**Abstract:** Vermiculite two-dimensional mixed-layer interstratified structures are a very attractive material for catalysis and photocatalysis. The iron-containing vermiculite from the Palabora region (South Africa) and its samples, which calcined at 500 and 700 °C, were studied in comparison with the α-Fe₂O₃ nanoparticles/vermiculite composites for the first time as photocatalysts of methanol decomposition, which is an organic pollutant and an efficient source for hydrogen production. The aim of the work was to characterize their structural properties using X-ray fluorescence, X-ray diffraction, infrared spectroscopy, nitrogen physisorption, diffuse reflectance UV-Vis spectroscopy and photoluminescence spectroscopy to explain the photocatalytic effects. The photocatalytic test of the samples was performed in a batch photoreactor under UV radiation of an 8W Hg lamp. The photocatalytic activity of vermiculite–hydrobiotite–mica-like layers at different water hydration states in the interstratified structure and the substitution ratio of Fe(III)/Al in tetrahedra can initiate electrons and h⁺ holes on the surface that attack the methanol in redox processes. The activity of α-Fe₂O₃ nanoparticle photocatalysts stems from a larger crystallite size and surface area. The hydrogen production from the methanol–water mixture in the presence of vermiculites and α-Fe₂O₃ nanoparticles/vermiculite composites was very similar and higher than the yield produced by the commercial TiO₂ photocatalyst Euronik P25 (H₂ = 1052 μmol/gcat). The highest yield of hydrogen was obtained in the presence of the Fe/V–700 composite (1303 μmol/gcat after 4 h of irradiation).

**Keywords:** vermiculite; α-Fe₂O₃ nanoparticles; structural properties; photocatalytic activity; methanol–water decomposition

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1. Introduction

Many researchers have focused on raw phyllosilicates as ideal materials for the immobilization of photocatalytic active nanoparticles to avoid potential environmental and health risks (e.g., [1,2]). Vermiculites play a great role as the layered carrier in the synthesis of photocatalysts with high performance. The structural unit layer of the 2:1 TOT type consists of two tetrahedral (T) Si-O sheets and an octahedral (O) Al-O/Al-OH sheet between them. The central Si atoms in the tetrahedra substituted by Al, and/or the octahedral Al atoms by atoms of lower oxidation number such as Mg, generate the negative charges of the layers that are compensated by interlayer cations [3,4]. Iron in the amounts varying from traces to up to 30 mass % can also substitute for Si in tetrahedral or for Al in octahedral sites [5]. The redox reaction Fe(II)–Fe(III) occurs on the clay mineral edges and the basal surface [6].
Vermiculite formed as a weathering product of phlogopite when K\(^+\) in the interlayer was replaced by exchangeable interlayer cations such as Mg\(^{2+}\), usually coordinated between two layers of water [7]. The hydration state of vermiculite is defined by the number of water layers in the interlayer space of the different phases, such as zero-, one- and two-water layer hydration states (0-, 1- and 2-WLHS, respectively) of mica, hydrobiotite and vermiculite [8–10]. Vermiculite from the Palabora (South Africa) occurring in the phlogopite- and apatite-rich pyroxenites was designated as a reference clay mineral sample named hydrobiotite [11]. The vermiculitization process of phlogopite was supported by oxidation of the Fe(II) at the octahedral sites and migration and oxidation to the Fe(III) at the tetrahedral sites in the mixed–layer phase [12]. Cation replacement in octahedra and tetrahedra averages layer orientation of water hydrated to the interlayer cation [13].

From the catalytic point of view, the vermiculite two-dimensional layered structure is a very attractive material due to the cation substitutions in the tetrahedral position, which provide more Brønsted-type acid sites on the surface in comparison with other hydrous 2:1 layer minerals such as smectites. Contributors to Brønsted acidity are also water molecules coordinated to the exchangeable cations and hydroxyl groups at the crystal edges. The Lewis acid sites are associated with unsaturated central atoms at the crystal edges, and the exchangeable cations (especially transition metal ions) and acidity resulting from the dissociation of water molecules in the hydration shell around the exchangeable cations in the interlayer (e.g., [14]).

Vermiculite (VMT) has been investigated as carrier of oxide nanoparticles, such as photocatalysts: TiO\(_2\)/VMT [15], NiO/VMT [16], CeO\(_2\)/VMT [17,18] and Fe\(_2\)O\(_3\)/VMT [19], or mixed oxides, such as Mn\(_2\)O\(_3\)-Fe\(_2\)O\(_3\)/VMT [20], MnO\(_x\)-CeO\(_2\)-Fe\(_2\)O\(_3\)/VMT [21] and CeO\(_2\)-Fe\(_2\)O\(_3\)/VMT [22] catalysts. Reli et al. [22] confirmed that the CeO\(_2\)-Fe\(_2\)O\(_3\) nanoparticles deposited on vermiculite only at 35%–70% showed slightly lower photocatalytic activity than stand-alone CeO\(_2\)-Fe\(_2\)O\(_3\) nanoparticles.

Unlike many reports synthesizing photocatalytically active catalysts and clay minerals used as catalyst supports, the number of reports utilizing the catalytic effects of natural vermiculite materials is still very rare. Vermiculites for the first time were successfully used to effectively split water for hydrogen generation under visible light [23] and were employed as a catalyst on thermal depolymerization products of polystyrene [24].

The clay-based photocatalyst has been recognized as one of the most popular solid photocatalysts for hydrogen production and wastewater degradation due to its special layered structure, large specific surface area and high adsorption capacity, which are suitable for the capture of organic pollutants. Interfacial water molecules associated with the clay surface (anionic director) were observed mainly in montmorillonites, where cation isomorphous substitution occurs only in the octahedral layer. Clay/clay base materials applied in the photo-reforming/water-splitting reaction for hydrogen production from water were attapulgite [25] and Hangjin2# clay [26].

The oxides are currently used as optically active materials because of their chemical stability and low cost. Among them, hematite \(\alpha\)-Fe\(_2\)O\(_3\) as a narrow-band semiconductor can convert about 40% of solar radiation into useful energy [27]. Therefore, it is generally assumed that the impregnation of clay with Fe\(_2\)O\(_3\) will increase the conversion of light into chemical energy. Until recently, there were few articles on photocatalytic water reduction using an \(\alpha\)-Fe\(_2\)O\(_3\)/clay-based heterosystem, and to the best of our knowledge, \(\alpha\)-Fe\(_2\)O\(_3\)/natural illitic clay was used for the first time [28].

