**Studies on highly dense pure YIG polycrystalline ceramics fabricated by tape-casting method**

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**Abstract**

Pure phase $Y_3Fe_5O_{12}$ (YIG) ceramics was successfully produced by tape-casting forming process and one-step solid-state reaction method. With the sintering temperature above 1100 °C, the pure phase YIG ceramics was synthesized with no YIP or Fe$_2$O$_3$ phase in XRD patterns. YIG ceramic sintering at 1400 °C for 10 h showed a clear grain structure with an obvious grain boundary, and no pores were observed in the SEM images. YIG ceramics in this paper has a high relative density which was 99.8% and the saturation magnetization was 28.2 emu/g at room temperature. The hysteresis loss at temperatures of 230-360 K was smaller than 10 mJ/kg. The $\tan\delta_e$ was nearly zero at 6~7 GHz and 11~12 GHz, showing that it can be used as a good material for microwave applications. In
addition, the low values of $\tan \delta_e$ and $\tan \delta_\mu$ indicates that it may have a good electromagnetic wave absorption ability.

**Keywords**
YIG ceramics; tape-casting; microwave; hysteresis loss; magnetodielectric ceramics

1. Introduction
Yttrium iron garnet, $Y_3Fe_5O_{12}$ (YIG) has a bcc cubic structure and belongs to the space group $Ia-3d$ (230). Furthermore, per unit cell (1895.6 Å$^3$) contains 8 formula units, where $Y^{3+}$ ions occupy the eight-coordinate (O$^2-$) dodecahedral sites, while $Fe^{3+}$ ions partially take up six-coordinate octahedral and three-coordinate tetrahedral sites. YIG materials and its modifications by doping as soft ferrites are suitable for magnetic microwave devices with high frequency and low microwave loss [1,2]. Microwave filters and oscillators based on YIG have a wider electric tuning range in frequency than other candidates including ferroelectric and varactors [3]. Multiferroic magnetoelectric materials developed on YIG exhibit both ferroelectricity and ferromagnetism, and typically yield much larger magnetoelectric coupling response than natural multiferroic single-phase compounds, making them suitable for magneto-electric applications [4]. Faraday rotators with substituted YIG films or bulk crystals own large Faraday Rotation capacity, low saturation magnetization and low optical absorption loss, showing high potential applications in magneto-optical devices [5,6].

Pure polycrystalline Bi-substituted YIG thin films were grown on Corning glass substrates by an original CVD technique and its Faraday rotation properties were studied [7]. Bi-substituted Y-Yb mixed iron garnet single crystals were grown by the flux method and the specific Faraday Rotation, optical absorption coefficient and the
saturation magnetization were investigated [5]. Nowadays, polycrystalline YIG (powder or bulk ceramic) are generally synthesized through conventional solid-state reaction (CSSR) method. For YIG ceramics, properties of density, purity and microstructure have great impact on the magnetic properties and therefore the performance of its microwave devices. However, some secondary phases (\(\text{Fe}_2\text{O}_3, \text{YFeO}_3, \text{YFe}_2\text{O}_4\)) are usually coexisted with YIG fabricated by CSSR method [8,9]. The growth kinetic and reaction mechanism of \(\text{Y}_2\text{O}_3\text{-Fe}_2\text{O}_3\) system were studied by different diffusion couples. And, the results showed that the formation of YIG phases were highly depended on pre-formed YIP phases and the diffusion of \(\text{Fe}^{3+}\) cations [10-12]. By adding excess 8-10 wt\% \(\text{Fe}_2\text{O}_3\), 99\% of YIG ceramic phase was achieved. What’s more, YIG with properly excess \(\text{Fe}_2\text{O}_3\) could be used for high frequency tunable dielectric resonator antenna (DRA) [13,14]. Recently, YIG ceramics with about 98-99\% of theoretical density were fabricated by solid-state reaction method [15,16]. While, new fabrication methods are pursued for much higher dense and purity fine-grained YIG ceramics.

