INVESTIGATION OF THERMAL AND STRUCTURAL PROPERTIES OF NATURAL AND ION-EXCHANGED ANALCIME

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ABSTRACT

In this study, the thermal and structural properties of natural analcime zeolite obtained from the Çanakkale-Küçükçay-Ayvacık region and its cationic forms (HA, FeA, AgA and CuA) were characterized using a range of methods. These included X-ray fluorescence (XRF), X-ray diffraction (XRD), thermogravimetric analysis (TG), differential thermogravimetric analysis (DTG), differential thermal analysis (DTA) and specific surface area measurement techniques. Cationic forms were prepared by using 1 M HCl, 1 M Fe(NO_3)_3·9H_2O, 1 M AgNO_3 and 1 M Cu(NO_3)_2·3H_2O solutions. Quantitative XRD analysis showed that the natural zeolite was mainly composed of analcime accompanied by quartz, feldspar, montmorillonite and amorphous substances. AgA and HA samples showed different thermal and structural behaviors due to their rearranged crystal structure and chemical composition. Adsorption properties of samples were determined by using adsorption isotherms and specific surface areas obtained by the Brunauer-Emmet-Teller (BET) method. The specific surface areas of the ion-exchanged analcime zeolites were found to be higher in comparison to natural analcime. Among all the modified forms, it was found that the H-form of the analcime sample has the largest surface area HA (41.88 m²/g).

Keywords: Zeolite, Analcime, N₂ Adsorption, XRD, TG–DTG–DTA

1. INTRODUCTION

Zeolites are defined as alumina silicates of alkali and earth alkali metals that have a crystal structure. The smallest structural unit of any zeolite crystal is the SiO₄ or AlO₄ tetrahedron. Natural zeolites are used in many fields like adsorption, catalysis, the building industry, in agriculture and earth enrichment [1]. Analcime, which has the chemical formula NaAlSi₂O₆·H₂O and a cubic crystal structure, is a hydrated, sodium aluminosilicate mineral [2, 3]. The crystal structure of analcime was determined by Taylor [2] and refined by Calleri and Ferraris [4], Knowles et al. [5], Ferraris et al. [6] and Pechar [7]. Mazzi and Galli [8] also demonstrated tetragonal and orthorhombic analcime structures. Analcime is also known as analcite. The name analcime comes from the Greek word ankis, which means “weak” or “without power”. Analcime includes trace amounts of potassium and calcium. Although they are classified as zeolite minerals, their structures and chemical properties are more similar to leucite-type feldspathoids. The Si/Al ratio changes between 1.8 and 2.8. Analcime can be found in basalt and other alkaline volcanic rocks. The crystal structure of analcime is composed of 4-member (Si,Al)–O rings, which are comprised of chains in three dimensions. Analcime (ANA) structures formed in this way include channels parallel to <110> that are connected to 8-member tetragonal rings. Unlike the big extra-framework positions and channel openings which are commonly found in many zeolites – like clinoptilolite – channels in analcime are pretty small, with openings ~ 1.6 Å × 4.2 Å [9, 10]. These small channels affect both the cation exchange behavior and dehydration/hydration behavior of analcime. This type of channel formation enables many structural changes, like structural deformation, channel collapse and ionic movements. Natural and synthetic analcime zeolites cannot be used as molecular sieves and catalysts because the channels in its crystal structure are pretty narrow. In zeolites with small pores, like analcime, only small molecules (e.g. H₂, He, H₂O) can penetrate the structure.
The physical and chemical properties of zeolites can be improved using several methods like acid treatment and ion exchange. Ion-exchange selectivity changes depending on the crystal structure of zeolite, the cation properties (ionic radius and hydration energy) and ion-exchange states [11, 12]. It was determined by Barrer [13] that analcime features two types of ion-exchange. The first type is the exchange of ions of at room temperature and the second type is the exchange at elevated temperatures. Since analcime has small pores, ion exchange happens more easily when the temperature rises [14]. Barrer and Hinds [14] showed that when Na-analcime is exchanged with transition metals, the amount of silica in zeolites increase. On the other hand, Co$^{2+}$, Ni$^{2+}$, Cu$^{2+}$ and Zn$^{2+}$ ions exchange to a lesser extent with sodium. If a pair of cations is exchanged with analcime, the cation-selection preference is determined by their size. Lower framework charge (high Si/Al) prefers the bigger cation and higher framework charge (low Si/Al) prefers the smaller cation. The exchange of the Li$^+$ atom with analcime is difficult due to Li$^+$ being very small and possessing high hydration energy. Barrer and Hinds [14] found that at high temperature, K$^+$, Ag$^+$, Ti$^4+$, NH$_4^+$ and Rb$^+$ cations exchange easily with Na-analcime, but determined that Sr$^{2+}$, Mg$^{2+}$, Co$^{2+}$ and Ni$^{2+}$ atoms exchange in very small amounts with Na-analcime [15]. The water content of analcime changes linearly with silica content. As the silica content increases, the number of cations decreases and correspondingly the number of water molecules also increases. Na$^+$ cations can be exchanged with alkali metal cations (Li$^+$, K$^+$), NH$_4^+$, Ag$^+$ and divalent cations (Ca$^{2+}$, Mg$^{2+}$) at high temperatures (225 °C). If the ions are too big, they occupy water locations. For example, when Na$^+$ in analcime is replaced with K$^+$ or Cs$^+$, the water molecule locations are occupied by alkali metal ions and H$_2$O molecules are removed from the structure. Therefore, the degree of hydration changes with the degree of ion exchange [16].

