Dielectric and Piezoelectric Studies of Dysprosium Doped BZT-BCNT Perovskite Ceramic System for Sensors and Actuators Applications

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Research Article

Keywords: Perovskite, BZT-BCNT, Dy-doping, Dielectric and Piezoelectric Studies, Ceramics, Sensors and Actuators

DOI: https://doi.org/10.21203/rs.3.rs-291438/v1

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Abstract

Perovskite structured Dy doped BZT-BCNT system has been investigated with variation of Dy doping through the solid-state reaction method. X-ray diffraction (XRD) and microscopic studies confirm the results in the rhombohedral stoichiometry without any secondary phases, and uniform grain growth, respectively. Tolerance factor and electronegativity also confirm the presence of Dy doping in BZT-BCNT ceramic system with a stable rhombohedral crystallinity forming vacancy sites for oxygen ions. Optimum piezoelectric properties in non-Pb piezoelectric ceramic systems were obtained due to trivalent donor Dy doping at A-site and pentavalent donor Nb at B-site are possible members for actuator and energy harvesters. Dielectric and piezoelectric studies signifying the rhombohedral perovskite behavior of this lead-free ceramic system confess that the proposed investigation could yield further potential piezoelectric sensors and actuators applicative futuristic studies.

1. Introduction

Piezoceramics engrosses the polycrystalline materials of perovskite form that are being explored in the contemporary times to explicate the Lead-free alternatives [1–3] with high dielectric constants [4], and piezoelectric (PE) response [5]. These materials are widely employed in various electronic devices due to their promising functional, structure, and microstructure (phase, lattice parameters, density, and grain size) properties that can be tailored through A-site or B-site engineering by soft or hard dopants, respectively [6, 7]. However, the efficacy of the system is dependent on the versatility of the sample fabrication, shape, sintering temperature, and dopant [1, 8–11]. Additionally, it is necessary to articulate the precise polarization rotation without having to overcome the large energy barrier [12–14] over the tetragonal (P4mm), and rhombohedral (R3m) phases at phase transition region where the orthorhombic (Amm2) and cubic (Pm3m) phases also coexist [15, 16]. (i) Perovskite (ABO₃) structured-titanates [17], (ii) Tungsten bronze (AB₂O₆) structured-niobates/tungstates [18], and (iii) Bismuth layer structured-oxides (BLSO) established by Aurivillius (A₄B₃O₁₂) family [19] possess adequate PE properties.

The dielectric and PE properties of the sintered ceramic powder are attributed to the volatile and precarious nature of the dopants, conjoining with the sintering conditions, microstructural, temperature dependent rhombohedral–tetragonal phase transition, surfactant effect on grain size, and composition ratios [20–22]. They are usually accounted for in terms of dielectric loss, tolerance factors, and electronegativity of the system. Enhanced PE coefficients are realized from the crystallinity and phase convergence nature of piezoceramic materials such as Barium Titanate (BaTiO₃ or BT), Barium Potassium Niobate (Ba₀.₅K₀.₅NbO₆ or BKN), Bismuth Potassium Titanate (Bi₀.₅K₀.₅TiO₃ or BiKT) and their modified compositions [23]. Notably, the electrical and mechanical energies of Barium-Zirconium-Titanate–Barium-Calcium-Titanate (BZT–BCT) is highly tunable [28–31] compared to the single phase BaTiO₃ owing to its binary-pseudo ferroelectric system [32]. The morphotropic phase boundary (MPB) separates the rhombohedral (Zr-rich near BZT site) and tetragonal (Ti-rich near BCT site) phases. In
summary, the dopants of soft and hard ceramics in Lead-free systems play a vital role in evincing the PE properties leveraging electronic structure and high polarizability [33].

Soft [34] and hard [35] electroceramics are constituted by doping donor (at A-site) and acceptor (at B-site) cations of similar/(higher, and lower) ionic radii, respectively. The soft ceramics hold good for sensing and actuating applications [27], while the hard ones are principally used in high voltage, mechanical loads, and power applications [36]. The effect of modified composition and donor/acceptor rare-earth ions on the MPB of BZT-BCT has been explored to yield the optimal PE properties [37–44]. Nevertheless, a very few investigations were available on Nb-doped BZT-BCT [41, 43, 44] (i.e., BZT-BCNT) out of which, in particular, Dy$^{3+}$ dopant system is not available to the best of our knowledge.

