Crystalline native defects in ZnO analyzed by photoluminescence applying Maxwell-Boltzmann statistics in the visible region

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Zinc oxide (ZnO) is prepared by Chemical Bath Deposition (green chemistry) technique at \( \sim 80.0 \pm 2^\circ \) C temperature. This manuscript continues with previous research examining the Photoluminiscence spectra situated at UV-Vis-region. In this investigation, three different molar concentrations of the progenitor reagent containing the \( \text{Zn}^{2+} \) cation are chosen in order to find the optimal conditions for crystal growth and determine the influence of this parameter on crystal growth. The average grain size and lattice strain using Scherrer’s equation and Williamson-Hall \((W - H)\) method are discussed. The normalized absorbance situated at UV-region, electronic transitions located at \( \sim 239 \text{ nm} (\sim 5.18 \text{ eV}) \) and \( \sim 283 \text{ nm} (4.38 \text{ eV}) \) associated with quantum confinement are appreciated. We consider that the inorganic nanomaterial has native defects and the strain is caused by point defects (vacancies and interstices). According to emission bands situated at Vis-region by means of Photoluminescence spectroscopic technique, green (GE) and yellow emission (YE) bands are discussed, which are associated with native defects. The Photoluminescence dependence with the trap density and the surface recombination velocity in the light of the Maxwell-Boltzmann theoretical model (MBM) results associated with the electronic transitions related to the native defects situated at Vis-region are investigated. The trap density \((\text{cm}^{-3})\) of nanocrystals located at range \( \sim 8.9 - 9.9 \times 10^{13} \) An approximate theoretical-experimental the kinetic model is shown, considering that the coordination complex ion is a key parameter in the crystal growth of ZnO nanocrystals. The optimized geometry of \([\text{Zn} (\text{NH}_3)_4]^{2+}\) molecule was obtained with the DFT method using the functional of H-GGA B3LYP.

Keywords: Green chemical; kinetic model; quantum confinement; ploluminescence; Maxwell-Boltzmann statistics.

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1. Introduction

The electron configurations of Zinc (Zn) and Zn\(^{2+}\) cation are \([\text{Ar}]3d^{10}4s^2\) and \([\text{Ar}]3d^{10}4s^0\), respectively. Zinc oxide (ZnO) is a II – VI semiconductor with a large band gap energy located at range 3.18–3.40 eV [1,2], inorganic material that possesses structural properties and requires careful examination to find the correlation with its morphological and optical properties. ZnO it is a semiconductor with important physical and chemical properties for wide application as a base material in the design of optoelectronic devices and biochemical applications [3]. Therefore, is very versatile inorganic material with applications on the optoelectronics devices, based on thin solid films, but also with a significant number of unresolved issues. Detailed studies of the key parameters in the kinetic chemistry of crystal growth for the synthesis of ZnO nanocrystals through theoretical experimental models [4], associated in the evolution of each of the stages that govern the origin in the surface morphology of this inorganic nanomaterial, continues to be a strong challenge for the scientific community [5]. Classical theoretical models propose that for kinetic formalism two mechanisms at crystal growth are: nucleation and autocatalytic growth and was considered to be the only theory of nucleation un-
til formulated an approach of constant slow nucleation followed by autocatalytic crystal growth [6]. The chemical and physical process of the nucleation and growth of nanocrystals have been investigated through the LaMer burst nucleation and following Ostwald ripening to describe the change in morphological characteristics, native crystalline defects and the particles size [7]. Considering the overwhelming synthesis techniques, it is found that ZnO is a very sensitive material to the synthesis technique, which leads to presenting different morphological, structural and optical properties. Another serious problem is the incorporation of different pollutants of different sizes are directly and indirectly that are inevitably incorporated into the matrix of this inorganic material [8]. Morphology is decisive and drastically governed by crystal growth parameters, in particular ZnO presents different, impressive and wonderful morphologies associated directly with the growth parameters as well as with the applied technique [9, 10]. It is of utmost importance to understand the wide range of theoretical and experimental chemical and physical techniques in the synthesis of nanocrystals and in particular of ZnO. In this way, the limitations in the synthesis promote a greater knowledge and with more clarity, from there it is possible to predict the properties that this nanocrystals possesses. Nanocrystals form the bulk of ZnO research trends due to their ease of chemical and physical using a wide array of methods enabling the synthesis with many shapes and sizes; nanospheres, nanocactus, nanonanoplates, nanorods, nanodendrites, nanotubes, nanoshells, nanoneedles, nanoribbons, nanobelts, nanosheets, nanopyramids, nanotowers, nanocombs, nanorings, nanosprings, nanowires, nanocages, nanopencils, etc. [9]. The chemical phenomenon is explained considering that is often driven by both interparticle electric interactions and the influence of the ionic environment, and it here different from bulk due to electric interactions with the nanocrystals surface. The environment of nanocrystals can be sensitively influenced by local ions and/or ligands, with effects already occurring at low concentrations of progenitor reagents. The ionic electrical interaction of particles present in the aqueous solution, is an important parameter which is associated with the typical short and long range van der Waals electrical forces. On the other hand, the growth of nanocrystals by oriented attachment is frequently reported as a method supplementary to the classical growth by Ostwald ripening process [11]. The classical model and non-classical crystallization pathways, in which particles: ions, cations, coordination complex cations, ionic clusters, amorphous precursors, poorly crystalline nanoparticles and nanocrystals, among others, have been investigated [12]. It is found that the crystalline morphology is different and interesting (in some cases they are amorphous materials) and the origin of this morphological behavior, is currently and since the past in deep investigation by the scientific community [13]. We present in this manuscript, a chemical kinetic model that allows us to understand in a simple way the key stages in the chemical synthesis of the crystal growth of ZnO nanocrystals. The inorganic nanomaterial is synthesized in a thin solid film format applying the Chemical Bath Deposition (CBD) technique. We further note that in this study we used the experimental results previously published [14]. Using the spectra obtained through the Photoluminescence spectroscopic technique located at UV-Vis region, the emission bands are assigned to crystalline defects associated with vacancies, intertices, stacking faults, different pollutants, stoichiometry, etc., are examined to find the correlation of these emission signals with the native crystalline defects [15]. Applying the theoretical model of Maxwell-Boltzmann statistics (MBM), the native defects are associated with vacancies/interstices, which in a first approximation are considered as particles that present corpuscular behavior [16].