Dehydration–hydration behavior of natural zeolites is important for many application areas and can have many differences, with the zeolite type depending on the nature of interactions between H$_2$O molecules and, the Al-Si framework and extra-framework cations. While zeolites that have cavities with structurally large exchange areas like clinoptilolite hydrate easily, zeolites that have small cavities, like analcime, hydrate very slowly [17]. Very slow rehydration of analcime can be attributed to the kinetic diameter (2.6 Å) of cavities in its structure being smaller than the water molecules (2.65 Å) [18].

Tuff formations with very significant analcime content in Turkey are located in the areas of Bigadiç (Emirler), Emet (Köpenez), Kirka (Karaören), Urla, Bahçe Çayır, Küçük Kava, Mihalçık, Kalecik-Hasayaş-Şabanöz mü-Candır, Polatlı-Mülki-Öğlakçı-Ayaş, Keşan and Gördes [19–22]. But, there are very few works on the thermal and structural characterization of this natural zeolite.

The aim of this work is the determination of the thermal and structural properties of analcime zeolite samples and its cationic forms (HA, FeA, AgA and CuA) using XRD, XRF, TG–DTG–DTA and specific surface area measurements.

2. MATERIALS AND METHODS

2.1. Materials and Chemicals

The analcime sample used in this work was provided from the Çanakkale-Küçükkuay-Ayvacık area. The sample was ground and sieved to a size less than 45 µm, using ASTM standard sieves. Cationic forms of analcime were prepared with 1 M Fe(NO$_3$)$_3$.9H$_2$O (FeA), 1 M AgNO$_3$ (AgA), 1 M Cu(NO$_3$)$_2$.3H$_2$O (CuA) and 1 M HCl (HA) solutions in a shaker for 6 hours at 80 °C. After this process, solutions were filtered and the sample was washed seven times with deionized water at boiling point and then dried at room temperature. Before the experimental processes, all samples were dried for 16 hours in a 110 °C oven and then stored in a desiccator.

The organic chemicals like HCl, Fe(NO$_3$)$_3$.9H$_2$O, AgNO$_3$ and Cu(NO$_3$)$_2$.3H$_2$O were provided by Merck (Darmstadt, Germany), and all solutions were prepared using deionized water.
2.2. CHARACTERIZATION

The chemical composition of samples was determined using powdered samples fused with lithium tetraborate using X-ray fluorescence (XRF – Rigaku ZSX Primus device). X-ray diffraction (XRD) patterns were obtained with a D8 Advance Bruker AXS instrument, using CuKα radiation (λ = 1.54 Å) at intervals 40 kV and 20 mA, in the range 3 to 50 °C (2θ). TG–DTG–DTA curves were determined at a 10 °C min⁻¹ heating rate, for a sample of approximately 30 mg, using an alumina crucible with a Setsys Evutation Seteram device over a temperature interval of 30 °C to 1000 °C. Surface areas of the samples were measured at −196 °C with a vacuum at 30 °C for 7 hours. High purity nitrogen gas (99.99 %) was used in adsorption measurements.

The BET gas adsorption method is the most common standard method used for calculating the specific surface areas of solids, and it involves the use of the BET equation [23]:

$$\frac{1}{V[(P_0/P) - 1]} = \frac{1}{V_mC} + \frac{C - 1}{V_mC} \left( \frac{P}{P_0} \right)$$

Here $P_0$ is the saturated vapor pressure of the gas over the solid. $P/P_0$ is the relative pressure of the adsorbate molecule. $C$ is the BET constant. $V$ is the amount adsorbed gas at the relative pressure $P/P_0$ and $V_m$ is the monolayer capacity. The BET surface areas of natural and ion-exchanged zeolite samples were calculated at the relative pressure interval of 0.05 to 0.2, using adsorption isotherms.