In this study, Dy$^{3+}$ doped 0.48(BZT)-0.52(BCNT) series, has been synthesized by a solid-state reaction method. The stoichiometric composite structure effect of BZT-BCNT in the presence of donor-Dy at A-site over rhombohedral structure is evaluated. Furthermore, the stability of that unit cell corroborated by the calculated tolerance factor and oxygen vacancy formation is verified by the calculated electronegativity. The effect of the presence of Dy at A-site, and Nb at B-site over microstructure and functional properties is characterized by the XRD, apparent density, dielectric, and piezoelectricity confessing the optimum PE properties of the proposed structure of non-Pb PE electroceramic system. This work prescience the capabilities of suggested material for specified prospective piezoelectric energy harvesters, multi-layer capacitors, and actuator applications.

2. Experimental Details

Stoichiometric composition of Dy doped BZT-BCNT is given as

$$0.48 \left[ Ba_{1-x} Dy_x \left( Zr_{0.28} Ti_{0.72 - \left( \frac{x}{4} \right)} \right) O_3 \right] - 0.52[ (Ba_{0.72} Ca_{0.28}) (Nb_{0.05} Ti_{0.9375}) O_3] ,$$

where the samples were synthesized by conventional solid-state reaction route varying the Dy concentration ($x$) as 0, 0.2, 0.4, 0.6, 0.8, 1 mol%. Analytical reagent grade (99.99% purity) starting materials BaCO$_3$, Dy$_2$O$_3$, ZrO$_2$, TiO$_2$, CaCO$_3$ and Nb$_2$O$_5$ (Sigma Aldrich, USA, and Germany) were mixed according to the stoichiometric formula (Table 1), and ball-milled these respective batch systems by using zirconia balls and ethanol in a polyethylene jar for 12 h. These ball-milled powders were filtered and dried at 100°C for 24 h, followed by calcination at 1150°C for 4 h. The set powders were ball-milled by using zirconia balls and ethanol in a polyethylene jar for 12 h to obtain agglomerate-free powders that were filtered and dried at 100°C for 24 h. The batch calcined powders were mixed with 5 wt% Polyvinyl Alcohol (PVA) binder, and compacted into pellets (with 12 mm in diameter and 2–3 mm in thickness by using a steel-die and conventional hydraulic press with a uniaxial pressure of 700–900 kg/cm$^2$), and sintered the respective green bodies at 1450°C for 3 h with a heating rate of 5°C/min, and after the sintering process, the samples were cooled to room temperature in the furnace. Dy doped BZT-BCNT
powder ceramic systems were analyzed by using X-ray diffraction (XRD) with a Bruker Endeavor X-ray diffractometer model D4/MAX-Bat at a scanning rate of 0.02 °/min over a range of Bragg angles 2θ between 20 and 70°. The apparent densities of the respective sintered samples were measured through the Archimedes method. As sintered ceramic surfaces were analyzed with scanning electron microscopy (SEM; JEOL model JSM 840A). An air-dry silver paste was applied on the polished pellet surfaces to form the electrodes. The electrode samples were characterized for dielectric constant (ε), and dissipation factor (tan δ) using a 1260A Impedance Analyzer. The samples were electrically poled in a silicon oil bath at 100°C by applying a dc field of 20 kV/cm. After aging for 1 day, the poled samples were characterized for piezoelectric properties by using a Berlincourt piezo-d-meter.