2. Experimental

2.1. Kinetic model in the chemical synthesis of ZnO nanocrystals

The Chemical Bath Deposition (CBD) technique, has been successfully applied at chemical synthesis of nanocrystals because it is versatile and relatively inexpensive, in which the chemical parameters of crystal growth are easily experimentally controlled [17]. Previous report, we systematically presented experimental details on the chemical synthesis to prepare rare earth and oxide-hydroxide transition metal and semiconductor nanocrystals [14, 15, 18].

The key chemical and physical parameters in CBD technique are systematically investigated with the aim of locating optimal conditions in small ranges of reaction temperature, dilute concentrations of precursor reagents, pH, etc. [14, 15, 17, 18, 19]. The chemical synthesis, is carried out in small temperature range located at 20 – 90 °C, kinetic mechanisms proposed by us is based on the indirect chemical synthesis of coordination complex \( \text{[M} (\text{NH}_3)_4]^{2+} \) ion (\( M = \text{Pb}^{2+}, \text{Cu}^{2+}, \text{Ni}^{2+}, \text{Zn}^{2+}, \text{Bi}^{3+}, \text{Er}^{3+} \), etc.). The kinetic model we consider in crystal growth, present an interesting structural phenomenon associated with the molecular distortion Jahn-Teller effect (JTE), which favors the synthesis of nanocrystal [20]. We propose the following key stages (i) creation of [Zn (NH\(_3\))\(_4\)]\(^{2+}\) ion in aqueous solution, at alkaline medium in which Zn(OH)\(_2\)(aq) nanocolloids are generated. The step is simple by direct addition of salt containing [Zn(OH)\(_6\))]\(^{2+}\) cation and direct combination of KOH, which generates the alkaline medium (pH \(\sim\) 8.3), (ii) the white suspension turns colorless when adding NH\(_4\)NO\(_3\) solution, which contributes NH\(_3\) molecules and the indirect creation of the coordination complex [Zn (NH\(_3\))\(_4\)]\(^{2+}\) cation, under these reaction conditions due to incubation period of the nanocolloids and the slow release of NH\(_3\) molecule with the formation of ZnO nanocrystals in the final stage crystal growth. Next, we present the chemical reactions associated with the energetic changes of Gibbs free energy \( (\Delta \text{G}^0) \), according to the \( \Delta \text{G}^0 = -n \varepsilon \tau \) equation, where: \( \Delta \text{G}^0 \) represents the free energy of Gibbs, \( \varepsilon \)\(^0\) half-cell potential [21, 22], \( n \) the number of equivalents (electron exchange) and \( \tau \) is a numerical con-
shows the spatial distribution of the spontaneous precipitation of ZnO nanocrystals, we propose the following stages:

\[ \text{Zn}^{2+} + 2e^- \rightarrow \text{Zn} \]
\[ \Delta G^0 = +147.20 \text{ kJ}, \quad (1) \]

\[ \text{ZnO}_2^{2-} + 2\text{H}_2\text{O} + 2e^- \rightarrow \text{Zn} + 4\text{OH}^- \]
\[ \Delta G^0 = +234.49 \text{ kJ}, \quad (2) \]

\[ \text{ZnOH} + \text{H}^+ + 2e^- \rightarrow \text{Zn} + \text{H}_2\text{O} \]
\[ \Delta G^0 = +95.92 \text{ kJ}. \quad (3) \]

