Thermal Transformation and Characterization of Synthetic Al-Substituted Maghemites (γ-Fe$_{2-x}$Al$_x$O$_3$)

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ABSTRACT: Burning is a common practice in tropical areas and related changes in mineralogy might affect the chemical and physical behavior of soils. Maghemite is a common iron oxide in soils formed from basic rocks in tropical regions. This mineral and hematite are the main pigments in these soils and exhibit high magnetization stemming from the precursor magnetite formed during the weathering process of primary minerals. The objective of the present study was to analyze changes in color, magnetic susceptibility values, Fourier transform infrared spectroscopy (FTIR) spectra, and available Fe and Al contents extracted with 1 mol L$^{-1}$ KCl during the process of thermal transformation of synthetic Al-maghemites into Al-hematites. Synthetic substituted maghemites with different degrees of Al-substitution (0.0, 1.0, 2.0, 2.9, 3.8, 5.6, 6.7, 10.0, 12.0, and 17.1 mol% Al) were subjected to a temperature of 500 ± 10 °C for 0, 5, 10, 16, 64, 128, 192, 360, 720, 2160, 3600, 5040 and 6480 min. The color of the samples was analyzed by a Munsell system in a colorimeter. Mass-specific magnetic susceptibility ($\chi_{LF}$) was measured at low-frequency. Available Fe and Al contents were estimated by a 1 mol L$^{-1}$ KCl solution. Fourier transform infrared spectroscopy (FTIR) spectra were obtained through use of a Bruker Vertex 70X FTIR spectrophotometer at a spectral resolution of 4 cm$^{-1}$. Contents of Fe and Al extracted by 1 mol L$^{-1}$ KCl in Al-hematites were not detected. All samples analyzed exhibited YR hue. Hue proportion decreased with increased heating time, and color changed from brown to red. The increase in isomorphic substitution (IS) led to increased hue values from maghemite to hematite, and the latter then became yellower. The $\chi_{LF}$ values decreased with an increase in heating time, indicating transformation from a ferrimagnetic phase (maghemite) to an antiferrimagnetic phase (hematite). With increasing IS, the maghemite $\chi_{LF}$ values decreased. Bands of the initial members (time 0) in the FTIR spectra were indexed as maghemites. The end members after completion of the heat treatment were identified as hematites. The IS of Fe by Al in maghemite influenced the thermal transformation to hematite, as well as the color and $\chi_{LF}$ of the minerals. The $\chi_{LF}$ proved to be very efficient in detecting maghemites remaining after thermal processing. Fe and/or Al were not ejected from the hematite crystalline structure after heat treatment.

Keywords: Fourier Transform Infrared Spectroscopy, color, Mass Specific Magnetic Susceptibility, KCl.
INTRODUCTION

Iron oxides constitute an important assemblage most of the minerals that compose Brazilian soils. Generally, tropical soils basic volcanic rocks contain a high concentration of magnetic and non-magnetic Fe oxides (Resende et al., 1988). According to Souza Júnior et al. (2010), soils with high magnetic susceptibility are correlated with good agricultural potential, and they occupy 5 % of Brazilian territory. The most important ferrimagnetic minerals in soils and sediments are magnetite (Fe₃O₄) and maghemite (γ-Fe₂O₃) (Mullins, 1977; Jong et al., 2000; Grimley et al., 2004).

Maghemite is a secondary Fe oxide, and it is easily identified in a clay fraction with a hand magnet (Schwertmann and Taylor, 1989). Particles larger than 10 nm are completely organized in a magnetic way at ambient temperatures, whereas smaller particles are said to be superparamagnetic (Coey and Khalafalla, 1972; Dearing, 1994).

Many authors have observed the effect of isomorphic substitution on magnetic susceptibility in synthetic (Batista et al., 2008; 2010; 2013) and natural minerals (Costa et al., 1999; Silva et al., 2010). In general, increasing isomorphic substitution (IS) (i.e. Al³⁺) decreases mass-specific magnetic susceptibility. This property has been used to quantify and qualify magnetic minerals in the soil.

