## Structural and chemical analysis of Zn ion exchange in thermally modified zeolite A4

### Análisis estructural y químico del intercambio iónico de Zn en zeolita A4 modificada térmicamente

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#### Abstract

This work reports the analysis of Zn ion exchange in thermally modified zeolite A4 (ZA4) for the potential formation of ZnO nanoparticles. The methodology consists of two steps. Step 1 consisted of carrying out different thermal treatments on ZA4. Step 2 involved the ion exchange of Zn with different concentrations of the Zn ion precursor. XRD and FTIR analyses revealed a transformation in the crystal and molecular structure of ZA4 after heat treatment. This work has been limited to studying ion exchange; however, it is very interesting to study the thermal behavior of ZA4 because this could improve the surface area of the material. The results obtained in this work demonstrate that heat treatment and Zn ion concentration affect the crystallinity and molecular structure of ZA4.

Keywords: Zn, zeolite A4, heat treatment, structural analysis.

#### Resumen

Este trabajo reporta el análisis del intercambio de iones de Zn en la zeolita A4 modificada térmicamente (ZA4) para la potencial formación de nanopartículas de ZnO. La metodología realizada, consta de un proceso de 2 etapas. La etapa 1 consistió en realizar diferentes tratamientos térmicos en ZA4. La etapa 2 involucró el intercambio iónico de Zn con diferentes concentraciones del precursor de iones Zn. Los análisis por DRX y FTIR muestran una transformación en la estructura cristalina y molecular de ZA4 después de los tratamientos. Este trabajo se ha limitado al estudio del intercambio iónico, sin embargo, resulta muy interesante conocer el comportamiento térmico de ZA4, ya que esto, puede favorecer el área superficial del material. Los resultados obtenidos en este trabajo demuestran que el tratamiento térmico y la concentración de iones Zn afectan la estructura cristalina y molecular de ZA4.

Palabras clave: Zn, zeolita A4, tratamiento térmico, análisis estructural.

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# 1 Introduction

Zeolites consist of  $[SiO_4]^{4-}$  and  $[AIO_4]^{5-}$  tetrahedral frameworks connected at vertices by oxygen atoms, with group I and II elements as exchange ions. The high selective adsorption capacity of ions is their main characteristic (Eroglu, Emekci, & Athanassiou, 2017; Leal-Perez, Flores-Valenzuela, Cortez-Valadez, et al., 2022). In particular, Zeolite Linde Type A (Zeolite LTA or Zeolite A) was the first synthetic zeolite commercially available (Breck, Eversole, & Milton, 1956). The general formula of LTA zeolite is  $X_{12}[(AlO_2)_{12}(SiO_2)_{12}]$ ·27H<sub>2</sub>O, where X could be K, Na or Ca (for Ca, 12 is changed by 6 due to the valence electrons), which are referred to as zeolites A3, A4, and A5, respectively. Moreover, the ratio of structural silicon to aluminum (Si/Al) is 1, and sodium ions are exchangeable structural cations (Julbe & Drobek, 2016). The fact that LTA zeolite has a Si/Al ratio equal to 1 means that it potentially has a high cation exchange capacity (Collins, Rozhkovskaya, Outram, & Millar, 2020).

Ion exchange in zeolites is a metathesis reaction (double displacement reaction) between the zeolite and a specific salt, where a specific reaction occurs between the cations of both species (Hoveyda & Zhugralin, 2007; Leal-Perez, Flores-Valenzuela, Vargas-Ortíz, et al., 2022). One of the first ion exchange researchers studying zeolites was Eichhorn, who reported that this type of reaction was reversible (Eichhorn, 1858). However, no industrial applications of this process are available, and for several years, zeolites were considered to have simple mineralogical curiosities. However, around the 1950s, important deposits were discovered in Japan, Italy, and the United States. After that, zeolites (called ion exchange resins) became the main product for solving problems related to water softening and deionization within a few years (Colella, 1996; Gans, 1905).

