INTRODUCTION

Water contamination with persistent dyes discharged from the textile, pharmaceutical, and food industries is a stringent issue for environmental protection since it affects the water transparency and gas solubility in water hindering as well the growth of aquatic species and the photosynthesis process. Acute or chronic effects on the health of water living organisms were also noticed depending on the dye pollutant concentration and the exposure time. Due to the organic nature of dye pollutants, dye contaminated wastewaters have higher values of the chemical oxygen demand (COD) and the total organic carbon content (TOC). Usually, these dyes cannot be degraded or biodegraded in classical wastewater treatment [1]. Moreover, the partial degradation products of the organic dyes can be highly carcinogenic or mutagenic compounds, which can have even more negative effects on the organisms than the dye itself. Hence, advanced oxidation processes (AOP) are required for their mineralization [2, 3]. Lately, among the AOP that have aroused, there is a growing interest for those based on sono-catalysis which are considered cheaper than the photo-catalytic ones [4-6]. Disodium (2E)-3-oxo-2-(3-oxo-5-sulfonato-2,3-dihydroindole-1H-indol-2-ylidene)-2,3-dihydro-1H-indole-5-sulfonate also called indigo carmine (IC) is an indigoid water-soluble dye (Fig. 1) authorized as synthetic food color for a wide range of food categories with a maximum permitted levels between 50 and 500 mg/kg food [7]. It may become toxic leading to atrioventricular block and polymorphous supraventricular
bigeminy as well as potentially carcinogenic when exceeding doses of 500 mg/kg body weight/day [7-9].

**Fig. 1.** The chemical structure of indigo carmine (IC)

Indigo carmine mineralization by photocatalytic AOP using TiO\(_2\) as the photocatalyst was found to be difficult since there was no noticeable loss of the TOC value and release of inorganic ions even if the water coloration disappeared [10]. Recently, it has been proved that MnO\(_2\) could efficiently remove indigo carmine from polluted water by a combined process using both photo and ultrasonic activation [11]. It has been considered that the catalytic activity of MnO\(_2\) was due to its high affinity for O\(_2\) and its efficient adsorption of the dye. The degradation of indigo carmine with this catalyst was more efficient with peroxydisulfate (S\(_2\)O\(_8\)) as an oxidation agent rather than with hydrogen peroxide which was found to inhibit the process. However, the development of AOP using greener oxidants such as hydrogen peroxide would be of higher interest. Another important challenge would be the enhancement of the solid affinity for the organic substrate. In 2019, it was reported that mixed oxides obtained by thermal decomposition of Zn\(_x\)Al layered double hydroxides (x=2-4) exhibit good adsorption capacities for both indigo carmine and methylene blue dyes [12]. Layered double hydroxides (LDH) also called hydrotalcite (HT)-like compounds are anionic clays having the general formula [M\(_{II}\)\(_{1-x}\)M\(_{III}\)\(_x\)(OH)\(_2\)]\(^{2+x}\)[A\(^{n-}\)]\(_{3x/2}\)•zH\(_2\)O, where M\(_{II}\) is a bivalent metal cation, M\(_{III}\) is a trivalent metal cation, A\(^{n-}\) is a charge compensation anion which may be either inorganic or organic, x has a value between 0.2 and 0.4. LDH solids are formed of positively charged brucite-type layers containing the cations hexacoordinated with hydroxyl groups. Between two brucite-type layers, there is a space called the interlayer region containing the compensation anions and water molecules. These solids are renowned by their anion exchangability and by their so-called “memory effect” which consist in the recovery of the layered structure when a mixed oxide obtained by the thermal decomposition of an LDH precursor (at temperatures lower than 650°C) is immersed in an aqueous solution containing different anions [13]. Hence, the affinity of Zn\(_x\)Al mixed oxides for the anionic dye indigo carmine was considered to be a consequence of the memory effect exhibited by their LDH precursors. Ten years ago, graphene oxide (GO) is a good catalyst for facilitating oxidation and hydration reactions [14]. Since then, the researches on the catalytic activity of GO were continuously extended and quite recently the catalytic activity of GO-MnO\(_2\) nanocomposites for azo dyes (reactive black 5) degradation with hydrogen peroxide was discovered and explained by the synergetic effect of the GO which favors the adsorption of the organic substrate and the MnO\(_2\) which promotes the oxidation [15]. However, the limitation of this process is the low pH value (e.g. pH=3) necessary to reach high conversions. Based on this state of the art, our attention was drawn to the obtaining of new composites by combining two different 2D materials, GO and a Cerium modified LDH which could act as catalysts for the oxidative removal of indigo carmine using hydrogen peroxide as oxidant. Herein, we present the results of our researches on the synthesis, characterization, and catalytic activity of Mg\(_{2}\)Al\(_{0.75}\)Ce\(_{0.25}\) LDH-GO composites for the oxidation of indigo carmine with H\(_2\)O\(_2\), both under ultrasonic irradiation and under conventional stirring.
MATERIALS AND METHODS

