Evaluation of New Properties of Calcined Magnetite Doped with Zinc and Lanthanum Nanoparticles Prepared in Oxygen and Nitrogen Atmospheres

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Research Article

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Abstract

The aim of the present research is the preparation of magnetite nanoparticles doped with Zn and La of the general formula ZnLa(x)Fe_{(3-x)}O_4 where x = 0.0, 0.1, 0.15, 0.2, 0.25 and 0.3; which were prepared via co-precipitation method, separated and calcined at 800°C under oxygen and nitrogen atmospheres. The stoichiometric ratios of the elements in the given general formula had been proved and confirmed by elemental analyses using XRF. Various techniques have been used to analyze the prepared nanoparticles. X-ray diffraction (XRD) confirmed the formation of one single phase of nanoparticles components with average crystalline size ranged from 6 to 9 nm at oxygen atmosphere and from 6 to 8 at nitrogen atmosphere. These size values had been calculated by Debye-Scherrer equation. The optical properties carried out by Cary Eclipse fluorescence spectrophotometer. The morphology of the prepared magnetite nanoparticles was studied using field emission scanning electron microscope (FE-SEM) and high-resolution transmission electron microscope (HRTEM). Optical properties were demonstrated by UV-visible-NIR spectrophotometer. The high absorbance values of Zn La(x)Fe_{(3-x)}O_4 were found to 80-97 and 80-99% in the visible wavelength range of 400-800 nm for the prepared samples in presence of O_2 and N_2 gas at room temperature, respectively. These proved the lower values of energy band gap in both conditions. The lowering of energy band gap of 1.89 eV for nanoparticles prepared in O_2 gas and 1.78 eV for prepared in N_2 gas in most samples may be attributed to incorporation of Zn cation. These results are confirmed by photo luminescence spectra (PL) measurements of the prepared samples. Magnetic properties of the given samples obtained from vibrating sample magnetometer (VSM); showed that most of these samples exhibited almost superparamagnetic behavior. These magnetic values (32.87 and 24.79 emu/g) infer super saturation magnetization of the given samples prepared in O_2 and N_2 gas at room temperature, respectively. Examination of the magnetic properties revealed decrease in saturation magnetization with increasing La ion concentrations incorporation up to x = 0.3. The evaluated magnetic and optical properties of the novel prepared ZnLa(x)Fe_{(3-x)}O_4 nano-materials revealed their possible use in different industrial fields like bio-applications, in electronic components, cosmetics, antibacterial agents, and in solar cells. The antibacterial activity study of the prepared NPs revealed them highly efficient against different kinds of bacteria that increase with increase of La % doped in their entity.

