Radiation Induced Synthesis of In$_2$O$_3$ Nanoparticles - Part II: Synthesis of In$_2$O$_3$ Nanoparticles by Thermal Decomposition of Un-irradiated and γ-irradiated Indium Acetylacetonate

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Pure cubic phase, In$_2$O$_3$ nanoparticles with porous structure were synthesized by solid state thermal oxidation of un-irradiated and γ-irradiated indium acetyl acetate in presence and absence of sodium dodecyl sulphate as surfactant. The as- synthesized In$_2$O$_3$ nanoparticles were characterized by X-ray diffraction (XRD), fourier transformation infrared spectroscopy (FT-IR), scanning electron microscopy (SEM), transition electron microscopy (TEM) and thermogravimetry (TG). The shapes and morphologies of as- synthesized In$_2$O$_3$ nanoparticles were highly affected by γ-irradiation of indium acetyl acetate precursor and by addition of sodium dodecyl sulphate as surfactant. Calcination of un-irradiated indium acetyl acetate precursor to 4 hours of 600 °C leads to the formation of spherical- shaped accumulative and merged In$_2$O$_3$ nanoparticles with porous structure, whereas irregular porous architectures composed of pure In$_2$O$_3$ nanoparticles were obtained by using γ-irradiated indium acetylacetate precursor. The as- prepared In$_2$O$_3$ nanoproducts exhibit photoluminescence emission (PL) property and display thermal stability in a wide range of temperature (25-800 °C) which suggest possible applications in nanoscale optoelectronic devices.

Keywords: indium acetylacetonate, thermal decomposition, γ-irradiation, indium oxide nanoparticles

1. Introduction

In$_2$O$_3$ is an important n- type semi- conductor with a wide direct band gap of about (3.5-3.8 eV) of potential use in optoelectronic devices and gas sensing applications. According to the inorganic crystal structure data base, In$_2$O$_3$ is reported to exist on three different structures, two body- centered cubic (bcc) structure (Space group 199) and (Space group 206) as well as rhombohedral (rh) one (Space group 167). Besides the different Bravais lattice, (bcc) or (rh), these polymorphs posses different space groups: $I \overline{2} 3$ (space group 199), $I a \overline{3}$ (space group 206) and $R \overline{3} C$ (space group 167). Several routes such as chemical vapor deposition (CVD), Sol gel, solution phase growth, and solid state decomposition have been employed for preparation of In$_2$O$_3$ nanoparticles.

Radiation induced synthesis of nanostructured materials plays an important role in the investigation and production of well- shaped and mono- dispersed nanoparticles. Methods based on the interaction of high energy charged particles and γ- ray are widely used in making ion- track membrane, polymeric nanocomposite, and metal oxide nanoparticle. To the best of our knowledge, we are the first research group deals with γ-irradiation induced synthesis of In$_2$O$_3$ nanoparticles.

In our first submission, we stated with un-irradiated and γ-irradiated indium acetate with 10$^2$ kGy γ- ray absorbed dose as precursors for preparation of In$_2$O$_3$ nanoparticles by sol- gel methods. The results demonstrated different morphologies and particle sizes for the as- synthesized In$_2$O$_3$ nanoparticles as result of γ- irradiation.

In the present study, we will continue our investigation into using un-irradiated and γ-irradiated indium acetylacetone (InAcAc) as precursors for preparation of In$_2$O$_3$ nanoparticles in presence and absence of sodium dodecyl sulphate (SDS) surfactant by thermal decomposition process. The results will discuss in view of the role of γ- irradiation and addition of surfactant on the morphologies, sizes and shapes of as-synthesized In$_2$O$_3$ nanoparticles.

2. Experimental

2.1. Material

Indium (III) acetylacetone (AcAc) (≥ 99.99% trace metals basis, Sigma- Aldrich) and sodium dodecyl sulphate (SDS) (Bioultra, ≥ 99.99% (GC), Sigma- Aldrich) were used as received.