The chemicals necessary for the synthesis of the LDH phase, e.g. magnesium nitrate hexahydrate Mg(NO$_3$)$_2$ $\times$ 6H$_2$O, aluminum nitrate nonahydrate Al(NO$_3$)$_3$ $\times$ 9H$_2$O, cerium nitrate hexahydrate Ce(NO$_3$)$_3$ $\times$ 6H$_2$O, anhydrous sodium carbonate Na$_2$CO$_3$, and sodium hydroxide NaOH (pearls) were all of the chemical purity grades and were purchased from Merck.

For the preparation of the GO phase, graphite powder 325 mesh from Aldrich, sodium nitrate NaNO$_3$ and potassium permanganate KMnO$_4$ (chemical purity from Merck), H$_2$SO$_4$ (98%) and hydrochloric acid HCl 37% (from Merck) and hydrogen peroxide H$_2$O$_2$ 30% (from ChimReactiv) were utilized.

Indigo carmine (IC) from Fluka AG was used for the preparation of the dye contaminated water.

All the aqueous solutions were obtained using distilled water with a conductivity of 2.5-5 $\mu$S/cm.

Three Mg$_x$Al$_{0.75}$Ce$_{0.25}$ LDH-GO composites bearing different concentrations of GO in the range of 5-15 wt.% abbreviated as HT3Ce-xGO where x stands for the concentration of GO ($x=5,10,15\%$ wt. GO) have been prepared by the co-precipitation of the LDH phase in the presence of GO at constant pH 10. First, the GO was prepared using the well-known Hummers method [16] by contacting 10 g of graphite powder with 5 g NaNO$_3$, 230 mL H$_2$SO$_4$ (98%), and 30 g KMnO$_4$ at 5-30 $^\circ$C for 1 h, followed by the addition of 710 mL distilled water and heating of the resulting mixture at 98-110$^\circ$C for 15 minutes and then adding 50 mL of H$_2$O$_2$ (30%) to reduce the remaining Mn$^{7+}$, Mn$^{4+}$ species to Mn$^{2+}$ (as water-soluble manganese sulfate), thus enabling the formation of GO yellowish flakes. The solid phase is then separated by centrifugation, washed first with dilute HCl (obtained by mixing equal volumes of distilled water and HCl 37%) and then with hot distilled water until the conductivity of the liquid in the centrifuge vials becomes lower than 100 $\mu$S/cm. After reaching this point, all the content of the centrifuge vials was collected and diluted with distilled water in a 5 L volumetric flask. The concentration of GO in the suspension was determined using the gravimetric method, by weighing in Petri dishes 3 liquid samples of 100 mL before and after the evaporation of water under vacuum at 60$^\circ$C for 24 h. The amount of solid recovered from the 3 samples was 0.4001 g, 0.4005 g, and 0.3998 g which gave an average value of the GO concentration in the suspension of 4 g/L.

