Different approaches to the synthesis of ZSM-22 zeolite with application in n-heptane cracking

Diferentes abordagens de síntese da zeolita ZSM-22 com aplicação no craqueamento do n-heptano
Diferentes enfoques para la síntesis de zeolita ZSM-22 con aplicación al craqueo de n-heptano

Received: 01/23/2022 | Reviewed: 02/01/2022 | Accept: 02/06/2022 | Published: 02/12/2022

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Abstract
The synthesis of ZSM-22 zeolite has been extensively studied due to its properties of form selectivity, acidity and hydrothermal stability, which is applied in important reactions in the areas of petroleum refining and petrochemicals. In view of this, the present work studied different approaches of synthesis of ZSM-22 using: (i) 1-diaminohexane as a structure directing agent, (ii) methanol and seed crystals, (iii) aging of the synthesis gel with addition of polymer, surfactant and silane and (iv) starch, calcium carbonate and silanized silica with subsequent desilication. Thus, the effects of these methodologies on the textural properties, acidity and catalytic activity of the zeolites obtained were evaluated. The samples were characterized by X-ray diffraction (XRD), nitrogen adsorption-desorption, scanning electron microscopy (SEM), ammonia desorption at programmed temperature (NH3-TPD) and thermal analysis (TG/DTG). The catalytic activity and selectivity of the catalysts were evaluated in the model catalytic cracking reaction of n-heptane at 650 °C for 180 min. The synthesis route using methanol and seed crystals allowed obtaining ZSM-22 in 3 h of crystallization, drastically reducing the synthesis time compared to other methodologies. Despite this, the use of 1-diaminohexane led to the obtaining of zeolite with textural properties, acidity and catalytic activity superior to the other samples. The generation of mesoporosity was obtained through the use of silanized silica and subsequent desilication, leading to greater catalytic stability and less deactivation by coke. All catalysts showed similar selectivity to the formation of compounds in the range C2 to C4.

Keywords: ZSM-22; Zeolite; Different synthesis approaches; Catalytic cracking.

Resumo
A síntese da zeólita ZSM-22 tem sido extensivamente estudada devido às suas propriedades de seletividade de forma, acidez e estabilidade hidrotérmica, sendo esta aplicada em reações na área petroquímica. Diante disto, o presente objetivou estudar diferentes abordagens de síntese da ZSM-22 utilizando: (i) 1-diaminohexano como agente
La síntesis de la zeolita ZSM-22 ha sido ampliamente estudiada debido a sus propiedades de selectividad de forma, acidez y estabilidad hidrotermal, la cual se aplica en importantes reacciones en las áreas de refinación de petróleo y petroquímica. En vista de esto, el presente trabajo estudió diferentes enfoques de síntesis de ZSM-22 utilizando: (i) 1-diaminohexano como agente director de estructura, (ii) metanol y cristales semilla, (iii) envejecimiento del gel de síntesis con adición de polímero, surfactante y silano, y (iv) aplicando almidón, calcio carbonato y sílica silanizada con posterior dessilicación. Así, se evaluaron los efectos de estas metodologías sobre las propiedades texturales, acidez y actividad catalítica de las zeolitas obtenidas. Las muestras se caracterizaron por difracción de rayos X (DRX), adsorción-descorción de nitrógeno, microscopía electrónica de barrido (MEB), desorción de amoníaco a temperatura programada (NH₃-TPD) y análisis térmico (TG/DTG). La actividad catalítica y la selectividad de los catalizadores se evaluaron en el modelo de reacción de craqueo catalítico de n-heptano a 650 °C durante 180 min. La ruta de síntesis utilizando metanol y cristales semilla permitió obtener ZSM-22 en 3 h de cristalización, reduciendo drásticamente el tiempo de síntesis en comparación con otras metodologías. A pesar de esto, el uso de 1-diaminohexano condujo a la obtención de una zeolita con propiedades texturales, acidez y actividad catalítica superior a las otras muestras. La generación de mesoporosidad se obtuvo mediante el uso de sílica silanizada y posterior desilicación, lo que condujo a una mayor estabilidad catalítica y menor desactivación por coque. Todos los catalizadores mostraron una selectividad similar a la formación de compuestos en el rango de C2 a C4.

Palabras clave: ZSM-22; Zeolita; Diferentes enfoques de síntesis; Craqueo catalítico.

1. Introduction

ZSM-22 is a silicon-rich zeolite, formed by non-intersecting one-dimensional channels with an opening diameter of 0.57 and 0.46 nm, comprised of 5, 6 and 10 membered rings. The channels are directed to the longest dimension of the crystals (crystallographic direction c) (Del Campo, et al., 2018). Due to the narrow pore size, ZSM-22 exhibits shape selective effect in different reactions, such as hydroisomerization dewaxing (Gao, et al., 2020), propene oligomerization (Redekop, et al., 2020; Dagle, et al., 2020) and methanol-to-hydrocarbons (MTH) (Liu, et al., 2017).

Valyocsik reported in a 1984 patent the first synthesis of ZSM-22 zeolite using alkanediamines in the C2-C12 range as structure-directing agents. The ZSM-22 standard synthesis method published by the Synthesis Commission of the International Zeolite Association (Robson et al., 2001), adopts 1,8-diaminoethane as OSDA, with prior aging of the synthesis gel for 24 h at room temperature, and subsequent crystallization at 160 °C for 2 to 3 days, under vigorous stirring (400 rpm). In low-stirring (less than 400 rpm) or static synthesis systems, other phases such as the MEL zeolite (Valyocsik, 1984; Robson & Lillerud, 2001), ZSM-5 zeolite and cristobalite (Verduijn et al., 1996) are obtained.