On the other hand, the ion exchange of Zn in a zeolite can offer a promising precursor for the formation of ZnO nanoparticles. A controlled size and homogeneous distribution were demonstrated for other nanoparticles synthesized from zeolites (Leal-Perez, Flores-Valenzuela, Vargas-Ortíz, et al., 2022). Zinc oxide (ZnO) is a semiconductor with a band gap of 3.37 eV at room temperature. Different physical and chemical properties can be obtained depending on the nanostructure morphology (Reyes-Zambrano, Lecona-Guzmán, Luján-Hidalgo, & Gutiérrez-Miceli, 2024). For this purpose, ZnO is considered a technologically relevant material with a variety of applications, such as semiconductors, sensors, magnetic and actuator materials, and cosmetic ingredients (Hasanpoor, Aliofkhazraei, & Hamid Delavari, 2016). In addition, Zn has healing and bactericidal properties, which are enhanced by being in a matrix such as a zeolite with a large surface area (Rajendran, Kumar, Houreld, & Abrahamse, 2018).

The ion exchange of Zn in zeolites is a relatively simple process because Zn has a 2+ valence, allowing only a single oxidation state and a unique way of bonding to the zeolite structure. There are different works related to Zn ion exchange and the formation of nanoparticles (Almutairi, Mezari, Magusin, Pidko, & Hensen, 2012; Luzgin et al., 2008; Ostroski et al., 2009). However, no studies on the ion exchange of Zn in thermally modified zeolites have been reported.

In this work, the effect of Zn ion exchange in A4 zeolite modified by thermal treatment was studied to increase the surface area of the matrix (ZA4). The crystalline and molecular structures were defined by XRD and FTIR.

# 2 Materials and methods

Synthetic zeolite A4 (Sigma Aldrich, 98 %), zinc acetate  $(CH_3COO)_2Zn.2H_2O$  (Sigma Aldrich, 99 %), and deionized water were used as the hydrates.

### 2.1 Thermally modified zeolite A4

Three samples with 12 grams of zeolite A4 were prepared, and sample 1 was thermally treated at 500  $^{\circ}$ C for 1 h. The second sample was thermally treated at 750  $^{\circ}$ C for 1 h. The third sample was thermally treated at 1000  $^{\circ}$ C for 1 h. All the samples were heated and cooled at a rate of 5  $^{\circ}$ C/min. These materials were named ZA4-500, ZA4-750 and ZA4-1000, respectively.

### 2.2 Thermally modified zeolite A4

For ZA4-500, the material was divided into 4 containers with 3 g, and 50 ml of distilled water each was added for hydration for 24 h. Afterwards, 3 solutions of 100 ml of (CH<sub>3</sub>COO)<sub>2</sub>Zn.2H<sub>2</sub>O were prepared at molarities of 0.01, 0.05, and 0.1 M. Then, hydrated zeolite and (CH<sub>3</sub>COO)<sub>2</sub>Zn.2H<sub>2</sub>O solutions were put in a thermal bath to obtain an internal temperature of 50 °C. After that, a solution of 0.01 M (CH<sub>3</sub>COO)<sub>2</sub>Zn.2H<sub>2</sub>O was slowly added to the container with hydrated zeolite. A solution of 0.05 M (CH<sub>3</sub>COO)<sub>2</sub>Zn.2H<sub>2</sub>O was slowly added to the container with hydrated zeolite. A 0.01 M solution of (CH<sub>3</sub>COO)<sub>2</sub>Zn.2H<sub>2</sub>O was slowly added to a vessel with hydrated zeolite. The final samples were filtered and dried at room temperature for 24 h. ZA4-500-0.01, ZA4-500-0.05, and ZA4-500-0.1 were obtained.

This process was repeated for ZA4-750 and ZA4-1000 under the same conditions, and the order of the nomenclature of the samples was maintained.