For the preparation of the Ce-modified LDH compound by co-precipitation at constant pH 10 under low supersaturation conditions, the method described for the obtaining of other rare-earth modified LDH [17] was adapted. An aqueous solution A having a concentration of 1.5 M of cations (Mg$^{2+}$, Al$^{3+}$, and Ce$^{3+}$ while respecting the molar ratio Mg$^{2+}$/Al$^{3+}$/Ce$^{3+}$ of 3/0.75/0.25) and a solution B containing NaOH and Na$_2$CO$_3$ (1 M concentration of Na$^+$ ions, with a molar ratio NaOH/Na$_2$CO$_3$ of 2.5/1) were prepared by dissolving the corresponding amounts of chemicals in distilled water.

TIM854 Titration Manager from Radiometer analytical specialized in maintaining a constant pH value was utilized for the precipitation. The precipitation reactor filled with 100 mL distilled water was placed on the stirring plate of TIM854, the pH electrode was immersed in the reactor and the pH of the water was adjusted to 10 using B solution which was placed in the automatic burette of TIM854, then it was started the concomitant dropwise addition of solution A and solution B to achieve the precipitation at room temperature under vigorous stirring. Using this equipment the flow rate of solution B was automatically adjusted to maintain the constant pH 10. After consuming all solution A, the precipitation reactor was transferred on a heating stirring plate, a vertical condenser was mounted on its mouth and the gel was aged at 75$^\circ$C for 18 h. Afterward, the gel was filtered and washed with distilled water until the conductivity of the washing water decreased below 100 $\mu$S/cm, then the solid cake was dried in an oven at 90$^\circ$C for 24 h yielding Mg$_3$Al$_{0.75}$Ce$_{0.25}$ LDH.

For the preparation of Mg$_3$Al$_{0.75}$Ce$_{0.25}$ LDH-GO composites, the amounts of metal nitrates necessary to obtain 9.5, 9, and 8.5 grams of LDH phase respectively, were dissolved in an aqueous suspension of GO containing 0.5, 1, and 1.5 grams of GO, and distilled water was added to reach the 1.5 M concentration of cations (Mg$^{2+}$, Al$^{3+}$, and Ce$^{3+}$) as in the above-
described solution A utilized for the obtaining of Ce-modified LDH. Solution B and all the other stages of the synthesis were identical to the above-described procedure. The characterization of the solids included X-Ray diffraction of powders (XRD), diffuse reflectance infrared Fourier-transform spectroscopy (DRIFTS), Raman spectroscopy, XRD patterns in the 2θ range 5-80° were obtained with XRD 6000 Shimadzu, equipped with a lamp of Cu (λCuKα = 1.5418 Å, 40 kV, 40 mA) in continuous scanning mode with a step of 0.02°/min and an acquisition time per step of 4 seconds. The dimensions of the crystallites were calculated using Scherrer’s formula:

\[ D_{hkl}= \frac{K\lambda}{B_{hkl}\cos\theta_{hkl}} \]

where K is a geometric factor (e.g. 0.9), λ is the wavelength of the incident X-ray, \( B_{hkl} \) is the width at half-intensity of the \( hkl \) reflection, and \( \theta_{hkl} \) is the Bragg angle of the same reflection. DRIFTS spectra of the freshly prepared and spent solid samples were recorded with a Jasco FT/IR 4700 spectrometer having a PIKE accessory for diffuse reflectance. The spectra were collected in the spectral range 4000-400 cm\(^{-1}\), with a scanning speed of 128 scans/min and a resolution of 4 cm\(^{-1}\) using KBr as background. A high-resolution confocal Raman microscope (Renishaw system, Renishaw UK Ltd) equipped with two laser lines (514 nm and 785 nm, respectively) and a Leica DM2500 microscope was utilized for recording the Raman spectra of the solid samples, in extended mode using the 514 nm laser line, monitoring the shifts in the Raman band position narrower than 0.5 cm\(^{-1}\), and measuring the Raman bands in the range 100 - 3100 cm\(^{-1}\). SEM analysis of the samples was performed on Hitachi Ultrahigh Resolution Scanning Electron Microscope SU8010 with a semi-in-lens type objective lens and cold FE-gun with small energy spread.