Hematite is paramagnetic above 956 K (682.85 °C) (Cornell and Schwertmann, 1996). At ambient temperatures, it is weakly ferromagnetic, and at 260 K (-3.15 °C), it undergoes a phase transition to an antiferromagnetic state. The magnetic behavior of hematite depends on particle size, crystallinity, and isomorphic substitution (IS), mainly by Al, Ga, Cr, In, Mn, Sn, and Ti (Cornell and Schwertmann, 1996).

Another reliable indicator of the presence of Fe oxides is soil color (Resende, 1976; Schwertmann, 1993). In the Brazilian Soil Classification System (Santos et al., 2013), color is used as a differential property at the second categorical level. Color measures may be collected using different scales, but the Munsell scale is commonly used in Brazil.

Along with organic matter, Fe oxides are the most important color agents: red (α-Fe₂O₃), yellow (α-FeOOH), brown (γ-Fe₂O₃), and black (Fe₃O₄) (Costa and Bigham, 2009). Color may also vary according to the particle size and shape of minerals, chemical composition, and type and level of IS (Costa and Bigham, 2009).

Soils developed from mafic rocks in tropical regions tend to be more reddish (richer in hematite) (Resende et al., 2007). Maghemite is an Fe oxide common in soils developed from basic effusive magmatic rocks and tends to constitute up to 50 % of the Fe oxides present in soils developed from these rocks (Costa et al., 1999). Maghemite can transform into hematite by heating above 500 °C (Bigham et al., 2002).

Changes in the color of Fe oxide because of IS of Fe³⁺ by other metallic cations have been observed by some researchers (Scheinost et al., 1999; Batista et al., 2010). The color of Al-substituted maghemite (γ-Fe₂₋ₓAlₓO₃) observed by Batista et al. (2010) ranged from 4.8YR to 7.1YR. Torrent and Barrón (2003) observed that the color of synthetic and natural Al-substituted hematite ranged from 9.5P to 5.3YR. The presence of Mn in the structure of hematite leads to a darker color, close to black (Cornell and Schwertmann, 1996). In contrast, the presence of Al reduces the crystal size of hematite and goethite and increases their value in the Munsell system, and they become lighter in color (Barrón and Torrent, 1984; Kosmas et al., 1986). Red hematite heated to temperatures above 800 °C becomes purple, due to increasing particle size (Steinwehr, 1969).

Some authors found that thermal transformation of Fe oxides that were isomorphically substituted caused migration and/or ejection of some elements from the crystalline structure of the mineral (Sidhu et al., 1980; Landers et al., 2009). However, some elements remain in the mineral even after heating (Sidhu et al., 1980).
The isomorphic substitution and thermal reaction can cause changes in mineralogical and chemical attributes of maghemite and hematite. The objective of the present study was to evaluate the influence of IS of Fe by Al after the thermal transformation of Al-maghemites into Al-hematites by analyzing mass-specific magnetic susceptibility, color, available Fe and Al in 1 mol L⁻¹ KCl, and Fourier transform infrared spectroscopy (FTIR).

**MATERIALS AND METHODS**

**Maghemite synthesis**

Aluminous maghemites were prepared by the oxidation of Al-magnetites, synthesized by co-precipitation in alkaline KNO₃ solutions containing FeSO₄·7H₂O with stoichiometric amounts of Al₂(SO₄)₃·7H₂O, according to the procedure described by Schwertmann and Cornell (1991) and Batista et al. (2010). Magnetite synthesis was performed in a N₂ atmosphere over 2 h periods to assess the impact of equilibration time on Al substitution. Following the oxidation and co-precipitation reactions, a hand magnet easily and completely attracted all the magnetite formed. This material was washed several times with distilled water to remove excess salts. It was frozen using liquid nitrogen, and dried in a lyophilizer. The magnetite was then heated in a furnace at 250 °C for 4 h, where it was converted to yellowish red maghemite, which was also attracted by a magnet. In order to purify the maghemites, poorly crystalline materials were selectively removed by one 4 h treatment with acid (pH 3.0) ammonium-oxalate (2.0 mol L⁻¹) in the dark using a sample to solution ratio of 1:1000, according to the procedure described by McKeague and Day (1966).