3. Results And Discussion

3.1 X-ray Diffraction

Fig. 1(a) shows the powder XRD patterns of Dy doped 0.48BZT-0.52BCNT ceramics. The XRD peaks were indexed to a perovskite-type of rhombohedral structure, which shows a single rhombohedral phase as can be evidenced in the BZT-BCT phase diagram [47]. XRD patterns show sharp crystalline peaks that show the Dy diffusion in the BZT-BCNT lattice to form a solid-solution with the single-phase rhombohedral perovskite structure and no secondary phases are found that can be confessed by the tolerance factor in section 3.2. However, several reports confirm the rhombohedral-tetragonal phase transition with the variation in BZT:BCT ratio, and/or dopant modifications at MPB [48–50]. To verify, we have plotted the enlarged XRD peaks for (220), (101), and (200) planes as shown in Figs. 1(b-d), respectively. The absence of deconvolution in the planes also supports the clear existence of rhombohedral crystallinity for Dy doped BZT-BCNT. It is well known that the phase genesis and evolution from oxide precursors obtained by solid-state reaction method is temperature dependent. XRD results indicate the formation of rhombohedral symmetry throughout the series attesting the Dy$^{3+}$ ions at A-site and Nb$^{5+}$ ions at B-site resulting in the optimum diffused solid solutions into the BZT-BCT lattice, respectively. The compositional stoichiometry could be achieved due to the calcination and sintering process. The substitution of Ba$^{2+}$ by trivalent Dy$^{3+}$ at A-site in BZT lattice and pentavalent Nb$^{5+}$ at B-site in BCT lattice that has a deep effect on the structural and functional properties of the electroceramic system.

The rhombohedrality is justified by the intensified sharp peaks as the Dy concentration increases, in which the Dy$^{3+}$ ions substitute Ba$^{2+}$ and create vacancies at B-site. BZT phase (either tetragonal, orthorhombic, rhombohedral, or cubic symmetry) depends upon the BZT:BCT ratio in the binary system. The Dy-doping enhanced the rhombohedrality showing a predominant single rhombohedral phase. Furthermore, the diffraction peaks shift to the lower diffraction angles, signifying the stable large donor trivalent ions (r > 0.94 Å) would occupy A-site while the small donor trivalent ions (r < 0.87Å) positioned at B-site, and the rest (0.87 Å < r < 0.94 Å) could possibly occupy A-site or B-site [51].

3.2 Tolerance Factor
The tolerance factor (Table 2) measures the stability of the perovskite unit cell of the ABO₃ configuration. According to Goldschmidt, the ionic radii in ABO₃ must fulfill the following categories: (i) the most stable structure is tetragonal \((t \geq 1)\) (ii) cubic \((t \approx 1)\), and (iii) rhombohedral or orthorhombic symmetries \((t < 1)\), respectively [48–50,52].

\[
t = \frac{r_A + r_O}{\sqrt{2}(r_A + r_O)}
\]

Where \(t\) is the tolerance factor, \(r_A\) is the ionic radii of the cations at A-site, \(r_B\) is the ionic radii of cations at B-site, and \(r_O\) is the ionic radii of the oxygen at respective sites in the perovskite unit cell. Tolerance factor values are tabulated in table 2. The tolerance factor is ranged in rhombohedral symmetry, and the rhombohedrality is enhanced with Dy doping. It is well known that the trivalent ions could occupy B-site \((r < 0.87 \text{ Å})\) or A-site \((r > 0.94 \text{ Å})\), respectively. Dy³⁺ \((r = 1.083 \text{ Å})\) substitutes Ba²⁺ \((r = 1.61 \text{ Å})\), predominantly enhancing the rhombohedrality in the lattice. Furthermore, no secondary phases are evidenced from the XRD studies. Hence, it confirms the formation of stable rhombohedral (R3m) perovskite structure.

### 3.3. Electronegativity

Furthermore, the averaged electronegativity difference \((e)\) is an added significant parameter to weigh the stability and bonding in the perovskite.

\[
e = \frac{(\chi_A - \chi_O) + (\chi_B - \chi_O)}{2}
\]

where are \(\chi_A, \chi_B,\) and \(\chi_O\) the respective electronegativities of A-site, B-site cations and O²⁻ anions, respectively in the perovskite unit cell. It is evident from table 2 that there is no distortion noticed in rhombohedral Dy doped BZT-BCNT structure. The \(e\) of Dy doped BZT-BCNT ceramics equation can be written as

\[
e = \left\{ \left[ (0.48(1 - y)\chi_{Ba-O} + y\chi_{Dy-O}) + 0.52(0.72\chi_{Ba-O} + 0.28\chi_{Ca-O}) \right] + \left[ (0.48(0.28\chi_{Zr-O}) + (0.72 - (X/4)\chi_{Ti-O})) + (0.52(0.05\chi_{Nb-O} + (1 - (5/4) * y)\chi_{Ti-O})) \right] \right\} / 8
\]

