The Primary Origin of Excellent Dielectric Properties of (Co, Nb) Co-Doped TiO$_2$ Ceramics: Electron-Pinned Defect Dipoles vs. Internal Barrier Layer Capacitor Effect

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Abstract: (Co, Nb) co-doped rutile TiO$_2$ (CoNTO) nanoparticles with low dopant concentrations were prepared using a wet chemistry method. A pure rutile TiO$_2$ phase with a dense microstructure and homogeneous dispersion of the dopants was obtained. By co-doping rutile TiO$_2$ with 0.5 at. % (Co, Nb), a very high dielectric permittivity of $\varepsilon' \approx 36,105$ and a low loss tangent of tan$\delta \approx 0.04$ were achieved. The sample–electrode contact and resistive outer-surface layer (surface barrier layer capacitor) have a significant impact on the dielectric response in the CoNTO ceramics. The density functional theory calculation shows that the 2Co atoms are located near the oxygen vacancy, creating a triangle-shaped 2CoV$_o$Ti complex defect. On the other hand, the substitution of TiO$_2$ with Nb atoms can form a diamond-shaped 2Nb2Ti complex defect. These two types of complex defects are far away from each other. Therefore, the electron-pinned defect dipoles cannot be considered the primary origins of the dielectric response in the CoNTO ceramics. Impedance spectroscopy shows that the CoNTO ceramics are electrically heterogeneous, comprised of insulating and semiconducting regions. Thus, the dielectric properties of the CoNTO ceramics are attributed to the interfacial polarization at the internal insulating layers with very high resistivity, giving rise to a low loss tangent.

Keywords: TiO$_2$; DFT; SBLC; giant/colossal dielectric permittivity; loss tangent

1. Introduction

The intensive investigation of a novel giant dielectric oxide (GDO) showing a high dielectric constant ($\varepsilon'$) has fuelled research in the field of dielectric materials [1–6]. The loss tangent (tan$\delta$) is also an essential factor for capacitor applications. ACu$_3$Ti$_4$O$_{12}$ (A = Ca, Cd, etc.), the famous GDO, is a widely researched GDO because it can show a large $\varepsilon'$ of $\sim 10^4$ over a wide temperature range [3,7–10]. This unusual behavior is owing to the existence of a potential barrier at the grain boundaries (GBs); that is, the Schottky barrier. ACu$_3$Ti$_4$O$_{12}$ ceramics have an electrically heterogeneous microstructure consisting of low conductivity and high conductivity parts. The microstructure is usually analyzed by the brickwork layer model of an internal barrier layer capacitor (IBLC) structure. Therefore, the dielectric properties of many GDOs can be improved by enhancing the electrical properties of the grains and GBs. Although ACu$_3$Ti$_4$O$_{12}$ ceramics can exhibit a large $\varepsilon'$ response, their tan$\delta$ values are very large, which cannot be used in ceramic capacitors [11–17]. The low-frequency tan$\delta$ of ACu$_3$Ti$_4$O$_{12}$ and other GDOs can be decreased by increasing the resistivity at the GBs ($R_{gb}$) via doping ions or designing a small grain-sized microstructure [8,18–20].

Recently, a new promising GDO was reported. By co-doping TiO$_2$ with aliovalent ions into the rutile structure (i.e., Ti$_{1-x}$(Nb$_{0.5}$In$_{0.5}$)$_2$O$_2$), a very high $\varepsilon' > 10^4$ with low tan$\delta \approx 0.02$ can be obtained [21]. The giant dielectric properties, with a low-temperature coefficient...
of the $\varepsilon'$ from 80 to 450 K of the Ti$_{1-x}$\((\text{Nb}_{0.5}\text{In}_{0.5})_x\)O$_2$ ceramics, are much better than those of the ACu$_3$Ti$_4$O$_{12}$ [15,22,23]. The existence of electron-pinned defect dipoles (EPDD) was suggested to be the primary cause of the observed dielectric properties. In this EPDD model, the diamond-shaped ($A = \text{Ti}^{3+}/\text{In}^{3+}/\text{Ti}^{4+}$) and triangular-shaped defects were predicted to be closely correlated, resulting in a low tan$\delta$ and high $\varepsilon'$ with temperature stability. The former and latter defects can be produced by co-doping with acceptor dopant (e.g., Ga$^{3+}$ or In$^{3+}$) and donor dopant (e.g., Nb$^{5+}$ or Ta$^{5+}$) [21,24,25].