At the chemical synthesis, it is proposed the coordination complex \([\text{Zn(OH)}_6]^{2+}\) cation, have undergone a fast transformation into \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation, which is supported by chemical mechanisms that could explain the transformation: dissolution, in situ crystallization and/or transformation of solid–solid phase of ZnO product. On the other hand, the \(\text{ZnO}_2^{2-}\) ion, is dissolved in the aqueous solution to form the coordination complex \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation at basic solution (pH ~ 8.3), where these growth units dehydrate when they are incorporated in the crystal growth [24]. Generally, chemical balances are presented in a compact and straightforward way and it is possible to apply Hess’s law, which allows us to carry out in a convenient and adequate way; addition and/or subtraction, change of direction of chemical reactions, etc; [25].

According to Hess’s law, we proceed to arrive at the final product

\[ \text{ZnOH}^+ + \text{H}^+ + 4\text{OH}^- \rightarrow \text{Zn(OH)}_4^{2-} + \text{H}_2\text{O} \]
\[ \Delta G^0 = -288.15 \text{ kJ}, \quad (4) \]

\[ \text{ZnO} + \text{H}_2\text{O} + 2e^- \rightarrow \text{Zn} + 2\text{OH}^- \]
\[ \Delta G^0 = +243.18 \text{ kJ}, \quad (5) \]

\[ \text{Zn}^{2+} + 4\text{OH}^- \rightarrow \text{ZnO}_2^{2-} + 2\text{H}_2\text{O} \]
\[ \Delta G^0 = -87.29 \text{ kJ}. \quad (6) \]

The intermediate complex of coordination \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation, releases slowly NH3 molecules, presenting the following chemical equilibria:

\[ [\text{Zn} (\text{NH}_3)_4]^{2+} \Leftrightarrow \text{Zn}^{2+} + 4\text{NH}_3 \quad \Delta G^0 = +53.30 \text{ kJ}. \quad (7) \]

From Eq. (7) it is observed that \(\Delta G^0 > 0\), the slow release of \(\text{Zn}^{2+}\) ion is favored, this is a key parameter to control the spontaneous precipitation of ZnO product. Figure 1 shows the spatial distribution of NH3 ligands (L) of the complex intermediate \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation. The optimized geometry of \([\text{Zn} (\text{NH}_3)_4]^{2+}\) molecule was obtained with the DFT method using the functional of H-GGA B3LYP [26]. The basis set B2 (also called WH3f) is specifically optimized for the Zn atom by Amin et al. [27]. The molecular configuration of \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation with greater relative thermodynamic stability according to our theoretical calculations, corresponds to tetrahedral geometry. The \(\text{Zn}^{2+}\) cation is located at central part at tetrahedron, in which \(L = \text{NH}_3\), are situated at the vertices with a coordinated covalent bond according to the valence bond theory. It is important to mention that for the case of the \(\text{Cu}^{2+}\) complex cation, in which the most thermodynamically stable molecular configuration corresponds to the planar-square, which undergoes molecular distortion due to the JTE effect [20]. We propose the formation of \([\text{Zn} (\text{NH}_3)_4]^{2+}\) cation (intermediately generated coordination complex), according to the chemical equilibrium in Eq. (7) in which the NH3 molecules are released slowly, the \(\text{Zn}^{2+}\) cation also combines slowly to form O-Zn-O chemical bonds, which are thermodynamically stable to finally reach the ZnO product. The stability of metal complex increases with decrease in size of the metal cations. Ionic radii of \(\text{Zn}^{2+} \sim 0.60 \text{ Å} \) cation and \(\text{CO}_2^{2-} \sim 1.62 \text{ Å}, \text{S}_2^{2-} \sim 1.84 \text{ Å}\) ions [28]. These findings highlight the impact of the presence of \(\text{H}_2\text{O}\) on the interaction of \(\text{Zn}^{2+}\) cation and NH3 molecules, in aqueous ammonia solution, which is an limitation with the hard/soft Lewis acids and bases classification of \(\text{H}_2\text{O}\) being a hard L associated at \(\text{NH}_3\), which is attributed as being soft. Experimental and theoretical studies.
have been widely employed to explore the chemical properties of solvated ionic species, and in particular to study molecular structure, ion-solvent distance, coordination number and L exchange, conformational changes of the solvation complex [29].