Electronegativity (Table 2) evaluates the oxygen vacancies and shows the tendency of the atomic bonding depending on the electron-ionization energy. It is well known that the lanthanides tend to replace
Ba$^{2+}$ rather than Ti$^{4+}$ and make it a donor or acceptor doping depending on its ionic radii in the unit cell. Electronegativity is the difference between the sum of cations and anions that reflects the tendency of the bonded atom to attract shared electrons. Therefore, the greater the electronegativity, the greater the attraction of electrons, and the nature of bond is determined by the differences in the electronegativity. Electronegativity is caused as the chemical bonding between the atoms that are formed which could be positive or negative electrons that determines the bond type i.e., ionic, covalent, or metallic. If the electronegativity is below 0.5, then the bond is said to be a non-polar covalent. If the electronegativity is between 0.5 and 2.1, it is said to be a polar covalent. If the electronegativity is more than 2.1, it is said to be an ionic in nature. The electronegativity of Dy doped BZT-BCNT is in the range of 0.6354 to 0.6316, which attests the polar covalent bonding. The multiple cations in the BZT-BCNT rhombohedral perovskite ceramic system resulted in sharing of Dy$^{3+}$ and Ca$^{2+}$ ions in the place of Ba$^{2+}$ (A-sites) at BZT and BCNT, respectively, also, Nb$^{5+}$ replacing Ti$^{4+}$ at B-site of BCT resulted in a polar covalent bonding of the material, which can stabilize the piezoelectric (PE) perovskite phase [53, 54].

3.4. Scanning Electron Microscopy (SEM)

The microstructure of Dy doped BZT-BCNT ceramics is presented in fig. 2(a-c) for the 0.2, 0.8, 1 mol%. It is evident that, small concentration of pentavalent Nb enhances grain growth, while Dy doping supported the domain reorientation in aligning and densifying the coarse grains. The oxygen vacancies supported the charge balance in the BZT-BCNT ceramic system due to the sintering process that in turn enhanced the density. The sintering process (temperature and time) caused enhancement in grain growth with the increase in Dy content. The homogeneously dense grains attested by SEM influenced the net polarization up to 0.8 mol% and thereafter attained saturation.

3.5. Density studies

Fig. 3 depicts the variation of density with Dy concentration for sintered ceramic samples. It is well known that the composition of the materials dictates the optimum sintering temperature [55, 56]. Initially, the density intensifies with increase in Dy concentration up to 0.8 mol% and reduces thereafter. One of the prominent reasons for lowering the density is the incomplete removal of pore or trapped pore present in the grain. Further, the decrease in density for 1 mol% suggests the limit of addition of dopant to BZT-BCNT ceramics. Therefore, the density study results indicate that the sintering temperature of 1450 °C is optimum for 0.8 mol%.

3.6. Dielectric studies

Fig. 4 shows the dielectric properties of Dy doped 0.48BZT-0.52BCNT ceramics characterized at 1 kHz. It is observed in Fig. 4(a) that the dielectric constant of the perovskite structured BZT-BCNT ceramic system increases gradually with temperature and reaches dielectric maximum at 98 °C which is the Curie transition temperature ($T_c$), and with additional rise in temperature the dielectric constant declines for pure. It can be witnessed that for Dy doping (0.2 to 1.0 mol%) of BZT-BCNT exhibited a similar trend in
this ceramic system. In addition to that, it is observed that maximum dielectric constant is achieved at a lower temperature for Dy doped BZT-BCNT systems up to 0.8 mol%, compared to the pure sample and thereafter the trend reverses. The curie transition temperature in the series has witnessed a similar trend. There are many factors that contribute to the high dielectric constant like, modifications in the grain size and grain boundary, change in density, release of internal stress, and the defect and domain wall motion [57].

Fig. 4b shows the variation of dielectric loss of Dy doped BZT-BCNT ceramics with temperature. It is detected that at room temperature the dielectric loss decreases gradually with rising Dy doping up to 0.8 mol% (optimum) and thereafter increases at room temperature. Furthermore, it is seen that with the increase in temperature, the dielectric loss decreases and reaches minimum at ~ 80°C for all the Dy doping BZT-BCNT ceramic systems, which might be owing to the manifold cationic arrangements or saturation in the perovskite ceramic system.

