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# TUNING THE OPTICAL AND MECHANICAL PROPERTIES OF Y<sub>2</sub>O<sub>3</sub> CERAMICS BY THE INCLUSION OF La<sup>3+</sup> ION IN THE MATRIX FOR INFRARED TRANSPARENT WINDOW APPLICATIONS

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### **ABSTRACT**

Infrared transparent windows are generally used to protect highly delicate infrared sensor circuits from the harsh environments. In the present work a combustion technique was effectively used to incorporate La³+ ion in the yttria matrix. The crystallites were in the size limit of 20 nm. Powder characterization using X-ray diffraction, HRTEM and FTIR spectroscopy revealed that the La³+ ions were effectively replacing the Y³+ ion in the yttria matrix. There was a slight reduction in optical band gap with La³+ concentration. A novel sintering mechanism was used for sintering the samples by coupling definite proportions of resistive heating and microwave heating. The highly dense pellets showed better transmittance and hardness properties, which improved with La³+ concentration. The present study authorises that combustion synthesis of the samples followed by resistive coupled microwave sintering can effectively be used to tune the optical and mechanical, properties of infrared transparent ceramics. Improving the transmittance properties without deteriorating the mechanical properties through the resistive coupled microwave sintering is the major highlight of this novel work.

**Keywords:** Combustion synthesis, Hardness, Infrared window, Resistive coupled microwave sintering

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# 1. INTRODUCTION

Infrared transparent ceramics are materials which allow electromagnetic radiations in the infrared range to pass through them without being much scattered or absorbed. The studies on transparent ceramics was triggered by the invention of commercially successful translucent  $\alpha$ -

Al<sub>2</sub>O<sub>3</sub> which was used as the envelope of high pressure Na-vapour lamp by General Electric under the trade name Lucalox in 1966 [1]. In electronic industry it found applications as it is used to protect highly delicate infrared sensor electronic circuits from the harsh environments. Infrared inspection ports and infrared cameras employ the transmittance property of the infrared transparent ceramics in the mid infrared range [2]. Infrared transparent window is one among the indispensable parts of homing missiles.

Yttria is considered to be one of the most promising candidates among all infrared transparent windows. It has remarkable optical properties but the hardness is at a moderate level compared to other window materials [3,4]. It is reported that the substitution of heavier ions in the yttria matrix effectively improves the mechanical and transmittance properties [5].  $Y_2O_3$  and  $La_2O_3$  both have 16 formula units per unit cell and no lattice distortions are expected in  $La_xY_{2-x}O_3$  up to a value x=0.24 [6,7]. There are reports that even at ~1900 °C the cubic phase of yttria with 16 mole percent of La is stable [8,9]. The phase change for 9 mole percent lanthanum doped yttria occur only after 2325 K [6]. Moreover it is reported that  $La^{3+}$  is an effective sintering aid which enhance the transmittance properties of infrared transparent windows [10].

Sintering of the transparent ceramics to high density without much grain growth is a major challenge in the fabrication of infrared transparent windows used as infrared sensor protective dome in homing missiles [11]. For pure nano crystalline yttria powder, the sintering temperature is above 1600 °C. Recently our research team has developed a new sintering strategy which effectively couple the conventional resistive heating and microwave heating in definite proportion to reduce the sintering temperature and soaking duration [12]. The yttria pellets sintered using the novel method showed better transmittance properties in the UV-Visible and mid-infrared region. Even a reduction in sintering temperature and soaking duration reduces the grain size to a great extent which improves the hardness and transmittance properties. The presence of La<sup>3+</sup> ion in the yttria matrix is reported to reduce the sintering temperature to ~1500 <sup>o</sup>C [13]. La<sup>3+</sup> ions enhance the grain boundary mobility of yttria and it is the reason behind improved sinterability [14]. In addition to this the La<sup>3+</sup> ions easily dissolves in the cubic lattice of yttria and reduce the grain growth [5,15]. Two step sintering mechanisms and high pressure sintering mechanisms were effectively used for sintering La doped yttria, but the transmittance is less than 80% [16]. Recently Gan et al. used hot pressing sintering to achieve a density of 99.9 % in 12 atom percent La doped vttria ceramics which showed a transmittance of 68.9 % at 400 nm and 81.9 % at 1100 nm where as a pellet sintered without applying pressure achieved 99.8 % of the theoretical density [17,18]. But no much work have so far been reported in this system, may be due to the difficulty in the sintering of the samples at low temperatures with reduced grain size and porosity.