Different parameters affect the crystallization and physicochemical properties of a specific zeolitic phase, such as synthesis method, chemical composition of the initial gel, type of reagents, synthesis gel SiO₂/Al₂O₃ ratio, alkalinity, water content, inorganic cations, organic directing organic structure-directing agents, temperature, aging of the synthesis gel, stirring and use of seed crystals (Yu, 2007; Shin et al., 2019). Rollmann et al. (1999) describe in their study the relationship between the structure of small amines and the zeolitic phase obtained. For a SiO₂/Al₂O₃ ratio of equal to 200, certain acyclic amines
were specific for the formation of 10-ring zeolites, as ethylmethylamine and diethylamine produced ZSM-22, while ethylamine and isopropylamine resulted in ZSM-23 zeolite. Accordingly, larger molecular size cyclic amines such as piperazine have produced zeolites with a 12-ring pore system such as ZSM-12. For synthesis gels with SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} ratios below 40, different zeolites were produced, such as ZSM-35 and MCM-22, showing a specific relationship between the amine employed and the zeolite obtained.

Wen et al. (2014) presented a study of the synthesis of pure-silica ZSM-22 by the dry gel conversion method. The gel was prepared in an acidic medium to hydrolyze the silica precursor, and the ionic liquid 1,3-alkylimidazolium chloride was used as an organic structure-directing agent. The pure-silica ZSM-22 structure was obtained within 2 days of crystallization. Using a similar procedure, the synthesis of Al-containing ZSM-22 was studied by adding aluminum salt to the starting gel. Lopes et al. (2016) reported the synthesis of the Al-TON structure using 1-butyl-3-methylimidazolium chloride as OSDA. This organic cation showed greater efficiency and selectivity for the synthesis of TON zeolites under different reaction conditions compared to 1,8-diaminooctane (a traditional OSDA). The scanning electron microscopy technique indicated that zeolites with different crystal sizes were obtained under static or stirred synthesis (static method from 30 to 50 μm and stirred method less than 2 μm), which consequently affects the textural properties of the zeolites.

Jamil et al. (2016) described the crystallization of ZSM-22 through the microwave solvothermal synthesis method, evaluating the effect of the type and amount of different co-solvents, such as ethanol, 2-propanol, glycerol, ethylene glycol and the aging time of the synthesis gel. They found that a long aging time (72 h) was crucial to obtain smaller and more uniform crystals. The addition of a co-solvent resulted in more elongated crystals, and the crystallinity of the zeolite was sensitive to the type of co-solvent. In general, the use of a co-solvent led to the formation of ZSM-5 zeolite as an impurity, which is proportional to the concentration of the co-solvent in the reaction medium.

Sousa et al. (2017) studied the effect of crystallization time, use of seed crystals, methanol as solvent, variation of the SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} and OH/SiO\textsubscript{2} ratios, the addition of potassium and stirring on the properties of ZSM-22 zeolite. Under static condition and SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} ratio of 100, ZSM-22 was obtained after 6 h, in the presence of methanol, OH/SiO\textsubscript{2} ratio of 0.2 and without addition of potassium. With the SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} ratio of 50, the increase in alkalinity (OH/SiO\textsubscript{2} ratio = 0.4) or in the potassium content resulted in a decrease in the crystallinity of the zeolite. The stirred synthesis with SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} ratio of 100 and low methanol content increased the crystallinity and specific surface area of the zeolite, reducing the crystallization time to 3 h.

Jamil et al. (2014) reported the synthesis of nanometric ZSM-22 after 72 h of synthesis gel aging at 50 °C and crystallization by the microwave method for 12 h at 180 °C. The nonionic polyoxyethylene surfactant - Brij-76 was added to the reaction medium and led to a decrease in size (between 90 – 300 nm) and agglomeration of the crystals, as well as the formation of competing phases. A better catalytic activity in the n-hexane cracking was observed for the catalysts with shorter crystal length (90 nm) in comparison to the larger ones (300 nm). This fact was attributed to the higher external surface area and the smaller length of the crystals, which offers more accessibility to the pores of the zeolitic structure (Rownaghi, Rezaei & Hedlund, 2012).

Due to the microporous structure of zeolites, especially those with one-dimensional pore systems such as ZSM-22, there is a diffusional resistance to bulky reagents and products (kinetic diameter greater than 1.4 nm) to the internal acid sites of the zeolite structure. Therefore, different studies have been carried out with the aim of improving the textural properties of these materials with the introduction of a secondary porosity (Verboekend, et al., 2011; Dyballa, et al., 2016; Del Campo, et al., 2018; Wang et al., 2019). Silva et al. (2019) obtained the micro-mesoporous ZSM-22 by adding nanometric silica previously silanized with trimethoxyphenylsilane to the reaction mixture, followed by desilication. The samples showed a pore
size distribution in the range of mesopores between 40 and 60 nm.

Zhu et al. (2008) synthesized hierarchical silicalite-1 (mesopores in the range of 50 – 100 nm) using nanometric carbonate (CaCO3) as a hard template for secondary porosity. After crystallization (140 °C for 7 days) the material was washed with HCl solution to remove the carbonate and consequently generate the mesopores. Zhang, Liu & Yan (2016) described the synthesis of mesoporous ZSM-5 using carbonized soluble starch. The zeolite showed high thermal and hydrothermal stability, with irregular mesopores distribution due to the different dimensions of the carbonized starch. After crystallization (170 °C for 5 days), carbon was removed by calcination at 800 °C.

The catalytic cracking of hydrocarbons on acidic solid catalysts involves a large number of process parameters and variables related to the reaction conditions and the physicochemical properties of the catalyst and hydrocarbon molecules. The difficulties to optimize catalytic cracking processes in order to obtain specific products are mainly due to: (i) catalyst deactivation with residence time, via coke formation, and (ii) the presence of a large number of reaction steps and products (Corma, et al., 2005; Rahimi et al., 2011; Corma, et al., 2017).