1. Introduction

Nanotechnology is exactly meant any technology that carried out in the nanoscale and has many applications in the real world, where the size of the atoms for the matter is about 1 to 100 nm as mentioned by I. Amato, et.al (1999) [1]. The peerless physical and chemical properties of nanomaterials have a profound effect by B. Bhushan (2015) [2] on commercial applications and for novel performance that benefits society. Nanomaterials have distinct physicochemical and biological properties. Particularly, their small size and diffusion abilities, shape, chemical composition, surface structure and charge, aggregation and agglomeration, and solubility can affect their interactions with biomolecules, cells, and organs. So, it can get into the human body in various ways, such as skin penetration, inhalation, or injection by Rahmandoust, M. et.al (2019) [3]. Magnetic nanoparticles present many applications in
biomedicine which have been evaluated by S. Laurent, et al (2008)[4] in vitro and in animal experiments. Compared to non-metallic nano formulations of similar sizes, the higher density Metallic nano particles are more readily taken up by cells, providing a benefit for cancer vaccination strategies as studied by Barnaby, et. al (2014)[5]. Iron oxide nanoparticles (IONPs) include many attributes of high magnetic saturation, stability, biocompatibility, and interactive functions at the surface. The surface of IONPs could be adjusted by organic materials or inorganic materials, such as polymers, biomolecules, silica, metals, and others that improve from its properties by Wu, et. al (2015)[6]. Laurent, et al (2011)[7] suggested bio-applications based on magnetic nanoparticles (NPs) have received considerable attention because NPs offer unique advantages over other materials. Magnetite is a spin-polarized, Fe$^{2+}$-Fe$^{3+}$ mixed-valence metal with a DC conductivity at 23 °C of about 0.1% that of Cu metal. Its crystals appear black and absorb throughout the UV-VIS-IR spectrum as indicating by Tang, et al (2003)[8]. A variety of synthetic methods such as co-precipitation are given by Rao, et. al (2018)[9], while thermal decomposition are given by Ibrahim, et. al (2018)[10], and hydrothermal and solvo thermal syntheses, sol–gel synthesis are given by de Mello, et. al (2019)[11]. The variety of magnetic materials can be formed from iron atom which has four unpaired electrons in its 3d orbital. The magnetic behavior of the materials can be classified into six types, depending on the particular response of the materials in the presence of a magnetic field. The materials are categorized by Spaldin, et. al (2010)[12]; into diamagnetic, paramagnetic, ferromagnetic, antiferromagnetic and superparamagnetic, on the basis of net magnetic moment of their atoms. Due to the advanced properties of nanomaterials, it was widely used by Ma, et. al (2010)[13] and Nel, et. al (2006)[14] in electronic components, cosmetics, antibacterial agents, and solar cells. During the last few decades there has been an increasing interest in utilizing nanoparticles and nanotechnology in cancer diagnosis and treatment by Barkalina, et. al (2014)[15] and Hofmann, et. al (2015)[16]. M. A. Zayed, et.al (2016)[17] prepared herein, novel ferrofluid nanocomposites xFe$_2$O$_3$/($1 - x$)Fe$_3$O$_4$ with different ratios of x ($0.0 \leq x \leq 1.0$) from magnetite and hematite nanoparticles (NPs) were prepared via a sonochemical method, for hyperthermia purposes. Various analytical and characterization techniques have been used to analyze and characterize the prepared ferrofluid nanocomposites. M. A. Zayed, et.al (2016)[18] were used various analytical techniques to analyze and characterize the ferrous/ferric constituents of the prepared ferrofluid nanocomposites xFe$_2$O$_3$/($1 - x$) Fe$_3$O$_4$ with different ratio x ($0.0 \leq x \leq 1.0$), from magnetite and hematite nanoparticles (NPS). These ferrofluid composites were prepared via the sonochemical method, for the first time, for hyperthermia purposes. P. M. Zélis et al. (2013)[19] obtained Cubic-like shaped Zn$_x$Fe$_{3-x}$O$_4$ particles with crystallite mean sizes D between 15 and 117 nm by co-precipitation. Amir, M., et al. (2015)[20] synthesized the single spinel phase nano-structured particles of FeB$_x$Fe$_{2-x}$O$_4$ (x = 0.1, 0.2, 0.3, 0.4 and 0.5) by the glycol-thermal method and the effect of B$^{3+}$ substitution on structural and dielectric properties of Fe$_3$O$_4$ were studied by L. Hu, et.al (2012)[21] investigated the effect of cobalt substitution on the size evolution, crystal structure, and magnetic properties of Fe$_3$O$_4$ nanoparticles. Mono-disperse Co$_x$Fe$_{3-x}$O$_4$ nanoparticles were prepared, using a one-step method, by the direct heating process of iron (III) and cobalt (II) acetylacetonates in a high-boiling-point inert organic solvent. The magnetic moments of magnetite nanoparticles were dramatically enhanced by J. Byrne, et.al (2014)[22] through the addition of zinc in a microbiologically driven synthesis
Super-paramagnetic iron oxide nanoparticles (SPIONs) had been approved by R. Hergt, et al (2006)[23] for clinical use due to their salient super-paramagnetic properties and low toxicity. Zn$^{2+}$ doped SPIONs possess significantly higher magnetic susceptibility than that of conventional SPIONs. Here we evaluated the potential toxicity of Zn$^{2+}$ doped Fe$_3$O$_4$ nanoparticles (Zn$_{0.4}$Fe$_{2.6}$O$_4$ NPs) in the liver and kidney of mice after repeated intragastric administration for 30 days. E. Fantechi et al. (2015)[24] found that magnetite nanoparticles (NPs) are widely carried out for biomedical applications, particularly as contrast agents for Magnetic Resonance Imaging and as heat mediators in Magnetic Fluid Hyperthermia. Y. Zhang, et al. (2002)[25] stated that, Superparamagnetic magnetite nanoparticles were surface-modified with poly (ethylene glycol) (PEG) and folic acid, respectively, to improve their intracellular uptake and ability to target specific cells. A progress report by Q. Pankhurst, et al. (2009)[26] shows that, a selection of scientific, technological and commercial advances in the biomedical applications of magnetic nanoparticles since 2003. G. R. J. Allard et al. (2015)[27] had been prepared the nanoparticles via the co-precipitation route and masked with 3-phosphonopropionic acid (3-PPA). M. Bloemen et al. (2014)[28] reported that, rare earth atoms exhibit several interesting properties, for example, large magnetic moments and luminescence. Introducing these atoms into a different matrix could lead to a material that showed multiple interesting effects. M. T. FARID, et al (2015)[29] studied the effect of neodymium (Nd) contents on the structural, electrical and dielectric properties of Nickel cobalt ferrites in the ferrite series Ni$_x$Co$_{1-x}$Nd$_y$Fe$_{2-y}$O$_4$ (x = 0.00, 0.20, 0.40, 0.60, 0.80, 1.0 and y = 0.00, 0.02, 0.04, 0.06, 0.08, 0.1). This series was synthesized by the sol-gel method. A series of Gd$^{3+}$ doped nano-crystalline Co-ferrites CoGd$_x$Fe$_{2-x}$O$_4$ (x = 0.0 to 0.1) had been prepared by E. Pervaiz, et al. (2013)[30] using sol-gel auto combustion technique. La- and Ni-doped Fe$_3$O$_4$ nanocomposite particles with a high saturation magnetization were prepared by MA, J et al. (2009)[31] via a homogeneous precipitation method in aqueous solutions. Fe-doped La$_{0.77}$Sr$_{0.23}$Mn$_{1-y}$Fe$_y$O$_3$ nanoparticles had been synthesized by Shlapa, et al. (2016) [32] using the sol-gel method. B. Balusamy, et al. (2012)[33] presented study to evaluated the bacterial toxicity of lanthanum oxide micron and nano-sized particles using shake flask method against gram-positive (Staphylococcus aureus) and gram-negative (Escherichia coli, Pseudomonas aeruginosa) bacteria. It was speculated that lanthanum oxide produced this effect by interacting with the gram-positive bacterial cell wall. Furthermore, lanthanum oxide bulk particles were found to enhance the pyocyanin pigment production in Pseudomonas aeruginosa.