3. RESULTS AND DISCUSSION

3.1. Chemical Analysis

The chemical compositions of the analcime samples in terms of the percentage of oxides are given in Table 1. It is determined that this sample is rich in Al₂O₃ and SiO₂. It is also seen that Na⁺, Ca²⁺ and Mg²⁺ cations decrease in analcime zeolites exchanged with heavy metal cations. As shown in Table 1, Ag⁺ cations enter the structure to a large extent as a result of treatment the analcime sample with a 1 M silver nitrate solution. A decrease in the iron, sodium, magnesium and calcium cations, along with aluminum, was detected as a consequence of treatment the analcime with 1 M HCl acid. Acid treatment increased the SiO₂/Al₂O₃ molar ratio from 5.63 to 6.85 by removing aluminum from the zeolite structure. The first purpose of the acid treatment is the removal of impurities and substances like Fe₂O₃ from the pores and cavities of zeolite [24].

<table>
<thead>
<tr>
<th>Chemical Analysis (%)</th>
<th>A</th>
<th>CuA</th>
<th>FeA</th>
<th>AgA</th>
<th>HA</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>71.992</td>
<td>71.599</td>
<td>73.125</td>
<td>71.086</td>
<td>77.148</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>2.409</td>
<td>2.405</td>
<td>2.767</td>
<td>2.365</td>
<td>0.761</td>
</tr>
<tr>
<td>MgO</td>
<td>0.799</td>
<td>0.749</td>
<td>0.593</td>
<td>0.731</td>
<td>0.255</td>
</tr>
<tr>
<td>CaO</td>
<td>0.891</td>
<td>0.507</td>
<td>0.465</td>
<td>0.494</td>
<td>0.619</td>
</tr>
<tr>
<td>K₂O</td>
<td>5.686</td>
<td>5.690</td>
<td>5.778</td>
<td>5.389</td>
<td>5.619</td>
</tr>
<tr>
<td>Na₂O</td>
<td>1.652</td>
<td>1.621</td>
<td>1.489</td>
<td>0.796</td>
<td>1.281</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.203</td>
<td>0.206</td>
<td>0.235</td>
<td>0.238</td>
<td>0.276</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.066</td>
<td>0.060</td>
<td>-</td>
<td>0.051</td>
<td>-</td>
</tr>
<tr>
<td>CuO</td>
<td>-</td>
<td>0.669</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ag₂O</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3.519</td>
<td>-</td>
</tr>
<tr>
<td>LOI</td>
<td>3.532</td>
<td>3.524</td>
<td>3.447</td>
<td>2.835</td>
<td>2.776</td>
</tr>
</tbody>
</table>
3.2. X-ray Analysis

XRD patterns were obtained for natural analcime and its ion-exchanged forms (Figures 1–3). As shown in Figure 1, it was determined that the natural zeolite sample includes analcime (A) as the main mineral, and, in addition to that, they also include quartz (Q), feldspar (F), montmorillonite (M) and amorphous substances. The characteristic diffraction peaks of analcime are determined to be at crystal spacings, $d = 5.60\,\text{Å}, 4.85\,\text{Å}, 3.43\,\text{Å}$ and $2.92\,\text{Å}$ in the XRD pattern of natural analcime sample [25, 26]. No other zeolite type is determined using XRD analysis. Quartz peaks were found at $d = 4.26\,\text{Å}, 3.34\,\text{Å}, 2.99\,\text{Å}, 2.46\,\text{Å}, 2.28\,\text{Å}$ and $2.12\,\text{Å}$. Feldspar peaks were seen at $d = 3.77\,\text{Å}, 3.46\,\text{Å}$ and $3.22\,\text{Å}$ [27].