Nowadays, major global problems have arisen, namely, increasing energy consumption, depletion of fossil fuels, and environmental pollution caused by human activity. Hydrogen appears to be a very promising possibility to replace fossil fuels with minimal impact on the environment. Water splitting is a promising technology based on natural, sustainable and clean resources. Hydrogen generation from water splitting is energetically an uphill process. For that reason, a sacrificial agent (such as methanol) was used as an electron donor for water splitting [29–32].
This study is an ongoing work on the structural, textural, optical and photocatalytic properties of raw and heat-treated vermiculite from the deposit in Palabora [24]. Vermiculite and its samples calcined at 500 and 700 °C were studied in comparison with the α-Fe₂O₃ nanoparticles/vermiculite composite heterosystem for the first time as photocatalysts of methanol decomposition. The aim was to explain the photocatalytic effects based on the structural properties of photocatalysts using X-ray fluorescence, X-ray diffraction, infrared spectroscopy, nitrogen physisorption, diffuse reflectance UV-Vis spectroscopy and photoluminescence spectroscopy.

2. Materials and Methods

2.1. Materials and Samples Preparation

Vermiculite mined from the deposit in the Palabora region in the Northern Province of South Africa was supplied by the company Grena Co. (Veselí nad Lužnicí, Czech Republic). The hematite precursor was iron chloride hexahydrate (FeCl₃·6H₂O) and sodium hydroxide (NaOH), supplied by Lach-Ner Co., Czech Republic.

The vermiculite was pulverized in a ball mill to a powder and sieved with a particle size <40 µm to the sample denoted as Ver. The Ver was then calcined to the samples denoted as V–500 and V–700 in a muffle oven (NABERTHERM P 330, NABERTHERM Inc., New Castle, USA) at 500 and 700 °C using a temperature ramp of 10 °C min⁻¹ and an isothermal step of one hour at the final temperature.

The crystalline α–Fe₂O₃ nanoparticles were prepared by chemical precipitation method [33]. The nanoparticle precursor was iron(III) chloride hexahydrate (FeCl₃·6H₂O) salt and sodium hydroxide (NaOH) solution as precipitator. An amount of 4 g of FeCl₃·6H₂O in 100 mL of distilled water was stirred in bottle at 450 rpm for 30 min at 70 °C in Heidolph Reax overhead shaker (REAX 20/4, Heidolph Instruments GmbH & Co. KG, Schwabach, Germany), to which 2 M NaOH was added drop by drop, until pH 11 value was reached.

α–Fe₂O₃ nanoparticles/vermiculite composites were obtained from three water dispersions containing 4 g of each Ver, V–500 and V–700 in 100 mL distilled water, which were stirred in bottles at 450 rpm for 30 min at 70 °C in Heidolph Reax overhead shaker (REAX 20/4, Heidolph Instruments GmbH & Co. KG, Schwabach, Germany). The solution of 100 mL of α–Fe₂O₃ nanoparticles precursor prepared before was then added and mixed together for 30 min. The precipitates were denoted as Fe/Ver, Fe/V–500 and Fe/V–700, collected and centrifuged at 6000 rpm. The solid fractions were repeatedly washed to free chlorides. The samples were dried at 40 °C and calcined at 500 °C using a temperature ramp of 10 °C min⁻¹ and an isothermal step of one hour at the final temperature in a muffle furnace (NABERTHERM L9/11/SW, Nabertherm GmbH, Lilienthal, Germany) to obtain crystalline α–Fe₂O₃ nanoparticles sole and attached to the vermiculite support.

2.2. Methods

The elemental analysis was obtained using a SPECTRO XEPOS energy dispersive X-ray fluorescence (ED-XRF) spectrometer (Spectro Analytical Instruments, Kleve, Germany). Each sample (2 × 4 g) was mixed with wax (0.9 g) and pressed at 10 tons for 30 s using a manual hydraulic press (BSL-2, Brio, Czech Republic) and a standard 24 mm diameter die, resulting in a cylindrical pellet thick of about 4 mm. Analysis was performed in an inert atmosphere (He). The amount of Fe(II) in Ver, V–500 and V–700 samples was determined by redox titration with the 0.1 M solution of potassium dichromate (K₂Cr₂O₇), which was then subtracted from the total Fe concentration determined by XRF. The difference was the amount of Fe(III).

The scanning electron microscope (SEM) JEOL JSM 7610F+ (JEOL, Tokyo, Japan) with Schottky cathode was utilized to examine the samples prepared on stubs with carbon tape and coated with 20 nm layer of Pt. Elemental composition analysis and mapping were performed using energy-dispersive X-ray spectroscope (EDS) Aztec Ultima Max 65 (Oxford Instruments, Abingdon, UK).
X-ray powder diffraction (XRD) phase analysis was performed using Rigaku SmartLab diffractometer (Rigaku Corporation, Tokyo, Japan), equipped with a detector D/teX Ultra 250 under CoKα radiation. The sieved samples before analysis were further powdered using agate mortar and pressed using microscope glass into a rotational sample holder. The XRD patterns were obtained in the reflection mode (Bragg-Brentano geometry) at 40 kV and 40 mA, at 15 rpm/min, a step size of 0.01° at the speed of 0.5 deg/min. The XRD patterns were evaluated using PDFX2 software No. 2.4.2.0 (Rigaku Corporation, Tokyo, Japan) and compared with database PDF-2, 2015 (ICDD, Newton Square, USA). In order to calculate the average crystallite size of the formed α-Fe₂O₃ nanoparticles, Scherrer equation: \( D_{(012)} = \frac{0.9 \lambda}{\beta \cos \theta} \), was utilized. In this equation, the parameters \( \lambda \), \( \beta \), and \( \theta \) stand for the incident X-rays wavelength, the full width at half maximum (FWHM) of the 012 diffraction peak, and the diffraction angle, respectively.