In the present study, an composite has been made to synthesis La$^{3+}$ doped ZnFe2O4 nanoparticles by the two-step sol gel method. Here, we report the synthesis and characterization of ZnFe$_2$-xLaxO$_4$ through the sol–gel method. Besides, several experiments were performed in order to investigate the chracstric future of prepared composite that include XRD, XRF, TEM, FESEM and FTIR analysis. The antibacterial activity study of the prepared NPs against different kinds of bacteria studied with varying the La % doped in the ZnFe$_2$-xLaxO$_4$.

2. Experimental

2.1. Materials and reagents
The initial chemicals Tetra hydrated ferrous chloride FeCl$_2$·4H$_2$O (M.wt = 198.81 g mol$^{-1}$) was purchased from Research-Lab fine industries, Zinc acetate dihydrateZn(CH$_3$COO)$_2$·2H$_2$O (M.wt = 219.50 g) was purchased from Oxford lab chem, Hexahydrated ferric chloride FeCl$_3$·6H$_2$O (M.wt = 270.30 g mol$^{-1}$), Sodium hydroxide NaOH (M.wt =40 g mol$^{-1}$ ) and Lanthanum chloride LaCl$_3$·7H$_2$O (M.wt =371.37 g mol$^{-1}$ ) was purchased from Alpha chemika. Hydrochloric acid (HCl) (M.wt = 36.46 g mol$^{-1}$) with purity 30-34 %, Hydrofluoric acid (HF) was purchased from Adwic, Ethyl alcohol (ethanol) (M.wt = 46.07 g mol$^{-1}$) was purchased from International company for SUP and MED industries. The distilled water used in all preparations usually collected from all glass equipment.

### 2.2.1. Preparation of Zn-La doped magnetite nanoparticles

Aqueous solution of tetra hydrate ferrous chloride (0.4 M), zinc acetate dihydrate (0.2 M) and hexahydrate ferric chloride (0.2 M) was added in 2:1:1 ratio and the same volumes were taken from each solution (50 mL). After 10 mins of constant stirring, NaOH was added drop-wise to the mixture with stirring until the suspension becomes homogenous and the pH of the solution reached 11, that in presence of oxygen atmosphere (O$_2$) once and repeated again in the presence of nitrogen atmosphere, where (N$_2$) gas flow (with 99.9% of purity) for 20 minute. The particles washed with warmed distilled water several times [34]. Then the reaction is kept at 85-90°C for 4 hours. The zinc doped magnetite nanoparticles were magnetically collected with the formula ZnFe$_3$O$_4$, then the aqueous solutions of tetra hydrate ferrous chloride, zinc acetate dihydrate, hexahydrate ferric chloride were mixed by 50 mL of an aqueous solution of Lanthanum chloride 7.H$_2$O for different ratios (0.1, 0.2, and 0.3 M). Also, after 10 mins of constant stirring, NaOH was added drop-wise to the mixture with stirring until the suspension becomes homogenous and the pH of the solution reached 11. The reactions occur also in the presence of oxygen and the same in a nitrogen atmosphere. The filtered particles were washed with warmed distilled water several times. The reaction is kept at 85-90°C for 4 hours. Then Zn-La doped magnetite nanoparticles were magnetically collected; which has the proposed formula ZnLa$_{(x)}$Fe$_{(3-x)}$O$_4$ where, $x = 0.0, 0.1, 0.15, 0.2, 0.25$ and $0.3$. The dried collected precipitates obtained in nitrogen or oxygen atmospheres were ignited in special crucibles at 800°C for two hours. All prepared samples were kept in suitable plastic vials in dry atmosphere desiccators.

### 2.2.2. Analyses

The spectrophotometric measurements of Fe$^{2+}$ and Fe$^{3+}$ in nano-materials solutions using selective indicators were carried out using recording Spectrophotometer, UV-Vis range from 150-1100 nm model Unicam UV 300 and quartz cell of 1 cm optical length was used. Then total iron and zinc determination had been performed using atomic absorption spectrophotometer model AA.6300 SHIMADZU with an air-acetylene flame. X-ray diffraction analyses were carried out using (analytical-x’ Pertpro with Cuka1 target, $\lambda=1.5404$ Å, 45 kV, 40 mA, the Netherland) to identify the formation of the samples in pure single phase. Vibrating sample magnetometer (VSM) of (Model DMS 4HF) was used to measure the magnetic behavior of magnetic materials. The morphology of the various materials was disclosed using field
EM emission scanning electron microscopy (FE-SEM-QUANTA FEG 250) attached with EDX unit (Energy dispersive X-ray Analyses). HRTEM instrument of Zeiss Sigma 500 VP Analytical FE-SEM Accelerating voltage range 0.2 to 30 kV Variable pressure range: 2-133 Pa was used to characterize the particle size, shape, and grain size and to confirm the lattice parameter variation across the interfaces using selected area electron diffraction (SAED) pattern.