In the rhombohedral symmetry, as Dy doping supported the space charge polarization due to the donor Dy\(^{3+}\) dopant in BZT lattice and pentavalent Nb\(^{5+}\) in the BCT lattice by balancing the charge and oxygen vacancies in the present system [58,59]. The polarization of a dielectric material is influenced by the dipolar, electronic, ionic, and interfacial interactions in the unit cell. The incorporation of Dy in the BZT and Nb in BCT drastically influenced the polarization in attaining optimum electrical properties that resulted in the increased dielectric constant. The dielectric constant (\(e_{RT}=1342\)) at room temperature, dielectric maximum (\(e_{Tc}=10667\)) at the Curie Temperature \(T_c\) (81°C) is found to be optimum for 0.8 mol% at the frequency of 1 kHz. It was reported that there was a mixed trend of the dielectric loss and Curie temperature with Dy incorporation in BaTiO\(_3\) [60], similarly, we confirm the same trend in this perovskite BZT-BCNT ceramic system.

### 3.7. Piezoelectric studies

Fig. 5 shows the piezoelectric coefficient \(d_{33}\) as a function of Dy concentration. It can be observed that with the increasing Dy concentration, the piezoelectric constant \(d_{33}\) increases up to 0.8 mol% and attains a maximum value 368 pC/N and then decreases sharply with further Dy concentration. It is believed that the observed high piezoelectric properties should be ascribed to the phase, polarization due to multiple cations, dopants, enhanced grain size (possible net polarization from respective domains), dense ceramics [61]. Further, lattice distortion provoked by Dy doping content may be advantageous to ferroelectric domain reorientation during poling process have some bearing on the improvement of piezoelectric properties. However, with the increasing of Dy content to 1 mol%, Dy ions should occupy the A-site of Ba\(^{2+}\), which might cause deformation in the perovskite ABO\(_3\) lattice to make a phase transition completed, resulting in a reduction of piezoelectric coefficient \(d_{33}\) [9, 61]. This shows that a proper addition of Dy induces a more significant piezoelectric activity in these perovskites, which can be chiefly ascribed to nature of the dopant and its content, the densification and grain morphology of the ceramic system.
4. Conclusion

We report the perovskite structured Dy doped BZT-BCNT system resulting in the rhombohedral stoichiometry without any secondary phases being observed. Superior net polarization due to homogeneous dense coarse grains, which not only enhanced the electrical properties but also favoring the piezoelectric applicative investigation while exhibiting the rhombohedral phase of Dy doped BZT-BCNT ceramic system. Tolerance factor and electronegativity confirm the stable rhombohedral crystallinity in forming vacancy sites by oxygen ions and polar covalent bonding in the perovskite phase, respectively. Optimum dielectric and piezoelectric properties are observed at 0.8 mol% due to donor Dy$^{3+}$ doping at A-site and donor Nb$^{5+}$ at B-site in the non-Pb perovskite structured (BZT-BCNT) ceramic system. The obtained results could be helpful in developing sensors and actuators.

Declarations

Acknowledgements & Funding Information

This work is supported by the research grants funded by Fondecyt (Programa Fondo Nacional de Desarrollo Científico y Tecnológico (FONDECYT) de la Agencia Nacional de Investigación y Desarrollo, Chile) Regular (N°1110583), Fondecyt Post-doctoral (N°3200832), Proyecto FONDEF, CONICYT, Chile. (ID15I10312), Chile, and CORFO project (N° 15IPPID-45708). The authors would like to sincerely thank the University of Concepcion, Chile and Andhra University, India for their enormous support and facilities provided. This work is also supported by the Manipal Academy of Higher Education (MIT-2017, 2020), Karnataka, India. The authors also like to be grateful towards the efforts offered by Mr. Ranganathan, Mr. Krishnamurthy and Ms. C. N. Devi for their technical assistance and valuable suggestions extended during this work.

Conflicts of interest/Competing interests

Authors declare no conflict/competing interests as per the guidelines of the journal.