In this work, high-dense pure YIG polycrystalline ceramics were fabricated by tape-casting forming process and one-step solid-state reaction method. Phase purity, relative density and micro-morphology depends on sintering temperature and holding time were studied. The activation energy was calculated from the linear Arrhenius plots. XPS was performed to check valance states of Fe and O atoms, as well as EPR spectrum was introduced to check O vacancy. Thereafter, magnetic and dielectric properties were chraccterized for the sample sintered at 1400 °C for 8h.

2. Material and methods

Commercially available powders of \(\text{Y}_2\text{O}_3\) (99.99\%) and \(\text{Fe}_2\text{O}_3\) (99.99\%) were used as raw materials without further purifying. Powders were precisely weighted in accordance with
the stoichiometry of $Y_3Fe_5O_{12}$ and then added into the mixed solvent of ethanol and xylene. 
Besides, proper amount of TEOS (Tetraethyl Orthosilicate) as sintering aid and fish oil as dispersant were put into the mixture. Then, the mixed slurry was planetary ball-milled for 24 h. Subsequently, some binder (Polyvinyl Butyral), plasticizers (Polyalkylene Glycol & Butyl Benzyl Phthalate) and defoamers (n-butyl alcohol & ethylene glycol) were added and ball-milled for another 24 h.

Thereafter, the tape-casting forming process was carried out. The distance between the blade and tape was 500 μm and the casting speed was 500 mm/min. The prepared slurry on the tape was dried at room temperature, and then cut into slices. A proper number of layers of slices were stacked and compressed under 20 Mpa at 120 °C to form into green body. The green body was calcined at 800 °C in an oxygen atmosphere to remove organic additives, and then sintered at various temperatures in a furnace under air atmosphere. Finally, the obtained YIG ceramics were prepared for characterization.

The phase identification of the fabricated ceramics was performed on an XRD system (Bruker, Germany) using Cu $K\alpha$ radiation ($\lambda=1.5406$ Å). The surface micro-morphology was carried on a field-emission SEM (JSM-6700F, JEOL, Japan). The XPS was conducted by using an X-ray photoelectron spectrometer (ESCALAB 250X, Thermo Scientific, USA). The EPR was conducted by using an electron paramagnetic resonance instrument (A300-10/12, Bruker, Germany). The bulk density was measured using the Archimedes’ method. The magnetic hysteresis loops were obtained by a vibrating sample magnetometer (VSM, QD, USA). The magnetic and dielectric properties were determined with a microwave network analyzer (PNA-N5244A, Agilent, USA).

3. Results and discussion
3.1. XRD

Fig. 1 shows the XRD patterns of YIG ceramics fabricated by combining tape-casting forming process and solid-state reaction method at various sintering temperatures in air atmosphere. It can be clearly observed that the ceramics with sintering temperatures above 1100 °C are all YIG pure phases (PDF# 71-2150). According to Ali’s report, YIG ceramics prepared through dry-pressing forming process generally contain unwanted phase of YIP even with 1250 °C sintering temperature [10]. It is suggested that YIP is formed as an intermediate phase and will further react with Fe₂O₃ to produce single-phase YIG by diffusion process [12]. It is suggested that Fe³⁺ cations diffuse and initiate the Fe-Y atomic bonding. Then, with Fe³⁺ cations continuously diffusing, the YIP layer covered on the surface of Y₂O₃ particles turned to be YIG while the inner Y₂O₃ changed into YIP. Considering the saturated vapor pressure of Fe and Y calculated by Eq. (1) and listed in Table 1, it can be seen that Fe has a larger saturated vapor pressure than Y, which signifies that the iron is more volatile in this system. If Fe₂O₃ is inhomogeneous in this system, high concentration of Fe element means more evaporation at high temperature. So, the existence of YIP impurity is highly depended on the inhomogeneous and evaporation of Fe₂O₃. Compared with dry-pressing forming method, our tape-casting forming method provide much better homogeneous mixture of Y₂O₃ and Fe₂O₃ particles, thus leading to the highly pure YIG ceramics.

\[ \log(p/\text{atm}) = A + B/T + C \cdot \log(T) + D/T^3 \]  

(1)

Here, coefficients in the equation can be obtained from reference [17] and listed in Table 1.

