C, and calcined in an oven at 550 °C overnight, rising the oven temperature at 100 °C steps and keeping the La-FAU zeolite at each step for 1 h. The amount of La-FAU zeolite was 39.66 g (6.0% yield based on crystallization gel of Table 2, including water).

The catalysts were prepared with the active form of the zeolite, and the fraction between 0.5 and 1.0 mm was used for the fixed-bed experiments [10].

Characterization

XRD and XRF analyses were carried out on all samples to ascertain the chemical and mineralogical composition of these catalysts, and they are summarized in Tables 3 and 4. The mineralogical composition of the catalysts and their crystallinity have been determined by XRD following the standard method ASTM D3906-03 (2013), as the average ratio of X-ray counts of selected diffraction peaks in the samples and in a supposedly full crystalline Na-FAU zeolite taken as reference.

The BET surface and pore volume of the catalysts were analyzed by nitrogen adsorption–desorption isotherms in a Micromeritics ASAP-2010 instrument and the results are summarized in Table 5. Prior to the measurements, all samples were degassed at 300 °C under vacuum overnight.

The acid site strength was investigated by temperature programmed ammonia desorption (NH3-TPD) on an ASAP 2920 analyzer. Before the measurement, approximately 0.1 g dry samples were degassed under He flow at 500 °C for 1 h. After cooling to 120 °C, the sample was exposed to NH3 (10% v/v) − He gas flow for 30 min. Then, the sample was swept by He for 1 h to remove the physically adsorbed NH3, and the TPD was carried out from 120 to 600 °C using a heating rate of 10 °C min−1.

Alkylation reaction

The catalyst La-FAU was placed in the bottom section of the tubular reactor supported on rockwool and prior to each reaction it was activated in air at 538 °C during 2 h. Isobutane, isobutene and nitrogen as diluent were fed to the reactor in the mass flows required for each experiment, that lasted for about 8 h, at a weight hourly spatial velocity (WHSV) defined in Eq. 3:
where \( m_\text{iC4} \) was the mass flow of isobutene in g/h, \( m_\text{iC4} \) was the mass flow of isobutane in g/h and \( m_\text{catalyst} \) was the catalyst mass in g.

All experiments were carried out in vapor phase, and the reactor effluents were sampled with a 100 µL gas syringe through a septum placed at the reactor outlet and were immediately injected in the GC. A heating wire was installed at the reactor outlet to avoid condensation of any liquid products before sampling. After each experiment, the catalyst was purged with a nitrogen flow of 6 NL/h for 2 h at 500 °C to desorb all products occluded in the zeolite channels.

Isobutene conversion (isobutene was always the limiting reagent) was calculated from Eq. 4:

\[
\text{Isobutene conversion} = \frac{A_\text{isobut} - A_\text{exp}}{A_\text{isobut}} \times 100, 
\]

where \( A_\text{isobut} \) represent the peak area of isobutene detected in the chromatogram when the reaction system is run without any catalyst, and \( A_\text{exp} \) represent this same peak area in each experiment with catalyst.

The 2,2,4-TMP selectivity was calculated from Eq. 5:

\[
\text{Isooctane selectivity, } S_{\text{iC8}} = \frac{A_{\text{iC8}}}{A_{\text{iC8}} + A_{\text{others}}} \times 100, 
\]

where \( A_{\text{iC8}} \) is the peak area of isooctane and \( A_{\text{others}} \) is the peak area of other alkylation and isobutene oligomerization products.

### Results and discussion

#### Catalyst characterization

The chemical composition of zeolites prepared in this work was analyzed by XRF and it is summarized in Table 3. Faujasite-type zeolites with Si/Al ratio near one are usually denoted by X-zeolite, whereas, those with Si/Al ratio higher than two are usually denoted by Y-zeolite. The Si/Al molar ratio for all zeolites in this study is around 2.0, which means that it could be possible that some of the aluminium atoms are extra-framework (EFAL), and they could be exchanged to increase the acid strength of the catalyst. The ion exchange procedures greatly decrease the crystallinity of the zeolites, thus the Na-FAU prepared in this work has almost the same crystallinity (92%) as the reference Na-FAU, but the La-FAU retains only 16% crystallinity with respect to reference. Figures 2, 3, 4, 5, 6 show the diffractograms of the FAU zeolites, where the decrease of crystallinity with time on stream is evident. However, the BET surface, the total micropore volume and the acidity results remain practically unaffected by the ionic exchange, which differs from previously reported results [8]. The BET surface of this work catalyst (Table 5) is slightly higher than the value of 470 m²/g reported for other alkylation catalysts, and the pore volume somewhat lower than the values between 0.255 and 0.295 cm³/g also reported [6].
La-FAU catalyst activation

The La-FAU catalysts showed a quick deactivation in the first alkylation experiments after a few minutes on stream [3]. Thus, it was activated by treatment with boron trifluoride etherate, injecting this solution (0.06 mL/min) with the isobutane/isobutene feed (molar ratio \(iC_4/iC_4^\text{m} = 5.0\), temperature 90 °C, WHSV 0.2 h\(^{-1}\), \(iC_4\) flow 3.70 NL/h, \(iC_4^\text{m}\) flow 0.74 NL/h). This activated La-FAU catalyst showed an activity equal to the original catalysts, that remained after 112 h TOS. In fact, the effect of boron could be due to the substitution of the extra-framework (EFAL) aluminium atoms by boron atoms, since boron appears in the chemical analysis of the catalyst after all the alkylation reactions in 2.6% w/w, see Table 3.
Alkylation reaction