Duplicates of the powdered, purified materials (~200 mg) were weighed into 50 mL polypropylene tubes; 3 mL of concentrated HCl was then added and the samples were allowed to rest for ~48 h until complete dissolution was achieved. Total Al and Fe contents were subsequently determined using a GBC 932AA atomic absorption spectrometer. Observed Al and Fe contents were calculated based on the amounts of Al and Fe obtained in total chemical analysis. The isomorphic substitutions (IS) observed in synthetic substituted maghemites were 0.0, 1.0, 2.0, 2.9, 3.8, 5.6, 6.7, 10.0, 12.0, and 17.1 mol% Al.

**X rays diffraction (XRD)**

The powdered material was analyzed on a Shimadzu XRD-6000 diffractometer using CuKα radiation (35 kv, 20 mA) with a Ni filter in a step-scanning mode (0.01 °2θ s⁻¹). An internal standard was not added. Identification of the mineral species in the samples was made from X rays diffractions (XRDs) according to the position and intensity of the diffraction planes (Whittig and Allardice, 1986; Schwertmann and Taylor, 1989).

**Maghemite-to-hematite transformation**

Synthetic maghemites with different degrees of IS were placed on mullite rectangular boats in a muffle furnace pre-heated to a temperature of 500 ± 10 °C. This temperature was chosen based on differential thermal analysis (DTA) and on literature results (Sidhu, 1988). The mullite refractories were placed on a steel tray and then heated. The samples remained in the furnace for different times: 0, 5, 10, 16, 64, 128, 192, 360, 720, 2160, 3600, 5040, and 6480 min. After that, they were withdrawn and immediately cooled (the tray was placed on ice) to interrupt further reactions (Sidhu, 1988) and then kept in plastic containers.

**Available Fe and Al in 1 mol L⁻¹ KCl**

Approximately 0.40 g of the heated material was placed in 15 mL Falcon tubes. Then, 10 mL of 1 mol L⁻¹ KCl was added (Claessen, 1997). The tubes were shaken for 10 min at 180 rpm in a horizontal shaker and then centrifuged at 2,200 rpm for 5 min. Extracts were stored in polypropylene containers and Fe and Al contents were determined by Atomic
Absorption Spectroscopy (AAS) in a GBC 932 AA spectrometer. Iron was determined in acetylene flame and Al in acetylene-nitrous oxide flame.

**Mass-specific magnetic susceptibility (χ_{LF})**

Mass-specific magnetic susceptibility was determined in all samples using a Bartington MS2 magnetic susceptibility system coupled to an MS2B sensor (Dearing, 1994). This dual frequency meter exposed the sample to a weak alternating magnetic field of ~80 A m⁻¹. The MS2B sensor has both low- (0.47 kHz) and high-frequency settings (4.7 kHz). Powdered samples (10 cm³) were weighed into 20 cm³ plastic vials. Volumetric magnetic susceptibility (k) was measured only at low-frequency settings, and low-frequency mass susceptibility (χ_{LF}) was calculated as follows (Dearing, 1994): χ_{LF} = (10 k/ m) m⁻¹, where m is the mass (g) and k is the volumetric magnetic susceptibility measured by the equipment.

**Fourier transform infrared spectroscopy (FTIR)**

The infrared (IR) spectra were recorded using a Bruker Vertex 70X FTIR spectrophotometer at a spectral resolution of 4 cm⁻¹. Each spectrum was obtained after acquiring 128 scans. Approximately 2 mg of each sample, plus 200 mg of KBr, were weighed and ground with an agate mortar and pestle until a homogeneous mixture was obtained, which was then pressed into disc-shaped pellets. The FTIR spectra from the pellets were recorded over the range of 400-4,000 cm⁻¹ and then analyzed using Origin® 8.0 software. FTIR bands were identified as suggested by Waldron (1955), White and De Angelis (1967), (Wilson et al., 1981), Cornell and Schwertmann (1996), and Socrates (2001).