#### 2.3 Characterization

XRD analysis was carried out with a PHI5100 BRUKER AXS D8 ADVANCE diffractometer. The vibrational energy of the bonds present in the structures of the different samples was determined by an IRAffinity 1-S Shimadzu Infrared Spectrometer. Micrographs and EDS results were measured with a JEOL JSM-7401F field emission scanning electron microscope (FESEM).

## **3 Results and discussion**

Figure 1 shows the FT-IR spectra of ZA4 and ZA4-500, ZA4-750, and ZA4-1000. In ZA4, the peaks at 1655 and 3423 cm<sup>-1</sup> are attributed to the distinctive vibrational stretching modes of the -OH bonds present in both chemical and physical water. Additionally, the peak at 1000 cm<sup>-1</sup> is characteristic of the asymmetric stretching vibrations of Si-O-Si and Si-O-Al. The peak at 556 cm<sup>-1</sup> indicates the symmetric stretching vibrations of the Si-O-Si and O-Si-O bonds, while the peak at 670 cm<sup>-1</sup> corresponds to the asymmetric stretching vibrations of the Si-O-Al bonds (Mozgawa, Król, & Barczyk, 2011; Şen, Bardakçi, Yavuz, & Gök, 2008). For ZA4-500, the characteristic peaks of ZA4 are similar. However, the peak at 556 cm<sup>-1</sup> shows a slight shift and the possible presence of a second peak near 500 cm<sup>-1</sup>. This may be attributed to the effect of temperature on the modification of the molecular structure of ZA4; moreover, the peaks at 1655 and 3423 cm<sup>-1</sup> are very small, which is caused by thermal treatment (Ates & Hardacre, 2012). For ZA4-750, the peak at 556 cm<sup>-1</sup> shifted to 570 cm<sup>-1</sup>, and the peak shifted to less than 500 cm<sup>-1</sup>. In addition, peak formations are observed in the range of approximately 713 to 633 cm<sup>-1</sup>, indicating possible molecular deformation in ZA4 (Aronne, Esposito, Ferone, Pansini, & Pernice, 2002; Yörükoğullar, Yilmaz, & Dikmen, 2010). These shifts and peak formations are more evident as the treatment temperature increases in ZA4; moreover, the peaks at 1665 and 3413 cm<sup>-1</sup> disappear. For ZA4-1000, the peak at 558 cm<sup>-1</sup> disappeared, which may indicate the breakdown of the D4R secondary structure, which is associated with that peak (Mozgawa et al., 2011), as well as the shift of the peak at 672 to 687 cm<sup>-1</sup>, indicating possibly significant modification in the molecular structure of ZA4. Similarly, the peaks at 1665 and 3413 cm<sup>-1</sup> disappeared after heat treatment. It can be observed that each of the thermal treatments applied to ZA4 caused the breakdown of SBU D4R, as indicated by the peak at 556 cm<sup>-1</sup> (Flores-Valenzuela et al., 2023).



Figure 1. FT-IR spectra of ZA4, ZA4-500, ZA4-750, and ZA4-1000.



Figure 2. FT-IR spectra of ZA4-500, ZA4-500-Zn0.01, ZA4-500-Zn0.05, and ZA4-500-Zn0.1.

Figure 2 shows the FT-IR spectra of ZA4-500, ZA4-500-Zn0.01, ZA4-500-Zn0.05, and ZA4-500-Zn0.1. ZA4-500-Zn0.01 shows similar behavior to that of ZA4-500, with the presence of a new peak at 897 cm<sup>-1</sup> attributed to the stretching modes of Zn-O (Ashokkumar & Muthukumaran, 2014). In addition, the peaks at 512 cm<sup>-1</sup> for ZA4-500-Zn0.05 and ZA4-500-Zn0.1 are associated with the stretching modes of Zn-O to octahedral coordination and a possible increase in the degree of tetrahedral polymerization of SiO<sub>4</sub> (Ginting et al., 2019; Xiong, Pal, Serrano, Ucer, & Williams, 2006), as well as elongation of the peak at 897 cm<sup>-1</sup>, indicating a greater presence of Zn-O bonds (García-Molina et al., 2024). Another peak at 1141 cm<sup>-1</sup> attributed to the normal polymeric O-H stretching vibration of H<sub>2</sub>O in the Zn-O lattice is observed (Ashokkumar & Muthukumaran, 2014). These results provide evidence of alterations in the molecular structure of ZA4-500 due to changes in the Zn concentration.