The infrared (IR) spectra were obtained on a Nicolet 6700 FTIR spectrometer from Thermo Scientific (Mundelein, IL, USA), equipped with an IR source, KBr beamsplitter and DTGS detector. The KBr pressed disk technique (1 mg of sample and 200 mg of KBr) was utilized to measure spectra in the mid-IR region (4000–400 cm\(^{-1}\)). The pellets were overnight heated at 120 °C to remove water adsorbed on the sample and KBr. The spectra were analyzed by the Thermo Scientific OMNIC™ software package. To compare the intensities of the bands of the samples, the spectra were normalized to the same intensity of the Si-O band near 1000 cm\(^{-1}\). The exact position of the overlapping bands (marked with * in the figures) was found by the second derivative operation using the Savitsky–Golay derivative embedded in OMNIC software.

The nitrogen physisorption measurements were performed on the 3Flex physisorption set-up (Micromeritics, USA). The specific surface area, \( S_{BET} \), was evaluated according to Brunauer–Emmett–Teller (BET) theory at 77 K for the \( p/p_0 = 0.05–0.25 \) measuring overall the 11 points within this relative pressure range [34]. The total pore volume, \( V_{tot} \), was evaluated as the adsorbed nitrogen volume at maximum \( p/p_0 \sim 0.99 \) from the adsorption branch of nitrogen adsorption–desorption isotherm. The micropore volume (\( V_{micro} \)) and the mesopore surface area (\( S_{meso} \)), were verified using the t-plot method, applying the Broekhoff-de Boer standard isotherm [35]. The mesopore–macropore-size distribution was evaluated from the adsorption branch of the nitrogen adsorption–desorption isotherm via the Barrett–Joyner–Halenda (BJH) method [36], assuming the cylindrical-pore geometry characterized by the diameter \( d_p \) of the pores. The micropore-size distribution was evaluated from the adsorption branch of the nitrogen adsorption–desorption isotherm, applying the Horwath–Kawazoe method [37] assuming the slit pore-geometry characterized by the width \( w_p \) of the pores.

The UV-Vis diffuse reflectance spectra (UV-Vis DRS) were recorded using a Shimadzu UV-2600 Series spectrophotometer (Shimadzu Scientific Co., Tokyo, Japan) from 220 to 800 nm. The measurements of all spectra were carried out by external 2D detector, an average diameter of 60 mm of an integration sphere and using the BaSO₄ standard. Kubelka–Munk function [38] was obtained by calculation from reflectance spectrum. The direct band-gap energies (\( E_g \)) values were determined from Tauc plot for \( n = 2 \) (constant for direct band-gap energy) [39].

The photoluminescence (PL) spectra were recorded using a spectrometer FLSP920 Series (Edinburgh Instruments, Ltd., Livingston, UK) from 350 to 620 nm with a 450 W Xenon lamp (Xe900) and a R928P type PMT detector. All PL spectra were measured at fixed 325 nm excitation wavelength with the width of the excitation slit at 3.0 nm, emission slit at 8.0 nm and dwell time of 0.5 s.

The photocatalytic tests were performed in a homemade photoreactor. Detailed description, scheme and real photo of photocatalytic apparatus are in the Supplementary Materials.
3. Results

3.1. XRF and SEM Analysis

The chemical composition of elements obtained from XRF analysis was recalculated to the stoichiometric metal oxides concentrations (Table 1). Different amounts of alkali oxides K$_2$O, Na$_2$O and MgO in Ver, V–500 and V–700 vermiculite samples in comparison with the Fe/Ver, Fe/V–500 and Fe/V–700 composite samples indicated partial cation exchange Na$^+$ for K$^+$ during the preparation procedure with NaOH precipitator. The amount of Fe$_2$O$_3$ 9.17, 10.20 and 11.12 mass% in Ver, V–500 and V–700, respectively, increased in the Fe/Ver, Fe/V–500 and Fe/V–700 composite samples due to the deposited hematite nanoparticles by 28.2, 25.0 and 28.0 mass%, respectively.

Table 1. Chemical XRF analysis (mass%) of vermiculites and α-Fe$_2$O$_3$ nanoparticles/vermiculite composites and the statistical error with a confidence interval of 1 sigma.

<table>
<thead>
<tr>
<th>Sample</th>
<th>SiO$_2$</th>
<th>TiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>Fe$_2$O$_3$</th>
<th>FeO</th>
<th>P$_2$O$_5$</th>
<th>CaO</th>
<th>MgO</th>
<th>K$_2$O</th>
<th>Na$_2$O</th>
<th>L.O.I. $^1$</th>
<th>Sum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ver</td>
<td>35.53 ± 0.02</td>
<td>1.26 ± 0.01</td>
<td>7.59 ± 0.01</td>
<td>9.17 ± 0.01</td>
<td>0.53</td>
<td>1.16 ± 0.01</td>
<td>3.67 ± 0.01</td>
<td>23.02 ± 0.05</td>
<td>5.60 ± 0.01</td>
<td>0.08 ± 0.01</td>
<td>12.30</td>
<td>99.91</td>
</tr>
<tr>
<td>V–500</td>
<td>38.27 ± 0.03</td>
<td>3.34 ± 0.01</td>
<td>8.11 ± 0.02</td>
<td>10.20 ± 0.01</td>
<td>0.62</td>
<td>1.22 ± 0.01</td>
<td>3.74 ± 0.01</td>
<td>24.46 ± 0.05</td>
<td>5.92 ± 0.01</td>
<td>0.09 ± 0.01</td>
<td>5.53</td>
<td>99.50</td>
</tr>
<tr>
<td>V–700</td>
<td>37.39 ± 0.03</td>
<td>3.35 ± 0.01</td>
<td>7.73 ± 0.02</td>
<td>11.12 ± 0.01</td>
<td>0.13</td>
<td>1.61 ± 0.01</td>
<td>4.53 ± 0.01</td>
<td>23.36 ± 0.05</td>
<td>5.85 ± 0.01</td>
<td>0.07 ± 0.01</td>
<td>5.27</td>
<td>98.41</td>
</tr>
<tr>
<td>Fe/Ver</td>
<td>25.94 ± 0.01</td>
<td>0.97 ± 0.01</td>
<td>5.40 ± 0.01</td>
<td>37.38 ± 0.02</td>
<td>2.687</td>
<td>1.15 ± 0.01</td>
<td>3.33 ± 0.01</td>
<td>16.55 ± 0.05</td>
<td>3.86 ± 0.01</td>
<td>1.40 ± 0.01</td>
<td>3.87</td>
<td>99.85</td>
</tr>
<tr>
<td>Fe/V–500</td>
<td>27.26 ± 0.02</td>
<td>0.96 ± 0.01</td>
<td>5.82 ± 0.01</td>
<td>35.16 ± 0.02</td>
<td>1.97</td>
<td>1.28 ± 0.01</td>
<td>3.19 ± 0.01</td>
<td>17.12 ± 0.05</td>
<td>3.83 ± 0.01</td>
<td>1.41 ± 0.01</td>
<td>3.86</td>
<td>99.89</td>
</tr>
<tr>
<td>Fe/V–700</td>
<td>24.97 ± 0.02</td>
<td>0.92 ± 0.01</td>
<td>5.23 ± 0.01</td>
<td>39.15 ± 0.02</td>
<td>2.321</td>
<td>1.21 ± 0.01</td>
<td>2.36 ± 0.01</td>
<td>16.34 ± 0.05</td>
<td>3.60 ± 0.01</td>
<td>1.22 ± 0.01</td>
<td>3.87</td>
<td>99.78</td>
</tr>
</tbody>
</table>