In the present work we discuss in detail the synthesis of La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub> by a modified combustion route. Various powder characterisation techniques like X-ray diffraction, Transmission Electron Microscopy, Fourier Transform Infrared Spectroscopy and UV-Visible spectroscopy used to analyse the quality of the powder are discussed in detail. The optical and hardness properties of the samples sintered via resistive coupled microwave sintering are also described in detail.

# 2. MATERIALS AND METHODS

A single step auto-igniting combustion technique was used to synthesize nano structured  $La_xY_{2-x}O_3$ . Stoichiometric amount of high purity  $Y(NO_3)_3.6H_2O$  (99.99 %, Alfa Aesar, USA) dissolved in double distilled water was mixed with  $La_2O_3$  (99.99 %, Alfa Aesar, USA) dissolved in nitric acid, to make a clear solution. Citric acid was added to the precursor solution which act both as fuel and the complexing agent [11]. By adjusting the nitric acid and ammonium hydroxide the pH of the precursor solution is made  $\sim$ 7. Nitric acid in the precursor

solution reacts with the ammonium hydroxide forming ammonium nitrate which acts as the extra oxidant. The precursor solution was heated using a hot plate at 250 °C in a combustion chamber. On persistent heating, the solution foams and finally ignites giving voluminous and fluffy product of combustion.

X-ray diffractometer (X'pert pro, PANalytical, the Netherlands) with Cu Kα radiation in the range of 20-60° was used for the determination of crystalline structure and phase of the as prepared sample. The X-ray diffraction data was utilised for estimating the crystallite size via Scherrer's equation and Williamson-Hall plot. High resolution transmission electron microscopy (JEM-2100, JEOL, USA) operating at 200 kV was used for verifying the particulate properties and the crystallite size distribution. Fourier Transform Infrared spectrometer (Spectrum 2, Perkin-Elmer, Singapore) with diamond ATR in the range 400-4000 cm<sup>-1</sup> was used to verify the phase purity of the as prepared sample. The absorption spectrum of asprepared samples of were recorded using a spectrophotometer (UV-1700, Shimadzu, Japan).

The as prepared powder was uniaxially compacted into pellets in a 14 mm diameter steel die at 20 MPa using a hydraulic press. A novel resistive coupled microwave sintering technique was used for sintering the samples. The resistive coupled microwave furnace (VBCC/HMF/71, VB Ceramics Consultants, India) effectively couples the resistive heating and microwave heating in desired proportion. A pair of molybdenum disilicide heating elements was used for resistive heating and a pair of 2.5GHz magnetrons with power 1.1 kW each provides microwave heating support. The effective control over the heating rate and uniformly distributed fast heating cycles yield high quality infrared transparent ceramics with enhanced properties. By varying the percentage composition of the microwave and resistive powers the micro structure and the properties of the sintered samples can be tuned. In the present work, after a series of trials the resistive and microwave heating were effectively coupled in the ratio 60:40 below 1100 °C and there after it was fixed at 40:60.

Archimedes principle was used to find the experimental density of the sintered samples. Coarse grinding, lapping and fine polishing are effectively utilised for optical quality surface finish. For a higher quality surface finish more intermediate sizes of abrasives are used and more material might be removed from each step. Fine grade diamond lapping and polishing pastes and fluids of 5 µm, 1µm, 0.5 µm, 0.25 µm and 0.1 µm were mainly used for this. Finally a float polishing mechanism was employed in which the work piece was submerged and rotated in a slurry of extra fine quality polishing powder in deionized water. The well-polished pellets were used for transmittance studies. The transmittance of radiation in the UV-Visible and in the IR range were measured using UV-Visible and FTIR spectrometers.