Given the above, systematic studies on a laboratory scale are commonly carried out using some low molecular weight hydrocarbons, such as n-heptane, as a representative of the feed charge (vacuum diesel) of the fluidized catalytic cracking process, enabling the determination of kinetic parameters, development of reaction mechanisms, catalyst designs and validation, adjustment and/or optimization of cracking units (Blay, et al., 2017; Afroukhteh-Langaroudi, Tarighi et al., 2018; Xu, et al., 2018).

In this context, the objective of this work was to evaluate the synthesis of ZSM-22 zeolite by different methodologies, using: (i) 1-diaminohexane as a structure-directing agent, (ii) methanol and seed crystals, (iii) gel aging of synthesis with addition of polymer, surfactant and silane, (iv) starch, calcium carbonate and silanized silica, with subsequent desilication, and thus evaluate the effects of these methodologies on the textural properties, acidity and catalytic activity in the catalytic cracking of n-heptane of the obtained zeolites.

2. Methodology

2.1 Synthesis of the ZSM-22 zeolite

2.1.1 Synthesis using 1,6-diaminohexane as structure-directing agent

The reagents were added in stoichiometric proportions in order to obtain a mixture with the following molar composition: 27.0NH2(CH2)6NH2:13.5K2O:1.0Al2O3:90.0SiO2:3600H2O (Valyocsik, 1984). The preparation of the reaction mixture consisted of the following steps: (i) dissolution of potassium hydroxide (Sigama-Aldrich, 85%) in 30% of the total amount of water required for synthesis (Solution A), (ii) dissolution of 1,6-diaminohexane (Sigma-Aldrich, 98%) in 30% of the water required for synthesis (Solution B), (iii) solubilization of aluminum sulphate (Merck, 99%) in 30% of the water required for synthesis, with sulfuric acid being subsequently added dropwise (J.T. Baker, 98%) (Solution C) and (iv) dispersion of colloidal silica (Ludox-AS-40, Sigma-Aldrich, 40%) in 10% of the water required for synthesis (Solution D). Subsequently, solution B was mixed with solution A, under mechanical stirring at 200 rpm for 10 min. Then, solution C was added under mechanical stirring at 200 rpm for 10 min. Finally, solution D was added under mechanical stirring at 200 rpm for 30 min. The hydrothermal treatment of the gel was carried out at 160 °C in a 1 L Parr Instrument Company stirred reactor (model 4520), under agitation at 400 rpm for 17 h. The solid was recovered by vacuum filtration, washed until neutral pH, and then dried in an oven at 120 °C for 12 h (Z22-ADE Sample).
2.1.2 Synthesis using methanol as a co-solvent

The molar composition used to obtain the synthesis gel was 9.0CH₃OH:0.3Na₂O:1.0Al₂O₃:90.0SiO₂:20.0H₂O (Morimoto, Takatsu & Sugimoto, 1983). The gel was prepared through the following steps: (i) dilution of sulfuric acid (J. T. Baker, 98%) in 25% of the total amount of water required for synthesis (Solution A), (ii) solubilization of aluminum sulfate (Merck, 99%) in 25% of the water required for synthesis (Solution B), (iii) dilution of sodium silicate (Pernambuco Química PQ-4, 10.07% Na₂O, 32.22% SiO₂) in 25% of the water required for synthesis (Solution C) and (iv) dilution of methanol (Synth, 99 %) in 25% of the water required for synthesis (Solution D). Then, solution B was mixed with solution A, under mechanical stirring at 200 rpm for 10 min. Subsequently, solution C was added under mechanical stirring at 200 rpm for 60 min. Finally, solution D was added under mechanical stirring at 200 rpm for 60 min. The crystallization was carried out at 170 °C in a 1 L Parr Instrument Company stirred reactor (model 4520), under agitation at 400 rpm for 20 h. Subsequently, the final solid was separated by vacuum filtration, washed to neutral pH, and dried in an oven at 120 °C for 12 h (Z22-Me Sample).

2.1.3 Synthesis using seed crystals and methanol as co-solvent

The molar composition of the synthesis gel was 27.0NH₃(CH₂)₃NH₂:13.5K₂O:1.0Al₂O₃:90.0SiO₂:4.5H₂SO₄:3600H₂O + 10% of seed crystals (mass percentage in relation to the amount of silica) (Sousa, et al., 2017). The mixing steps were: (i) dilution of sulfuric acid (J. T. Baker, 98%) in 20% of the total amount of water required for synthesis (Solution A), (ii) solubilization of aluminum sulfate in 20% of the water required for synthesis (Solution B), (iii) dilution of sodium silicate in 20% of the water required for synthesis (Solution C), (iv) methanol dilution in 20% of the water required for synthesis (Solution D) and (v) dispersion of ZSM-22 seed crystals (sample Z22-ADE, previously synthesized) in 20% of the water required for synthesis (Solution E). In sequence, solution B was mixed with solution A, and subsequently the solutions C and D were added, with each step under mechanical stirring at 200 rpm for 10 min. Finally, solution E was added under mechanical stirring at 200 rpm for 30 min. The synthesis process was carried out at 170 °C in a 1 L Parr Instrument Company stirred reactor (model 4520), under agitation at 400 rpm for 3 h. After this period, the final solid was separated by vacuum filtration, washed to neutral pH, and oven dried at 120 °C for 12 h (Z22-Me-CS Sample).