In the present work, the properties of the synthesized magnetic nanoparticles were investigated. The characteristics of the nanocomposites depend upon their crystal sizes, shapes and structures; which were characterized by advanced and sophisticated methods of analyses using different spectroscopic techniques. So, samples were completely analyzed by UV-Vis spectrophotometry and their solid state structures were investigated by XRD, EDX, HTEM and FESEM. The magnetic properties of the prepared materials were determined by means of the magnetic susceptibility and VSM. Thermal gravimetric analysis (TGA) was performed using a SDT Q 600 V20.9 build 20 (TA Instruments) thermo analyzer with a heating rate of 10 °C/min, using a O₂ at a flow rate of 50 mL/min. The photoluminescence spectra of the prepared nano compounds were performed using Cary Eclipse Absolute PL Quantum Fluorescence QM-40 Spectrometer and a Ge photodetector coupled to a data acquisition system composed of a microcomputer-controlled SR530 lock-in amplifier. The 350.7 nm excitation wavelength of a krypton ion laser (Coherent Innova) was used, with the laser output kept at 200 mW. The biological activities nanocomposites had been studied.

2.3a. Biological anti-bacterial studies

The antimicrobial activities of the test samples were determined by means of a modified Kirby-Bauer disc diffusion method by BrabuBalusamy et al [35] under standard conditions using Mueller-Hinton agar medium (tested for composition and pH), as described by NCCLS Sakiyan et al 2004[36]. Plates inoculated with filamentous fungi as Aspergillus flavus at 25°C for 48 hours; Gram (+) bacteria as Staphylococcus aureus, Bacillus subtilis; Gram (-) bacteria as Escherichia coli, Pseudomonas aeruginosa they were incubated at 35-37°C for 24-48 hours and yeast as Candida albicans incubated at 30°C for 24-48 hours and, then the diameters of the inhibition zones were measured in millimeter by Mahdavi et al 2013[37].

3. Results And Discussion

3.1 Particle Size distribution and Morphology Effects

3.1.1 HR-TEM Comparison

Lanthanum oxides have low magnetic characters similar to magnesium oxides in nature that coming after magnetite. Lanthanum has flat or cylindrical pipes crystal shape but it is diffused when mixed with magnetite less than 0.2 %[38]. Therefore, these shapes increased in size or its diameter when La% increased from 0.1 to 0.3 % and the La flat shapes become more common than the cylindrical common shape of magnetite and the most common shape of nano-material becomes more flat shape as shown in
Fig. 1. For zinc in nano-material goes around magnetite and it takes nano-cylindrical or hexagonal together with fluctuation of their crystal shapes. It is also noticed that, the variation of oxygen into nitrogen atmosphere change crystalline structures the doped Zn-magnetite with different ratios of La especially when ignited at 800°C. At which in nitrogen atmosphere it takes collective cylindrical Nano pipes groups under the effect of its homogeneity with magnetite[39]. In case of oxygen atmosphere (Fig.2) La appear with magnetite as big layers within which magnetite appears as spots; this means that La oxides crystals are homogenized with some magnetite oxides via oxygen – oxygen bonding. Generally heating of nano-materials to 800°C leads to homogeneity of their crystalline structures and consequently increasing in their magnetic properties gradually with increasing ratio of La.

3.2 FE-SEM analysis

The typical morphologies of the products in each La ratio (0.0-0.3 %) were observed by FE-SEM. The morphology of the Zn La(x)Fe_{(3-x)}O_4 NPs was evaluated by FE-SEM method. Fig.3 clearly indicates that the Fe_3O_4 spheres of ca. 30-50 nm in diameter are composed of many fine primary magnetite nanocrystals and exhibit a relatively smooth surface. These Fe_3O_4 spheres were further used as cores for the growth of La shells to obtain the La_xZnFe3O4 NPs core–shell nanostructures. Figure 1 depicts FE-SEM micrographs and reveals that nanostructures show spherical geometry, with diameters near 20 nm, which form irregular agglomerates with diameters from 50 to 70 nm. This agglomerate formation indicates that the surface energy of the nanoparticles is relatively strong. Similar results are obtained by[40]. Fig.3 clearly showed the uniform distribution of Fe, O and La elements in the structure of Zn La(x)Fe_{(3-x)}O_4 NPs. The iron and oxygen presence is related with iron oxide core. La was also verified in the scanning and its presence is consistent with the La nanolayer (shell). The FE-SEM results are similar that obtained by[41]. Fig.1, 2 demonstrated a micrograph performed by FE-SEM and the chemical distribution obtained to Zn La(x)Fe_{(3-x)}O_4 NPs in N_2 atmosphere. Fig.4 confirms the agglomerate formation with dimensions near 100 nm to 300 nm that depict the presence of iron, oxygen and La elements, respectively. Compared to the ZnFe_3O_4 core, the outside surfaces became coarse after the growth of La layers on the surface process, as shown in Fig.1 in 0.1, 0.2 and 0.3 % La. We can say from the TEM images (Fig.1,2) indicates that the La are successfully loaded onto the surfaces of ZnFe_3O_4 spheres clearly demonstrating the formation of core–shell nanostructures. The average size of ZnFe_3O_4 and Zn La(x)Fe_{(3-x)}O_4 NPs even at O_2 or N_2 atmosphere (Fig.3,4) are 30 and 50 nm, respectively. Fig. 2 indicates that core–shell heterostructure is the predominant morphology of the products[42], which agrees well with the SEM observation (Fig 4). Obviously, the thickness of the outer shell with light contrast is about 10 nm, while the diameter of the inner core with dark contrast is about 30-25 nm. The magnified TEM image further confirms that the shell is porous and composed of nanoparticles (Fig. 2).