**Color determination**

Color was determined from powdered samples with Konica Minolta CR-400 equipment set up for the Munsell scale (hue, value, and chroma). The amount available of each sample was placed on gloss paper positioned on a workbench lined with black poster paper and then measured.

**RESULTS AND DISCUSSION**

**X rays diffraction (XRD)**

The XRD reflections showed only the presence of the two phases in all materials analyzed: maghemite, Al-maghemite, hematite and/or Al-hematite (data not shown). For example, figure 1 shows XRD spectra of Al-maghemitess with 3.8 mol% isomorphic substitution (IS) without heating and Al-hematite with IS of 3.8 mol% Al that was heated for 6480 min. Those samples were arbitrarily chosen.

**Available Fe and Al in 1 mol L⁻¹ KCl**

Iron and Al content extracted by 1 mol L⁻¹ KCl solution was not detected in Al-hematites in any degree of IS. Element migration in mineral structures and element ejection have been observed by different authors. Landers et al. (2009) observed Ni migration in natural goethite samples because of thermal treatment. In contrast, Sidhu et al. (1980) emphasized that during thermal transformation of maghemite into hematite, element ejection mainly occurs with divalent ions that exhibit a lower valence and larger ionic radii than Fe³⁺, a situation not observed in this experiment.

**Color**

Differences in particle size, morphology, and chemical composition may cause variations in mineral structure and behavior (Cornell and Schwertmann, 1996). The maghemite hue values (Figure 2) tend to increase with increasing isomorphic substitution (IS) but without any significant correlation (p>0.05). Hematite hue values also increased with increasing IS,
and this relationship was best represented by an exponential regression model ($R^2 = 0.88$). Therefore, hematite showed greater changes in color at lower degrees of IS, becoming yellower. Color stabilization in hematite samples was achieved at hue 2.76YR.

**Figure 1.** X rays diffraction of Al-maghemite (not heated) and Al-hematite (heated for 3,680 min) with 3.8 mol% Al.

**Figure 2.** Relationship between YR hue values of synthetic Al-substituted maghemites (a) and hematites (b) samples. a.u.: arbitrary unit.
The Fe-to-Al isomorphic substitution effect on mineral color is difficult to detect because Al substitution reduces particle size and changes chemical composition, which might also change mineral color (Batista et al., 2010). Maghemite and hematite hues ranged from 5.1YR to 7.1YR, and from 0.3YR to 2.9YR, respectively. Hue values observed for maghemite were higher than the values suggested by Bigham et al. (2002) (2.5YR to 5YR). The hue values of hematite were also different from those proposed by Cornell and Schwertmann (1996) (5R to 2.5YR). Bhuiyan et al. (2007) verified that hematite substituted by Nd³⁺ exhibited a brighter yellowish-red color compared to the pure phase.

The behavior of chroma and value data did not show significant correlation (p>0.05) with the different degrees of IS. For Barrón and Torrent (1984) and Kosmas et al. (1986), structural Al seems not to significantly influence the hue and chroma of synthetic Al-hematites, although the crystals became lighter (increasing value).

Increased heating time led to a reduction in the proportion of the YR hue for all the degrees of substitution studied (Figure 3); i.e., the samples became redder. The Munsell hue values were higher for the highest Al contents. As previously observed, change in color with heating time is due to the maghemite-to-hematite transformation; i.e., the color changes from brown (maghemite) to red (hematite). The samples that were a mix of maghemite and hematite showed intermediate values between maghemite with no substitution and maghemite with 17.1 mol% Al. Al-free maghemites showed an abrupt change in color compared to Al-substituted maghemites. Thus, a higher IS led to a slower transformation rate. In other words, the higher the content of chemical elements, the higher the energy necessary to eject or redistribute them in the mineral crystalline structure. When samples with different IS are exposed to the same temperature, the sample that has higher IS will transform slower (Sidhu et al., 1980).