$^1$ L.O.I. — Loss on ignition at 1000 °C.

The metal oxgens from the XRF chemical analyses of Ver, V–500 and V–700 (Table 1) were recalculated based on the 22 negative charges per formula unit (pfu) to the cation content for the half-unit cell composition O$_{10}$(OH)$_2$ according to Foster [3] (Table 2).

Table 2. Cations on the basis 22$^-$/O$_{10}$(OH)$_2$ in vermiculite samples.

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Si$^4+$</th>
<th>Al$^{3+}$</th>
<th>Fe$^{3+}$</th>
<th>Tet.ch.</th>
<th>Al$^{3+}$</th>
<th>Fe$^{3+}$</th>
<th>Fe$^{2+}$</th>
<th>Mg$^{2+}$</th>
<th>Ti$^{4+}$</th>
<th>Oct.ch.</th>
<th>Tot.ch.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ver</td>
<td>2.887</td>
<td>0.726</td>
<td>0.387</td>
<td>−1.113</td>
<td>0.00</td>
<td>0.173</td>
<td>0.063</td>
<td>2.687</td>
<td>0.077</td>
<td>+0.327</td>
<td>−0.786</td>
</tr>
<tr>
<td>V–500</td>
<td>2.900</td>
<td>0.723</td>
<td>0.377</td>
<td>−1.100</td>
<td>0.00</td>
<td>0.204</td>
<td>0.039</td>
<td>2.681</td>
<td>0.076</td>
<td>+0.366</td>
<td>−0.734</td>
</tr>
<tr>
<td>V–700</td>
<td>2.902</td>
<td>0.707</td>
<td>0.391</td>
<td>−1.098</td>
<td>0.00</td>
<td>0.258</td>
<td>0.010</td>
<td>2.653</td>
<td>0.079</td>
<td>+0.416</td>
<td>−0.682</td>
</tr>
</tbody>
</table>

Tet.ch. — Tetrahedral charge; Oct.ch. — Octahedral charge; Tot.ch. — Layer unit charge.

SEM images of vermiculite flakes Ver, V–700 and Fe/V–700 observed in the backscattered electron mode (Figure 1) showed discontinuities, marginal undulations and bumps of various sizes. Striking aggregates of flakes were observed in the Ver sample (Figure 1a) and more compact and larger flakes in V–700 (Figure 1b). Calcination temperature influenced micro-morphological textural variations in surface observed as small humps, marginal rolling and discontinuity of the layers. The Fe/V–700 composite contained large aggregates (clusters) of randomly oriented vermiculite flakes, with no hematite nanoparticles observed (Figure 1c). The EDX mapping images confirmed a less homogeneous distribution of Fe on the surface of Ver in comparison to calcined V–700. The EDX elemental spectra of Ver and V–700 are very similar and demonstrate hematite nanoparticles by a higher amount of Fe on the surface of Fe/V–700.

3.2. XRD Phase Analysis

The XRD patterns from 2 to 50° 2θ (Figure 2a,b) of Ver, V–500 and V–700 show vermiculite (marked as V, JCPDS card no. 00-076-0847), hydrobiotite (marked as Hb, JCPDS card no. 00-049-1057) and phlogopite (marked as Ph, JCPDS card no. 01-074-3145) and only Ph in V–700 (Figure 2a). The XRD patterns of Fe/Ver, Fe/V–500 and Fe/V–700 composite samples (Figure 2b) contained only Ph and hematite (H) (Fe$_2$O$_3$, JCPDS card no. 01-073-2234).
The Fe/V–700 composite contained large aggregates (clusters) of randomly oriented vermiculite flakes, with no hematite nanoparticles observed (Figure 1c). The EDX mapping images confirmed a less homogeneous distribution of Fe on the surface of Ver in comparison to calcined V–700. The EDX elemental spectra of Ver and V–700 are very similar and demonstrate hematite nanoparticles by a higher amount of Fe on the surface of Fe/V–700.

Figure 1. SEM images and EDX scanning distribution of Fe (red points) on the surface with the EDX elemental spectra of: (a) Ver, (b) V–700 and (c) Fe/V–700.

Hydroxyapatite (marked as HA, JCPDS card no. 00-064-0738) was a mineral residue from the parent phlogopite-apatite-rich pyroxenites occurring in all samples. The chemical composition of HA corresponds to the nearly identical CaO/P₂O₅ ratios of 3.00 ± 0.04 mass% and 2.61 ± 0.08 mass% in all vermiculite samples and in Fe/vermiculite composite samples, respectively (Table 1).

Characterization of the mixed layers of hydrobiotite [40] occurring in the interstratification domains of vermiculites is performed according to the basal space values, which are given for each peak in the range from 2 to 15° 2θ (Figure 2c,d). The basal peak series in vermiculite samples (Figure 2c) suggest mixed layering of vermiculite–mica-like layers, containing the interlayer cations at different water layer hydration states (WLHS). Ver is composed of hydrobiotite, Hb (1-WLHS, d = 1.216 nm), which predominates over vermiculite, V (2-WLHS, d = 1.469 nm), and a very small peak of phlogopite, Ph (0-WLHS, d = 1.025 nm). The hydrobiotite-type layer sequence [41] produced a broad peak with d = 2.65 nm of Hb in the various mixed-layer stacking sequences in the long-range order [11]. The peak with d = 1.269 nm was assigned to the transitional phase, which can be formed due to an interstratification between the 2-WLHS of V and a less hydrated Hb. The peak with d = 1.025 nm of Ph suggests 0/1-WLHS in the interlayer space [10]. Vermiculite (V) in V-500 was partly dehydrated to the Hb and Ph of the similar peak intensities. In the V-700, a small peak of V (d = 1.469 nm) and only Ph (d = 1.025 nm) can be observed.