Vickers indenter method (HMV 2TAW, Shimadzu, Japan) was used for studying the hardness of the well sintered pellets. Various indentation loads P of 0.98, 1.96, 2.94 and 4.90 N were used for a duration of 14 s to study the load dependent hardness of the samples. If the hardness of the sample decreases with increase in load it is called positive indentation size effect (ISE) [19]. For detailed analysis a semi empirical relation called modified proportional specimen resistance (MPSR) model is used for studying the ISE behaviour in the sintered samples [20]. The hardness of the sample for different indentations were estimated using the relation

$$P = a_0 + a_1 d + a_2 d^2$$

Where P is the applied load, 'd' is the indentation size, ' $a_0$ ', ' $a_1$ ' and ' $a_2$ ' are parameters obtained from curve fitting of the experimental results. Here ' $a_0$ ' is the term related to the residual stresses generated in the specimen surface,  $a_1$  is a constant related to the proportional resistance of the specimen which is directly proportional to the Young's modulus and ' $a_2$ ' is a constant related to load independent micohardness. The load independent hardness called true hardness  $H_T$  can be obtained from ' $a_2$ ' by the relation

$$H_T = ka$$

Where k is a constant equal to 1.8544 for Vicker's indenter [21].

### 3. RESULTS AND DISCUSSION

X-ray diffraction technique was effectively used to analyze the phase purity of the sample. X-ray diffraction patterns of  $Y_2O_3$  and  $La_xY_{2-x}O_3$  (where  $x=0.05,0.10,0.15,\,0.20$  and 0.25) are shown in Fig. 1. All the peaks correspond to the cubic phase of yttria and are indexed using JCPDS file no-895591. No additional peaks were observed in the XRD pattern, which clearly indicate that the  $La^{3+}$  ions are substituting  $Y^{3+}$  ions in the yttria cubic structure. The ionic radius of  $La^{3+}$  (1.032  $\overset{\circ}{A}$ ) is larger than that of  $Y^{3+}$  (0.90  $\overset{\circ}{A}$ ) ions [22]. As the  $La^{3+}$  concentration increases the diffraction peaks shift towards shorter 20 side which attributes to the increase in d-spacing due to the substitution of bigger  $La^{3+}$  ion in the  $Y^{3+}$  sites [16, 23].

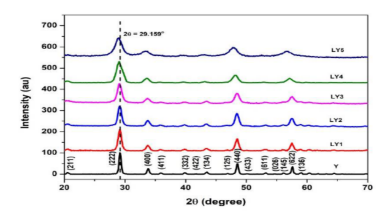


Figure 1 XRD pattern of  $La_xY_{2-x}O_3$  (where x= 0, 0.05,0.10,0.15,0.20 and 0.25)

Table 1 shows the variation in d-spacing and cell parameters for the (222) peak with increasing La<sup>3+</sup> ion concentration. For convenience sample codes were assigned for each concentration and the crystallite sizes were calculated from the high intensity peaks using Scherrer formula.

The extent of line broadening in the diffraction pattern due to the strain, which is caused by the non-uniform displacement of the atoms with respect to their reference lattice position, could be identified by Williamson-Hall (W-H) method [24]. In this method, the reciprocal peak width ( $\beta \cos \theta$ )/ $\lambda$  is plotted against the reciprocal lattice distance ( $4sin\theta$ )/ $\lambda$ , and the resulting plot is fitted linearly. Fig. 2 shows the Wiliamson-Hall plot for the  $La_xY_{2-x}O_3$  nanoparticles. The lattice strain is estimated from the slope of the graph and the reciprocal of Y-intercept gives the crystallite size. The lattice strain and crystallite sizes for  $La_xY_{2-x}O_3$  with different  $La^{3+}$  ion concentration are calculated and included in table 1. It is noticed that as growth of  $La_xY_{2-x}O_3$  is restricted by the substitution of  $La^{3+}$  ion leading to relatively small crystallite size as compared to pure yttria. Similar behavior is observed by many researchers in oxide systems on incorporation of bigger ions in the crystal lattice. The positive slope indicates that the strain developed is tensile in nature [25].