RESULTS AND DISCUSSION
The diffraction pattern of Mg\(_3\)Al\(_{0.75}\)Ce\(_{0.25}\) LDH in Fig. 2 shows besides the typical reflections (003, 006, 009, 012, 015, 018, 110, and 113) of a lamellar hydrotalcite-like structure (Mg\(_3\)Al-ICDD card no 00-054-1029), those characteristic to CeO\(_2\) cubic phase (111, 200 and 220) with fluorite structure (ICDD card no 00-034-0394) and low-intensity lines of cerium hydroxycarbonate (marked with * on Fig. 2) (Ce(CO\(_3\))\(_2\)O\(_2\)H\(_2\)O -ICDD card 00-044-0617). It seems that Ce\(^{3+}\) fails to enter in the brucite-type layer due to its larger ionic radius (1.01 Å) compared to those of Mg\(^{2+}\) (0.72 Å) and Al\(^{3+}\) (0.535 Å). Therefore, the LDH-modification with Ce leads mainly to an alteration of the structure explaining the ill crystallized HT-phase observed in the XRD patterns of the dried Mg\(_3\)Al\(_{0.75}\)Ce\(_{0.25}\) LDH. The XRD pattern of GO (ICDD card 00-065-1528) shows the characteristic (001) diffraction line for layered hexagonal crystals which appears in the same region as the most intense line (003) of the LDH. The XRD patterns of all the composite materials HT3Ce-xGO had more intense and better-defined diffraction lines characteristic to the lamellar hydrotalcite-like

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structure than the neat $\text{Mg}_3\text{Al}_{0.75}\text{Ce}_{0.25}$ LDH or the neat GO. Besides, only the specific lines of $\text{CeO}_2$ are noticed while those corresponding to the cerium hydroxycarbonate are missing. The formation of the $\text{CeO}_2$ phase was favored by the increase of GO content in HT3Ce-xGO composites. These facts suggest that the precipitation of the LDH in the presence of GO suspension promotes the crystallization of the LDH phase along with the embedding of fine $\text{CeO}_2$ particles having Scherrer dimensions around 3 nm.

![Fig. 2. XRD patterns of the solid samples](image)

![Fig. 3. SEM microphotographs of the composite structures HT3Ce-xGO (15.0kV, x 4.5k, 10\(\mu\)m)](image)

The composite structures, analyzed by SEM in powder form, showed for all the samples well-defined grains, with different sizes and surface morphology when closely inspected. As emphasized by XRD structural analysis, the grains composition contains a well-defined LDH structure decorated with a distinct $\text{CeO}_2$ phase. In this context, Ce ions involvement should be noticed, since they could play a key role with the GO layers during the LDH synthesis. GO layers can offer plenty of sites, left non-oxidized during GO synthesis. Therefore the GO interaction during LDH synthesis should not be ignored, from both physical (nucleating) and chemical reason (interaction with Ce ions). Moreover, it was already reported that the ceria phase has an affinity for attachment onto GO [18]. The grain surface, revealed the transition from a relatively smooth surface towards a rougher polymorphic surface as the GO concentration increased. At high GO content, better accessibility towards the smaller elementary particles was observed. The elementary particles consist either in layered polymorphic particles with edges or in „ovoidal” composite particles (Fig. 3). The presence of „ovoidal” particles on the surface of the grains (marked with red circles in Fig. 3) drastically increases with GO concentration. These results are in good agreement with XRD.
data which are suggesting a certain involvement of the GO in the composite HT3Ce-xGO phase arrangements. A possible explanation of the overall mechanism for the occurrence of the „ovoidal” composite particles and their positioning towards the surface could be related also to the complex role played by Ce ions. On one hand, they can influence the oxidant-reductive balance and act on GO sites (not completely oxidized) and by another, they could induce a basic character in the LDH reaction loci, leading to a spherical association of the hexagonal LDH platelets [19].

The presence of GO in all the composites HT3Ce-xGO was clearly evidenced by Raman spectra of the samples which are displayed in Fig. 4. The spectrum of HT3Ce shows several bands characteristic to the LDH structure e.g. a broad Raman band at 137 cm\(^{-1}\) (assigned to the LDH lattice vibrations), a band at 548 cm\(^{-1}\) (symmetric stretching vibration of the hydrogen bonds formed between interlayer water and carbonate where the two hydrogen atoms of the H\(_2\)O molecule are bridged to the two oxygen atoms of the carbonate anion) and another band related to the \(v_1\) symmetric stretch of A1' symmetry of carbonate anion at 1069 cm\(^{-1}\) [20].