A 90 % color change in Al-free and in 17.1 mol% Al maghemites was observed after heating for 10 and 768 min, respectively. This increasing resistance to color change with increasing IS and the differences in the initial (maghemite) and final (hematite) colors follow the same reasoning used to explain phase alteration. It is evident that

![Figure 3. Relationship between YR Hue values of synthetic Al-maghemites transformed into hematites throughout heating period at 500 °C. a.u.: arbitrary unit.](image-url)
higher temperatures (Sidhu, 1988) or longer heating periods are necessary for the solid state maghemite-to-hematite transformation when the minerals have undergone isomorphic substitution.

Burning, which is still widely used for land clearing in tropical regions, may also change soil color. The maximum temperature and duration of the exposure are important indicators of fire severity (Wells et al., 1979). Low severity fires (short exposure to <250 °C) have shown to temporarily affect biological and chemical properties of soil. More severe burning may alter fundamental properties as texture, mineralogy, and cation exchange capacity (Sertsu and Sanchez, 1978; Ulery and Graham, 1993; Ulery et al., 1996; Ketterings et al., 2000).

Some properties of Indonesian soils were studied by Ketterings and Bigham (2000) and they observed that surface burning tends to reduce the value and chroma, observing that above 600 °C, a pronounced reddening may occur that would indicate the thermal conversion of goethite (yellow) to maghemite (reddish brown) and to hematite (red). Lugassi et al. (2009) observed this reddening at 300 °C in Israeli soils (loess). Therefore, the temperature of change in soil colors is dependent on the mineralogy.

**Mass specific magnetic susceptibility**

For the Al-substituted maghemites, $\chi_{LF}$ values ranged from 44,000 (0.0 mol% Al) to 52,968 (2.9 mol% Al) $\times 10^{-8}$ m$^3$ kg$^{-1}$, and for the Al-substituted hematites, $\chi_{LF}$ values ranged from 166 (2.0 mol% Al) to 1,270 (17.1 mol% Al) $\times 10^{-8}$ m$^3$ kg$^{-1}$. Dearing (1994) established values ranging from 41,000 to 44,000 $\times 10^{-8}$ m$^3$ kg$^{-1}$ for maghemite (ferrimagnetic) and from 27 to 169 $\times 10^{-8}$ m$^3$ kg$^{-1}$ for hematite (antiferrimagnetic) (Figure 4); i.e., our values are higher than those registered by Dearing (1994) and similar to those obtained by Batista et al. (2010; 2013) for synthetic maghemites.

![Figure 4. Mass specific magnetic susceptibility ($\chi_{LF}$) values to synthetic Al-substituted maghemites (a) and hematites (b) samples. (○) Outlier.](image)

$\hat{y} = 172.90 + 58.75x$

$R^2 = 0.91$ p<0.01

$\hat{y} = 51708.39 + 286.29x$

$R^2 = 0.74$ p<0.01
With increasing IS, $\chi_{LF}$ values decreased (Figure 4b) since $\text{Al}^{3+}$ is a paramagnetic ion (Cullity, 1972; Wolska, 1990) and has a similar dilution effect on magnetization, as observed previously for Zn (Batista et al., 2008). The dilution effect might be associated with i) partial relocation of the vacancies from octahedral to tetrahedral sites in the mineral structure (Wolska, 1990; Takei and Chiba, 1966) and ii) breaking down the order of both cation and vacancy distributions in the octahedral sub-lattice (Gillot et al., 1982). However, this decrease is not linear (Gillot et al., 1982).