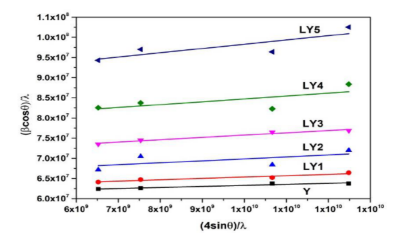


Figure 2 Williamson-Hall plot for La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub> with different La<sup>3+</sup> ion concentration

Table 1 Variation in d-spacing and cell parameter with La<sup>3+</sup> ion concentration

La <sub>x</sub> Y <sub>2-x</sub> O <sub>3</sub>	Sample Code	d-spacing (Å) for (222) plane		Crystallite size		_
		JCPDS file 895591	Present Sample	Scherrer Formula	W-H plot	Micro strain
$\mathbf{x} = 0$	Y	3.0615	3.0601	$14.16 \pm 0.20$	16.30	2.17 x10 <sup>-4</sup>
x = 0.05	LY1	3.0615	3.0639	$13.20 \pm 0.14$	16.1	$2.50 \times 10^{-4}$
x = 0.10	LY2	3.0615	3.0657	$12.16 \pm 0.23$	15.4	4.91 x10 <sup>-4</sup>
x = 0.15	LY3	3.0615	3.0693	$11.71 \pm 0.25$	14.2	$5.73 \times 10^{-4}$
x = 0.20	LY4	3.0615	3.0722	$10.68 \pm 0.24$	12.8	$7.15 \times 10^{-4}$
x = 0.25	LY5	3.0615	3.0767	$9.74 \pm 0.23$	11.4	$1.08 \times 10^{-3}$

There is a linear relationship between the La<sup>3+</sup> concentration and lattice constant in the yttria matrix as shown in Fig. 3. A graph is plotted with the lattice constant (a) with the concentration of La<sup>3+</sup> concentration (C) and is linearly fit according to the relation a = 10.602 + 0.0105C. It is found that lattice constant changes from 10.613 to 10.658 Å as the La<sup>3+</sup> concentration changes from 1 mole % to 5 mole %.

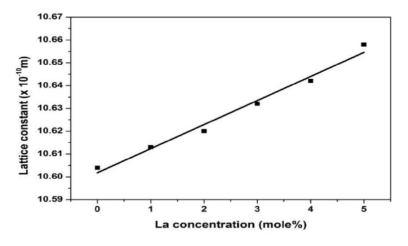


Figure 3 Variation in cell parameter with La<sup>3+</sup> ion concentration

The crystallite size distribution from the TEM image of the as prepared LY3 sample which is shown in the Fig. 4 supports the XRD results. The crystallite size are found to be in the range 7 - 24 nm. Maximum number of crystallites lie in the size range 10 -12 nm and the average

crystallite size for the entire distribution is calculated as  $11.58\pm0.33$  nm. Fig. 5 shows the HRTEM image of the as prepared LY3 powder sample. The crystallographic planes are well defined and they are in good agreement with the XRD data.

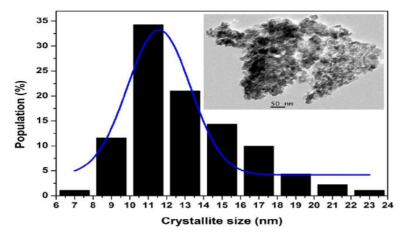


Figure 4 Crystallite size distribution of as prepared LY3. TEM image is shown inset.

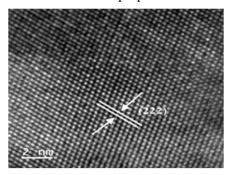
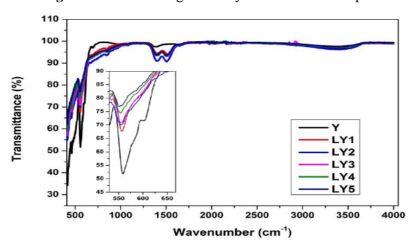


Figure 5 HRTEM image of as synthesized LY3 sample



**Figure 6** FTIR spectrum of as synthesized La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub>. The characteristic peak for different compositions are shown in the inset