**Table 1.** Vapor pressure of metallic elements Fe and Y in the solid state at about 1800 K.

| Element | A     | B       | C      | D      | T(melt)/K | p/atm       |
|---------|-------|---------|--------|--------|-----------|-------------|
| Fe      | 7.1   | -21723  | 0.4536 | -0.5846| 1808      | $3.652 \times 10^{-4}$ |
Fig. 1. The XRD patterns of YIG ceramics sintered under various temperatures in air atmosphere.

3.2. SEM and XPS

Figs. 2(a-f) show the SEM images of YIG ceramics prepared at various sintering temperatures for 10 h and the evolution of morphologies is in accord with Ostwald ripening process. Along with the sintering temperature increasing from 900 to 1400 °C, driven by the sintering kinetics, little grains gradually adhere to each other with grain boundaries and grow to be big grains to reduce the interfacial area, as a result, gaps and pores between grains are gradually eliminated at the same time. Samples sintered at 1400 °C exhibit no pores existed in both grains and boundaries, indicating a quite high densification, as
discussed in Section 3.3. Grain size in Fig. 4(f) is non-uniform and varies from several microns to several dozens of microns, which can be ascribed to the asynchronous growth of grains at different regions and thereof migration of grain-boundaries, as shown in Figs. 2(a-e).

![Fig. 2. SEM images of YIG ceramics sintered at (a) 900 °C, (b) 1000 °C, (c) 1100 °C, (d) 1200 °C, (e) 1300 °C and (f) 1400 °C for 10 h.](image)

According to Chen’s report, Fe$^{2+}$ ions and oxygen vacancies may exist in ceramics fabricated by high temperature sintering process due to the low oxygen conditions [16], and this can be verified by XPS measurement. Fig. 3 shows XPS spectra of YIG ceramics sintered at 1400 °C for 10 h. The XPS spectrum confirmed the existence of Fe 2p$_{1/2}$, Fe 2p$_{3/2}$, Fe 3s, Fe 3p, O 1s, O 2s, Y 3s, Y 3p$_{1/2}$, Y 3p$_{3/2}$, C 1s, Y 3d, Y 4s and Y 4p signals for the sample as shown in Fig 3(a). In Fig. 3(b), based on the Lorentzian-Gaussian fitting, Fe 2p$_{3/2}$ peak and Fe 2p$_{1/2}$ peak can both be divided into Fe$^{3+}$ and Fe$^{2+}$ peaks which are centered at 711.6 eV, 726.4 eV, 710.2 eV and 724.2 eV. Meanwhile, O 1s spectrum in Fig. 3(c) was divided into two peaks centered at 529.9 eV and 531.6 eV. And the latter peak
indicated the existence of oxygen vacancy which can also be proved by the EPR spectrum, as shown in the illustration in Fig. 3(c).

![EPR spectrum and XPS spectra](image)

**Fig. 3.** XPS spectra in (a) the survey range, (b) Fe ion and (c) O ion in YIG ceramics sintered at 1400 °C for 10 h.

### 3.3. Relative density and activation energy

The sintering kinetics, namely densification kinetics, of YIG ceramics could be characterized by the relationship between the relative density and sintering conditions (temperatures and holding times). Densities of YIG ceramics sintered at various
temperatures and holding times were measured by Archimedes’ method and were plotted in Fig. 4(a). Obviously, higher sintering temperature and longer holding time did lead to higher relative density. Besides, the relative density increased significantly along with the temperature below 1375 °C, while it increased slowly when the temperature further raised. For samples sintered at 1450 °C for 8 h, the relative density reached the highest value of 99.8%, which was bigger than that values reported in references [15] and [18]. The high relative density indicated extremely low residual porosity and highly dense microstructure. Densification kinetics can be interpreted by Arrhenius equation: [10,18,19]