The basal peak series in α-Fe₂O₃/vermiculite samples (Figure 2d) corresponds to the Ph (d = 1.025 nm, 1/0-WLHS). However, Fe/V–500 contains interstratification domains composed of mixed layers of Ph and other dehydrated phases: the transient 1-WLHS phase (d = 1.159 nm) and mixed layered 1/0 intermediate space (d = 1.246 nm) The
interstratifications between these contracting and non-contracting forms were observed in many vermiculites [41].

Figure 2. XRD patterns from 2 to 50° 2θ of: (a) vermiculites (Ver, V–500 and V–700) and (b) α-Fe₂O₃ nanoparticles/vermiculite composites (Fe/Ver, Fe/V–500 and Fe/V–700). Peaks are marked: V = vermiculite, Hb = hydrobiotite, Ph = phlogopite, H = hematite and HA = hydroxyapatite; and XRD patterns from 2 to 15° 2θ with d-values of peaks of: (c) vermiculites and (d) α-Fe₂O₃/vermiculite composites.

Diffractions of hematite (H) and the Ph are mostly overlapped. Therefore, the single reflection from the H(012) plain (marked H₀₁₂ in Figure 2b) was used to calculate the mean coherent crystallite size \( D(012) \) [42] (Table 3). The obtained mean size of Fe₂O₃ crystallites, generated as a result of deposition from the Fe(OH)₃ precursor (Table 3), indicates that the mean crystallite size is about 30 nm and the deviation from the mean value is about 3 nm.

Table 3. The mean coherent crystallite size \( D(012) \) of hematite in \( \alpha \)-Fe₂O₃ nanoparticles sole and in \( \alpha \)-Fe₂O₃/vermiculite composites.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Fe–500</th>
<th>Fe/Ver</th>
<th>Fe/V–500</th>
<th>Fe/V–700</th>
</tr>
</thead>
<tbody>
<tr>
<td>( D(012) ) (nm)</td>
<td>29</td>
<td>31</td>
<td>27</td>
<td>34</td>
</tr>
</tbody>
</table>

3.3. Infrared Spectroscopy Analysis

The IR spectra of Ver, V–500 and V–700 were analyzed to obtain information on the changes in the composition of Ver upon heating (Figure 3a). In contrast to XRD, the identification of the individual minerals in vermiculites is less straightforward because IR spectroscopy provides evidence, primarily on the nearest environment of the OH groups, i.e., on the occupancy of the central positions in the trioctahedral sheet. Therefore, for the
example, this method is not able to detect unambiguously hydrobiotite but only biotite and vermiculite. The spectrum of Ver shows a broad complex band in the OH stretching region (Figure 3A). The absorption band at 3715 cm\(^{-1}\) related to the stretching vibration (\(\nu\)) of the Mg\(_2\)OH groups, is characteristic of phlogopite, while a band at 3666 cm\(^{-1}\) is ascribed to the vibrations of Mg\(_2\)Fe\(^{2+}\)OH groups, which can be present also in the biotite or vermiculite [43–45]. The inflection at 3565* cm\(^{-1}\) (revealed by the second derivative) and a weak band at 3537 cm\(^{-1}\) probably correspond to vMgFe\(^{3+}\)OH and vFe\(^{3+}\)Fe\(^{3+}\)OH, respectively, of biotite and/or vermiculite. The broad band at 3404 cm\(^{-1}\) belongs to the stretching vibrations of water molecules. Heating to 500 °C only slightly modifies the shape of the complex OH stretching band; more pronounced changes are recognized after vermiculite heating at 700 °C (Figure 3A). The intensity of the broad band of water significantly decreased alike those of the vMgFe\(^{3+}\)OH and vFe\(^{3+}\)Fe\(^{3+}\)OH bands. The bands of phlogopite at 3712 and 3661 cm\(^{-1}\) remained unchanged, which is in agreement with XRD results indicating phlogopite as the dominant phase in V–700.

![Figure 3. IR spectra of: (a) vermiculite (Ver) and samples heated at 500 °C (V–500) and 700 °C (V–700), (A)–OH stretching region; (B)–Si-O and OH bending region. * indicates the position obtained by the second derivative operation and (b) Ver, Fe/Ver and Fe–500.](image)

In the 1200–400 cm\(^{-1}\), the bands related to the OH bending vibrations are overlapped with the bands resulting from stretching and bending (\(\delta\)) vibrations of the tetrahedral sheets (Figure 3B). The low position of the dominant band of this region near 1000 cm\(^{-1}\), related to the stretching vibrations of the central atoms of the tetrahedral sheet, confirmed the presence of Al or Fe in addition to Si in tetrahedra. This broad Si-O bond was shifted from 999 cm\(^{-1}\) in Ver to 1002 cm\(^{-1}\) in V–500 and 1004 cm\(^{-1}\) in V–700. The bands at 818, 727 and 685 cm\(^{-1}\) are due to Al\(^{IV}\)-O\(_{ap}\), Al\(^{IV}\)-O-Si and Si-O\(_{ap}\), respectively. The band at 676 cm\(^{-1}\) is related to \(\delta\)Mg\(_3\)OH of vermiculite, while the position at 605 cm\(^{-1}\) is assigned to \(\delta\)Mg\(_3\)OH of phlogopite [43]. The weak band at 637 cm\(^{-1}\) corresponds to \(\delta\)Mg\(_2\)Fe\(^{2+}\)OH [45]. The spectra of V–500 and V–700 show only minor changes compared to the unheated sample. Only a decrease in the 637 cm\(^{-1}\) band intensity signalized changes in the trioctahedral sheet occupancy (Figure 3B).