Figure 3 shows the FT-IR spectra of ZA4-750, ZA4-750-Zn0.01, ZA4-750-Zn0.05 and ZA4-750-Zn0.1. The FT-IR spectra showed no apparent effect on the characteristic peak for ZA4-750; the peak at approximately 690 cm<sup>-1</sup> was more intense; however, the FT-IR study was carried out qualitatively. Furthermore, the peak at 556 cm<sup>-1</sup> disappears when ion exchange is carried out. Therefore, we can affirm



Figure 3. FT-IR spectra of ZA4-750, ZA4-750-Zn0.01, ZA4-750-Zn0.05 and ZA4-750-Zn0.1.



Figure 4. FT-IR spectra of ZA4-1000, ZA4-1000-Zn0.01, ZA4-1000-Zn0.05, and ZA4-1000-Zn0.1.

that the molecular structure of ZA4 thermally treated at 750 °C is affected by nanoparticle formation, ionic clusters or simple Zn ion exchange (Flores-Valenzuela et al., 2023; Ginting et al., 2019).

Figure 4 shows the FT-IR spectra of ZA4-1000, ZA4-1000-Zn0.01, ZA4-1000-Zn0.05, and ZA4-1000-Zn0.1. Under experimental conditions, the FR-IR spectra did not significantly change during ion exchange; therefore, we can conclude that ion exchange does not affect the molecular structure of ZA4 thermally treated at 1000 °C.

Figure 5 shows the X-ray diffraction patterns of ZA4, ZA4-500, ZA4-750 and ZA4-1000. The characteristic Miller indices of ZA4 are indicated by the solid line, indexed with card JCP2:01-089-5423. For ZA4-500, the same Miller indices characteristic of ZA4 were observed; however, a slight decrease in the intensity of the (220), (222) and (420) peaks was observed because of the heat treatment process applied to the zeolite. For ZA4-750, the characteristic peaks of ZA4 are observed, and new peaks at 21.3°, 24.6°, 29.4°, and 35.3° are also observed. According to the Miller indices, (003), (01-2), (01-3) and (01-4) belong to the "nepheline" group (Na23.55Al24Si24O96), which has a monoclinic structure and is indexed with PDF 96-901-0481. Nepheline is a porous tectosilicate with a chemical formula similar to that of ZA4. It presents



Figure 5. X-ray diffraction patterns of ZA4, ZA4-500, ZA4-750 and ZA4-1000.



Figure 6. X-ray diffraction patterns of ZA4-500, ZA4-500-Zn0.01, ZA4-500-Zn0.05 and ZA4-500-Zn0.1.

a slight deficit of the element Na, creating a new structure. For ZA4-1000, only peaks for nepheline are observed, which may indicate that ZA4 changes to a monoclinic nepheline structure at high temperatures.

Figure 6 shows the X-ray diffraction patterns of ZA4-500, ZA4-500-Zn0.01, ZA4-500-Zn0.05 and ZA4-500-Zn0.1. For ZA4-500-Zn0.01, a similar behavior to that of ZA4-500 is observed, which may indicate that ion exchange at this concentration does not modify the crystalline properties of ZA4-500. For ZA4-500-Zn0.05, a similar behavior to that of ZA4-500 is observed; however, a decrease in the relative intensity of the peaks associated with Miller indices (220), (222) and (420) is evident. However, this peak could not be indexed to Zn<sup>0</sup> or ZnO; therefore, we may consider the formation of a forbidden plane in the zeolitic structure (He et al., 2015; Williams & Carter, 1996; Yan, Luo, Schaffer, & Qian, 2012). For ZA4-500-Zn0.1, a low peak resolution is observed, which can be interpreted as a tendency to transform the sample into an amorphous material, possibly caused by a high concentration of Zn in solution (Šponer, Sobalík, Leszczynski, & Wichterlová, 2001). Additionally, the crystallographic behavior is similar to that of ZA4-500-Zn0.05, where new peaks can be seen at 13.7° and 17.9°, which are not associated with any possible formation of nanoparticles. This is interpreted as the formation of forbidden planes. An increase in peak intensity is also observed at 22.9°