\[ \ln K = \frac{-E_a}{RT} + C \]

where, \( E_a \) is the activation energy for YIG ceramic densification, \( R \) is the gas constant (8.314 J/mol/K), \( T \) is the absolute temperature, \( C \) is the constant, and \( K \) is the rate constant depending on temperature. Densification rate versus reciprocal temperature was plotted in Fig. 4(b), where rate constant \( K \) was related to relative density and calculated from Fig. 4(a). By linear fitting, the value of \( E_a \) was determined to be 183.81 kJ/mol, which was comparable with the activation energy of 169 kJ/mol for lattice diffusion or grain boundary formation [18] and much smaller than the energy of 484 kJ/mol for YIG formation from YIP [10]. It can be drawn that the mechanism of YIG phase formation and ceramics densification was the result of homogeneous diffusion probably assisted by the sintering aids of SiO₂ (products of TEOS decomposition) located at grain boundaries.
Fig. 4. (a) Relative densities of YIG ceramics sintered at various temperatures and holding times. (b) The activation energy analysis for YIG ceramics.

3.4. Intrinsic magnetic property

The magnetic properties of YIG ceramics sintered at 1400 °C were studied by the static hysteresis loops under various temperatures, as shown in Fig. 5(a). The well-defined hysteresis loop can be clearly observed at 30 K, while turned to be narrower along with the increase of temperature. The saturation magnetization \(M_s\) and hysteresis loss (area of the hysteresis loop) depending on temperature were plotted in Fig. 5(b). The value of \(M_s\) decreased from 38.9 emu/g down to 25.3 emu/g as the temperature increased from 30 K up to 360 K. And, the trend could be ascribed to the enhanced thermal vibrations of atoms and lattices under high temperatures, and hence magnetic dipoles were difficult to be aligned [20]. It was worth mentioning that the room-temperature (300 K) value of \(M_s\) was 28.2 emu/g, which was a little higher than that value (27.4 emu/g) reported in previous literature [15]. The result indicated that our YIG ceramics exhibited much higher relative density and more compact microstructures. Hysteresis loss, characterized by the area of the hysteresis loop, generally decreased with temperature. However, in this work, there was an upturn
point at 230 K (4 mJ/kg), and the mechanism needed to be further studied. Besides, hysteresis loss at 30 K was as high as 238.8 mJ/kg, while values of that at temperatures of 230-360 K was smaller than 10 mJ/kg.

Fig. 5. (a) Magnetic hysteresis loops and (b) $M_s$ and hysteresis loss of YIG ceramics sintered at 1400 ºC for 8 h at various test temperatures.

3.5. Electromagnetic parameters

Fig. 6 shows zero-field complex permittivity properties of home-made YIG ceramic circle (the fabricated YIG ceramics were grinded into powder, homogenously mixed with paraffin by a mass ratio of 3:7, and thereafter formed into a 2 mm thick circle with an inner diameter of 3 mm and an outer diameter of 7 mm). Along with the frequency increasing from 2 to 18 GHz, the changing trend of real part $\varepsilon'$ (the ability for the storage of electrical energy) and the imaginary part $\varepsilon''$ (the dissipation capacity of electrical energy) is just the opposite, and some relaxation peaks are clearly observed as a result of multiple relaxation processes, as shown in Fig.6 (a). [21] In addition, the dielectric loss tangent $\tan\delta_\varepsilon$ determined by $\varepsilon''/\varepsilon'$ fluctuates with the frequency, and interestingly, $\tan\delta_\varepsilon$ at about 6~7 GHz and 11~12 GHz is extremely low suitable for practical microwave devices, as shown
in Fig. 6(b). As we know, the dielectric loss is ascribed to conduction loss and polarization relaxation loss. And, polarization relaxation loss can be characterized by the Cole-Cole semicircle, that is, a simple semicircle in the $\varepsilon'\varepsilon''$-coordinate plane means a polarization relaxation process. As shown in Fig. 6(c), there are three semicircles suggesting that the dielectric loss is determined by multiple dielectric relaxation losses, and this result is in accordance with relaxation peaks in Fig. 6(a). Future work will be focused on dielectric properties of YIG ceramics under applied magnetic field in a wide range of frequency.