2.1.4 Synthesis through gel aging using polymer, surfactant and silane

The synthesis was carried out using a two-step procedure: (i) aging of the synthesis gel and (ii) crystallization. The molar composition 27.0NH₃(CH₂)₃NH₂:13.5K₂O:1.0Al₂O₃:90.0SiO₂:4.5H₂SO₄:3600H₂O:2.5(X or Y or Z) (where X = polymer, Y = surfactant and Z = silane) and methodology of the synthesis gel preparation for the aging stage was similar to those used for the Z22-ADE sample (topic 2.1.1). The synthesis gel was aged for 20 h at 120 °C, under stirring at 200 rpm in a 300 mL Parr Instrument Company stirred reactor (model 4566). Then, the gel was divided into equal parts, to which were added, under mechanical stirring at 200 rpm for 30 min, Pluronic P123 polymer (Sigma-Aldrich, 99%), Brij93 surfactant (Sigma-Aldrich, 99%), and vinyltrimethoxysilane silane (Sigma-Aldrich, 98%). Each gel was transferred to the same reactor as in the aging step. Crystallization was carried out under stirring at 400 rpm at a temperature of 160 °C for 20 h. After crystallization, the final solid was recovered by vacuum filtration, washed to neutral pH, and oven dried at 120 °C for 12 h (Z22-P123, Z22-Brij93 and Z22-Vynil Samples).

2.1.5 Synthesis using soluble starch

The molar composition used was similar to that of the Z22-ADE sample plus a percentage by mass of starch in relation to the amount of silica: 27.0NH₃(CH₂)₃NH₂:13.5K₂O:1.0Al₂O₃:90.0SiO₂:4.5H₂SO₄:3600H₂O + 1.5 Starch/SiO₂ (ratio of
1.5 times the amount of starch to silica, by mass) (Zhang, Liu & Yan, 2016). The synthesis gel preparation procedure was carried out in the following steps: (i) dissolution of aluminum sulphate (Merck, 99%) in 25% of the water required in the synthesis, followed by dropwise addition of sulfuric acid (J. T. Baker, 98%) (Solution A), (ii) dispersion of the soluble starch (Sigma-Aldrich, 99%) in 25% of the water required for synthesis. Solution A was mixed with solution B, under stirring at 200 rpm for 10 min. Then, colloidal silica (Ludox-AS-40, Sigma-Aldrich, 40%) was added to the previous mixture, under stirring at 200 rpm for 10 min, obtaining a paste in which the starch is highly dispersed.

This paste was transferred to a 100 mL teflon vessel, which was placed in a 300 mL Parr Instrument Company stirred reactor (model 4566), and heated at 60 °C for 4 h under agitation at 300 rpm. After the heat treatment, a (black) solid was obtained which was ground. Then, the resulting solid was carbonized in a muffle oven at 170 °C for 17 h. Subsequently, (iii) dissolution of 1,6-diaminohexane in 30% of the water required for synthesis (Solution C) and (iv) dissolution of KOH in 20% of the water required for synthesis (Solution D). Solution D was added to solution C, under stirring at 200 rpm for 10 min. The carbonized solid was then added to the previous mixture, under stirring at 200 rpm for 30 min, and this new mixture was submitted to hydrothermal treatment, in the same Parr reactor described above, at 80 °C for 2 h. The final solid was washed to neutral pH and separated by vacuum filtration, and then oven dried at 120 °C for 12 h (Z22-Starch-Cal sample).

### 2.1.6 Synthesis using calcium carbonate

The stoichiometric molar composition of the synthesis gel 9.0CH$_3$OH:0.3Na$_2$O:1.0Al$_2$O$_3$:90.0SiO$_2$:20.0H$_2$O + 0.16 CaCO$_3$/SiO$_2$ (ratio of 0.16 times the amount of CaCO$_3$ to silica, by mass) (Zhu, et al., 2008), and the preparation procedure adopted were similar to that described for the Z22-Me sample (topic 2.1.2). Calcium carbonate was added at the end of the preparation of the reaction mixture, under mechanical stirring at 200 rpm for 30 min. Crystallization was carried out at 170 °C for 20 h. Then, the resulting solid was submitted to an acid treatment with a 0.5 mol L$^{-1}$ hydrochloric acid solution (Neon, 37%), under mechanical stirring at 100 rpm for 30 min, to remove the CaCO$_3$. The ratio of 3 g of solid per 30 mL of solution was used. Finally, the material was washed to neutral pH by vacuum filtration and dried at 120 °C for 12 h (sample Z22-CaCO3-HCl).

### 2.1.7 Synthesis using silanized silica

The molar composition of the synthesis gel 27.0NH$_2$(CH$_2$)$_6$NH$_2$:13.5K$_2$O:1.0Al$_2$O$_3$:90.0SiO$_2$ (-20% by mass):4.5 H$_2$SO$_4$:3600H$_2$O + 20% silanized silica (percentage by mass, in relation to SiO$_2$) (Silva, et al., 2018) was similar to that described for the Z22-ADE sample (topic 2.1.1). 20% of the colloidal silica (Ludox-HS-40, Sigma-Aldrich, 40%) required for the synthesis were previously separated, which was mixed with a solution of 2.5 g of silane in 40 mL of water, and this mixture was then dried in an oven for 12 h at 110 °C. After this time, the solid obtained was added to the synthesis gel, under mechanical stirring at 200 rpm for 10 min. The hydrothermal treatment of the gel was carried out at 160 °C in a 1 L Parr Instrument Company stirred reactor (model 4520), under agitation at 400 rpm for 20 h.

Subsequently, the resulting solid was submitted to an alkaline treatment with a sodium hydroxide solution (Merck, 98%) of 0.1 mol L$^{-1}$ at 80 °C for 2 h. The ratio of 1 g of solid to 80 mL of alkaline solution was used (Del Campo, et al., 2018). Finally, the material was washed to neutral pH, separated by vacuum filtration and dried at 120 °C for 12 h (Z22-HS40-Des sample).