Besides the quasi-intrinsic effect of the complex defect dipoles [21,26], extrinsic effects based on interfacial polarization have been widely proposed, such as the surface barrier layer capacitor (SBLC) [27] and IBLC models [28,29]. Although the non-Ohmic contact at the sample–electrode interface (i.e., sample–electrode, SE effect) can have a remarkable impact on the dielectric properties [27], a significant increase in $\varepsilon'$ is usually accompanied by an enormous tan$\delta$ value [24,30]. According to the SBLC and IBLC effects, the insulating layers in co-doped TiO$_2$ polycrystalline ceramics can be formed by doping with an acceptor dopant. At the same time, the semiconducting part can be produced by doping with a donor dopant [29,31]. Thus, the origin of the giant dielectric response in co-doped TiO$_2$ is still under discussion.

Until recently, many co-doped TiO$_2$ systems have been intensively investigated to develop a new co-doped TiO$_2$ system that can exhibit a high $\varepsilon'$ > $10^4$ with low tan$\delta$, such as (Ga, Nb), (Ag, Nb), (In, Nb), (Sc, Nb), (In, Ta), and (Y, Nb) co-doped TiO$_2$ systems [4,25,31–34]. To the best of our knowledge, the preparation, characterization, dielectric properties, and formation of defect dipoles of (Co, Nb) co-doped TiO$_2$ (CoNTO) ceramics have never been reported. The aims of this study were to prepare and characterize the CoNTO ceramics using a wet chemistry method for obtaining a new co-doped TiO$_2$ system that can exhibit a high $\varepsilon'$ and low tan$\delta$, and to clarify the primary contribution of the giant dielectric properties in CoNTO ceramics.

In the present study, the synthesized CoNTO ceramics with low levels of co-doping concentrations were systematically studied to obtain a high $\varepsilon'$ and low tan$\delta$, and to clarify the primary contribution of the giant dielectric properties in CoNTO ceramics. X-ray photoelectron spectroscopy (XPS), scanning electron microscope (SEM), Raman scattering spectroscopy, and X-ray diffraction (XRD) techniques were used for the characterization of the sintered CoNTO ceramics. Impedance spectroscopy and first-principle calculations were used to evaluate the possible origin of the dielectric response in the CoNTO ceramics. The intrinsic and extrinsic effects were discussed in detail.

### 2. Results and Discussion

Figure 1 shows the XRD patterns of the 0.5% CoNTO and 1% CoNTO powders and sintered 0.5% CoNTO and 1% CoNTO ceramics. A main phase of the rutile TiO$_2$ (JCPDS 21-1276) with a tetragonal structure is clearly observed in all samples with no impurity phase. The lattice parameters ($a$ and $c$ values) of all the samples calculated by a Rietveld refinement method are summarized in Table 1. Both values are nearly the same as the $a$ (4.593 Å) and $c$ (2.959 Å) values for the rutile TiO$_2$ structure. These calculated values are comparable to those reported in the (In, Nb) co-doped TiO$_2$ ceramics 21. The lattice parameters of the 0.5% CoNTO and 1% CoNTO powders are slightly changed. This may be due to the fact that, after the calcination process, the (Co, Nb) dopants did not, or only faintly, substitute into the rutile TiO$_2$ structure. The lattice parameters of sintered ceramics had slightly decreased compared to those of the powders with the same doping content. This result may be due to the substitution of rutile TiO$_2$ with (Co, Nb) dopants into the structure. On the other hand, the observed increase in the lattice parameters of the 1% CoNTO ceramic compared to those of the 0.5% CoNTO ceramic may be associated with the different ionic radii between the dopants and the host Ti$^{4+}$ ions.
CoNTO ceramic compared to those of the 0.5% CoNTO ceramic may be associated with the different ionic radii between the dopants and the host Ti4+ ions.