The chemical and structural information for identifying and characterizing the nanopowders are further studied using FTIR spectra. The FTIR spectra shown in Fig. 6 confirms that the  $La_xY_{2-x}O_3$  powder transmits in the mid IR range. The absorption peak at 559 cm<sup>-1</sup> is assigned to Y-O stretching in the case of pure yttria [26]. In the present sample there is a slight shifting of this peak to longer wavelength range as reported in the literature [27]. But there is no additional peaks. From the XRD pattern it is clear that in the present composition  $La^{3+}$  ions

dissolves in the yttria matrix and there is no change in the phase, but there is a slight increase in the lattice parameter. ie as La<sup>3+</sup> ion concentration increases the cell volume increases. With the increase in the cell volume the IR absorption peak shifts towards longer wavelength range [28]. The FTIR spectrum in figure 6 corroborate the results obtained from the XRD. Fig. 7 shows the variation in absorption wave number with cell volume.

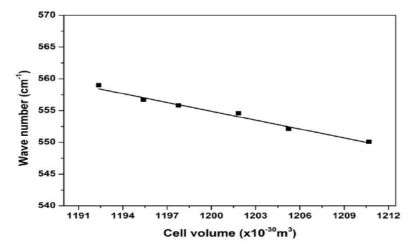


Figure 7 Variation in absorption wave number with cell volume

The high energy edge of a transparent ceramics is determined by the energy gap between the conduction and valence orbital band which is called the energy gap  $E_{\rm g}$  [29]. The UV-Visible absorption spectrum was recorded in the range 200-800 nm and is shown in Fig. 8. The sample showed maximum absorption in the UV region and decreases towards the visible region. The wavelength corresponding to maximum absorption for the samples shifts towards longer wavelength range as  $La^{3+}$  ion concentration increases.

The measured reflectance is converted into Kubelka-Munk (K-M) function  $F(R) = (1-R)^2/(2R)$ , where R is the reflectance. The K-M function is proportional to the absorption coefficient  $\alpha$  in the Tauc equation [29]. Fig. 9 shows Tauc's plot in the form of Kubelka-Munk function of the optical absorption spectrum of yttria nanopowder. A graph is plotted with  $(F(R)*hv)^2$  against the energy hv of the incident photon and the optical band gap energy obtained from the intercept of the linear region with the energy axis at  $(F(R)*hv)^2 = 0$ . It is observed that with La<sup>3+</sup> addition the band gap gradually decreases 5.65eV of pure yttria to 5.51eV of sample LY5.

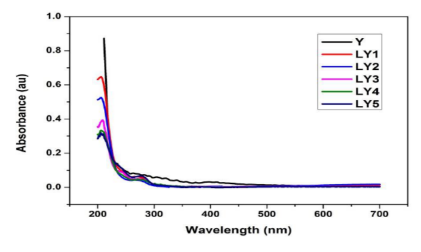
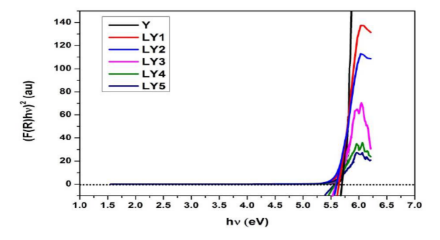


Figure 8 UV-Visible absorption spectrum of as synthesized La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub>



**Figure 9** Tauc's plot of as synthesized La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub> in terms of Kubelka Munk function.

Table 2 Density achieved by the	e pellets of different composition
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Sample	Sintering	Maximum relative
Code	Temperature	density achieved
	(±10°C)	(%)
$Y_{RM}$	1450	$99.2 \pm 0.01$
$LY1_{RM}$	1440	$99.2 \pm 0.02$
$LY3_{RM}$	1430	$99.2 \pm 0.02$
$LY5_{RM}$	1430	$99.1 \pm 0.02$

The La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub>nanopowder is uniaxially compacted into pellets in a 14 mm diameter steel die at 20 MPa using a hydraulic press. All the pellets were sintered using resistive coupled microwave heating showed high relative density >99% of the theoretical density. For a comparative study the samples Y, LY1, LY3 and LY5 sintered via resistive coupled microwave sintering were coded as Y<sub>RM</sub>,LY1<sub>RM</sub>, LY3<sub>RM</sub> and LY5<sub>RM</sub> respectively and tabulated in table 2.