On the other hand, the equilibrium constant \( k_{eq} \) of \([\text{Zn} (\text{NH}_3)_4]^{2+}\) ion, provides us with semi-empirical information regarding the thermodynamic stability of cation and is related to the concentrations of the ligands (L). In this particular case are: \( L = \text{NH}_3 \) is molecule monodentate according to the following expression [30]

\[
k_{eq} = \frac{[\text{Zn} (\text{NH}_3)_4]^{2+}}{[\text{Zn}^{2+}] [\text{NH}_3]^4} \sim 4.6 \times 10^{17}. \tag{8}
\]

As mentioned above, in our conditions the following chemical equilibria are proposed:

\[
[Zn (NH_3)_4]^{2+} + 4OH^- \rightleftharpoons ZnO_2^{2-} + 2H_2O + 4NH_3
\]

\[
\Delta G^0 = -28.97 \text{ kJ}. \tag{9}
\]

Adding the chemical (1) and (5) equilibria

\[
Zn^{2+} + 2OH^- \rightleftharpoons ZnO + H_2 \quad \Delta G^0 = -95.98 \text{ kJ}. \tag{10}
\]

The precipitated white powder corresponds in a first stage, which synthesis chemical was carried out with Zn\(^{2+}\) cation (in aqueous solution), obtaining hydrozincite \([\text{Zn}_5 (\text{CO}_3)_2 (\text{OH})_6]\), which stoichiometrically blends \([\text{Zn}(\text{OH})_6]^{2+}\) (hexahydrated cation) and \(\text{ZnCO}_3\) [31]. The process chemical might incorporate many chain reactions leading \(\text{ZnO}\) formation in solid state

\[
[Zn (NH_3)_4]^{2+} + \text{ZnO}_2^{2-} \rightleftharpoons 2\text{ZnO}_s + 4\text{NH}_3 + 2e^-
\]

\[
\Delta G^0 = +7.13 \text{ kJ}. \tag{11}
\]

Hydrolysis of thiourea at room temperature (RT), provides \(\text{CO}_3^{2-}\) and \(\text{HS}^-\) ion, which are identified by FTIR studies [32,33]

\[
\text{SC} (\text{NH}_2)_2 + 3\text{H}_2\text{O} \rightleftharpoons \text{CO}_3^{2-} + \text{HS}^- + \text{NH}_3. \tag{12}
\]

On the other hand, according to the experimental results of XRD previously reported in the structural investigation of \(\text{ZnO}\) product [14], the crystalline \(\text{Zn}_5 (\text{CO}_3)_2 (\text{OH})_6\) product is identified in quantities not quantified by XRD technique. The chemical equilibria presented by the formation of the pollutant is shown below

\[
[Zn (NH_3)_4]^{2+} + 5\text{ZnO}_2^{2-} + \text{CO}_3^{2-} + 5\text{OH}^- \rightleftharpoons \text{ZnO} + 4\text{NH}_3 + 2e^- + \text{Zn}_5 (\text{CO}_3)_2 (\text{OH})_6. \tag{13}
\]

The synthesis of the green chemistry CBD of inorganic materials has been previously reported [14]. However, we will briefly point out some of the strategic steps that we applied in the chemical synthesis of samples.

The CBD technique is reduced to preparing three solutions at \(\text{MC} = \sim 1.0 \text{ M, } \sim 2.0 \text{ M and } \sim 3.0 \text{ M of } \text{Zn(NO}_3)_2 \cdot 5\text{H}_2\text{O} \text{ salt provided by } \text{Zn}^{2+} \text{ cation. All precursor reagents were analytically pure and used as received without any further purification were of } \sim 99.9\% \text{ purity (Baker). The following progenitor reagents are held constant: } \text{KOH} (0.5 \text{ M}, \text{NH}_4\text{NO}_3 (1.5 \text{ M) and SC} (\text{NH}_2)_2). \text{ The time of deposition is } \sim 1.0 \text{ h while maintaining the } \text{T} \sim 80 \pm 2^\circ \text{C temperature. The glass substrates were previously immersed in } \text{HCl}/\text{H}_2\text{O for 2.0 h; after which they were rinsed in distilled water and dried in a clean hot-air flow. The substrate cleaning was carried out by immersing them into an acid-chromium mixture } \text{K}_2\text{Cr}_2\text{O}_7/\text{HCl}/\text{H}_2\text{O for 24.0 h and then rinsing them in deionized water. The thin solid films were labeled: } \text{ZnO} - A, \text{ZnO} - \text{B and ZnO} - \text{C samples, according to } \text{MC} \sim 1.0 \text{ M, } \sim 2.0 \text{ M and } \sim 3.0 \text{ M, respectively. All the films are white and the porosity decreases with the increase at } \text{MC. The samples are repeatedly washed with deionized water (} \sim 1.8 \text{ M} \text{O}) \text{ to remove impurities that are generally suspended in the solution and these adhere weakly on the crystals. The Photoluminescence (PL) spectra was characterized by a main peak, under optical excitation provided by an Ar+ laser beam, with a pump power of 10, 350 nm as excitation, using a Science-Tech model 9040 apparatus.}

3. Discussion

In this manuscript, we continue to investigate the systematically obtained experimental results of \(\text{ZnO}\) synthesized by the CBD technique, to investigate the correlation of morphological, structural and optical properties with the aim of systematically applying the Maxwell-Boltzmann theoretical model (MBM), considering how first approximation that the different crystalline defects (vacancies and intertices) behave as free particles.