Precipitation procedure and calcination temperature of 500 °C of hematite nanoparticles onto vermiculite modified the shape of the spectrum in the 700–400 cm\(^{-1}\) region, i.e., in the area where vibrations of the Fe\(_2\)O\(_3\) occur (Figure 3b). The contribution of the hematite bands at 530 and 444 cm\(^{-1}\) increases the absorption in this region; however, the positions of the bands observed for Ver remained almost unchanged for Fe/Ver, i.e., near 1000 cm\(^{-1}\) (vSi-O), 818 cm\(^{-1}\) (Al\(^{IV}\)-O\(_{ap}\)), 733 cm\(^{-1}\) (Al\(^{IV}\)-O-Si), 686 cm\(^{-1}\) (Si-O\(_{ap}\)) and 602 cm\(^{-1}\) (Mg\(_3\)OH). The IR patterns of the Fe/V–500 and Fe/V–700 were identical to that
of Fe/Ver; therefore, they are not presented. Unfortunately, the IR spectroscopy cannot distinguish whether hematite particles are interacting with the vermiculite surface or if the samples consist of two separate phases of vermiculite and hematite.

3.4. Textural Properties $S_{\text{BET}}$ and $V_{\text{tot}}$

Measured nitrogen adsorption–desorption isotherms (Figure 4a,b) and evaluated pore-size distributions (Figure 4c,d) of Ver and its heated samples revealed any significant changes within the meso- and macroporous structure, arising from particle-to-particle interactions [46]. Vermiculites, Ver, V–500 and V–700, comprise minor smaller mesopores ($d_p$ about 3 nm) corresponding to the basal interlayer space and dominant large mesopores and macropores ($d_p > 40$ nm) corresponding to the slit-shaped pores between particles (Figure 4c).

![Figure 4. Nitrogen adsorption–desorption isotherms of: (a) Ver, V–500 and V–700 and (b) Fe/Ver, FeV–500 and Fe/V–700 and pore size distributions of: (c) Ver, V–500 and V–700 and (d) Fe/Ver, Fe/V–500 and Fe/V–700.](image)

The surface area of 23 m$^2$/g and the total pore volume of 85 mm$^3_{\text{liq}}$/g of V–500 were slightly smaller in comparison with Ver (25 m$^2$/g and 86 mm$^3_{\text{liq}}$/g). The heating of Ver at 700 °C increased in V–700 the total pore volume to 100 mm$^3_{\text{liq}}$/g in the mesoporous–macroporous structure while the size of the surface area did not change (Table 4).
Table 4. Textural properties $S_{\text{BET}}$ and $V_{\text{tot}}$ of vermiculites, $\alpha$-$\text{Fe}_2\text{O}_3$ nanoparticles and $\alpha$-$\text{Fe}_2\text{O}_3$ nanoparticles/vermiculite composites.

<table>
<thead>
<tr>
<th></th>
<th>Ver</th>
<th>V–500</th>
<th>V–700</th>
<th>Fe–500</th>
<th>Fe/Ver</th>
<th>Fe/V–500</th>
<th>Fe/V–700</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_{\text{BET}}$ ($m^2/g$)</td>
<td>25</td>
<td>23</td>
<td>24</td>
<td>36</td>
<td>58</td>
<td>53</td>
<td>63</td>
</tr>
<tr>
<td>$V_{\text{tot}}$ ($mm^3liq/g$)</td>
<td>86</td>
<td>85</td>
<td>100</td>
<td>197</td>
<td>145</td>
<td>143</td>
<td>161</td>
</tr>
</tbody>
</table>

The Fe–500 nanoparticles sample showed a surface area of 36 $m^2/g$ and a total pore volume of 197 $mm^3liq/g$. Precipitation of nanoparticles on Ver, V–500 and V–700 and heating at 500 °C to the Fe/Ver, Fe/V–500 and Fe/V–700 composites caused an increase in the specific surface area of about 2.5, 2.2 and 2.6 $m^2/g$ times and a total pore volume about 1.7 $mm^3liq/g$ times (Table 4). Moreover, the mesopores with the pore diameter of about 10–11 nm formed mainly in Fe/V–700 composite can be beneficial to the transformation, diffusion and adsorption of molecules [47].

3.5. UV-Vis Diffuse Reflectance (DR) Spectroscopy

Vermiculites have a certain capacity for adsorbing light in the UV-Vis region on the absorption edge around 450 nm [47] (Figure 5). The optical ability of the samples is performed by the UV-Vis diffuse-reflectance spectra (DRS) (Figure 5a). The reflectance recalculated into the absorption intensities of the direct band energy gap ($E_g$) [39] can then be estimated in the Tauc plot (Figure 5b) using the intercept of the tangent to the plot of the optical adsorption coefficient $\alpha$ and photonic energy $h\nu$.

Figure 5. DRS spectra (a) and Tauc plots (b) of vermiculites, $\alpha$-$\text{Fe}_2\text{O}_3$ nanoparticles and $\alpha$-$\text{Fe}_2\text{O}_3$/vermiculite composites.

The first $E_g$ values 1.64 eV in Ver, 1.70 eV in V–500 and 1.86 eV in V–700 were assigned to the iron bound in the vermiculite structure, as it was also documented by IR spectra above. The second $E_g$ value of 3.03 eV in Ver decreasing to the 2.50 eV in V–500 and to 2.36 eV in V–700 was attributed to the vermiculite [47,48] (Figure 5a; Table 5).

Table 5. The band-gap energies $E_g$ of vermiculite samples, $\alpha$-$\text{Fe}_2\text{O}_3$ nanoparticles and $\alpha$-$\text{Fe}_2\text{O}_3$ nanoparticles/vermiculite composite samples.

<table>
<thead>
<tr>
<th></th>
<th>Ver</th>
<th>V–500</th>
<th>V–700</th>
<th>Fe–500</th>
<th>Fe/Ver</th>
<th>Fe/V–500</th>
<th>Fe/V–700</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_g$ (eV)</td>
<td>1.64</td>
<td>1.70</td>
<td>1.86</td>
<td>1.99</td>
<td>1.55</td>
<td>1.50</td>
<td>1.35</td>
</tr>
</tbody>
</table>
The Fe–500 nanoparticles showed characteristic nearly constant reflectivity in the range of 350–550 nm, a shoulder near 620 nm, and a maximum near 750 nm (Figure 5a), in a good agreement with the spectrum of hematite powder [49] and the Eg value of 1.99 eV (Figure 5b). The Eg value of the iron oxide (Fe2O3) semiconductor material is about 2.3 eV [50]. The Eg values of α-Fe2O3 nanoparticles prepared using different methods were reported in the range from 1.88 to 1.93 eV [51].