### 2.2 Heat treatment and ion-exchange

All synthesized samples were calcined at 550 °C for 6 h in a muffle furnace, at a heating rate of 2 °C min$^{-1}$, under an air flow of 100 mL min$^{-1}$. All calcined materials were submitted to three consecutive ion exchange steps with 0.5 mol L$^{-1}$...
ammonium nitrate solution (Sigma-Aldrich, 99%) at 60 °C for 2 h, to obtain zeolite in its acidic form. The ratio of 1 g of solid to 75 mL of solution was used. After the ion exchanges, the materials were oven dried at 100 ºC for 12 h. Finally, a new heat treatment was carried out, under a synthetic air flow of 100 mL min⁻¹ at a heating rate of 2 °C min⁻¹ up to 550 ºC for 6 h.

2.3 Characterization

The X-ray diffraction analysis (XRD) was performed in a Shimadzu XRD-6000 diffractometer, CuKa (λ = 0.1542 nm), Ni filter, 40 kV voltage and 30 mA current. The data was collected in the 20 range between 3 and 40º, with goniometer velocity of 2º min⁻¹ and step of 0.02º. The area of the diffraction peaks located in the 20 regions of 10.9º - 19.67º and 23.6º - 25.0º (Sousa, et al., 2017) were used to calculate the crystallinity of the materials.

Nitrogen adsorption-desorption measurements were performed using a Micromeritics ASAP 2020 equipment at -196 ºC, in the relative pressure range (P/P₀) between 0.01 and 0.99. The solids were previously degassed at 350 ºC for 12 h, under vacuum of 2 µmHg. The specific surface area (S_BET) was calculated using the BET method (Brunauer, Emmett & Teller, 1938). The microporous area (S_Micro), external surface area (S_total) and microporous volume (V_Micro) were determined by the t-plot method (Lippens & De Boer, 1965). The total pore volume (V_Total) was measured using the single-point BET method (Dollimore & Spooner, 1974) at P/P₀ = 0.995. The mesoporous volume (V_Meso) was obtained by the difference between the total pore volume and the micropore volume (V_Meso = V_Total - V_Micro) (Silva, et al., 2019). The pore size distribution was obtained from the adsorption branch of the isotherm by the BJH method (Barrett, E. P., Joyner, L. G. & Halenda, 1951).

Temperature-programmed desorption of ammonia (NH₃-TPD) was conducted in a multipurpose reaction system (SAMP3 model equipment, Termolab - Brazil) equipped with thermal conductivity detector. In these measurements, a sample of approximately 100 mg was submitted to a pretreatment at 400 ºC for 1 h, under helium atmosphere with a flow rate of 30 mL min⁻¹. Then, the temperature was reduced to 100 ºC and the sample was kept in contact with the ammonia stream at a flow rate of 30 mL min⁻¹ for 40 min. The next step of the analysis was the removal of the physisorbed NH₃ molecules at 100 ºC for 1 h, under a helium flow rate of 30 mL min⁻¹. Finally, the NH₃ desorption curves were obtained in the temperature range between 100 and 800 ºC, with a heating rate of 10 ºC min⁻¹ and under helium flow rate of 30 mL min⁻¹.

Scanning electron microscopy (SEM) was carried out using a Shimadzu SSX-550 and Mira3 Tescan electron microscopes. Thermal analyzes (TG/DTG) were used to determine the coke content of the partially deactivated samples after the n-heptane catalytic cracking reaction, and were carried out in a Shimadzu DTG-60H thermobalance, using alumina crucibles, with sample mass of approximately 6 mg. The catalysts were previously treated under a nitrogen atmosphere with a flow rate of 50 mL min⁻¹ for 1 h at 100 ºC, to eliminate volatile compounds, and then submitted to a synthetic air atmosphere with a flow rate of 50 mL min⁻¹, under a heating rate of 10 ºC min⁻¹ in the range of ambient temperature to 800 ºC.

2.4 Catalytic tests

The catalytic tests were performed in a Termolab TCAT-3 catalytic evaluation unit. The HZSM-22 samples that, based on the NH₃-TPD results, showed higher values of acid site densities were evaluated in the n-heptane cracking model reaction (Sigma-Aldrich, 99%). The tests were carried out using 100 mg of zeolite in a fixed bed tubular microreactor at 650 ºC, under atmospheric pressure for 180 min, WHSV of 2 h⁻¹ and nitrogen was used as carrier gas with a flow rate of 20 mL min⁻¹. The reaction products were analyzed on a Shimadzu GC-2014 chromatograph equipped with a FID detector.
3. Results and Discussion

3.1 X-ray diffraction

Figure 1 shows the X-ray diffractograms of the synthesized samples and the pattern for the TON structure (Treacy & Higgins, 2007). All samples presented diffraction peaks referring to the ZSM-22 zeolite (JCPDS PDF 38-197), without the presence of impurities compared to the simulated standard diffractogram. The samples crystallinity followed the order: Z22-Me-CS (100%) > Z22-ADE (98%) > Z22-P123 (66%) > Z22-Vynil (66%) > Z22-Me (63%) > Z22-Brij93 (34%) > Z22-CaCO3-HCl (31%) > Z22-Starch-Cal (24%) > Z22-HS40-Des (22%).

Figure 1. X-ray diffractograms of the ZSM-22 samples.