From the XRD analysis examination of ion-exchanged analcime zeolites, some changes in the peak intensities are seen (Figures 2 and 3). These changes in the peak intensities can be attributed to changes in the chemical composition of the mineral due to the ion-exchange process (Table 1). While a considerable decrease in the intensity of the peak at $d = 5.60\,\text{Å}$ is seen in the XRD pattern of HA sample, it is seen that the intensities of the quartz peaks at $d = 4.26\,\text{Å}$ and $d = 3.34\,\text{Å}$ are largely increased (Figure 2). This was likely related to both dealumination and the partial destruction of the analcime structure. It is also seen that insoluble impurities like quartz and feldspar are left in the structure after the acid process [28]. It is seen however, that while the intensities of the quartz peaks at $d = 3.34\,\text{Å}$ of FeA and CuA samples increase by a considerable amount, the analcime peak at $d = 5.60\,\text{Å}$ from the AgA sample disappears completely and its quartz peak at $d = 3.34\,\text{Å}$ has decreased by a considerable amount (Figures 2 and 3). XRD analysis exhibited that considerable change was observed in the crystallinity of natural zeolite after Ag-exchange treatment. This can be related to cation migration and the elimination of water molecules from the structure channels due to heating. Natural zeolites have a good selectivity for silver ions [29].

![Figure 1. X-ray diffraction pattern of natural analcime sample. (A: Analcime; K: Quartz; F: Feldspar; M: Montmorillonite)](image-url)
Figure 2. X-ray diffraction pattern of HA and CuA samples

Figure 3. X-ray diffraction patterns of FeA and AgA samples
3.3. Thermal Analysis

Dehydration of analcime is a relatively simple single-step process, since the water molecules occupy at a single distinct location in the crystal structure [8]. Hydration–dehydration thermodynamics of analcime minerals have been examined by many researchers [30–39]. No phase transition has been observed during the dehydration of cubic structured analcime at high temperature [35].

The TG–DTG–DTA curves of analcime samples are shown in Figure 4. The decomposition temperatures of AgA and HA samples shifted toward lower temperatures compared with natural analcime. An endothermic peak is observed in the DTA curve of the HA sample at 77 °C. On the other hand, in the DTA curves of natural analcime A, CuA, FeA and AgA samples, two endothermic peaks are observed. The first endothermic peak is observed at temperatures of 70 °C (AgA) and 83 °C (CuA, FeA and A). A second endothermic peak is observed at temperatures of 466 °C (FeA), 472 °C (AgA), 480 °C (CuA) and 481 °C (A). The first endothermic peak, which is observed up to 200 °C in the DTA curves of the analcime samples was higher and corresponds to the elimination of physically adsorbed water from the structure. The second endothermic peak, which was observed in a broad temperature interval from 200 °C to 500 °C shows a loss of water that is more tightly bound to the structure. Dehydroxylation was observed through the greater loss of water at high temperatures.

![Figure 4. TG–DTG–DTA curves of natural and modified analcime samples](image-url)
The temperature intervals and mass loss values determined by TG–DTG analysis are given in Table 2. It was observed that for a series of heavy metal cation-exchanged forms the thermal stability and water retentivity of natural analcime increased in the order as Ag < Fe < Cu. It was determined that the total mass loss of CuA, A, FeA, HA and AgA analcime samples at 1000 °C after TG analysis was 3.46 %, 3.30 %, 3.23 %, 2.94 % and 2.73 %, respectively. It is seen that the total mass loss of natural and modified analcime samples are very close to the values of loss ignition (LOI) (Table 1) of the same samples. It was determined that greater loss of water occurs for the zeolites that are exchanged with divalent cations. In addition to that, it is seen that zeolite, which is exchanged with large, monovalent cations show less water loss (Table 2).

Table 2. (% Mass loss of analcime samples at different temperature intervals)

<table>
<thead>
<tr>
<th>Sample</th>
<th>30–200 °C</th>
<th>200–370 °C</th>
<th>370–750 °C</th>
<th>750–1000 °C</th>
<th>Tot. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.98</td>
<td>0.76</td>
<td>1.48</td>
<td>0.08</td>
<td>3.30</td>
</tr>
<tr>
<td>AgA</td>
<td>0.66</td>
<td>0.63</td>
<td>1.37</td>
<td>0.06</td>
<td>2.73</td>
</tr>
<tr>
<td>CuA</td>
<td>0.97</td>
<td>0.90</td>
<td>1.50</td>
<td>0.10</td>
<td>3.46</td>
</tr>
<tr>
<td>FeA</td>
<td>1.06</td>
<td>0.64</td>
<td>1.41</td>
<td>0.12</td>
<td>3.23</td>
</tr>
<tr>
<td>HA</td>
<td>1.03</td>
<td>0.52</td>
<td>1.23</td>
<td>0.16</td>
<td>2.94</td>
</tr>
</tbody>
</table>