3.3 Particle Size distribution
The hydrodynamic diameter provides information about inorganic core along with functionalized/coated material and the solvent layer attached to the particle as it moves under the influence of Brownian motion. Usually, the magnetic properties of nanomaterials can be varied by increase of surface to volume ratio of particles and variation of their size. Decreasing of particles size in nanometer scales led to creation of magnetic death layer on the particles surface due to deviation of magnetic moments orientation, formation of disordering configuration of moments that are caused to develop superparamagnetic behavior (below specific critical size)\[43\], making a difference in normal cationic distribution in crystalline structure, and finally variation of some surface properties like ability of water absorption on surface. Furthermore, decreasing of particle size and increasing of their surfaces are very useful for biological and medical applications. In contrast, when particles size is decreased less than 10 nanometers (for many of materials), the crystallinity of particles is also decreased, so the amount of saturation magnetization is dropped off\[44\].

Particle size was determined by atomic force microscopy. The average size derived from the height scan was below 40 nm. The distribution profile of sizes from the TEM height scan is shown in Fig.5a. A section of the AFM scan is included in the inset. The results from the TEM correspond to the distribution acquired from DLS (Fig.5b). The hydrodynamic diameter distribution of the Zn La(x)Fe_{3-x}O_4 NPs was of 36 nm and volume ratio of 16.9%. The hydrodynamic diameter by DLS is slightly larger than that determined by AFM, as expected. A similar study has been performed elsewhere and it was observed that DLS was limited for accurate particle sizing and mostly DLS provides higher values. Fig.5 indicates that the optimized co-precipitation method effectively produces a well-dispersed and narrower distribution of particle diameters\[45\]. The high stability of Zn La(x)Fe_{3-x}O_4 NPs is given by their very low electrostatic potential.

The hydrodynamic sizes of (N_2) Zn La(x)Fe_{3-x}O_4 NPs and (O_2) Zn La(x)Fe_{3-x}O_4 NPs were 1248 and 900 nm, respectively. The increase in the hydrodynamic size of functionalized Zn La(x)Fe_{3-x}O_4 NPs is due to an increase in La or LaOmoieties onto the surface of the magnetite Zn La(x)Fe_{3-x}O_4 NPs. This demonstrates a more hydrophilic nature of Zn La(x)Fe_{3-x}O_4 NPs than Fe_3O_4 NPs. The size obtained from DLS was large as compared with TEM because in DLS dispersed NPs moves through a liquid medium, and thin layer of electric dipole of the solvent adheres to its surface\[46\].

### 3.4 X-RD and XRF analysis

The crystalline structures of the Zn Fe_3O_4 before and after doping with La were investigated by XRD (Fig. 6a).

The framework peaks were weakened or vanished after La was added onto Zn Fe_3O_4. Obvious peaks of Fe_3O_4 located at 24°, 33, 35.6°, 54.1°, 57.3° and 62.4° were detected on sample ZnLa(x)Fe_{3-x}O_4, indicating the successful loading of Fe_3O_4. No peaks corresponding to lanthanum compound was observed on the XRD pattern of Zn Fe_3O_4 (Fig.6a&b). The samples of different La ratio (0.1-0.3 %) having the La dispersed onto structure of Zn La(x)Fe_{3-x}O_4 samples. The diffraction peaks of Zn La(x)Fe_{3-x}O_4.
x_jO_4 samples are highly crystalline, which is consistent with the standard pattern of Fe_2O_4 (JCPDS: 01–073-1964)[47]. Furthermore, the XRD patterns of Zn La(x)Fe_{3-x}O_4 samples differ significantly from the Zn Fe_3O_4 peaks. The major diffraction peaks at 2θ = 22.67°, 32.24°, 39.74°, 46.25°, 57.48°, and 67.57° could be indexed to (100), (110), (111), (200), (211), and (220) for lanthanum iron oxide [JCPDS: 01–075-0541][48]. In addition, a small portion of Fe_2O_3, with peaks at 2θ = 24.35°, 35.71°, 62.43°, and 72.68°, was obtained. This was thought to have originated from the transformation of unreacted iron species into Fe_2O_3 at 850 °C owing to tube furnace heating. The XRD pattern of Zn La(x)Fe_{3-x}O_4 samples exhibits the corresponding diffraction peaks with lower intensity owing to the size effect and domination of the Zn Fe_3O_4 heterojunction.

As the La ratio was increased from 1 and 0.3%, no obvious differences were observed, and the Zn Fe_3O_4 were delaminated at high La ratio. Fig. 2 shows the XRD patterns of the Zn La(x)Fe_{3-x}O_4 samples calcinations in N_2 atmosphere. The La_xZnFe_3O_4 samples calcinations in N2 atmosphere were not decomposed even at high ratio of La, which confirmed the stability of the material owing to the chemical interaction of the La particles with the Zn Fe_3O_4. Sharp diffraction peaks of La nanoparticles were obtained (Fig. 6b), which indicated the crystalline nature of the synthesized La_xZnFe_3O_4 samples. Broadening of the peaks indicates that the particles formed were in the nanometer scale. Particle size were estimated from half width of the observed diffraction peaks with (1 0 1), (1 1 1), (1 2 1), (1 1 0), (1 1 1), (2 3 0), (0 2 0), (2 0 2), (0 2 2), (2 2 0), (3 1 1), (0 0 4) Bragg reflection. The average particle size of the synthesized nanoparticles was 60 nm, which was obtained using the Debye–Scherrer equation[49].