The behavior of hematites was opposite to maghemites (Figure 4a). An increase in Al substitution causes an increase in $\chi_{LF}$ values. However, this result is contrary to the results of Murad and Schwertmann (1986). Increasing $\chi_{LF}$ values may be due to remnant maghemite components that maintain magnetism, although XRD data only points to the presence of hematite. Increasing Al IS changes the temperature of the solid phase thermal transformation from maghemite to hematite (Sidhu, 1988); therefore even long exposure of maghemites to 500 °C temperature was not able to completely transform maghemite to hematite, and this effect was more pronounced as IS increased.

If $\chi_{LF}$ values were used to calculate maghemite content in the samples (Figure 4a), there would be less than 2 %, with this value hardly being detectable by the XRD technique using a standard configuration (0.06 °2θ, 0.6 s).

With increasing heating time, $\chi_{LF}$ values of synthetic Al-maghemites decreased exponentially, indicating a solid state thermal transformation of a ferrimagnetic phase (maghemite) into an antiferrimagnetic one (hematite) (Figure 5). Equations concerning the adjustments are presented in table 1. To have a 90 % reduction in $\chi_{LF}$ values, it would be necessary to heat for 23 and 1774 min for Al-free and 17.1 mol% Al, respectively.

Maghemites with a low degree of crystallinity and with IS (especially for Al, Ga, Cr, Mn, Sn, and Ti) exhibit modified magnetic behavior (Cornell and Schwertmann, 1996). These two factors are interdependent, and it is hard to separate the substitution effect from the crystallite size effect on $\chi_{LF}$. Hematite synthesis, for example, influences both factors. Therefore, low crystallinity
and Al substitution allow the coexistence of ferrimagnetic and antiferrimagnetic phases above specific temperatures in an apparent homogeneous sample (Murad and Johnston, 1987).

**Fourier transform infrared spectroscopy (FTIR) spectra**

Bands of the initial members (time 0) of the FTIR spectra in figure 6 have been indexed as maghemites by the interpretation proposed by Waldron (1955) and White and De Angelis (1967). The end members after heat treatment (360 and 3600 min times, figures 6a and 6b).

<table>
<thead>
<tr>
<th>mol% Al</th>
<th>Equation</th>
<th>$R^2$</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>$\hat{y} = 801.9 + 47088.8\exp(-(x-(0.31))/3.6)$</td>
<td>0.99</td>
<td>0.01</td>
</tr>
<tr>
<td>1.0</td>
<td>$\hat{y} = -473.5 + 54684.0\exp(-(x-(0.25))/5.8)$</td>
<td>0.91</td>
<td>0.05</td>
</tr>
<tr>
<td>2.0</td>
<td>$\hat{y} = 211.5 + 52166.3\exp(-(x-(0.3))/6.65)$</td>
<td>0.91</td>
<td>0.05</td>
</tr>
<tr>
<td>2.9</td>
<td>$\hat{y} = 1548.8 + 53832.6\exp(-x/8.5)$</td>
<td>0.90</td>
<td>0.01</td>
</tr>
<tr>
<td>3.8</td>
<td>$\hat{y} = 907.8 + 53145.6\exp(-x/9.06)$</td>
<td>0.90</td>
<td>0.01</td>
</tr>
<tr>
<td>5.6</td>
<td>$\hat{y} = 927.4 + 30419.9\exp(-x/(4.8))/155.2 + 51632.5\exp(-x/(4.8))/4.8$</td>
<td>0.99</td>
<td>0.01</td>
</tr>
<tr>
<td>6.7</td>
<td>$\hat{y} = 634.9 + 45447.2\exp(-x/(4.2))/5.6 + 28889.1\exp(-x/(4.2))/199.5$</td>
<td>0.99</td>
<td>0.01</td>
</tr>
<tr>
<td>10.0</td>
<td>$\hat{y} = 516.4 + 24573.1\exp(-x/(1.6\times10^{-7}))/389.7 + 24573.1\exp(-x/(1.6\times10^{-7}))/389.7$</td>
<td>0.99</td>
<td>0.01</td>
</tr>
<tr>
<td>12.0</td>
<td>$\hat{y} = 578.9 + 24002.1\exp(-x/(2.9))/311.2 + 21002.1\exp(-x/(2.9))/311.3$</td>
<td>0.98</td>
<td>0.01</td>
</tr>
<tr>
<td>17.1</td>
<td>$\hat{y} = 2459.9 + 23498.8\exp(-x/(25.4))/342.5 + 23498.8\exp(-x/(25.5))/342.5$</td>
<td>0.98</td>
<td>0.01</td>
</tr>
</tbody>
</table>