2.2. Synthesis of In$_2$O$_3$ nanoparticles

Four samples were prepared for the experiment. The first sample contains 0.0024 mole of un-irradiated indium acetylacetone (sample A). The second sample contains 0.0024 mole of un-irradiated indium acetylacetone and...
1 mL of 0.1 M sodium dodecyl sulphate solution (sample B). The third sample contains 0.0024 mole of γ-irradiated indium acetylacetonate of 10^2 kGy absorbed dose (sample C). The fourth sample contains 0.0024 mole of γ-irradiated indium acetylacetonate of 10^2 kGy absorbed and 1 mL of 0.1 M sodium dodecyl sulphate solution (sample D). The four samples were encapsulated into four separate stainless steel cells and allowed to stand in a muffle furnace. The temperature was raised at a heating rate of 10 °C min⁻¹ to 600 °C and kept constant for four hours. The reaction took place under the autogenic pressure of the encapsulated materials. At the end of the reaction, the containers were gradually cooled (5h) to room temperature, and after opening, yellow powders nanoparticles were collected in clean and dry containers and subjected to characterization.

2.3. Characterization

X-ray powder diffraction patterns (XRD) were recorded on Siemens D 5000 X-ray diffract meter with CuKα radiation (λ=1.54 Å) and operating at 40 KV/ 40 mA. The scan mode was continuous speed of 2.0 deg. / min. TG measurements were recorded on Perkin-Elmer TG A7 thermogravimetric analyzer in the temperature range of 30-1000 °C. The sample weight was 10.0 ± 0.1 mg with heating rate of 10 °C/min. FT-IR measurements were recorded as KBr pellets in the range of 200-4000 cm⁻¹ on Perkin-Elmer FT-IR spectrophotometer (spectrum 1000). SEM and TEM images were captured using the models (SEM, JSM-6360 ASEM, JEOL, Japan) and (TEM, JEM-2100F, JEOL, Japan) electron microscopes. Photoluminescence spectra of as-synthesized In₃O₄ nanoparticles were recorded using Perkin-Elmer Luminescence Spectrophotometer with a 450 W Xenon lamp as the excitation source (550 B) at room temperature.

For irradiation, samples were encapsulated under vacuum in glass vials and exposed to successively increasing doses of radiation at constant intensity. A Co-60 γ-ray source model gamma cell 220 from MDS (Nordion, Canada) was used for irradiation of the samples. The source was calibrated against a Fricke ferrous sulphate dosimeter and the absorbed doses in the irradiated samples were calculated by applying appropriate corrections on the basis of photon mass attenuation and the energy absorption coefficients for the sample and the dosimeter solutions. The dose rate was 9.83 kGy h⁻¹ and the transient dose was estimated to be 12.07 Gy. All of the irradiations were conducted at 25 °C. After irradiation, the samples were stored at room temperature for 24 h before analysis.

3. Results and Discussion

Figure 1 shows typical TG/ DTG/DTA curves for the thermal decomposition of un-irradiated and γ-irradiated indium acetylacetonate in air atmosphere. The thermograms collected for γ-irradiated InAcAc display some modifications in their thermal behaviors compared to those obtained using un-irradiated sample. The major differences in the thermal behavior between un-irradiated and γ-irradiated samples are listed in Table 1.

Figure 2 show XRD patterns of the as-synthesized In₃O₄ nanoparticles obtained under the different experimental conditions mentioned in the experimental section. All the diffraction lines are indexed undisputedly to pure phase of bcc In₃O₄ structure without indication of other crystalline by-products. The peaks are quite broad, which demonstrates the nanometric grained nature of the products. The average crystallite size calculated by Debye-sherrer formula from the (222) reflections are listed in Table 2.