The broad XRD diffraction peaks of ZnLa(x)Fe_{3-x}O_4 samples at 13.04°, 22.82°, and 29.54° are assigned to the lanthanum ZnFe_3O_4 structure (PDF No. 00-054-0212)[50]. The intensity of the Zn Fe_3O_4 signals decreases as a result of the growth of crystalline La_xZnFe_3O_4 nanocomposites as interlayer spacers on the surface of ZnFe_3O_4 nanoparticles. This result indicates the successful deposition of La on the ZnFe_3O_4 nanoparticles surface. Generally the sharp diffraction peaks as well as high intensity implied the good crystallinity of ZnLa(x)Fe_{3-x}O_4 that calcinated in O_2 or N_2 atmosphere (Fig.6a&b). Also, in the XRD pattern of ZnLa(x)Fe_{3-x}O_4 samples, the intensity of the above peaks decreased. This result might be attributed to the increases of La content of ZnLa(x)Fe_{3-x}O_4 samples which were dispersed onto the surface of ZnFe_3O_4 nanoparticles.

The chemical composition of the nanoparticles sample was examined by XRF analysis. The peak clearly indicated the presence of three essential constituents Zn, La, O and Fe with La occupying different percentage (Table. 1). The different La (0.1- 0.25 %) elemental ratios was here excellent agreement with the molecular composition of ZnLa(x)Fe_{3-x}O_4 nanoparticles samples. This clearly indicates that ZnLa(x)Fe_{3-x}O_4 nanoparticles have been successfully synthesized. The increase in La element percentage alongside peaks of iron and oxygen are shown in Table .1. The % of oxygen is relatively high due to the contribution of oxygen from LaO used in coating the surface of ZnFe_3O_4 nanoparticles. Further
functionalization with La element is shown in Table 1. The % of oxygen is dramatically increased due to contribution of oxygen from LaO[50].

**Table 1** XRF analyses of ZnLa(x)Fe(3-x)O4 nanoparticles.

| Samples          | molecular formula |
|------------------|-------------------|
| Key              | Fe    | Zn | La | O   |
| 0 % La in O₂ atmosphere | 1.67  | 1  | 0  | 3.67 |
| 0 % La in N₂ atmosphere | 1.18  | 1  | 0  | 3.18 |
| 0.1 % La in O₂ atmosphere | 0.98  | 1  | 0.085 | 2.98 |
| 0.1 % La in N₂ atmosphere | 1.48  | 1  | 0.108 | 3.48 |
| 0.2 % La in O₂ atmosphere | 1.4   | 1  | 1.7 | 3.5  |
| 0.2 % La in N₂ atmosphere | 1.07  | 1  | 0.09 | 3.07 |
| 0.3 % La in O₂ atmosphere | 1.02  | 1  | 0.23 | 3.02 |
| 0.3 % La in N₂ atmosphere | 1.9   | 1  | 0.26 | 4    |

**3.5 Thermal Analyses (TGA and DSC)**

Thermogravimetric analysis (TGA) was employed to study the thermal stability of ZnLa(x)Fe(3-x)O₄ magnetic NPs that calcinated in O₂ and N₂ atmosphere (Fig. 7a&b). As shown in the TG plot, the magnetic composite supported a 40.09 % weight loss at approximately 200-344 °C and 242-353°C for 0.3La calcinated in N₂ and O₂ atmosphere, respectively, which was ascribed to the which demonstrated the content of the LaO coating. For the magnetic composite, the second degradation step ranges between 443-672 °C and 434-672 °C for 0.3La calcinated in N₂ and O₂ atmosphere with weight loss of 1.1% and 2.76%, respectively. The degradation step indicate the decomposition of magnetite core of the ZnLa(x)Fe(3-x)O₄ magnetic NPs[50]. Differential scanning calorimetric analysis (DSC) curve of ZnLa(x)Fe(3-x)O₄ shows two endothermic main peaks at 168,165 and 684, 648 °C[51] for 0.3La calcinated in N₂ and O₂ atmosphere, which suggest the removal of La shell and decomposition of ZnLa(x)Fe(3-x)O₄ NPs respectively.

**3.6 Zeta optional of nanoparticles**

Zeta-potential measurements were further used to confirm the presence of La on the surface of ZnFe₃O₄ NPs. As shown in Fig. 8a&b, the zeta potential of (N₂) ZnFe₃O₄ NPs is increased with increasing La
percentagedue to presence of La metal not La oxide on the surface, while the zeta potential of \((O_2)\) ZnFe\(_3\)O\(_4\) NPs is decreases derived from the -ve charge of surface LaO as shown in Fig.8a. The \((O_2)\) La\(_{0.3}\)ZnFe\(_3\)O\(_4\) NPs shows a higher zeta potential of -13.2 mV derived from the -ve charge of LaO. Calcinations of Zn La\((x)\)Fe\((3-x)\)O\(_4\)NPs in N\(_2\) atmosphere (Fig.8b), lead to a increases of zeta potential, which is due to the fact that the La have +ve not LaO (-ve) charge crystal[50, 51]. So the biomedical activity of ZnFe\(_3\)O\(_4\) NPs calcinated in N\(_2\) atmosphere is more effective in killing bacteria with respect to O\(_2\) calcinated samples.