Besides these, there is also a band-specific to nanocrystalline CeO\(_2\) at 465 cm\(^{-1}\) and weak Raman shifts in the high-frequency region between 800 and 900 cm\(^{-1}\), specific to peroxide (O\(_2^2^-\)) stretching vibrations at the oxidized and defective ceria surfaces [21].

The Raman spectrum of GO shows the D band which is the dominant sp\(^2\) Raman signature of disorder in nanocrystalline carbonic structures at 1321 cm\(^{-1}\) and the band G at 1586 cm\(^{-1}\) (characteristic to planar carbonic structures with sp\(^2\) hybridized C atoms) [22], while the ratio between the intensities of the two bands is \(I_D/I_G=1.316\). Besides the band at 137 cm\(^{-1}\) characteristics to the LDH lattice vibrations, in the Raman spectra of the HT3Ce-xGO composites, the characteristic features of the GO component overlap the bands specific to the LDH component. Only the spectrum of HT3Ce-5GO shows deformed weak bands in the region 200-800 cm\(^{-1}\) which could be caused by the LDH component of the nanocomposite. In the spectra of all composites, the positions of the GO bands D and G are slightly shifted to 1326 and 1593 cm\(^{-1}\) respectively while the ratio \(I_D/I_G\) increases with the GO content in the sample.
(5%, 10% 15%) to 1.357, 1.424, and 1.473 respectively, indicating an increased disturbance of the GO layer. DRIFT spectra presented in (Fig. 5a) showed that GO has the characteristic absorption peaks for C=O carbonyl stretching at 1731 cm\(^{-1}\), 1640 cm\(^{-1}\) (\(\delta_{\text{H}_2\text{O}}\)), 1427 cm\(^{-1}\) (\(\nu_{\text{as cm}}\)), 1264 cm\(^{-1}\) (\(\nu_{\text{C-OH}}\)), 1160 cm\(^{-1}\) (edge OH), 1065 cm\(^{-1}\) (\(\nu_{\text{C-O}}\)), as well as the band at 3640 cm\(^{-1}\) characteristic to isolated –OH groups and the doublet around 2800 cm\(^{-1}\) characteristic to methylene groups [23]. The spectrum of HT3Ce displayed a broad asymmetric absorption band in the region of hydroxyl groups with a maximum centered at 3597 cm\(^{-1}\), the band-specific for interlayer water bending at 1653 cm\(^{-1}\), the one for carbonate ion vibration at 1435 cm\(^{-1}\), a broad asymmetric absorption band with the maximum at 946 cm\(^{-1}\) which is due to the overlapping of different vibration modes and a doublet of weak bands at 505 and 580 cm\(^{-1}\) characteristic to Ce-O and Al-O vibrations [24]. The spectra of the HT3Ce-xGO composites indirectly showed a perturbation of the LDH interlayer region under the influence of the GO component caused by noticeable vibrations of hydrogen bonds formed between water molecules and the anions located in the interlayer region (shoulder at 3450 cm\(^{-1}\)) and the shifting of the band corresponding to \(\nu_{\text{OH}}\) to higher wavelengths (3625 cm\(^{-1}\) compared to 3597 cm\(^{-1}\) in HT3Ce). The occurrence of the bands characteristic for edge OH found in GO at 1264 and 1160 cm\(^{-1}\) (which are more visible in the spectrum of HT3Ce-15GO) has indicated that a decrease in size of the particles found in the powder form is correlated with the increase of the content in GO component, thus confirming the observations from SEM analysis [25].