### Table 1. Lattice parameters of (Co0.33Nb0.67) x Ti1-x O2 powders and ceramics.

| Sample            | Lattice Parameter (Å) |
|-------------------|-----------------------|
|                   | a         | c         |
| 0.5% CoNTO powder | 4.596     | 2.962     |
| 1% CoNTO powder   | 4.596     | 2.962     |
| 0.5% CoNTO ceramic| 4.593     | 2.960     |
| 1% CoNTO ceramic  | 4.595     | 2.961     |

Figure 1. XRD patterns of: (a) 0.5% CoNTO; (b) 1% CoNTO powders and sintered ceramics; (c) 0.5% CoNTO; and (d) 1% CoNTO ceramics.

Figure 2a,b show the surface morphologies and the elemental mapping in the 0.5% CoNTO and 1% CoNTO ceramics, respectively. It was found that the dopants were homogeneously dispersed in the microstructure. A dense microstructure consisting of grains and grain boundaries is observed. The average grain size of the 0.5% CoNTO (~16.5 µm) and 1% CoNTO (~7.4 µm) ceramics was significantly changed by variations in the co-doping concentrations. The decrease in the mean grain size of the 1% CoNTO ceramic was likely due to the solute drag mechanism associated with the formation of a space charge at the grain boundaries [32,35,36]. The relative densities of the 0.5% CoNTO and 1% CoNTO ceramics were 94.83% and 97.64%, respectively. Elemental mapping images show homogeneous dispersion of the (Co, Nb) dopants and the major elements of Ti and O throughout the microstructure, without segregation of the dopants at any specific region.

Figure 3 shows Raman spectra of the 0.5% CoNTO and 1% CoNTO ceramics compared to the undoped TiO2 ceramic. The peak positions of the Raman shift of the E_g mode were the same position at 445.6 cm⁻¹. The peak positions of the A_1g mode of the undoped TiO2, 0.5% CoNTO, and 1% CoNTO ceramics were 610.5, 610.0, and 610.0 cm⁻¹, respectively. Generally, the E_g mode in the Raman spectrum of the rutile TiO2 is attributed to the vibration mode of oxygen along the c axis, which can be correlated with the presence of oxygen vacancies in the structure. The A_1g model is related to the vibration of the Ti–O bond [24,28]. The unchanged Raman shift of the E_g mode and the A_1g model mode may be due to a small doping concentration.
Figure 2. Field emission scanning electron microscope (FE-SEM) images and mapping images of all elements of (a) 0.5% CoNTO and (b) 1% CoNTO ceramics.

Figure 3. Raman spectra of TiO$_2$, 0.5% CoNTO, and 1% CoNTO ceramics.

The 0.5% CoNTO and 1% CoNTO ceramics were characterized using XPS techniques to clarify the influences of (Co, Nb) co-dopants on the formation of defects in the rutile TiO$_2$ structure. XPS spectra of the 0.5% CoNTO and 1% CoNTO ceramics are shown in Figure 4. Two peaks of Nb 3$d$ electrons were detected, corresponding to 3$d_{3/2}$ and 3$d_{5/2}$, respectively, which is entirely consistent with those observed in Nb$^{5+}$ single-doped TiO$_2$ materials [21]. Generally, doping a rutile TiO$_2$ ceramic with Nb$^{5+}$ can produce free electrons and eventually cause a reduction in Ti$^{4+}$ to Ti$^{3+}$, following the relationship [21]:

$$2\text{TiO}_2 + \text{Nb}_2\text{O}_5 \rightarrow 2\text{TiO}_2 + 2\text{NbO}_5 + 8\text{O}_2 + \frac{1}{2}\text{O}_2,$$  \hspace{1cm} (1)