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Tables
Table 1: Stoichiometric compositions of the Dy doped BZT-BCNT ceramic system.

| Compositions | 1 - z [Ba_{1-z}Dy_{z} (Zr_{0.28}Ti_{0.72})O_{3}] - z[(Ba_{0.2}Ca_{0.2}Zr_{0.7}Ti_{0.3})O_{3}] | Formulae |
|--------------|-----------------------------------------------------------------|----------|
| Stoichiometric Formula | 0.48 [Ba_{1-z}Dy_{z} (Zr_{0.28}Ti_{0.72-0.48})O_{3}] - 0.52[(Ba_{0.2}Ca_{0.2}Zr_{0.7}Ti_{0.3})O_{3}] | Where Dy_{z} = 0, 0.2, 0.4, 0.6, 0.8, 1 mol%, and Nb_{y} = 5 mol% |

BZT-BCNT

0.48 [BaCO_{3} + (0.28ZrO_{2} + 0.72TiO_{2})] + 0.52 [(0.72BaCO_{3} + 0.28CaCO_{3}) + (0.025NbO_{2} + 0.9375TiO_{2})]

0.2% Dy + 0.48 BZT - 0.52 BCNT:

0.48 [(0.98 BaCO_{3} + 0.01 Dy_{0.2}O_{3}) + (0.28 ZrO_{2} + 0.715 TiO_{2})] + 0.52 [(0.72 BaCO_{3} + 0.28 CaCO_{3}) + (0.025 NbO_{2} + 0.9375 TiO_{2})]

0.4% Dy + 0.48 BZT - 0.52 BCNT:

0.48 [(0.96 BaCO_{3} + 0.02 Dy_{0.4}O_{3}) + (0.28 ZrO_{2} + 0.710 TiO_{2})] + 0.52 [(0.72 BaCO_{3} + 0.28 CaCO_{3}) + (0.025 NbO_{2} + 0.9375 TiO_{2})]

0.6% Dy + 0.48 BZT - 0.52 BCNT:

0.48 [(0.94 BaCO_{3} + 0.03 Dy_{0.6}O_{3}) + (0.28 ZrO_{2} + 0.705 TiO_{2})] + 0.52 [(0.72 BaCO_{3} + 0.28 CaCO_{3}) + (0.025 NbO_{2} + 0.9375 TiO_{2})]

0.8% Dy + 0.48 BZT - 0.52 BCNT:

0.48 [(0.92 BaCO_{3} + 0.04 Dy_{0.8}O_{3}) + (0.28 ZrO_{2} + 0.700 TiO_{2})] + 0.52 [(0.72 BaCO_{3} + 0.28 CaCO_{3}) + (0.025 NbO_{2} + 0.9375 TiO_{2})]

1% Dy + 0.48 BZT - 0.52 BCNT:

0.48 [(0.90 BaCO_{3} + 0.05 Dy_{1.0}O_{3}) + (0.28 ZrO_{2} + 0.695 TiO_{2})] + 0.52 [(0.72 BaCO_{3} + 0.28 CaCO_{3}) + (0.025 NbO_{2} + 0.9375 TiO_{2})]

Table 2: Tolerance factor and Electronegativity of Dy doped BZT-BCNT ceramic system.

| Compositions | Ionic Radii/ Electronegativity of A-site cations | Ionic Radii/ Electronegativity of B-site cations | Tolerance Factor | Electronegativity |
|--------------|-----------------------------------------------|-----------------------------------------------|-----------------|-----------------|
| BZT-BCNT     | Ba^{2+}: 1.61Å / 0.89                         | Zr^{4+}: 0.72Å / 1.33                         | 0.8450          | 0.6354          |
| Dy 0.2mol % +| Dy^{3+}: 1.083Å / 1.22                       | Ti^{4+}: 0.605Å / 1.90                        | 0.8445          | 0.6347          |
| BZT-BCNT     | Ca^{2+}: 1.34Å / 1.00                         |                                               |                 |                 |
| Dy 0.4mol % +| O^{3-}: 1.42Å / 3.44                         | Nb^{5+}: 0.74Å / 1.84                         | 0.8440          | 0.6339          |
| BZT-BCNT     |                                               | O^{3-}: 1.42Å / 3.44                         | 0.8435          | 0.6331          |
| Dy 0.6mol % +|                                               |                                               | 0.8431          | 0.6324          |
| BZT-BCNT     |                                               |                                               |                 |                 |
| Dy 0.8mol % +|                                               |                                               |                 |                 |
| BZT-BCNT     |                                               |                                               |                 |                 |
| Dy 1mol %    |                                               |                                               | 0.8426          | 0.6316          |