The nanocrystals labeled by the \(\text{ZnO} - A, \text{ZnO} - \text{B and } \text{ZnO} - \text{C} \text{ symbology, were investigated applying systematically SEM, XRD, optical absorption and Photoluminescence (PL) technique previously reported [14]. The percentage composition expressed in } \% \text{ of atoms (stoichiometry), is quantified in } \text{ZnO} - A : \text{ Zn } \sim 67.65, \text{ O } \sim 32.35, \text{ ZnO} - \text{B : } \text{ Zn } \sim 44.18, \text{ O } \sim 76.38 \text{ and } \text{ZnO} - \text{C : Zn } \sim 49.70, \text{ O } \sim 50.30 \text{ samples. Morphological images were obtained by Scanning Electron Microscopy (SEM) of } \text{ZnO-A, ZnO-B and ZnO-C thin solid films. Here, we analyze the different morphologies and their correlation with the optical properties that these materials present. The } \text{ZnO-A sample presents spongy morphology with cavities of different sizes. ZnO-B film has well defined crystals of long micro-javelins and } \text{ZnO-C sample, morphological images of tetrapod is presented. It is shown that the } \text{MC} \text{ of } \text{Zn(NO}_3)_2 \text{ in our experimental conditions of precursor reactants, temperature, stirring are constant, with different } \text{MC therefore, it is a key parameter that significantly modifies the stoichiometry and morphology of thin solid films. According to (JCPDS card No. 36-1451) standards the hexagonal phase is identified.}

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However, some structural differences are observed and compared with each other. Zn-A and Zn-B samples, broadening of crystalline planes with respect to Zn – C sample, is observed. The characteristic thing in Zn – A and Zn – B samples, is the emergence of a crystalline plane at angular position located at $2\theta \sim 32.85^\circ$, corresponds to Zinc-hydroxyl-carbonate $\text{Zn}_5(\text{CO}_3)_2(\text{OH})_6$ (Zinc hydrocincite) according to (No. 04-013-7572) standards. The MC increase in ZnO – A and ZnO – B samples, produces a decrease at relative intensity of the (002) reflection plane, reaching a relative maximum at Zn-C sample. A reflection associated with the crystalline plane of $\text{Zn}_5(\text{CO}_3)_2(\text{OH})_6$ material, is observed. Structural behavior, with increase of $\sim 3.0$M of the salt containing the $\text{Zn}^{2+}$ cation, favors the thermodynamic stability of the nanocoloids and the crystalline growth of the free $\text{ZnO}_{(s)}$ of Zinc-hydroxyl-carbonate pollutant. The grain size (GS) presented the following numerical values: Zn – A $\sim 33.4$ nm, Zn – B $\sim 36.5$ nm, and Zn – C $\sim 34.2$ nm, respectively. The addition of thiourea, the mixture could result in strong electrostatic interaction with the polar surfaces of growing ZnO nanocrystals thus resulting in decreasing the energy of polar surfaces and hence slowing down the growth rate of the polar planes being the exposed basal surface of the nanocrystals which grows slowly with well developed facets, the intrinsically anisotropic growth of ZnO along the (002) orientation is substantially suppressed and crystal growth then proceeds sideways, which results in the formation of Zn-A and Zn-B spongy morphology with cavities of different sizes [35]. In crystalline growth they are identified as pollutants and generally adhere to the surface and in general to the volume of nanocrystals in quantities not yet quantified. Structural parameter called strain ($\varepsilon$) is evaluated by applying the experimental of XRD results. Williamson-Hall method was applied. The grain strain is related to the measured FWHM ($\beta$) of the diffraction plane by following $\beta \cos \theta = \lambda/GS + \varepsilon \lambda \sin \theta$ equation, $\lambda$ and $\theta$ are wavelength of the X-ray source and Bragg’s angle, respectively [36]. The plot of $\varepsilon$ vs. $\sin \theta$ for ZnO – A, ZnO – B and ZnO – C films, the slope of the plot gave the amount of residual $\varepsilon$. The lattice constant $c$ of ZnO – C films is larger than the ZnO – A and ZnO – B and all the films exhibit tensile $\varepsilon$. ZnO – A (002) plane position was shifted to higher angle ($2\theta$) as well as the increase in the intensity of the crystalline planes with increasing at MC. The variation of $c$ with increasing $MC$ suggests that the compressive stress is generated during deposition due to MC which according to the low MC induces structural defects; thickness, roughness, grain boundaries, and stacking faults are parameters associated with stress [37]. The increasing MC increases the atomic mobility and reduces the structural defects, and thus a relaxation of ZnO – C film. Thus, the extrinsic stress will not be present and the total estimated stress values must be to dominantly intrinsic. Also, the decrease in residual stress might be brought about by release of defects, such as interstitials.