Hematite nanoparticles attached to the vermiculite surface changed the shape of the spectra and shift the maximum reflectance to about 750 nm. The Eg value, 1.99 eV, of Fe–500 nanoparticles decreased to 1.55 eV in Fe/Ver composite and then to 1.50 eV in Fe/V–500 and to 1.35 eV in Fe/V–700 composites (Figure 5b; Table 5). The reduction in the band-gap Eg value of the Fe2O3 photocatalyst can be ascribed to a combination effect of vermiculite negative layer charge.

3.6. Photoluminescence (PL) Study

Hematite nanoparticle has a local collapse of the Fe magnetic moment and strong local lattice distortion in the strong antiferromagnetic background, which can generate luminescence after photoexcitation [52]. Photoluminescence (PL) spectra (Figure 6) in the wavelength range of 350–600 nm show a UV emission band (374–380 nm) and a green-yellow emission band (410–500 nm). The position of the UV band at 380 nm in Ver, V–500 and V–700 was found to shift towards a lower wavelength at 374 nm in Fe/Ver, Fe/V–500 and Fe/V–700 composites. This position was attributed to the electron transition from the donor level formed by oxygen vacancies to the acceptor level [53]. The sole Fe–500 nanoparticles exhibited a relatively weak emission band and the intensity of the spectra increase in Fe/Ver, Fe/V–500 and Fe/V–700 composites. However, α-Fe2O3 nanoparticles attached to the interstratified structure of the Fe/V–500 composite (Figure 2c,d) exhibited the most intensive PL spectrum. The emission band at 413 nm and 438 nm can be attributed to emission deep level defects associated with oxygen vacancies or at the particle boundaries [53,54].

![Figure 6. PL emission spectra of vermiculites, α-Fe2O3 nanoparticles and α-Fe2O3/vermiculite composites.](image-url)

The emission band at about 489 nm in Ver, V–500 and V–700 was assigned to the charge transfer transition between the iron ions [55], whose intensity corresponds to the Fe2+/Fe3+ ratios of 0.36, 0.19 and 0.04 in the octahedral positions, respectively (Table 2). The emission bands at about 576 nm and 599 nm were assigned to the Fe3+ ligand field transition [56] of...
the recombination of electrons trapped at oxygen vacancy near the conduction band edge with holes in deep Fe-related surfaces [53].

3.7. Photocatalytic Hydrogen Generation from Methanol–Water Mixture

Photocatalytic hydrogen generation from the methanol–water mixture in the presence of vermiculite and Fe/vermiculite composite samples at 254 nm irradiation was performed during the time (0, 1, 2 and 4 h) and the yields of products of hydrogen (divided by 10), methane and carbon monoxide after 4 h (Figure 7). Methane and CO₂ are typical intermediates of photocatalytic methanol decomposition [57]. The generation of hydrogen yields from the photocatalytic decomposition of methanol–water mixture with the presence of vermiculites, α-Fe₂O₃/vermiculite composites and commercial photocatalyst TiO₂-P25 showed the dependence on the time of irradiation 0–4 h (Figure 7a). The highest yields of hydrogen were obtained in the presence of Fe/V–700 composite (1303 µmol/g<sub>cat.</sub>). All investigated photocatalysts produced higher yields of hydrogen than commercial TiO₂ photocatalyst Evonik P25 (1052 µmol/g<sub>cat.</sub>) (Figure 7b).

![Figure 7](image_url)

**Figure 7.** Photocatalytic hydrogen generation from the methanol–water mixture in the presence of vermiculite and Fe/vermiculite composite samples at 254 nm irradiation: (a) Time dependence on the yields of hydrogen and (b) Yields of H₂, CH₄ and CO after 4 h of irradiation; P25 = commercial photocatalyst TiO₂ Evonik P25; H₂/10 = yield divided by 10.

4. Discussion

The mixture of hydrated vermiculite–hydrobiotite–mica-like layers in the interstratified structure of vermiculite transformed on the thermal treatment to the dehydrated phases. The specific surface area and the total pore volume within the mesoporous–macroporous texture of Ver and V–500 were very similar, but a higher total pore volume was formed in V–700 (Table 4).

The similar yields of hydrogen produced from the mixture of methanol–water and vermiculites Ver (1197 µmol/g<sub>cat.</sub>), V–500 (1202 µmol/g<sub>cat.</sub>) and V–700 (1183 µmol/g<sub>cat.</sub>) did not support the knowledge of improving the photocatalytic properties of the catalysts by higher annealing temperature [47,48,58].

In general, photocatalytic degradation of methanol takes place in two ways [29,59,60]. One way is the direct hole transfer reaction, where photo-generated holes serve as oxidation sites for adsorbed methanol molecules. The following possible decomposition reactions...
in Equations (1)–(4) indicate the release of CH₂O (formaldehyde) and CO as intermediate products and H⁺ as the final decomposition product:

\[
\begin{align*}
\text{CH}_3\text{OH} + h^+ & \rightarrow \cdot\text{CH}_2\text{OH} + H^+ \\
\cdot\text{CH}_2\text{OH} & \rightarrow \text{CH}_2\text{O} + H^+ + e^- \\
\text{CH}_2\text{O} + 2O & \rightarrow \cdot\text{CHO} + \text{HO}_2\cdot \\
\cdot\text{CHO} + 2O & \rightarrow \cdot\text{CO} + \text{HO}_2\cdot
\end{align*}
\]

The indirect hole transfer reactions occur by the hole photo-induced products while the presence of HO• radicals to decompose the molecules is essential, e.g., in Equations (5)–(7):

\[
\begin{align*}
\text{H}_2\text{O} + h^+ & \rightarrow \text{HO}• + H^+ \\
\text{CH}_3\text{OH} + \text{HO}• & \rightarrow \cdot\text{CH}_2\text{OH} + \text{H}_2\text{O} \\
\text{CH}_2\text{OH} + \text{HO}• & \rightarrow \cdot\text{CH}_2\text{O} + \text{H}_2\text{O} \\
H^+ + e^- & \rightarrow \text{H}• \\
\text{H}• + \text{H}• & \rightarrow \text{H}_2
\end{align*}
\]

The holes are strong oxidizing agents and responsible for the splitting of water to form H₂ according to Equations (8) and (9).