The unit cell parameter (a) for cubic In₃O₄ was calculated using the following Equation:

\[ d = \frac{a}{\sqrt{h^2 + k^2 + l^2}} \]  

Where \( d \) is the d-spacing, \( h, k, \) and \( l \) are miller indices. By selecting (222) plane, the values of unit cell parameter (a) of the obtained In₃O₄ nanoparticles under the given experimental conditions were calculated with standard deviation of ± 0.001 and the results are listed also in Table 2.

From Table 2, it could be seen that the In₃O₄ unit cell obtained using γ-irradiated InAcAc precursor calcined in presence of 1 mL SDS (sample D) gave the smallest value of the parameter (a) due to the radiation damage induced by γ-irradiation in the host lattice of InAcAc as we will discuss in the section “role of irradiation”.

Figure 3 shows SEM micrographs of In₃O₄ nanoparticles prepared by calcination of the four samples of InAcAc. All the micrographs display the nanocrystalline nature of the as-synthesized nanoparticles. However, different morphologies were obtained. In sample (a), the surface is composed of porous, nearly spherical-shaped accumulative and merged particles. In the case (b), the irregular porous architecture composed of pure In₃O₄ was observed. In the case (c), the SEM image display aggregates of irregular-shaped In₃O₄ porous microsphere. For case (d), the surface is covered by porous layer of ill-defined In₃O₄ polyhedron.

TEM images of the as-prepared In₃O₄ nanoparticles are shown in Figure 4. For samples (a), (b) the TEM images display monodisperse In₃O₄ nanoparticles, with uniform size and shape. In case (c) the TEM image display porous structure of In₃O₄. For sample (d), the TEM image show formation of dark spots of agglomerated crystallites (quantum dots). There is a random distribution in the size and shape of these crystallites. The average particle size calculated by a simple computer algorithm for samples a, b and c were found to be around 48, 38 and 18.5 nm in good agreement with the values obtained from XRD measurements. The average diameter calculated by computer program in several grids is ~ 3.5 nm [18].

If we compare the morphology of In₃O₄ nanoparticles synthesized by sol-gel method in our previous publication with that reported in the present investigation using solid state thermal decomposition process, different morphology obtained. In case of sol-gel method the SEM and TEM images of In₃O₄ nanoparticles synthesized using un-irradiated indium acetate precursor display agglomerated and ill-defined spherical nanoparticles and nanocubes respectively. In the present investigation, the SEM and TEM images display uniform structure of well-defined and monodispersed nanoparticles. In sol-gel method, γ-irradiation afforded coral-like structure (SEM image) and blocks of agglomerated nanocubes (primary of noncrystallites) with big holes between
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Figure 1. TG/DTG/DTA curves of: a) Un-irradiated InAcAc; b) γ-irradiated InAcAc (10$^2$ kGy).

Table 1. TG/DTG/DTA data of indium acetylacetonate (In$_2$O$_3$).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Un-irradiated InAcAc</th>
<th>γ-irradiated InAcAc</th>
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<tbody>
<tr>
<td>No weight loss was detected up to 190 °C.</td>
<td>There is a minor weight loss (~5%) in the temperature range of (120-160 °C). The corresponded DTA curve reveal broad and small endothermic peak maximized at 138 °C attributed to the loss of adsorbed water on the high surface area of InAcAc created by formation of points defects and lattice imperfections as result as of γ-irradiation$^1$.</td>
<td>The main decomposition process for the release of acetylacetone moieties show thermal event in the temperature range of (180-260 °C) with small broad three endothermic peaks centered at 188, 233, 250 °C attributed to step wise release of three coordinated acetyl acetone molecules. The experimentally calculated weight loss percentage (29.8%) was found to be in good agreement with that calculated theoretically (31.6%).</td>
</tr>
<tr>
<td>The minor weight gain for the oxidation to In$_2$O$_3$ was detected in the temperature range of 280-320 °C with small exothermic peak at 309 °C.</td>
<td>The corresponding decomposition behavior contains three thermal events. The DTA curve reveal two sharp endothermic peaks maximized at 188 and 266 °C and one minor and broad endothermic peak centered at 222°C. There is also good agreement between the experimental (29.9%) and theoretical values (31.6%) assuming the release of three acetyl acetone molecules.</td>
<td></td>
</tr>
<tr>
<td>The TG curve shows the weight loss gain process to occur at a higher and narrow region (310-320 °C). The DTA curve reveal broad exothermic peak monitored at 338 °C.</td>
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</table>
these blocks (TEM image). In the present investigation γ-irradiation afforded irregular porous architecture (SEM) image and mesoporous structure of In$_2$O$_3$ nanoparticles. Further discussion on the role of γ-irradiation on the morphology, size and shape of the as-synthesized nanoparticles will be published in a separate review after complete publication of all series of this project.