**Figure 6.** FTIR spectra of synthetic Al-substituted maghemite with 0.0 (a) and 12.0 (b) mol% Al in the range 370 - 800 cm$^{-1}$. a.u. = arbitrary unit.
6b, respectively) were identified as hematite (Wilson et al., 1981). In the FTIR spectra for the intermediate heating times, a mixture of maghemite and hematite characteristic bands was observed. The samples with shorter and longer heating times were similar to the spectra of hematite and maghemite, respectively.

Fourier transform infrared spectroscopy bands that shifted to higher frequency indicate an increase in the bond energies within the maghemite structure. Therefore, if the observed shifts are not artifacts of the method, then Al substitution introduces a significant strain into the octahedral and tetrahedral sites associated with the Fe-O vibrations, as reported by Batista et al. (2010) for Zn-substituted maghemites.

With increasing IS, the maghemite bands shifted to higher wavelengths, except the 558.0 cm\(^{-1}\) band, which shifted in the opposite way. Another trend was observed for the 480.3 and 395.0 cm\(^{-1}\) bands, which shifted to shorter wavelengths initially (5.6 mol% Al) and then to longer wavelengths (12.0 mol% Al). The 583.4 and 421.8 cm\(^{-1}\) bands disappeared with increasing IS (Figure 7a). The FTIR bands of the hematite samples shifted to shorter wavelengths initially (5.6 mol% Al) and then to longer wavelengths (12.0 mol% Al), except the 378.0 cm\(^{-1}\) band, which shifted to shorter wavelengths with increasing IS (Figure 7b).

The strongest bands in hematite occur due to light scattering by the crystallites (Fysh and Fredericks, 1983). They identified two bands (550 and 470 cm\(^{-1}\)) that show some change in position with increasing Al content. Accordingly, in this study, the 470 cm\(^{-1}\) band changed its position with IS (Figure 7b). The IR spectrum of hematite depends on

![Figure 7. FTIR spectra of synthetic Al-substituted maghemite (a) and Al-hematite (b) in the range 370 - 800 cm\(^{-1}\). a.u.: arbitrary unit.](image-url)
particle shape (Serna et al., 1982). According to Barrón et al. (1984), Al substitution produces shifts in the frequencies by as much as 10 to 15 cm$^{-1}$.

CONCLUSIONS

The Fe and Al contents analyzed by 1 mol L$^{-1}$ KCl solution did not suggest Al migration to hematite borders or ejection from the mineral structure.

The values of the Munsell YR hue decreased with increasing heating time, changing from brown (5.1-7.1YR) to red (0.3-2.9YR) colors.

Increasing IS led to increasing hematite and maghemite hue values; thus, the former became yellower.

The $\chi_{LF}$ values decreased with increasing heating time, indicating the transformation of a ferrimagnetic phase (maghemite) into an antiferromagnetic one (hematite). With increasing IS, maghemite and hematite $\chi_{LF}$ values decreased and increased, respectively. In maghemites, the $\chi_{LF}$ values ranged from 44,000 (0.0 mol% Al) to 52,968 × 10$^{-8}$ m$^3$ kg$^{-1}$ (2.9 mol% Al), whereas for hematites, these values ranged from 166 (2.0 mol% Al) to 1,270 (17.1 mol% Al) × 10$^{-8}$ m$^3$ kg$^{-1}$.

Bands of the initial members (time 0) in the FTIR spectra have been indexed as maghemites. The end members after completion of the heat treatment were identified as hematite. In the FTIR spectra for the intermediate heating times, a mixture of maghemite and hematite characteristic bands was observed.

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