$$\text{Ti}^{4+} + e^- \rightarrow \text{Ti}^{3+} \hspace{1cm} (2)$$

As shown in Figure 4, the binding energies of Ti$^{3+}$ and Ti$^{4+}$ for the 0.5% CoNTO ceramic were about 457.52 and 458.54, respectively, while the binding energies of Ti$^{3+}$ and Ti$^{4+}$ for the 1% CoNTO ceramic were about 457.67 and 458.42 eV, respectively. The oxidation states of Co were Co$^{2+}$ and Co$^{3+}$. As shown in Figure 4 for the XPS spectra of Co 2$p_{1/2}$ and Co 2$p_{3/2}$
were observed at about 779.15–780.19 eV and 794.55–794.86 eV, respectively, confirming the existence of the Co\(^{3+}\). Additionally, a small peak at relatively higher binding energies was observed. This can be ascribed to the Co\(^{3+}\). The XPS spectrum of O1s profiles was measured. Three peaks were obtained from the fitted data. The XPS peaks at 529.84, 532.43, and 531.15 eV were ascribed to the oxygen lattice in the bulk ceramic, oxygen lattices of other cation–oxygen bonds, and oxygen vacancies, respectively [21,37]. The substitution of TiO\(_2\) with acceptor Co\(^{2+}\) dopant can result in the existence of oxygen vacancies due to charge compensation, following the relation:

\[
\text{CoO} \rightarrow \text{Co}^{2+} + VO + O_2.
\] (3)

The dielectric properties as a function of frequency at room temperature for the 0.5% CoNTO and 1% CoNTO ceramics are illustrated in Figure 5a,b and its insets. The as-fired samples exhibited ultra-high \(\varepsilon\)' values of \(\sim 10^3–10^4\), over the frequency range of 40–10\(^7\) Hz. At 1 kHz, the \(\varepsilon\)' values of the 0.5% CoNTO and 1% CoNTO ceramics were \(3.6 \times 10^4\) and \(3.6 \times 10^3\), respectively, while the tan\(\delta\) values were 0.039 and 0.079, respectively.

Figure 4. XPS spectra of 0.5% CoNTO and 1% CoNTO ceramics.
The dielectric properties as a function of frequency at room temperature for the 0.5% CoNTO and 1% CoNTO ceramics were attributed to the high-frequency behavior of many giant dielectric oxides is associated with the DC conduction of the long-range motion of free charge carriers. Removing the resistive outer surface layer results in an increase in conductivity, leading to the observed increase in low-frequency $\tan\delta$. The increased $\varepsilon'$ values of the polished 0.5% CoNTO and 1% CoNTO ceramics were attributed to the non-Ohmic sample–electrode contact.

After the dielectric properties of both polished samples were measured, the electrodes and outer surface layer were removed. In this step, the sample thickness was reduced by 0.12 mm. The polished samples were painted by Ag paste and heated in air at 600 °C for 0.5 h. After that, the dielectric properties of the annealed samples were remeasured. It was found that the $\varepsilon'$ of the annealed samples significantly increased compared to that of the as-fired samples, especially in low-frequency range. Furthermore, it was found that the $\tan\delta$ values of the polished samples significantly increased in frequencies below $10^4$ Hz, as shown in the insets of Figure 5a,b. These results indicated that the outer surface layers of the 0.5% CoNTO and 1% CoNTO ceramics are the insulating layers. Generally, a low-frequency $\tan\delta$ of many giant dielectric oxides is associated with the DC conduction of the long-range motion of free charge carriers. Removing the resistive outer surface layer results in an increase in conductivity, leading to the observed increase in low-frequency $\tan\delta$.
and $\tan \delta$ values of the annealed 1% CoNTO sample were significantly reduced compared to those of the polished 1% CoNTO sample. At 1 kHz, the $\varepsilon'$ values of the annealed 0.5% CoNTO and 1% CoNTO ceramics were $3.3 \times 10^4$ and $7.6 \times 10^3$, respectively, while the $\tan \delta$ values were 0.032 and 0.043, respectively. These results indicated the important role of the insulative outer surface layer that contributed to the dielectric properties of the 0.5% CoNTO and 1% CoNTO ceramics. During the annealing process at 1200 °C, the surfaces of the polished samples were re-oxidized by filling oxygen vacancies on the surface and along the GBs. Free charges on the surfaces and the GB regions were reduced by filling oxygen ions during the annealing process [27,32,38]. Therefore, the SBLC and IBLC mechanisms can be used to explain the dielectric properties of the (Co, Nb) co-doped TiO$_2$ ceramics. Thus, it is clearly shown that the SBLC and IBLC effects have a significant impact on the dielectric properties of the 0.5% CoNTO and 1% CoNTO ceramics.