Figure 5 shows FT-IR spectra of the as-synthesized In$_2$O$_3$ nanoparticles. Four main intensive bands centered at 416, 539, 566 and 604 cm$^{-1}$ were observed and characteristic for the cubic In$_2$O$_3$ phase. The chemisorbed H$_2$O and CO$_2$ molecules on the surface of In$_2$O$_3$ could be easily detected by recording the bands appeared in the range of 3400-3500 cm$^{-1}$ and 1700-1600 cm$^{-1}$ attributed to ν$_{OH}$ and ν$_{C=O}$ respectively.

As shown from Figure 5 no significant changes were recorded in the IR-spectra of the investigated samples as a result of γ-irradiation and surfactant addition.

### 3.1. PL analysis

Photoluminescence (PL) is the spontaneous emission of light from a material under optical excitation. The excitation energy and intensity are chosen to probe different regions and excitation concentrations in the sample. PL investigations can be used to characterize a variety of materials parameters and provides electrical characterization. It is a reflective and extremely sensitive probe of discrete electronic states. It is well known from literature that bulk In$_2$O$_3$ cannot emit light at room temperature. However, PL emissions of the nanocrystalline In$_2$O$_3$ samples are possible due to the effect of oxygen vacancies and recently to indium inertial and their associated complex defects.

Figure 6 shows the RT-PL spectra of the as-synthesized In$_2$O$_3$ nanoparticles. While exciting the sample with 266 nm, strong emission at 383 nm and a weak emission at 650 nm were recorded. In general, emission spectra can be divided into two broad

<table>
<thead>
<tr>
<th>Precursor</th>
<th>a, Å</th>
<th>average diameter, nm</th>
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<tr>
<td>Un-irradiated InAcAc (A)</td>
<td>10.086</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>γ-irradiated InAcAc (B)</td>
<td>10.032</td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>Un-irradiated InAcAc + SDS (C)</td>
<td>10.041</td>
<td>20</td>
<td>10.114 Å</td>
</tr>
<tr>
<td>γ-irradiated InAcAc + SDS (D)</td>
<td>10.031</td>
<td>5</td>
<td></td>
</tr>
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</table>
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Figure 3. SEM micrographs of the as-synthesized In$_2$O$_3$ nanoparticles (a-d). a) Un-irradiated AcAc; b) γ-irradiated AcAc (10$^2$ kGy); c) Un-irradiated AcAc and SDS; d) γ-irradiated AcAc (10$^2$ kGy) and SDS.

Figure 4. TEM micrographs of the as-synthesized In$_2$O$_3$ nano-particles (a-d). a) Un-irradiated AcAc; b) γ-irradiated AcAc (10$^2$ kGy); c) Un-irradiated AcAc and SDS; d) γ-irradiated AcAc (10$^2$ kGy) and SDS.
categories, the near-band edge (NBE) emission and deep level (DL) emission. The peak at 383 nm is due to near band edge emission owing to excitonic transition. The PL peak at 650 nm could be attributed to imperfections and defects due to oxygen vacancies. However, regarding the results of FT-IR, the surface of nanoparticles is highly active as confined by the presence of hydroxyl groups. In high active surface several types of defects are expected to give rise to oxygen vacancies and other point defects in the host lattice of cubic indium oxide. These oxygen vacancies induce the formation of new energy levels in the band gap which facilitates the electronic transitions between valence band and conduction band. The as-obtained four samples of In$_2$O$_3$ display the same PL spectra without any significant changes as a result of γ-irradiation and surfactant free addition and only one representative spectrum will be displayed.