![Fig. 5. DRIFT spectra of the freshly prepared solids (a) – GO and HT3Ce; (b) HT3Ce-xGO composites](image)

The results of the catalytic tests performed in a batch system at room temperature using 1 wt.% catalyst in simulated wastewater containing 420 ppm indigo carmine (IC) and H\(_2\)O\(_2\) (5.5 wt.%) as the oxidation agent are displayed in Table 1 and Table 2. The results in Table 1 highlight, without any doubt the activation effect of the ultrasonic irradiation which allowed doubling the DR\% in the first 30 minutes of reaction time and reaching after 2 h to the same DR\% as in the tests performed with LDH containing catalysts under continuous stirring for 4 h. The results of the blank tests performed without any catalyst have indicated also the role played by US irradiation in H\(_2\)O splitting into radicals which have increased the amount of oxidizing species in the reaction mixture [6]. Under both testing conditions, the composite catalysts HT3Ce-xGO were more active than the neat components and their activity increased along with their GO content. This fact was well correlated to the structural analysis of the composites which showed their higher
crystallinity compared to the neat components, as well as the beneficial effect of GO content increase on the distribution of the small-sized particles, which could lead to an enhancement of the catalytic activity due to better accessibility to the active sites.

Table 1. The extent of dye removal (DR%) after the oxidation of IC with H$_2$O$_2$ in the tests performed under ultrasonic irradiation (UST) and under stirring (400 rot/min) (ST)

| Catalyst     | HT3Ce   | HT3Ce-5GO | HT3Ce-10GO | HT3Ce-15GO | GO     | blank |
|--------------|---------|-----------|------------|------------|--------|-------|
| UST 30 min   | 48.0    | 57.1      | 61.7       | 78.1       | 49.9   | 12.7  |
| 1 h          | 55.4    | 68.6      | 73.2       | 84.6       | 52.2   | 12.9  |
| 2h           | 70.3    | 81.5      | 85.6       | 97.3       | 58.2   | 13.3  |
| ST 30 min    | 20.6    | 27.2      | 28.6       | 33.5       | 15.1   | 4.4   |
| 1 h          | 35.6    | 40.5      | 44.1       | 58.5       | 20.6   | 4.7   |
| 2 h          | 57.9    | 65.3      | 69.7       | 76.5       | 32.3   | 5.1   |
| 3 h          | 66.2    | 72.0      | 75.6       | 88.2       | 41.2   | 5.7   |
| 4 h          | 70.5    | 81.6      | 85.4       | 97.0       | 54.1   | 6     |
| 5 h          | 80.2    | 87.5      | 90.2       | 97.6       | 60.4   | 6.2   |
| 6 h          | 86.0    | 93.2      | 96.0       | 98.0       | 64.5   | 6.7   |
| 24 h         | 90.0    | 96.3      | 97.7       | 99.5       | 68.3   | 7.5   |

Table 2. The values of COD and TOC after the oxidation of IC with H$_2$O$_2$ in the tests performed under ultrasonic irradiation (UST) for 2 h and under stirring (400 rot/min) (ST) for 4 h, and 24 h (the initial values of COD and TOC were 574.7 mg O$_2$/L, and 174.2 mg C/L, respectively)

| Catalyst     | HT3Ce   | HT3Ce-5GO | HT3Ce-10GO | HT3Ce-15GO | GO     | blank |
|--------------|---------|-----------|------------|------------|--------|-------|
| UST 2 h      |         |           |            |            |        |       |
| COD (mg O$_2$/L) | 256    | 205       | 186        | 132        | 312    |       |
| COD decrease (%) | 55.3   | 64.3      | 67.6       | 77         | 45.6   |       |
| TOC (mg C/L)  | 108     | 86        | 78         | 56         | 131    |       |
| TOC decrease (%) | 37.5   | 50.0      | 54.6       | 67.7       | 23.9   |       |
| ST 4 h       |         |           |            |            |        |       |
| COD (mg O$_2$/L) | 255.8  | 204.7     | 187.2      | 133.8      | 331.2  |       |
| COD decrease (%) | 55.5   | 64.4      | 67.4       | 76.7       | 42.4   |       |
| TOC (mg C/L)  | 107     | 86        | 79         | 56         | 139    |       |
| TOC decrease (%) | 37.7   | 50.0      | 54.4       | 67.4       | 19.3   |       |
| ST 24 h      |         |           |            |            |        |       |
| COD (mg O$_2$/L) | 166    | 137       | 130.6      | 122.3      | 265.9  |       |
| COD decrease (%) | 71.1   | 76.2      | 77.3       | 78.7       | 53.7   |       |
| TOC (mg C/L)  | 70      | 58        | 55         | 51         | 112    |       |
| TOC decrease (%) | 59.6   | 66.6      | 68.2       | 70.2       | 35.2   |       |

For all the catalysts, the values of COD and TOC determined after the tests performed with ultrasonication for 2 h, and after the tests performed under stirring for 4 h and 24 h are presented in Table 2.