3.4. Surface Area

Surface area data, correlation coefficients and BET constant C for natural and modified analcime samples are shown in Table 3. As seen in Figure 5, the adsorption isotherms of analcime samples are type II according to International Union of Pure and Applied Chemistry (IUPAC) classification [40]. It is seen that the specific surface areas of modified analcime samples are bigger than the surface areas of natural zeolite samples (Table 3). The BET surface areas decreased in the sequence HA > FeA > CuA > AgA > A. The specific surface area of the natural chabazite increased from 10.40 m²/g to 25.28 m²/g after treatment 1 M iron nitrate solution. The considerable increase in the BET surface area of FeA was ascribed to decationation and dealumination. It is also seen that the smallest surface area belongs to the AgA (14.60 m²/g) sample among the analcime samples that are exchanged with heavy metal cations. This decrease in the surface area can be attributed to the partial blockage of the channels in the structure by the Ag⁺ (1.26 Å) cations after the analcime was modified by treatment with silver nitrate solution [41].

It is seen that the analcime sample processed with a 1 M HCl acid solution had the biggest BET specific surface area with respect to the others. The surface area of the natural analcime sample has increased from 10.40 m²/g to 41.88 m²/g (Table 3). The acid treatment causes the Al³⁺ atoms to remove from the zeolite structure and zeolite cations to be exchanged with H⁺. As a result of the removal of aluminum from the structure, important changes occur in the adsorption properties of zeolite. Acid treatment also causes the amorphous substances inside the channels within the structure to remove and new pores to form. If the intensity of the acid process is too high, it may cause partial or total damage to the structure of the zeolite crystal [24, 42–44].

Table 3. Surface areas of natural analcime and its ionic forms

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>Multi Point BET Surface Area (m²/g)</th>
<th>BET Constant C</th>
<th>Correlation Coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>10.40</td>
<td>59.18</td>
<td>0.999</td>
</tr>
<tr>
<td>HA</td>
<td>41.88</td>
<td>248.60</td>
<td>0.999</td>
</tr>
<tr>
<td>FeA</td>
<td>25.28</td>
<td>174.70</td>
<td>0.999</td>
</tr>
<tr>
<td>CuA</td>
<td>17.66</td>
<td>84.60</td>
<td>0.999</td>
</tr>
<tr>
<td>AgA</td>
<td>14.60</td>
<td>48.34</td>
<td>0.999</td>
</tr>
</tbody>
</table>
4. CONCLUSION

Natural and ion-exchanged analcime samples were characterized using the XRF, XRD, TG–DTG–DTA and nitrogen adsorption methods. It was determined that the natural zeolite has analcime as the main mineral but also includes quartz, feldspar, montmorillonite and amorphous substances. XRD analysis showed that important changes occurred in the crystal structure of the Ag- and H-forms of analcime. While the analcime peak of AgA disappears completely, the quartz peak intensity decreased by a considerable amount. On the other hand, the analcime peak in the HA sample decreased significantly, while the quartz peak increased by a significant amount.

DTA curves of ion-exchanged analcime zeolites showed that these samples exhibited thermal stability up to high temperatures. However, it is found that the Ag- and H- forms of analcime show different thermal properties due to the change in their crystal structure and chemical composition. It was determined that the thermal decomposition temperatures of Ag-exchanged analcime shifted towards lower temperature values compared to natural analcime. This shows that the AgA sample had less thermal stability compared to natural zeolite. On the other hand, the amount of water molecules in the AgA sample decreased significantly due to the decrease in cations in the analcime structure after the Ag-exchange. Therefore, it was seen that AgA has the least amount of mass loss among the ion-exchanged samples. Also, the removal of cations and aluminum from the analcime structure, after treatment with the acid solution, caused a decrease in the hydrophilic nature of HA. This resulted in the H-form of analcime having both a lower endothermic peak temperature, and less mass loss, compared to natural analcime.

It was determined that the surface areas of ion-exchanged analcime zeolites are increased in comparison to natural analcime. It was found that while AgA (14.60 m²/g) has the smallest surface area, HA (41.88 m²/g) has the largest surface area of these samples. The decrease in the surface area of the AgA sample can be attributed to the blockage of analcime pores by the Ag⁺ cations during the modification process with the silver nitrate solution. On the other hand, the increase in the surface area of the HA sample can be attributed to the changes in the pore structure by the removal of amorphous substances and aluminum from the analcime structure when treatment with the acid solution.

Consequently, it was seen clearly that the structural, thermal and adsorption properties of analcime samples are affected by the ion-exchange process. The fact that natural zeolite are very cheap, abundant and easily obtained, makes these materials attractive in many application areas. Therefore, the thermal and structural characterization of natural and modified zeolites has great importance.
REFERENCES