The materials synthesized using methanol and seed crystals (Z22-Me-CS) and 1,6-diaminohexane (Z22-ADE) showed diffraction peaks with greater definition and intensity. The use of 1,6-diaminohexane as a structure-directing agent is described in several studies that aimed to obtain ZSM-22, either for structure modification or for application (Liu, et al., 2015; Niu, et al., 2017). The Z22-Me-CS sample obtained the expected result for zeolite synthesis using seed crystals. In this case, it was possible to crystallize ZSM-22 in a short time (3 h) in the absence of structure-directing agents. Wang et al. (2014) and Sousa et al. (2017) reported a similar effect on the use of seed crystals in the reaction mixture, which act as pre-formed nuclei, thus reducing the crystallization time. The Z22-Me sample showed intermediate crystallinity, which is probably associated with the weak affinity of the OH group of the alcohol with the zeolitic structure, compared to other organic compounds such as amines (Qian, et al., 2001), resulting in a lesser effect of pore filling and structure directing. Jamil et al. (2016) described the importance of alcohol as a co-solvent, being responsible for promoting the control of crystal growth and inhibiting the
formation of competing phases.

There is a small crystalline organization along the amorphous phase in the synthesis gel aged for 20 h, which can be attributed to some intermediate species of silicate (Mintova & Valtchev, 2002). Jamil et al. (2014) reported an increase in the crystallinity of ZSM-22 from 52% to 98% with the addition of a small amount of the brij76 surfactant (Brij/Al = 2.5). However, when adding a greater amount of surfactant (Brij/Al = 10) there was a decrease in crystallinity or amorphous material was formed. Although the amount adopted in this work was Brij/Al = 2.5, the crystallinity of the material synthesized with Brij93 may be related to the higher molecular weight of this compound (Brij76 MW = 314.6 g mol\(^{-1}\) and Brij93 MW = 356.58 g mol\(^{-1}\)) resulting in an excess amount in the medium reaction, and in this way, slowing down the formation of the ZSM-22 zeolite. According to Aguado, Serrano & Rodríguez (2008), the presence of the silane group in the reaction mixture would have the function of inhibiting the growth and agglomeration of the zeolite crystals. The same effect observed for the material obtained with Brij93 occurred in the cases of samples synthesized with pluronic P123 and vinyltrimethoxysilane.

The material synthesized using starch (JCPDS PDF 39-1911) showed an amorphous region due to the presence of this compound in the sample. After the calcination process, the ZSM-22 phase became better defined (Z22-Starch-Cal), suggesting the effective removal of starch from the zeolite structure. The sample synthesized with carbonate exhibited a mixture of the ZSM-22 and CaCO\(_3\) phases (JCPDS PDF 47-1743). The carbonate was removed after the acid treatment and the characteristic peaks of the TON phase (Z22-CaCO\(_3\)-HCl) were the only observed. Both treatments to remove starch and CaCO\(_3\) led to an increase in the crystallinity of the zeolites (Starch from 18% to 24% and CaCO\(_3\) from 18% to 31%). The material obtained with the addition of silanized silica denoted the formation of pure ZSM-22, with a decrease in the intensity of the peaks after desilication (Z22-HS40-Des), due to the extraction of structural silicon atoms (Silva, et al., 2018).

### 3.2 Nitrogen adsorption-desorption measurements

Figure 2 shows the nitrogen adsorption-desorption isotherms of the ZSM-22 zeolites. Practically all samples presented isotherms with a type I profile, exhibiting low adsorbed amounts in the range of P/P\(_0\) < 0.1, related to the typical microporosity of the TON structure (He, et al., 2021), with a small increase in adsorption in the range P/P\(_0\) < 0.8, indicative of the intercrystalline voids of nanorod clusters. The exception was the Z22-Me-CS sample, which showed a combination of isotherms with types I and IV profiles, with the formation of a hysteresis at relative pressures between 0.6 < P/P\(_0\) < 1.0, denoting a micro-mesoporous material. The BJH distribution exhibits an "apparent" mesoporosity for the samples Z-ADE, Z22-Me-CS and Z22-Brij93, possibly due to the filling of irregular pores formed by the intergrowth between the crystals or particles that form these materials (Carvalho & Urquieta-Gonzalez, 2015). Z22-HS40-Des and Z22-Starch-Cal samples had pores distribution in the micro-mesopore range, which is effect of the post-synthesis treatment carried out on the solids.

The textural properties obtained from the nitrogen adsorption-desorption analyzes are listed in Table 1. The results for the Z22-ADE sample are similar to the values described in the literature (Lazareva, et al., 2019), indicating the effectiveness of 1,6-diaminohexane as OSDA to synthesize the TON structure. The Z22-Me sample had lower area and volume values, denoting a lower “filling” effect of the zeolite pores by methanol in relation to the amine. The use of crystallization seeds (Z22-Me-CS) provided an increase in the external area and, together with the Z22-ADE sample, resulted in a pseudo mesoporosity, due to the interstices between the nanorod clusters (Sousa, et al., 2017). The materials prepared with the aging of the synthesis gel and subsequent polymer surfactant and silane addition (Z22-P123, Z22-Brij93 and Z22-Vynil) showed an increase in the external area in comparison to Z22-ADE sample, as they are formed by smaller crystals (Jamil, et al.; 2014). The sample obtained with silanized and desilicate silica (Z22-HS40-Des) effectively showed an increase in S\(_{\text{ext}}\) and generation of V\(_{\text{Mes}}\) as described in the BJH distribution.
Figure 2. Nitrogen adsorption-desorption isotherms of ZSM-22 samples. For better visualization, the following isotherms were shifted: Z22-20h-Brij93 (40 cm$^3$ g$^{-1}$) and Z22-Vynil (130 cm$^3$ g$^{-1}$).

Source: Authors.