![Fig. 6](image)

**Fig. 6.** Frequency dependence of (a) $\varepsilon'$ and $\varepsilon''$, (b) $\tan\delta_{\varepsilon}$, and (c) Cole-cole semicircle of YIG ceramics sintered at 1400 °C for 8 h.

For complex permeability, the real part $\mu'$ (the ability for the storage of magnetic energy) and the imaginary part $\mu''$ (the dissipation capacity of magnetic energy) have the same changing trend along with the rise of frequency from 2 to 18 GHz, as shown in Fig. 7(a). Apparently, $\mu'$ and $\mu''$ exhibit three distinct resonance peaks located at about 6.2 GHz, 11.6 GHz and 16.6 GHz in turn, meaning perfect magnetic loss ability at these frequencies, which is well agree with the result of magnetic loss tangent $\tan\delta_{\mu}$ determined by $\mu''/\mu'$, as shown in Fig. 7(b). Generally, the magnetic loss contains exchange resonance, natural resonance and eddy current loss in the investigated frequency from 2 GHz to 18 GHz.
According to previous report, if the eddy current loss is the only reason for the magnetic loss, frequency dependence of $C_0 = \frac{\mu''}{\mu'\omega f}$ should be a constant. [23] In Fig. 7(c), $C_0$ is nearly constant with little fluctuations at the range of 5~18 GHz, suggesting that the eddy current loss dominates the magnetic loss but also the exchange resonance may exist. As for lower frequency between 2 GHz and 5 GHz, the natural resonance will always dominate. The electromagnetic wave (EMW) dissipation is both rely on the dielectric loss and magnetic loss. For our YIG ceramics, values of tan$\delta_e$ and tan$\delta_\mu$ were pretty low, indicating that this material may have a good EMW absorption ability, on which future work will be focused.

![Fig. 7](image.png)

**Fig. 7.** Frequency dependence of (a) $\mu'$ and $\mu''$, (b) tan$\delta_\mu$, (c) $C_0$ of YIG ceramics sintered at 1400 °C for 8 h.

**4. Conclusion**

In this study, pure phase YIG ceramics were successfully produced by tape-casting forming process and one-step solid-state reaction method. The microstructure of the YIG ceramics sintering at 1400 °C for 10 h showed a clear grain structure with an obvious grain boundary, and no pores were observed in the SEM images. At 1450 °C with a holding time of 8 h, the relative density of the YIG ceramic was nearly 99.8%. The activation energy was calculated to be 183.81 kJ/mol. The saturation magnetization was 28.2 emu/g at 300 K.
And, hysteresis loss at 30 K was as high as 238.8 mJ/kg, while values of that at temperatures of 230-360 K was smaller than 10 mJ/kg. The $\tan\delta_e$ was nearly zero at 6~7 GHz and 11~12 GHz. The low values of $\tan\delta_e$ and $\tan\delta\mu$ indicates that it may have a good electromagnetic wave absorption ability. This work provided a new fabrication method for much higher dense and purity fine-grained YIG ceramics. And the future work will be focused on the study of the uptown point (230 K) on the hysteresis loss chart, the dielectric properties under applied magnetic field in a wider range of frequency and the EMW absorption of the dense YIG ceramics.

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