Optical propertie were investigated through the refractive index $n(\lambda)$ from the reflection coefficient and the optical extinction data, following the Fresnel equation [39]. Figure 3 displays the real of $n(\lambda)$ spectrum of ZnO – A, ZnO – B and ZnO – C nanocrystals. The refractive was calculated using Eq. (5)

$$n = \frac{1 + R}{1 - R} + \sqrt{\frac{4R}{(1 - R^2)}} - k^2. \quad (14)$$

To determine the values of extinction coefficient ($k$) we used the following equation:

$$k = \frac{\alpha \lambda}{4k}. \quad (15)$$

The data were recorded in the wavelength range $\sim 200 – 400$ nm absorption, and then the optical data were used to calculate the $n(\lambda)$ using an angle of incidence of $30^\circ$.

The electronic transitions of greater relative intensity are seen at ZnO-A and ZnO-B nanocrystals. A plausible explanation is proposed; ZnO – A and ZnO – B samples has a high concentration of native defects. The observed absorption bands can, in principle, be justified by the optical and structural phenomena and electronic transitions locted at UVVis, native defects as vacancies, grain boundaries, stacking faults, stoichiometric and quantum confinement effect [40]. It has been ascribed to a slight deviation of the local symmetry of the Zn$^{2+}$ ion induced by the $\text{CO}_3^{2-}$ and $\text{OH}^-$ ions modifiers for O$^{2-}$ ion, such differences in the optical properties arising.

![Normalized absorbance spectra of ZnO-A, ZnO-B, and ZnO-C thin solid films][1]

**Figure 2.** Normalized absorbance spectra of of ZnO-A, ZnO-B and ZnO-C thin solid films $[\text{Zn(NH}_3)_4]^{2+}$ ion.

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from the ZnO – A and ZnO – B electronic transition. The absorption bands intensity was found to increase with the concentration of the –OH. The presence of –OH groups induces a greater relative intensity in the absorption bands and the values agree with those already reported [41]. It can be related to the decrease in GS causes more atoms to be closer to the surface and thereby increasing the rate of trapping of photogenerated holes (h+) at the surface, which in turn enhances the emission intensity. The n(λ) exhibits anomalous dispersion in the near-UV region, associated with the steep onset of absorption. In the Vis-region, the values are comparable for all three samples, i.e. they range situated at range ~ 2.2 – 2.3 eV. These are compatible with earlier results for pure ZnO film [42,43]. We can observe, that the n(λ) for all samples increase when the λ increase. This phenomenon is attributed to light scattering and to the increase of absorbance.

Urbach’s energy (E_u) has a strong experimental theoretical form for the study of crystal lattice defects, these are generally associated with crystalline disorder (which can be ionic, atomic and/or molecular). Analyzing the structural behavior, it has been found that for a wide variety of inorganic solid crystals, the energies quantify the static, structural disorder causing localized exponential-tail states, and dynamic disorder from electron-phonon (e−–h+) scattering. The theoretical model E_u is powerful and allows us to examine the degree of of optical absorption generated by crystal defects. It has been found that that sub-gap absorption due to singlet excitons is universally dominated by thermal broadening at low photon energies and the associated Urbach energy equals the thermal energy, regardless of static disorder [44]. The studies of the E_u parameters allow us to extract optical information associated with the crystalline disorder, stacking faults, stoichiometry, and disordered crystalline grain boundaries, GS, orientation of crystalline planes, among other parameters related to crystal growth, resulting in the formation of confined situations into the E_g energy. The absorption spectra, α(v) are used to define the E_u corresponding to the electronic intratransitions between the extended states of the valence band (VB) and the localized states of the conduction band (CB). In the low energy photon (hv) regime, it is assumed that the spectral dependence of absorption edge follows the empirical E_u given by α(v) = α_0 exp(hv/E_u) equation [45], where α_0 is a constant, E_u denotes an energy which is constant or weakly dependent on temperature and is often interpreted as the width of the tail of localized states in the E_g. Figure 4 displays the ln(α) as a function of energy (eV) spectra of ZnO-A, ZnO-B and ZnO-C films.