Table 1. Textural properties obtained from nitrogen adsorption-desorption analyses.

| Sample           | $S_{BET}$ (m$^2$ g$^{-1}$) | $S_{Micro}$ (m$^2$ g$^{-1}$) | $S_{Ext}$ (m$^2$ g$^{-1}$) | $V_{Total}$ (cm$^3$ g$^{-1}$) | $V_{Micro}$ (cm$^3$ g$^{-1}$) | $V_{Meso}$ (cm$^3$ g$^{-1}$) |
|------------------|-----------------------------|-----------------------------|---------------------------|-------------------------------|-------------------------------|-------------------------------|
| Z22-ADE          | 265                         | 243                         | 20                        | 0.23                          | 0.1                           | 0.13                          |
| Z22-Me           | 229                         | 198                         | 31                        | 0.11                          | 0.08                          | 0.03                          |
| Z22-Me-CS        | 254                         | 185                         | 71                        | 0.23                          | 0.08                          | 0.16                          |
| Z22-P123         | 259                         | 224                         | 35                        | 0.15                          | 0.09                          | 0.06                          |
| Z22-Brij93       | 263                         | 223                         | 40                        | 0.20                          | 0.09                          | 0.11                          |
| Z22-Vynil        | 276                         | 230                         | 46                        | 0.17                          | 0.09                          | 0.08                          |
| Z22-Starch-Cal   | 162                         | 129                         | 33                        | 0.13                          | 0.05                          | 0.08                          |
| Z22-CaCO3-HCl    | 202                         | 175                         | 27                        | 0.14                          | 0.07                          | 0.07                          |
| Z22-HS40-Des     | 282                         | 192                         | 90                        | 0.39                          | 0.08                          | 0.31                          |

Source: Authors.
3.3 Scanning electron microscopy

The micrographs (Figure 3) in general did not show differences in the morphology of the materials, that are composed of agglomerates of crystals with nanorods shape, characteristic of ZSM-22 (Li, et al., 2021), with size in the 1-2 μm range. The interstices between the crystals provided the characteristic of a pseudo mesoporosity, for purely microporous materials, obtained from nitrogen adsorption-desorption analyses. The exception was the Z22-Vynil sample, which exhibited crystals with a higher agglomeration degree, taking the form of practically a single particle. Due to the peculiar morphology of the ZSM-22, it was not observed cracks in the crystals of the Z22-HS40-Des sample caused by the desilication process, as reported by Verboekend et al. (2011). The effect of micro-mesoporosity generation was better highlighted by the nitrogen adsorption-desorption analyses.

Figure 3. Micrographs of the ZSM-22 samples: (a) Z22-ADE, (b) Z22-Me, (c) Z22-Me-CS, (d) Z22-P123, (e) Z22-Brij93, (f) Z22-Vynil, (g) Z22-HS40-Des, (h) Z22-CaCO3-HCl and (i) Z22-Starch-Cal.

Source: Authors.

3.4 Temperature-programmed desorption of ammonia

Figure 4 shows the ammonia desorption profiles of the ZSM-22 zeolites. All samples provided two main peaks of ammonia desorption, one approximately between 100 and 320 °C and the other in the range of 320 and 550 °C, indicating the presence of Lewis and Brønsted acidity, respectively (Li, et al., 2017; Silva, et al., 2020). The first low temperature desorption event represents the weak acidic sites and the second high temperature event is attributed to the strong acidity sites (Gao, et al., 2020; Quintela, et al., 2021). The exceptions were the Z22-Vynil and Z22-CaCO3-HCl samples, which exhibited a third less intense peak between 550 and 620 °C. This fact is linked to a process of dehydroxylation of the structures (Lok, Marcus & Angell, 1986).
Assuming the stoichiometry of one ammonia molecule per acid site, the acid site densities can be obtained from the Gaussian deconvolution of the TPD curves (Lok et al., 1986). The results from the NH$_3$-TPD analyzes are listed in Table 2. Nearly all synthesized samples had higher densities of weak acidic sites compared to strong acidic sites, which is a negative effect, since Brønsted acidity plays a role of greater importance in the catalytic cracking of hydrocarbons (Gao et al., 2020). The exception was the Z22-P123 sample, which showed similar densities between the two desorption events.

Hayasaka et al. (2007) reported that through X-ray excited photoelectron spectroscopy (XPS) analysis they observed that a large part of the aluminum was located on the external surface of the ZSM-22 crystals, having a very siliceous internal structure. Recent investigations on aluminum location in TON type zeolites by NMR techniques revealed a preferential population of T3 and T4 sites (Camblor et al., 1998). Especially the T4 sites located in lobes of the main because they are prime candidates for aluminum positions (Hayasaka et al., 2007). This implies a higher proportion of acid sites near pore mouths (Thaker et al., 2016), which favour undesirable bimolecular reactions, such as alkylation, hydrogen transfer and isomerization, because of less spatial restrictions (Rahimi & Karimzadeh, 2011). Primary products (such as ethylene and propene) can undergo further reactions at external acid sites, leading to the formation of coke precursors (benzene, toluene and xylenes) and coke deposits (Javaid et al., 2015; Inagaki et al., 2016).