For the most active catalyst, HT3Ce-15GO, the COD after 30 minutes reaction time under ultrasonic irradiation was 221 mg O$_2$/L, while TOC was 93 mg C/L marking a decreased percentage of 61.6% for COD and 46.2% for TOC. The values of COD and TOC displayed in Table 2, showed that the total mineralization of the indigo carmine was not reached even when the dye removal exceeded 95%. This fact was also noticed by other authors when they performed the photocatalytic oxidation on TiO$_2$ [10]. However, the values obtained for COD using HT3Ce-15GO were very close to the
maximum admissible value for the direct discharge in water bodies of the treated wastewater sample (e.g. 125 mg O₂/L) [26]. For all the other catalysts, the treated wastewater fulfilled the requirements for discharge in a water treatment plant (e.g. COD < 500 mg O₂/L) [27].

The DRIFT spectra of the catalysts recovered after the catalytic tests performed during 2 h under ultrasonic irradiation and after 24 h under stirring conditions are presented in Fig. 6.

The spectra of the composites used in UST (Fig. 6a) indicate that the ultrasonic radiation and the contact with the dye and H₂O₂ affected similarly all three composites, mainly by decreasing the intensity and the broadness of the bands located in the region 4000-3000 cm⁻¹ where the vibrations characteristic to hydroxyl groups and interlayer interactions through hydrogen bonds appear. This fact correlated with the shifting of the band at 1650 to 1662 cm⁻¹, and the appearance of two new bands at 1210 and 1111 cm⁻¹ may be due to the incorporation of indigo carmine or some of its degradation products in the interlayer region of the solids. The spectra of the composites used in ST (Fig. 6b) show a similar effect in the mid-infrared region accompanied also by a decrease of the intensity and the broadness of the bands in the region 4000-3000 cm⁻¹, but this effect is more pronounced for the samples HT3Ce-5GO and HT3Ce-10GO. The spectrum of HT3Ce-15GO suggests that most probably because this solid had smaller particles, the incorporation of guest species in its interlayer region was more intense since the intensity of the band at 3270 cm⁻¹ is twice as big compared to the other two composites.

**CONCLUSIONS**

Based on the above-presented results, it may be concluded that the co-precipitation of Mg₃Al₇₅Ce₀₂₅ LDH in the presence of GO suspension leads to HT3Ce-xGO composites with increased crystallinity of the LDH-phase compared to the synthesis of Mg₃Al₀₇₅Ce₀₂₅ LDH using only aqueous solutions when the LDH phase was impurified with Cerium hydroxycarbonate and CeO₂. This fact suggests that the presence of GO nanosheets with an increased number of defect sites on the surface, as it was indicated by the I_D/I_G ratio of 1.316 determined by Raman spectroscopy, could both promote the in-situ formation and immobilization of LDH on GO, and inhibit the agglomeration of LDH nanosheets during the growth process while orientating the crystallization of the LDH phase. The XRD patterns of the composites indicated that besides the CeO₂ phase there were no other impurities. Raman spectroscopy highlighted clearly the existence of GO in the HT3Ce-xGO composites. The appearance of CeO₂ nanocrystallites in the structure of the composites played the main role in promoting the catalytic oxidation of indigo carmine with
H$_2$O$_2$. Structural analyses indicated that the dimensions of the nanocomposite particles have decreased with the increase of GO content, thus leading to an enhanced dispersion of the catalytic active sites. The results of the catalytic tests for the oxidation of indigo carmine suggested a synergetic effect of GO and HT3Ce which led to increased catalytic activity, and its highest value was noticed for the composite HT3Ce-15GO, mostly under ultrasonic irradiation.

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