Figure 7. X-ray diffraction patterns of ZA4-750, ZA4-750-Zn0.01, ZA4-750-Zn0.05 and ZA4-750-Zn0.1.



Figure 8. X-ray diffraction patterns of ZA4-1000, ZA4-1000-Zn0.01, ZA4-1000-Zn0.05, and ZA4-1000-Zn0.1.

and  $36.7^{\circ}$ , and the peak at  $35.7^{\circ}$  could be associated with the (101) plane of the ZnO nanoparticles (Borda, Torres, & Lapidus, 2022; Raoufi, 2013). However, this peak may be shielded by the ZA4 peak. For this sample, evidence of alteration in the crystal structure of ZA4-500 by ion exchange is observed.

Figure 7 shows the X-ray diffraction patterns of ZA4-750, ZA4-750-Zn0.01, ZA4-750-Zn0.05 and ZA4-750-Zn0.1. For these samples, the behavior of the crystalline structure is very similar and without apparent change for different concentrations. Therefore, we can determine that ion exchange does not affect the crystalline structure of ZA4-750; moreover, no evidence of characteristic peaks of nanoparticles is observed. Therefore, the formation of nanoparticles is absent or simply does not satisfy the diffraction criteria. (Warren, 1941; Whittig & Allardice, 2018).

Figure 8 shows the X-ray diffraction patterns of ZA4-1000, ZA4-1000-Zn0.01, ZA4-1000-Zn0.05, and ZA4-1000-Zn0.1. For those samples, the crystal structure behavior is very similar and without apparent variation for all concentrations, maintaining the nepheline structure. For this reason, we can reach the same conclusion as for ZA4-750, where we predict that ion exchange does not affect the crystalline structure of ZA4-1000 and that there is no evidence of characteristic peaks of nanoparticles.

Figure 9 shows the SEM and EDS micrographs for a) ZA4-500-Zn0.01, b) ZA4-500-Zn0.05 and c) ZA4-500-Zn0.1. For each sample, the morphology is similar and has no significant impact. However, EDS analysis shows that the atomic percentage of Zn for a) is 2.4 %, b) is 4.2 % and c) is 5.9 %. This is consistent due to the increasing Zn concentration in each sample. However, the Al/Si ratio was 1:1. However, variations in the molar concentration of the ion exchange result in changes in the Zn/Al ratio. Moreover, the samples for ZA4-750 and ZA4-1000 showed a low presence of Zn, lower than 0.5 atomic %, and therefore, this analysis was not included.



Figure 9. SEM and EDS images of a) ZA4-500-Zn0.01, b) ZA4-500-Zn0.05 and c) ZA4-500-Zn0.1.

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# Conclusion

FT-IR and XRD analyses showed significant molecular and crystalline structure modifications in zeolite due to heat treatment, as well as evidence of changes in each of the samples due to ion exchange. SEM micrographs showed no significant evidence of morphological variation. Additionally, the EDS analysis showed that heat treatment at 500 °C contributed to a greater ion exchange performance of Zn, which increased with increasing molar concentration. Furthermore, we conclude that heat treatments at 750 °C and 1000 °C cause phase transformations in the zeolite to nepheline, which is not suitable for ion exchange.

Importantly, the study of the structural and chemical properties of the thermally modified zeolite allows the prediction of different technological applications. This could be the formation of ZnO nanoparticles to be used as coatings in feed ovens where the elimination of bacteria harmful to the human body is required.

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