It has been recently reported that the E_u does not correlate with the topological disorder in ZnO nanocrystals [46]. The absorption edge fluctuations are linked to the variations of the E_g, the width of Urbach tails. From these spectra, it can be seen that the optical absorption edge shifted to higher λ(μm). This was attributed to decrease in GS, preferred orientation and stoichiometry. The defects and disorders may lead to forming a delocalized state near the band level and the enhancement of the E_u value. The structural motif for the localization of the mid-gap states is a crystalline-like atomic environment within the amorphous network. Therefore, these mid-gap states trap an extra electron spontaneously, creating deep traps in the E_g [47]. Urbach energy was calculated as E_u ~ 11.2 eV for Zn – A, E_u ~ 11.3 eV and Zn – C ~ 11.4 eV, respectively. In our opinion, E_u increase is associated with increase in native crystalline defects and impurities, resulting in a nanomaterial with major disorder and major density of localized states. Research related to the
optical behavior of the E, originates from E, fluctuations attributable to chemical composition variations at nanoscale, while electrostatic fluctuations contribute significantly.

The photoluminescence (PL) spectroscopy technique, is applied in research of the band edge electronic transition levels of a material have been performed by many research groups [48]. We present a previous study applying the technique of PL located at UV-Vis region with the aim of investigating the green (GE) and yellow (YE) emission bands, respectively [14]. Generally, native defects located at UV-Vis region by means of PL technique and recorded at range \( \sim 1.5 - 3.0 \text{ eV} \) was carried out by deconvolution, is observed that different emission signals are overlapping with other energy levels and PL measurements indicate GE band at \( \sim 495 \text{ nm (} \sim 2.50 \text{ eV)} \) [49]. Shallow acceptor levels are created at \( \sim 0.3 - 0.4 \text{ eV} \) above the top of the valence band (VB) due to zinc vacancy (V\(_{\text{Zn}}\)) and oxygen interstitial (O\(_i\)), respectively [50].

Figure 5 shows the theoretical and experimental PL spectra of (a) Zn-A, (b) Zn-B and (c) Zn-C thin solid film. The distribution of particles is reflected by the experimental PL spectra of (a) Zn-A, (b) Zn-B and (c) Zn-C nanocrystals. The experimental emission band recorded here, they are asymmetric implies different signals overlapping each other. We performed an analysis of nanocrystal emission bands located at UV-Vis region, applying the statistical theoretical model based on Maxwell-Boltzmann (MBM) distributions, in this context, several detailed studies have been carried out [15, 16, 51]. The emission band applying the MBM is indicated by a sky blue stripe. The emission band at \( \sim 2.25 \text{ eV} \) may be related to deep level defects [52]. The emission band situated at \( \sim 607 \text{ nm (} \sim 2.04 \text{ eV)} \), is known YE band in ZnO [53]. PL results from the recombination of a photogenerated hole \((h^+)\) with the electron \((e^-)\) occupying oxygen vacancies site. Oxygen, in general, exhibits three types of charge states of oxygen vacancies. GE band has been proved to be an outcome of singly oxygen vacancies in the sequence of \( (V_{O^0}^0, V_O^+ \text{ and } V_O^{2+}) \) located below the bottom of the conduction band (CBD). The emission band is produced by the recombination of delocalized electrons close to the CB with deeply trapped holes in the \( O^+ \) point defect level. The deep level emission is known to be related to intrinsic \((V_{\text{Zn}}, V_O, \text{ or } Zn_i)\) and extrinsic (acceptor) point crystal defects. Vis-emission bands is also reported from ZnO nanocrystals owing to various crystal defects. Zinc interstitial \((Zn_i)\) produces a shallow donor level at \( \sim 0.5 \text{ eV} \) below the bottom of CB. In our cases, we observed PL emission in the Vis-region which indicates that the emission band in this case is governed by the crystal defects related deep level emission over the band edge UV-emission [54]. The Vis-emission band at \( \sim 485 \text{ nm (} \sim 2.55 \text{ eV)} \) arises due to electronic transition between interstitial zinc \((Zn_i)\) and zinc vacancy \((V_{Zn})\) level arises due to transition between \( Zn_i \) and \( V_{Zn} \) level [50]. The emission band \( \sim 527 \text{ nm (} \sim 2.3 \text{ eV)} \) can be related to singly ionized oxygen vacancy \((V_O)\). It is also reported that the defect-related PL emission dominates for the nanorods of high-aspect ratio as compared to that of bulk because of more number of surface states and incorporation of pollutant. According to these experimental results we observed that our material presents contribution of defects associated with GE and YE bands. This is confirmed by the spectrum of optical absorption and morphological images obtained by SEM. The emission band associated at the amount of interstitial oxygen \((O^+)_i\), and according to the studies obtained by SEM technique, it is appreciated that samples ZnA and Zn-B present a lower amount of oxygen with respect to \( Zn - C \), which is almost stoichiometric \((Zn/O_1)\). Therefore, the emission bands must show a significant change as seen here. GE and YE bands is observed at \( \sim 600 - 450 \text{ nm (} \sim 2.06 - 2.75 \text{ eV)} \) which is attributed to the amount of non-stoichiometry, producing intrinsic defects in the material that may originate from the \( V_{Zn} \) and anti-site defect in the ZnO. However, \( V_O \) would be the dominant intrinsic defect under both Zn-rich and O-rich conditions and it is a deep double donor. The red emission (RE) band has been related to the
Zn$_i$ which causes a lattice disorder along c-axis, introducing shallow donor levels [4]. It seems that the band could originate from a donor acceptor pair transition. Chemical pollutants may also cause a surrounding of structural defects by distorting the lattice as well as the surface of a nanocrystal. The trap-states may be caused by the thermal fluctuations of the molecules, which are expected to result in shallow trap states within the $E_g$ [55]. The UV emission for the bulk ZnO is detectable at very low temperature but for the tetrapod-shaped ZnO (Zn-A sample) it was detectable even at room temperature owing to mainly two reasons (i) first being the high quality and lesser impurities and structural defects in ZnO tetrapods and (ii) is related to the quantum confinement effect in nanocrystals [48]. A remarkable difference in response from a group of randomly oriented tetrapods and a single tetrapod oriented that one of its arm is aligned parallel to the incident laser beam. The reason for the broad Vis-emission band is that ZnO tetrapod have a large surface to volume ratio. This difference in optical behavior is reasoned on the basis that the vertical leg of the aligned tetrapod behaves like a Fabry-Perot resonant cavity, the essential reason for high-directivity antenna with different superstrates can be revealed in terms of the Fabry-Perot resonant theory [56].