Figures 6 and 7 show the temperature dependences of the $\varepsilon'$ and $\tan \delta$ values of the 0.5% CoNTO and 1% CoNTO ceramics. The relaxation peak of $\tan \delta$ was observed in the 0.5% CoNTO and 1% CoNTO ceramics in the temperature range from −60 to 40 °C. The dielectric relaxation was likely attributed to the electrical response of internal insulating interfaces between grains [39]. The dielectric relaxation in a low temperature range is similar to that observed in ACu$_2$Ti$_4$O$_{12}$ ceramics. This result may be due to the Maxwell–Wagner polarization relaxation at the insulating GBs [40,41]. The relaxation peak of $\tan \delta$ shifts to high temperatures as the frequency increases, indicating a thermally activated relaxation mechanism.

![Figure 6](image_url)

**Figure 6.** Temperature dependences of (a) $\varepsilon'$ and (b) $\tan \delta$ of 0.5% CoNTO ceramic.

By using impedance spectroscopy analysis, the electrical properties of the grains and GBs can be characterized [8]. As shown in Figure 8 and the inset (1), only parts of large arcs are observed in a high-temperature range (100–200 °C) for the 0.5% CoNTO and 1% CoNTO ceramics, indicating the electrical response of the insulating parts (GBs and resistive outer surface layer). The nonzero intercept of the impedance spectra on the $Z'$ axis was observed for all the ceramics (not shown), confirming the existence of semiconducting grains with the grain resistance of $R_g \approx 40$ and 25 Ω.cm for the 0.5% CoNTO and 1% CoNTO ceramics, respectively. Therefore, the 0.5% CoNTO and 1% CoNTO ceramics are electrically heterogeneous and comprised of insulating and semiconducting parts. Furthermore, the diameter of the large arc decreased with increasing temperature, indicating the decrease in total resistance of the insulating parts ($R_i$). Thus, the colossal permittivity in the 0.5% CoNTO and 1% CoNTO ceramics prepared by a wet chemical process method may be primarily caused by extrinsic factors resulting from the IBLC and SBLC effects. The result indicates that the dielectric properties are associated with the electrical responses of grain and grain boundaries. As shown in the inset (2) of Figure 8, although the arc of the 0.5% CoNTO and 1% CoNTO ceramics cannot be seen at 200 °C, it is clearly observed that the arc of the 1% CoNTO ceramic was much larger than that of the 0.5% CoNTO ceramic.
This result indicates that the $R_i$ of the 1% CoNTO ceramic was larger than that of the 0.5% CoNTO ceramic. As shown in Figure 2a,b, the mean grain size of the 1% CoNTO ceramic was smaller than that of the 0.5% CoNTO ceramic. Thus, the number of insulating GBs per volume (GB density) of the 1% CoNTO ceramic should be higher than that of the 0.5% CoNTO ceramic, resulting in the increased $R_i$.