The samples that were submitted to a post-synthesis treatment, Z22-HS40-Des and Z22-CaCO3-HCl, possibly suffered a certain degree of structure dealuminization. Verboekend et al. (2011) reported that due to the peculiar morphology of the ZSM-22 crystals in the form of "needles" and the higher concentration of Al on the external surface, alkaline and/or acid treatments, despite generating inter- and intra-crystal mesoporosity, can lead to the leaching of part of the aluminum atoms of the zeolitic structure and even the breakdown of the finest crystals, which can influence the catalytic properties of these materials.
Table 2. Results obtained from the NH$_3$-TPD analyses.

| Sample            | 1st Peak | | 2nd Peak | | ∑C (μmol g$^{-1}$) |
|-------------------|----------|-----------------|----------|-----------------|
|                   | $T_{\text{Max}}$ (°C) | C (μmol g$^{-1}$) | $T_{\text{Max}}$ (°C) | C (μmol g$^{-1}$) |                  |
| Z22-ADE           | 224      | 264             | 457      | 119             | 383              |
| Z22-Me            | 217      | 102             | 436      | 40              | 142              |
| Z22-Me-CS         | 224      | 291             | 454      | 125             | 416              |
| Z22-P123          | 233      | 121             | 449      | 123             | 244              |
| Z22-Brij93        | 204      | 244             | 448      | 117             | 360              |
| Z22-Vynil         | 198      | 197             | 437      | 66              | 263              |
| Z22-HS40-Des      | 203      | 265             | 443      | 56              | 321              |
| Z22-CaCO3-HCl     | 161      | 150             | 444      | 73              | 223              |
| Z22-Starch-Cal    | 202      | 121             | 576      | 17              | 138              |

Source: Authors.

3.5 Catalytic performance

From the textural properties and total density of acidic sites results, the Z22-ADE, Z22-Me-CS, Z22-Brij93 and Z22-HS40-Des samples were selected to evaluate the catalytic performance in the n-heptane cracking reaction.

The conversions and coke content of ZSM-22 zeolites for the catalytic cracking of n-heptane are shown in Figure 5. It was found that the Z22-ADE and Z22-Brij93 samples showed higher initial conversions, followed by a strong decrease in catalytic activity. This led to a higher coke content deposited on these microporous materials. The Z22-Me-CS and Z22-HS40-Des samples initially provided lower conversions, showing greater stability during the reaction time. The Z22-Me-CS sample, despite having strong acid site density similar to Z22-ADE and Z22-Brij93, does not exhibit the same catalytic activity behavior, indicating that for these one-dimensional materials the acidity is related not only to the SiO$_2$/Al$_2$O$_3$ ratio, but also with the coordination of aluminum atoms in the crystal structure (Silva, et al., 2019). The lower activity showed by the Z22-HS40 sample is related to a lower density of strong acid sites, resulting from a certain degree of dealumination of the structure after desilication (Verboekend, et al., 2011). On the other hand, the generation of mesopores led to a greater stability and lower coke content compared to the microporous zeolites.
The coke content was measured by thermal analysis (Figure 6), where it was found that all catalysts had two mass loss events, with the first occurring at temperatures below 550 °C, due to thermal degradation of light coke (aliphatic compounds), and the second at temperatures above 550 °C, attributed to the combustion of heavy coke (aromatic compounds) (Zhang, et al., 2018). The deactivation process of zeolites occurs mainly due to the formation of coke, which is produced through secondary reactions that occur on the catalyst surface, which can lead to the blocking of pore openings and deactivation of active sites. Therefore, the mesopores generated in the Z22-HS40-Des sample allowed an increase in molecular diffusion, and with this, the micropores were more slowly blocked due to the greater accessibility to the interior of the zeolite structure (Silva, et al., 2022).
The products distribution obtained in the n-heptane catalytic cracking is listed in Table 3. There was no significant change in the products percentage, in which all catalysts showed selectivity to the formation of compounds in the range C2 to C4 in similar proportions. The reaction temperature used favors the formation of ethene, as it is formed by primary carbonium ions that require high activation energy (Wang, et al., 2015).

Table 3. N-heptane catalytic cracking products distribution and coke content after 180 min of reaction.

| Sample          | Product distribution (%) |      |      |      |      |
|-----------------|--------------------------|------|------|------|------|
|                 | C1           | C2   | C3   | C4   | C5+  |
| Z22-ADE         | 1.0          | 10.0 | 32.0 | 32.0 | 25.0 |
| Z22-Me-CS       | 1.0          | 12.0 | 33.0 | 30.0 | 24.0 |
| Z22-Brij93      | 1.0          | 13.0 | 40.0 | 36.0 | 11.0 |
| Z22-HS40-Des    | 1.0          | 11.0 | 34.0 | 33.0 | 21.0 |

Source: Authors.

4. Conclusion

ZSM-22 zeolite was successfully obtained from all the studied synthesis routes. The use of methanol together with seed crystals led to a reduction in crystallization time (3 h), with the Z22-Me-CS sample being the most crystalline (100%). The acidity, specific area and volume values were close to those obtained from conventional synthesis (Z22-ADE) that uses 1,6-diaminohexane as an organic structure-directing agent. The synthesis using only methanol (Z22-Me) resulted in the crystallization of ZSM-22 with inferior characteristics in comparison to Z22-ADE, indicating a low interaction of alcohol with the zeolitic structure. The aging of the synthesis gel with subsequent addition of polymer, surfactant and silane did not cause significant changes in the properties of ZSM-22. Among the synthesis routes using calcium carbonate, starch and silanized silica followed by alkaline post-treatment, only the latter generated the micro-mesoporous ZSM-22.

The optimization of the ZSM-22 synthesis is not easy to perform, because, as reported in the literature, this zeolite has a formation process sensitive to any slight change in the synthesis parameters, which may lead to undesired physicochemical characteristics or even contamination by other crystalline phases. For further research, we propose the study of the synthesis of ZSM-22 with the simultaneous use of different amines, to evaluate the effect of different kinetic diameters and affinity of these compounds on the crystallization, textural properties and acid density of the zeolite. Another proposed study is the application of amino acid salts as possible mesoporosity agents, in order to obtain micro-mesoporous structures to improve the diffusivity in the ZSM-22 zeolite.

Acknowledgments

The authors are grateful for the financial support of the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES), through a research grant and the Petrobras.

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