For a comparative study the transmittance of the pellets in the UV-Visible range with different La  $^{3+}$  ions sintered via resistive coupled microwave heating are used. The maximum transmittance of 80.5% is shown by the sample LY5  $_{RM}$ . The comparison of transmittance of different samples in the UV-Visible region are shown in Fig. 10.

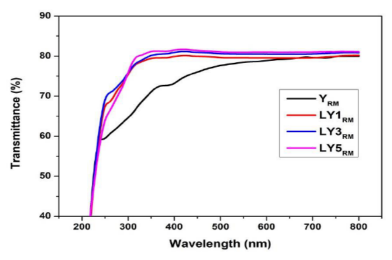


Figure 10 Transmittance percentage in the UV-Visible range in different pellets sintered using resistive coupled microwave heating

The transmittance properties of the pellets in the mid infrared range is studied. The upper IR cut off or the long wavelength edge of a transparent ceramics is determined by the energy absorption used for transition between the vibration states of the lattice called phonons. Usually the overtones of the principal lattice vibration define the upper IR cut off [11].

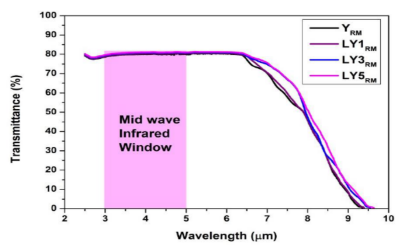


Figure 11 Transmittance percentage in the mid IR range in different pellets

The transmittance of radiations through the pellets with different  $La^{3+}$  ion concentration and sintered via resistive coupled microwave heating are studied. It is shown in Fig. 11. From the figure it is clear that as the concentration of  $La^{3+}$  ion increases there is a slight increase in the infrared cut off. Since ionic size of  $La^{3+}$  ion is greater than that of  $Y^{3+}$ , La-O fundamental phonon vibrational frequency is less than that Y-O so the phonon absorbing edge shifts towards the longer wavelength range [11]. Compared to pure yttria sintered via resistive coupled microwave heating,  $La_xY_{2-x}O_3$  having a slight increase in the percentage transmittance. Fig. 12 shows the variation in dielectric constant with frequency for different samples.

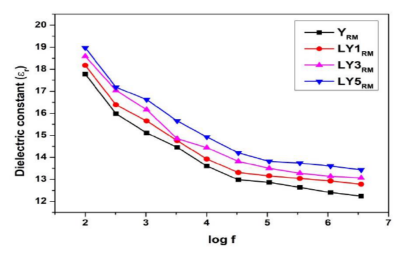


Figure 12 Variation in dielectric constant with frequency for different pellets

For the sample  $Y_{RM}$  the dielectric constant is found to be 12.23 and that for the sample LY5<sub>RM</sub> it is 13.48. The increase in transmittance in the UV-Visible region and mid-IR region may be due to the change in dielectric properties of the yttria matrix with La<sup>3+</sup> ions. The variation of dielectric loss with frequency for different pellets are shown in Fig. 13. As in the case of pure yttria all other samples show a decrease in dielectric loss with temperature [30]. But it is worth to note that the dielectric loss increases with the increase in La<sup>3+</sup> concentration,

which may be the reason behind the improved sinterability of La substituted yttria. In resistive coupled microwave sintering the green pellet absorbs the microwave energy generated by the magnetrons and convert it into heat within. The absorbed microwave power is proportional to the magnitude of electric field distribution in the sample. As the dielectric loss increases the microwave absorption in the sample increases [31]. In microwave sintering, the densification commences at a relatively low temperature by enhancing the diffusion mechanisms. So the sintering temperature and the soaking time can be reduced considerably. The microwave ponderomotive force induced mass transport is considered to be the reason behind the high densification in microwave sintering [32]. Even in low dielectric loss materials as temperature increases the dielectric loss factor increases and at higher temperatures they can effectively absorb the microwaves [33].