![Figure 6](image1)

**Figure 6.** Temperature dependences of (a) $\varepsilon'$ and (b) $\tan\delta$ of 0.5% CoNTO ceramic. $\varepsilon'$ and $\tan\delta$ for 0.5% CoNTO and 1% CoNTO ceramics, respectively. Therefore, the 0.5% CoNTO and 1% CoNTO ceramics are electrically heterogeneous and comprised of insulating and semiconducting parts. Furthermore, the diameter of the large arc decreased with temperature ($100–200 \, ^\circ C$) for the 0.5% CoNTO and 1% CoNTO ceramics. Inset (1) shows the impedance spectra of 1% CoNTO ceramic at different temperatures and inset (2) shows the comparison of impedance spectra at 200 °C for 0.5% CoNTO and 1% CoNTO ceramics.

![Figure 7](image2)

**Figure 7.** Temperature dependences of (a) $\varepsilon'$ and (b) $\tan\delta$ of 1% CoNTO ceramic.

![Figure 8](image3)

**Figure 8.** Impedance spectra at different temperatures ($100–200 \, ^\circ C$) for 0.5% CoNTO ceramic. Inset (1) shows impedance spectra of 1% CoNTO ceramic at different temperatures and inset (2) shows the electron hopping mechanism between Ti $^{3+}$ and Ti $^{4+}$ ions occurred inside the semiconducting grains of the 0.5% CoNTO and 1% CoNTO ceramics. This result indicates that the Ri of the 1% CoNTO ceramic was larger than that of the 0.5% CoNTO ceramic. As shown in Figure 200 °C, it is clearly observed that the arc of the 1% CoNTO ceramic was much larger than that of the 0.5% CoNTO ceramic. Thus, the number of insulating GBs per volume (GB density) of the 1% CoNTO ceramic was larger than that of the 0.5% CoNTO ceramic. Consequently, the dielectric properties are associated with the electrical responses of grain and grain boundaries. As shown in the figure, the mean grain size of the 1% CoNTO ceramic was smaller than that of the 0.5% CoNTO ceramic. Therefore, it is reasonable to suggest that the colossal dielectric properties of the 0.5% CoNTO and 1% CoNTO ceramics are electrically heterogeneous and comprised of insulating and semiconducting grains with the grain resistance of $R_g \sim 40$ and $25 \, \Omega \cdot \text{cm}$. The nonzero intercept of the impedance spectra on the Z-axis indicates that the dielectric properties of the insulating parts (GBs and GB densities) can be characterized [8]. As shown in Figure 8 and the inset (1), only parts of large arcs are observed in a high-temperature range ($100–200 \, ^\circ C$) for the 0.5% CoNTO and 1% CoNTO ceramics, indicating the electrical response of the insulating parts (GBs and GB densities). The 2CoVoTiO2 and 2NbTiO2 structures simultaneously in three configurations, for structures 1–3, as shown in Figure 7.
To clarify the possible origin of the colossal dielectric properties in 0.5% CoNTO and 1% CoNTO ceramics, we calculated the most stable configurations of 2CoV_oTiO_2 and 2NbTiO_2. To investigate the lowest energy configuration of the 0.5% CoNTO and 1% CoNTO ceramics, the 2CoV_o triangular and 2Nb diamond defects were placed into the TiO_2 structure simultaneously in three configurations, for structures 1–3, as shown in Figure 9. The stable structure of 2NbTiO_2 is structure 1 due to the lowest total energy. This result indicates that the 2CoV_o triangular defect does not prefer to be close to the 2Nb diamond defect. For the formation of defect clusters of EPDDs, these two types of defects must be close together. However, according to DFT calculations, the lowest total energy can be obtained when the 2CoV_o triangular and 2Nb diamond defects are far away. Therefore, it is reasonable to suggest that the colossal dielectric properties of the 0.5% CoNTO and 1% CoNTO ceramics were attributed to the SBLC and IBLC effects, in which the electron hopping mechanism between Ti^{3+} and Ti^{4+} ions occurred inside the semiconducting grains of the 0.5% CoNTO and 1% CoNTO ceramics.