Thermal stability of as-synthesized In$_2$O$_3$ nanoparticles was tested by recording their TG/ TGA/ DTA curves in the temperature range (25-400 °C) and the results are shown in Figure 7. The TG curve display thermal stability with minor weight loss (∼2%) in the temperature range of (50-350 °C). The DTA curve shows multiple peaks maximized at 50, 180, 216, 281 and 311 °C. These thermal events are due to release of adsorbed water, organic and in-organic species from the active surface of the as-synthesized nanoparticles. Broad exothermic peak was detected at 369 ºC, probably due the formation of non-stoichiometric composite In$_2$O$_3$.

3.2. Role of irradiation

In solid sample, the radiation effect is dominated by direct ionization of the material, whereas for aqueous solutions the reaction with radical species, such as OH$^-$ or solvated electrons is the dominant mechanism for damage to a solute. Upon irradiation with Co-60 γ-ray source, the Compton scattering is the main mode of interaction of γ-ray with indium and oxygen atoms and multiple ionization occurs. This event leads to the formation of electron-hole pairs, lattice imperfection and extended defects. From crystallographic point of view, when unit cell is exposed to γ-irradiation, the atoms may be disturbed from the equilibrium positions. Accordingly, the scattering action changes so that the structure factor becomes different and the lattice dimensions become smaller. Increasing of the dose absorbed by the precursor, the radiation effect may be significantly remarkable and contraction of the unit cell occurs. If the contraction is below a certain limit, broadening of the powder diffraction results.

As a result of these effects, In$_2$O$_3$ nanoparticles synthesized using γ-irradiated precursors are smaller in size and displayed different morphology compared to the nanoparticles obtained from un-irradiated indium acetate. In the case (D), the trapped electrons in the host lattice of InAcAc may react with surfactant moiety to create another source of points and line defects in InAcAc due to the damage of SDS on the surface of the new phase formed by In$_2$O$_3$ nanoparticles as a result of γ-irradiation. This damage may increase the rate of nucleation of the nanoparticles to have another impact on morphologies and shapes of the as-synthesized nanoparticles.

It is worthy mentioned that the cell parameter, a of In$_2$O$_3$ calculated using γ-irradiated indium acetate (10.05 Å) by sol-gel method in our previous work was found to be higher than the calculated value by solid state thermal decomposition of indium acetylacetonate of the same adsorbed dose (10.03 Å). This behavior could attribute to high concentration of the trapped electrons and holes generated by γ-irradiation in the solid state and leads to high degree of lattice imperfection, and extended defects. Part of the trapped electrons in the case of sol-gel method reacts with benzyl alcohol and the gel leading to the formation of intermediate species of very short half-life.
4. Conclusion

This study reports the synthesis of high purity porous In$_2$O$_3$ crystallites with different morphologies by thermal decomposition of un-irradiated and γ-irradiated anhydrous indium acetylacetonate (InAcAc) precursor in presence and absence of surfactant for 4 h at 600 °C. Thermal oxidation of γ-irradiated indium acetylactonate precursor with $10^2$ kGy in presence of sodium dodecyl sulphate leads to the formation of nanocrystalline of cubic In$_2$O$_3$ with porous structure and having average diameter in the range of 1-3 nm. The as-synthesized In$_2$O$_3$ nanoparticles show PL property due to the presence of singly ionized oxygen vacancies. The as-synthesized In$_2$O$_3$ nanoparticles display thermal stability in the temperature range of (25-400 °C). The porous nature of the as-synthesized In$_2$O$_3$ nanoparticles suggests the potential application of these particles as gas-sensing material.

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