3.7 Photoluminescence spectroscopy analysis

The separation of between electron and hole is one of the important parameters influencing photocatalytic performance. In this context, photoluminescence spectroscopy was employed in order to evaluate the recombination rate for the photo generated electron–hole pairs for the ZnLa\((x)\)Fe\((3-x)\)O\(_4\) at N\(_2\) and O\(_2\) calcinations conditions (see Fig. 9).

The concentration of La ion in ZnLa\((x)\) Fe\((3-x)\)O\(_4\) synthesis is one of the most important factor that can affect the optical properties of ZnLa\((x)\)Fe\((3-x)\)O\(_4\). It is well illustrated that the excitation is generated by the photon absorption, which leads to band-edge and \(Zn^{2+}\) ion-related luminescence processes. The effect of concentration of \(La^{3+}\) ion as dopant was studied from 0.1% to 0.3% in the synthesis of ZnLa\((x)\)Fe\((3-x)\)O\(_4\). Fig. 9 shows the PL spectra of the ZnLa\((x)\)Fe\((3-x)\)O\(_4\) at N\(_2\) and O\(_2\) calcinations conditions.

The lowest intensity for PL spectrum of the ZnLa\((x)\)Fe\((3-x)\)O\(_4\) calcinated in O\(_2\) (Fig 9a) atmosphere reveals that recombination rate of the charges in this nanocomposite is the lowest in comparison with the Zn La\((x)\)Fe\((3-x)\)O\(_4\) calcinated in N\(_2\) atmosphere (Fig 9b). For both composites enhancement in intensity was observed with increasing La concentration (Fig. 9) and this enhancement was larger for C1 confirming more effective interaction with Fe\((3-x)\)O\(_4\). In fact, low intensity in PL spectrum of ZnLa\((x)\)Fe\((3-x)\)O\(_4\) calcinated in O\(_2\) atmosphere indicates that separation between the charge carriers is enhanced, leading to contribution of more electrons and holes in the oxidation and reduction reactions. The PL spectra appeared in the region between 700 and 800 nm, being identical for all the samples. In this way, the peaks around 733 nm arise from defect state luminescence. Besides, the bands at around 423 nm can also be related to recombination of free excitations. Therefore, the electron–hole separation efficiency can be improved, resulting in the increase in photo catalytic activity. The spectrum of ZnLa\((x)\)Fe\((3-x)\)O\(_4\) calcinated in O\(_2\) and N\(_2\) atmosphere have the peak at 423 nm which is originated from defects caused by missing of some Zn ion in ZnLa\((x)\)Fe\((3-x)\)O\(_4\) due to replacement of Zn with La in the crystal structure, and the peak at 733 nm which is originated from the \(4T1\rightarrow6A1\) transition in 3d5 electronic layer of Zn (II) ion. However the intensity of the peak at 423 nm of ZnLa\((x)\)Fe\((3-x)\)O\(_4\) calcinated in O\(_2\) atmosphere is lower than that calcinated in N\(_2\) atmosphere. These results indicate that ZnLa\((x)\)Fe\((3-x)\)O\(_4\) nanoparticles contain La ions and have similar PL properties with Zn Fe\((3-x)\)O\(_4\) nanoparticles in visible region, which can be used for labeling application in biomedicine.
3.8 Biological Activity

Fig. 10 show the results of descriptive indices for values of inhibition zone diameter and growth numbers of bacterial colonies at different weight percents G1 to G13 of calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs against different kind of bacteria. Where samples G1 to G13 are given in Table 2.

Table 2. Samples Key of different weight percent as G1 to G13 of calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs against different kind of bacteria.

| Key | La and Zn in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
|-----|--------------------------------------------------|
| G1  | La 0.0% in O\(_2\) atmosphere, in calcined ZnFe\(_3\)O\(_4\)NPs |
| G3  | La 0.0% in N\(_2\) atmosphere, in calcined ZnFe\(_3\)O\(_4\)NPs |
| G4  | La 0.1% in O\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G5  | La 0.3% in O\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G6  | La 0.10% in N\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G7  | La 0.30% in N\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G8  | La 0.15% in O\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G9  | La 0.15% in N\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G10 | La 0.20% in O\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G11 | La 0.20% in N\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G12 | La 0.25% in O\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |
| G13 | La 0.25% in N\(_2\) atmosphere in calcined ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs |

Based on the results presented in Fig. 10, the highest inhibition zone diameter was 25 mg/mL at 0.2% La doped magnetite O\(_2\); while the lower inhibition zone diameter of E. coli K88 was 15 mg/mL which obtained at the two nanoparticles concentration of from 0.1% to 0.3% La-doped magnetite (N\(_2\)). Diameter zone inhibition of E. coli bacteria was 17, 18, 19, 20, 22 and 23 mg/mL comparison to 100% of control at nanoparticle concentrations nanoparticle samples of La 0.1% in O\(_2\), La 0.2% in N\(_2\), La 0.3% in O\(_2\), and La 0.1% in O\(_2\) respectively Fig. 10. Results for E. coli in Fig. 10 was observed that number of E. coli K88 growth colonies values in different concentrations of ZnLa(x)Fe\(_{(3-x)}\)O\(_4\)NPs was significant different (p<0.05) at nanoparticles of G\(_{10}\), G\(_5\), G\(_4\), G\(_1\), G\(_{12}\), G\(_8\), G\(_6\) & G\(_3\) and G\(_9\). And had non-significant different (p<0.05) of bacterial growth no. colonies at nanoparticles of G\(_{11}\) and G\(_{13}\). The bacterial growth no. colonies in
ascending order was 37, 38, 41, 43, 46, 55, 65, 66, 68, 71 and 73\% at nanoparticles concentrations of G\textsubscript{10}, G\textsubscript{5}, G\textsubscript{4}, G\textsubscript{1}, G\textsubscript{12}, G\textsubscript{8}, G\textsubscript{7}, G\textsubscript{6} & G\textsubscript{3}, G\textsubscript{9}, G\textsubscript{11} and G\textsubscript{13} mg/mL.

Data in Fig. 11 showed that Laincreasing inhibitory effects and decreasing bacterial growth numbers of different kinds of positive and negative bacteria as the concentrations increased Lain ZnLa(x)Fe\textsubscript{(3-x)}O\textsubscript{4} NPs may distort and damage bacterial cell membrane, resulting effect on bacterial cellular contents and death of bacterial cells.

4. Conclusion

The present research involved preparation of magnetite nanoparticles doped with Zn and La of the general formula ZnLa(x)Fe\textsubscript{(3-x)}O\textsubscript{4} where x =0.0, 0.1, 0.15, 0.2, 0.25 and 0.3; which were prepared via co-precipitation method, separated and calcined at 800\textdegree C under oxygen and nitrogen atmospheres. The stoichiometric ratios of the elements in the given general formula had been proved and confirmed by elemental analyses using XRF. Various techniques have been used to analyze the prepared nanoparticles. X-ray diffraction (XRD) confirmed the formation of one single phase of nanoparticles components with average crystalline size. The morphology of the prepared magnetite nanoparticles was studied using field emission scanning electron microscope (FE-SEM) and high-resolution transmission electron microscope (HRTEM). Magnetic properties of the given samples obtained from vibrating sample magnetometer (VSM); showed that most of these samples exhibited almost superparamagnetic behavior. The evaluated magnetic and optical properties of the novel prepared Zn La\textsubscript{(x)}Fe\textsubscript{(3-x)}O\textsubscript{4} nano-materials revealed their possible use in different industrial fields like bio-applications, in electronic components, cosmetics, antibacterial agents, and in solar cells. The antibacterial activity study of the prepared NPs revealed them highly efficient against different kinds of bacteria that increase with increase of La \% doped in their entity.

Declarations

Credit author statement

1- Mohamed A. Zayed. - Suggested the idea, supervised the work, revised the manuscript and provided essential advice. Close supervision of all experimental work, characterizations and analyses, made amendments and corrections of the first draft and submit the manuscript for publication.

2 – Mahmoud Fathy Mubarak: Performed most of experimental work, characterizations and analyses of this work and wrote the first draft of the manuscript.

3- Gehad M. Abdel-maksoud. - Performed some of the experimental work, and preparation of magnetite nanoparticles.

Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Figures**

![Figure 1]
HRTEM of Fe3O4 nano-powders co-doped with Zn2+ ions and different La3+ ion molar ratio synthesized via co-precipitation route under N2 atmosphere and annealing at 800 oC for 2 h

Figure 2

HRTEM of Fe3O4 nano-powders co-doped with Zn2+ ions and different La3+ ion molar ratio synthesized via co-precipitation route under O2 atmosphere and annealing at 800 oC for 2 h
Figure 3

FESEM of Fe3O4 nano-powders co-doped with Zn2+ ions and different La3+ ion molar ratio synthesized via co-precipitation route under O2 atmosphere and annealing at 800 oC for 2 h
Figure 4

FESEM of Fe3O4 nano-powders co-doped with Zn2+ ions and different La3+ ion molar ratio synthesized via co-precipitation route under N2 atmosphere and annealing at 800 oC for 2 h
Figure 5

Size distribution of the magnetite nanoparticles at different calcinations atmospheres: (a) N2. (b) O2
Figure 6

XRD patterns of Fe3O4 nano-powders co-doped with Zn2+ ions and different La3+ ion molar ratio synthesized via co-precipitation route under A) O2 and B) N2 atmosphere and annealing at 800 oC for 2 h
Figure 7

TGA and DSC analysis of 0.3 Zn La(x)Fe(3-x)O4 calcinated in a) N2 and b) O2 atmosphere
Figure 8

Zeta potential of Zn-La(x)Fe(3-x)O4NPs calcinated in A) N2 and B) O2 atmosphere
Figure 9

The photoluminescence (PL) spectra of the ZnLa(x)Fe(3-x)O4 at N2 and O2 calcinations conditions.
Figure 10

The diameter of inhibition zone of the studied bacteria at different concentrations of calcined ZnLa(x)Fe(3-x)O4NPs at 800oC.

Figure 11

Antibacterial micro-dilution test of Zn La(x)Fe(3-x)O4 NPs