![Figure 9. Energy-preferable structure of the 2CoV_o triangular defect and 2Nb diamond with different configurations.](image)

3. Materials and Methods

(Nb_{2/3}Co_{1/3})_xTi_{1-x}O_2 (CoNTO) powders with x = 0.5% (0.5% CoNTO) and 1% (1% CoNTO) were prepared by a wet chemistry method. Co(NO_3)_2•6H_2O (Kanto chemical, >99.5%), Diisopropoxynitriamion bis(acetylacetonate) (C_{16}H_{26}O_6Ti, Sigma–Aldrich), NbCl_5 (Sigma–Aldrich, >99.9%), deionized water, and citric acid were used as the starting raw materials. First, Co(NO_3)_2•6H_2O and NbCl_5 were dissolved in an aqueous solution of citric acid under constant stirring at ~25 °C (solution A). Second, a C_{16}H_{26}O_6Ti solution was dropped into solution A at 130 °C until a viscous gel was obtained. Third, a viscous gel was heated at 350 °C in an oven for 1 h to form dried porous precursors. Then, the resulting dried precursors were ground and calcined in air at 1000 °C for 12 h to produce the rutile phase in the 0.5% CoNTO and 1% CoNTO powders. Next, the obtained 0.5% CoNTO and 1% CoNTO powders were carefully ground. After that, the powders were pressed into pellets of ~1.0 mm in thickness and ~9.5 mm in diameter. Finally, the pellets were sintered at 1450 °C for 5 h. The heating and cooling rates were 2 °C/min and ~10 °C/min, respectively.

The prepared powders and sintered ceramics were characterized using XRD (PANalytical, EMPYREAN), SEM (FEI, QUANTA 450), XPS (AXIS Ultra DLD, UK), and Raman (Horiba Jobin-Yvon T64000) techniques. The densities of the sintered samples were measured using an Archimedes’ method. For the measurement of the dielectric properties, the top and bottom surfaces of the sintered samples with thickness < 1 mm were painted with Ag paste and heated in air at 600 °C for 0.5 h to make good electrode contact. The dielectric properties of the as-fired 0.5% CoNTO and 1% CoNTO ceramics were tested using a KEYSIGHT E4990A impedance analyzer over the frequency range from 40 to 10^7 Hz.
using an oscillation voltage of 0.5 V. The dielectric properties as a function of temperature were measured using a step increase of 10 °C from −60 to 200 °C.

The stable configuration of the (Co, Nb) co-doped rutile TiO$_2$ was investigated using the density functional theory (DFT) implemented in the Vienna ab initio simulation package (VASP). According to the pseudopotential used in this work, the projector augment wave approach and the Perdew–Burke–Ernzerhof (PBE) form of exchange–correlation functional was chosen. The 600-eV plane-wave energy cutoff and $3 \times 3 \times 3$ k-point samplings with Monkhorst–Pack scheme were successfully tested. The conjugate–gradient algorithm was carried out, and the force acting on each ion was calculated by the Hellmann–Feynman theorem.

4. Conclusions

(Nb$_{0.67}$Co$_{0.33}$)$_x$Ti$_{1-x}$O$_2$ ceramics with different co-dopant concentrations were successfully prepared using a wet chemistry method. A highly dense microstructure was obtained in all ceramics. High $\varepsilon'$ $\approx 10^{3}–10^{4}$ and very low $\tan\delta$ $\approx 0.032–0.079$ at 1 kHz were achieved. The XPS analysis indicated that the substitution of TiO$_2$ with Co$^{2+}$ and/or Co$^{3+}$ caused the existence of V$_O$ for charge compensation, while doping TiO$_2$ with Nb$^{5+}$ could cause the existence of free electrons, giving rise to the electron hopping mechanism between Ti$^{4+}$ and Ti$^{3+}$ in the semiconducting grains. Examination of the possible formation of defect structures was performed using a DFT calculation. The 2Nb diamond did not correlate with the 2CoV$_O$ triangular shapes, indicating that there was no EPDD. According to the impedance spectroscopy and DFT calculation, it can reasonably be suggested that the origins of the colossal dielectric properties are attributable to the IBLC and SBLC effects.

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