This reaction of isobutane with isobutene was carried out as a model reaction for the alkylation of isobutane with the C₄ cut from FCC, where isobutene is the most reactive component of this C₄ fraction. The product analyzed was 2,2,4-trimethylpentane (2,2,4-TMP, isooctane, iC₈), although obviously other branched C₈ and higher molecular weight compounds Cₙ⁺ are produced in this alkylation. The C₈ fraction is composed of other trimethylpentanes apart from 2,2,4-TMP (2,3,3-TMP and 2,3,4-TMP), dimethylhexanes (DMH) (2,4-DMH, 2,5-DMH and 3,4-DMH), C₈ olefins, and other C₈ alkanes. Isooctane, iC₈, 2,2,4-TMP, is the C₈ hydrocarbon with the highest research octane number (RON) and motor octane number (MON) of 100 points. These other C₈ alkylation and higher molecular weight products Cₙ⁺ have been quantified in the chromatograms, and their areas have been taken into account to calculate the
The isooctane selectivity after Eq. 5. The effect of reagents molar ratio, temperature and WHSV on the isobutene conversion and isooctane selectivity has been studied.

**Effect of the molar ratio isobutane/isobutene \(iC_4/iC_4^e\)**

The effect of the molar ratio \(iC_4/iC_4^e\) was investigated at 90 °C and WHSV 0.05 h⁻¹, and it is summarized Fig. 7.

The isooctane selectivity remained practically constant, but the isobutene conversion reached a maximum of 65% between molar ratios 10 and 15, in good agreement with the molar ratio of 13 for this reaction on La-FAU catalyst, recently reported by Yang et al. [8]. This high molar ratio of isobutane to isobutene is necessary to minimize the isobutene oligomerization reaction, although it means that a great amount of isobutane need to be recycled to make the process economically acceptable.

**Effect of the temperature**

The effect of temperature was investigated at molar ratio \(iC_4/iC_4^e\) of 15 and WHSV 0.05 h⁻¹, in the range 70–150 °C, and the results are summarized in Fig. 8.

The highest isooctane selectivity was achieved at 130 °C but at lower isobutene conversion, 50%. As it was expected, when increasing the conversion to 65% at 90 °C, the isooctane selectivity decreased. The optimal temperature found in our experiments maximizing the isooctane selectivity was 130 °C, in disagreement with previously reported results of 80 °C [6], which gave more importance to the butenes conversion.

**Effect of the WHSV**

The effect of the WHSV was investigated at 130 °C and molar ratio \(iC_4/iC_4^e\) of 15, and the results are summarized in Fig. 9.

Thus, it could be estimated that the optimal reaction conditions for this La-FAU catalyst are 130 °C, molar ratio \(iC_4/iC_4^e\) 15 and WHSV of 0.10 h⁻¹.

Figure 10 shows the thermogram of the La-FAU catalyst after one of the alkylation experiments. This thermogram shows one sharp peak at 156 °C, which corresponds to the desorption of the C₈ products, isooctane among them, and another much broader peak between 456 and 475 °C, that would represent the desorption of higher molecular weight products, C₉⁺. The adsorption behavior of these catalysts has been reported recently in the literature [15].

The crystallinity of the La-FAU catalyst after 56 h TOS was slightly lower (13%) than the that of the new catalyst (16%), both referred to the Na-FAU taken as crystalline reference. However, after 112 h TOS, the crystallinity has decreased to 5.3%, although the La-FAU mass loss of catalyst was negligible.

The mechanism of this alkylation reaction could be schematized in Fig. 11 [5]:

The isooctane selectivity reached the highest value of 15.1% at WHSV of 0.10 h⁻¹, but with the lowest isobutene conversion of 48.1%.
An isobutene $iC_4$ molecule is adsorbed on the Brönsted acid sites and one $H^+$ adds to the double bond, giving an adsorbed $terc$-butyl carbocation. A second isobutene $iC_4$ molecule adsorbs and react with the adsorbed $terc$-butyl carbocation, giving an adsorbed iso-octyl carbocation. The hydride transfer $H^-$ from an adsorbed isobutane molecule

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Fig. 10  Thermogram of the La-FAU catalyst after one of the alkylation experiments

Fig. 11  Mechanism of alkylation of isobutane $iC_4$ with isobutene $iC_4^-$ on La-FAU catalyst (F$^-$ represents zeolite framework)
$iC_4$ (present in great excess in the vapor phase) produces the isooctane final molecule that desorbs from the catalyst surface, leaving an adsorbed $terc$-butyl carbocation coming from the isobutane $iC_4$ that repeats the mechanistic cycle.

These preliminary alkylation results on La-FAU catalyst show the feasibility of trying this catalyst in a pilot scale plant, avoiding the use of dangerous liquid mineral acids and their environmental problems associated.

**Conclusions**

La-FAU Catalyst has been synthesized by hydrothermal method and ion exchange with $La^{3+}$ solutions, and it has been successfully checked in the alkylation reaction of isobutane with isobutene. The rapidly deactivated original La-FAU catalyst has been reactivated by co-injection of a boron trifluoride etherate during an alkylation experiment, and this catalyst reactivation lasted for at least 112 h of time on stream.

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