The vermiculite structure contained ions that can promote the formation of electron e⁻/holes h⁺ pairs. The created h⁺ can directly react with the methanol, OH⁻ in the medium and also with the hydroxyl groups at the edges and surface of vermiculite to form HO• radicals. The presence of water on the vermiculite photocatalyst can initiate the indirect hole transfer process via the generation of HO• radicals, which are responsible for the methanol oxidation and the similar intermediates as of direct hole transfer [60].

Hydrogen yield on V–500 was slightly higher in comparison with the yield on Ver and V–700 while not dependent on a specific surface (Figure 8a), but dependency on the substitution of Fe(III) and Al in tetrahedra expressed as the Fe(III)/Al ratio in tetrahedra of Ver, V–500 and V–700 is 0.533, 0.521 and 0.553, respectively, (Table 2; Figure 7b).

The crystallite size is generally related to the photoactivity of materials [61,62]. The positive photocatalytic effect of α-Fe₂O₃ nanoparticles containing larger crystallites was observed on the most active photocatalyst Fe/V–700 composite with the highest surface area (63 m²/g) (Figure 8a). The largest crystallite size (34 nm) provided a larger number of active sites (Figure 8c) to produce the highest yield of hydrogen (1303 μmol/gcat.) from all the tested samples.

The Fe/V–500 composite of the smallest surface area (53 m²/g) (Figure 8a) and the nanoparticles’ smallest crystallite size (27 nm) (Figure 8c) produced the lowest yield of hydrogen (1094 μmol/gcat.) from all the tested samples.

The recombination rate of electrons and holes can be determined from PL spectra (Figure 6). A lower PL intensity indicates a lower recombination rate of photo generated charge carriers. The α-Fe₂O₃ nanoparticles in the Fe/V–500 composite significantly increased photoluminescence in contrast to the other samples indicating the highest electron-hole recombination rate resulting in the few electrons and holes that can participate in the redox reactions.

It is also known that the hematite surface in an aqueous solution can become hydroxylated and the surface chemistry indicated a different relationship between energy levels of the conduction band edge on the surface and hydrogen reduction [63]. Therefore, both direct and indirect transfer mechanisms can work competitively over the Fe/Ver, Fe/V–500 and Fe/V–700 composite surfaces on photocatalytic methanol decomposition.
Figure 8. The yields of hydrogen in the presence of investigated photocatalysts dependency on: (a) the surface area ($S_{\text{BET}}$), (b) Fe(III)/Al ratio in tetrahedral position of vermiculite samples and (c) the mean crystallite size $D_{\text{012}}$ of $\alpha$-$Fe_2O_3$ nanoparticles anchored in nanocomposite samples.

5. Conclusions

The aim of this study was a detailed characterization of the structural and textural properties of raw and heat-treated iron-containing vermiculite in comparison with their use as carriers in $\alpha$-$Fe_2O_3$/vermiculite composites to explain their photocatalytic activity under 254 nm light irradiation on the hydrogen production from methanol–water mixture.

The results of the XRF, XRD and IR analyses, physisorption and optical analysis allow a comparison of the vermiculite properties after calcination at temperatures of 500 and 700 °C as follows:

- Calcination decreased LOI = 12.3 mass% in Ver to 5.5 and 5.3 mass% in V–500 and V–700, respectively.
- Change in the hydration state of vermiculite was observed in the interlayer space of the V-Hb-Ph phases. The intensity of the basal peaks allows the representation of V-Hb-Ph (%) phases in Ver: V(30)-Hb(60)-Ph(10), in V–500: V(35)-Hb(37)-Ph(28) and in V–700: V(6)-Hb(0)-Ph(94) to be estimated.
- Infrared spectroscopy, confirmed the calcination effect on the presence of Al or Fe in addition to Si in the tetrahedral sheet.
- Calcination at 700 °C increased the pores volume with the formation of slit-like pores between the particles.
- Calcination supported increasing the amount of Fe(III) bound in the structure, which increases the direct band energy gap (Eg), while reducing the Fe$^{2+}$/Fe$^{3+}$ ratio in octahedra and PL intensity.
The photocatalytic degradation of methanol–water mixture takes place on the surface of vermiculite by the action of iron ions that can promote the generation of $e^−/H^+$ pairs, which play a role in the redox reaction. The similar proportion of one- and two-water layers in the interstratified structure of V–150 initiated higher photocatalytic activity due to the indirect hole transfer process via the generation of HO• radicals.

The photocatalytic activity of α-Fe$_2$O$_3$/vermiculite composites increased both with increasing specific surface area and increasing crystallite size of α-Fe$_2$O$_3$. The experimental results showed that all the investigated photocatalysts exhibited higher photoactivity compared to the commercial TiO$_2$ photocatalyst Evonik P25.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/min12050607/s1, Figure S1. Scheme (a) and real photo (b) of photocatalytic experiments. Figure S2. Repeated photocatalytic experiments in presence Fe/V-500 (a) and Fe/V-700 photocatalysts (b).

Author Contributions: Conceptualization, M.V. and K.K. (Kamila Kočič); methodology, K.K. (Kamila Kočič); J.M., L.M. and J.P.; validation, M.V., K.K. (Kamila Kočič), J.P. and L.M.; formal analysis, K.K. (Kateřina Klemencová) and B.T.B.; investigation, K.K. (Kateřina Klemencová) and B.T.B.; resources, K.K. (Kamila Kočič); data curation, K.K. (Katerina Klemencová); J.P., J.M. and L.M.; writing—original draft preparation, M.V., K.K. (Kamila Kočič), J.M. and L.M.; writing—review and editing, M.V. and K.K. (Kamila Kočič); visualization, M.V., K.K. (Kamila Kočič), J.M. and L.M.; supervision, M.V. and K.K. (Kamila Kočič); project administration, K.K. (Kamila Kočič); funding acquisition, K.K. (Kamila Kočič) and L.M. All authors have read and agreed to the published version of the manuscript.

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References
3. Foster, M.D. Interpretation of the composition of vermiculites and hydrobiotites. Clays Clay Miner. 1963, 10, 70–89. [CrossRef]


33. Lassoued, A.; Dkhil, B.; Gardi, A.; Ammar, S. Control of the shape and size of iron oxide (α-Fe2O3) nanoparticles synthesized through the chemical precipitation method. Results Phys. 2017, 7, 3007–3015. [CrossRef]