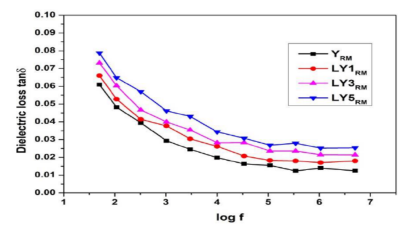


Figure 13 Variation in dielectric loss with frequency for different pellets

To analyse the load dependent hardness of the samples  $Y_{RM}$ ,  $LY3_{RM}$  and  $LY5_{RM}$  MPSR model is used. Variation in indentation size with applied load for different  $La^{3+}$  concentration is shown in Fig. 15.

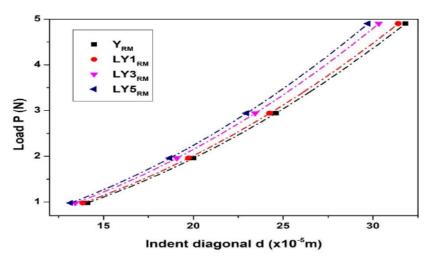


Figure 15 Variation in indentation size with applied load for different La<sup>3+</sup> concentration

A P versus d graph is plotted for different compositions as in Fig 15 and the parameters obtained are tabulated in table 3. The results reveals that as the presence of  $La^{3+}$  ions in the matrix increases the hardness increases and maximum hardness of 10.05 GPa is shown by the sample LY5<sub>RM</sub>.

Samples	a <sub>0</sub> (N)	a <sub>1</sub> ( <b>N/μm)</b>	$a_2 = \frac{P_c}{d_0^2}$ (N/ $\mu$ m <sup>2</sup> )	H <sub>T</sub> (GPa)
$Y_{RM}$	0.00064	0.00227	0.00477	8.85
$LY1_{RM}$	0.01011	0.00262	0.00487	9.03
$LY3_{RM}$	0.00182	0.00238	0.00525	9.73
$LY5_{RM}$	0.01638	0.00447	0.00542	10.05

# 3. CONCLUSIONS

The comprehensive analysis based on XRD and FTIR spectroscopy reveals that by the modified auto igniting combustion technique La<sup>3+</sup> ion effectively substitutes Y<sup>3+</sup> ions in the yttria matrix without changing the body centred cubic phase. In nanostructured La<sub>x</sub>Y<sub>2-x</sub>O<sub>3</sub>( where x = 0, 0.05, 0.10, 0.15, 0.20 and 0.25) powders the crystallites size analysis using Scherer formula reveals that at La<sup>3+</sup> concentration increases crystallite size decreases. The TEM image analysis support the estimation of crystallite size and d-spacings obtained from XRD data. The crystallite size calculated for LY3 sample using XRD is  $11.71 \pm 0.25$  nm and TEM is  $11.58\pm0.33$  nm. Williamson Hall plot is used to deconvolute the size and strain effect in XRD peak broadening which indicates that there is tensile strain in the crystallites which may contribute to peak broadening. As La<sup>3+</sup> ion concentration increases the lattice strain increases due to difference in ionic size of La<sup>3+</sup> and Y<sup>3+</sup> ions. As La<sup>3+</sup> ion concentration increases the cell volume increases but no phase changes are observed which attributes to the shift in absorption peak towards longer wavelength range in the FTIR spectrum. No additional peaks are observed in the spectrum which indicates that no impurities are present in the sample. The powder characterization conclude that nano crystalline phase pure yttria synthesized by the single step auto igniting combustion is stable at relatively high temperatures and transmit electromagnetic radiations in the UV-Visible and mid IR range. Pellet sintered via resistive coupled microwave heating showed high density, reduced grain size and minimum porosity, which attributes to the improved transmittance of 80% in the Uv-visible and 81% in the mid IR range. The studies on dielectric properties of the sample reveal that as La<sup>3+</sup> ion concentration increases the dielectric constant and dielectric loss of the samples increase which attributes to the increased sinterability and improved transmittance. The microhardness of LY5<sub>RM</sub> is 10.05 GPa which is found greater than that of pure yttria. Thermal conductivity of the samples decreases with La<sup>3+</sup> concentration. The results clearly indicate that the novel resistive coupled microwave sintering can effectively and economically be used to fabricate high quality infrared transparent windows and the presence of La<sup>3+</sup> ion in the matrix considerably improves the hardness.

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