$$I(E) \propto \alpha(E)E^2 \exp \left[ \frac{(E - E_g)}{K_B T} \right]. \quad (16)$$

Maxwell Boltzmann theoretical model (MBM), shown in Fig. 6. Table I presents the trap density (cm$^{-3}$)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Density traps (cm$^{-3}$)</th>
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<tbody>
<tr>
<td>ZnO-A</td>
<td>$8.9 \times 10^{13}$</td>
</tr>
<tr>
<td>ZnO-B</td>
<td>$9.3 \times 10^{13}$</td>
</tr>
<tr>
<td>ZnO-C</td>
<td>$9.9 \times 10^{13}$</td>
</tr>
</tbody>
</table>

These experimental results are generally concerned with high lifetimes [15,16]. The presence of a detectable PL emission bands in the nanocrystal, characterized by the presence of ZnO, demonstrates that Zn$^{2+}$ cation are already optically active in this crystalline configuration. Surface recombination varies widely even in the high-lifetime regime because it depends on the surface’s state, i.e., bare, passivated, contaminated, polarized of π-cloud, etc. The Zn-A and Zn-B samples show a small shift towards lower photon energy ($h\nu$), which is associated with a higher concentration of native defects as well as impurities identified by XRD and SEM. In other words, this phenomenon is associated with the hybridization ($sp$) of orbitals generating the π-cloud of delocalized electrons, phenomena has not been properly appreciated [57]. The molecular configuration of these inorganic materials presents the criteria that are based for the existence of π-delocalized electrons (delocalization of π-electron cloud) associated with the conjugated bonds external electromagnetic radiation produces a strong distortion of the polarization of the π-electronic cloud, which also generates significant changes in the electronic transitions in this solid nanomaterial [58]. The interesting part of inorganic material associated with this planar configuration focuses on the short-range van der Waals electrostatic interactions produced by the distortion in the π-electronic cloud of the functional group of Zinc-hydroxylcarbonate (> C=O ··· H-O-C-Zn-) pollutants. One approach to address these issues consists of growing high-quality single crystalline bulk and thin films in which the concentrations of impurities and intrinsic defects are controlled. A very interesting study has been published in which they apply the theory of doping and native defects in ZnO based on density-functional calculations, discussing the stability and electronic structure of native point defects and impurities and their influence on the electrical conductivity and optical properties of ZnO [59].

### 4. Conclusions

The search for the optimal conditions to prepare ZnO in a direct and simple way, present a greater scientific advance every day. Applying the appropriate technique in the synthesis of ZnO, the main objective is to have interesting morphological, structural and optical properties for its possible application. Considering the optical property, it is found that the PL signals recorded in the Vis-region of the spectra are associated with native defects. These are in turn associated with
It is well known that the origin of the aforementioned PL emission bands is still in a deep scientific debate. However, from the multiple reports on ZnO made by various research groups, they are identified and carefully examined through theoretical-experimental studies. We found a drastic and marked difference in the theoretical-experimental PL emission signals, with respect to those recorded by the MBM. In this theoretical model, the effect of other defects was not considered. However, it is applied here as a first approximation.

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