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Influence of Non-Stoichiometry on the Structure and Properties of Ba(Zn$_{1/3}$Nb$_{2/3}$)O$_3$ Microwave Dielectrics. IV. Tuning $\tau_f$ and the Part Size Dependence of $Q \times f$

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Abstract
High $Q$ ceramics of Ba$_3$W$_2$O$_9$ (BW)-substituted Ba(Zn$_{1/3}$Nb$_{2/3}$) O$_3$ (BZN) were prepared with a zero $\tau_f$ through the partial substitution of Zn by Ni and Co. The small concentrations of B-site vacancies introduced by the substitution of BW accelerated the kinetics and stability of the cation ordering and lowered the sintering temperature. Dense, zero $\tau z_f$ ordered solid solutions such as 0.99Ba(Zn$_{0.3}$Co$_{0.7}$)$_{1/3}$Nb$_{2/3}$O$_3$−0.01BW with $\varepsilon_r$=34.4 and $Q \times f$=82 000 at ~8 GHz could be obtained after sintering at 1380°C for 5 h and annealing at 1300°C for 24 h. Partially ordered ceramics in the Zn/Co and Zn/Ni solid solutions show a large gradient in the ordering throughout the pellets, which produces a resonant frequency dependence of their $Q \times f$ value. The ordering gradient is associated with the increased constraints on the growth of the 1:2 ordered structure within the interior of larger and thicker pellets and can be minimized by extended annealing.

Comments

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High Q ceramics of Ba$_3$W$_2$O$_9$ (BW)-substituted Ba(Zn$_{1/3}$Nb$_{2/3}$)O$_3$ (BZN) were prepared with a zero $\tau_f$ through the partial substitution of Zn by Ni and Co. The small concentrations of B-site vacancies introduced by the substitution of BW accelerated the kinetics and stability of the cation ordering and lowered the sintering temperature. Dense, zero $\tau_f$ ordered solid solutions such as 0.99Ba(Zn$_{0.3}$Co$_{0.7}$)$_2$Nb$_3$O$_9$-0.01BW with $\varepsilon_r = 34.4$ and $Q \times f = 82\,000$ at $\sim 8\,\text{GHz}$ could be obtained after sintering at $1380^\circ\text{C}$ for 5 h and annealing at $1300^\circ\text{C}$ for 24 h. Partially ordered ceramics in the Zn/Co and Zn/Ni solid solutions show a large gradient in the ordering throughout the pellets, which produces a resonant frequency dependence of their $Q \times f$ value. The ordering gradient is associated with the increased constraints on the growth of the 1:2 ordered structure within the interior of larger and thicker pellets and can be minimized by extended annealing.

I. Introduction

In a previous paper, the substitution of small concentrations of Ba$_3$W$_2$O$_9$ (BW) were shown to accelerate the kinetics of cation ordering, increase the stability of the order, and improve the sintering of Ba(Zn$_{1/3}$Nb$_{2/3}$)O$_3$ (BZN). As a result, BZN–BW ceramics show some of the highest $Q \times f$ values (at $Q \times f \sim 118\,500$) reported for a niobate perovskite and approach those of their “super Q” tantalate counterparts. The improvements in the ordering from the surface to interior and their limited (typically $\leq 1\%$ or less), it again has a very large effect on the extent and stability of the B-site order and the sintering, and enables the formation of high $Q$ ceramics with $Q \times f = 110\,000$. While both of these studies focused on optimizing the structure and dielectric losses, all of the BZN ceramics had a non-zero value of the temperature coefficient of resonant frequency ($\tau_f = 20–25\,\text{ppm}/\text{C}$). This paper focuses on the formation of high $Q$ BW-substituted ceramics with a zero $\tau_f$ through the partial substitution of Zn by Ni and Co. During the study, partially ordered ceramics were found to contain a gradient in the ordering from the surface to interior and their $Q \times f$ values were dependent upon the part size and resonant frequency. The lowering of $Q \times f$ with frequency is well known to occur in super $Q$ tantalate ceramics such as BZT and has limited their use for 2 GHz base-station applications. For the Ni- and Co-substituted BZN–BW systems studied in this paper, when the gradient in the order was removed, the dependence of $Q \times f$ on frequency was almost eliminated. Based on this finding, new methodologies are proposed to limit the degradation of $Q \times f$ in large ceramic pucks.

The tuning of the temperature coefficient of resonant frequency ($\tau_f$) of BZN to a zero value has been explored in systems with a variety of different chemistries. For example, the partial replacement of Ba by Sr in solid solutions of (1–$x$)BaZn$_{1/3}$Nb$_{2/3}$O$_3$–Sr(Zn$_{1/3}$Nb$_{2/3}$)O$_3$ results in a zero $\tau_f$ at $x = 0.7$,

with frequency is greatly reduced. The lowering of $\tau_f$ has also been studied in the (1–$x$)BZN–($x$)Ba(Ni$_{1/3}$Nb$_{2/3}$)O$_3$ (BNN) and (1–$x$)BZN–($x$)Ba(Co$_{1/3}$Nb$_{2/3}$)O$_3$ (BCO) systems. BCoN undergoes a transformation from a 1:2 ordered to disordered structure at $\sim 1425^\circ\text{C}$, slightly higher than pure BZN ($\sim 1375^\circ\text{C}$). Therefore, ordered solid solution phases are typically formed in BZN–BCoN after prolonged annealing at 1300–1400$^\circ\text{C}$, although there is disagreement on the zero-$\tau_f$ composition with values ranging from $x = 0.4$ to 0.7 being reported. For Ba(Ni$_{1/3}$Nb$_{2/3}$)O$_3$, the literature contains contradictory reports for the disordered temperature, which varies from 1400 to $> 1500^\circ\text{C}$. The X-ray diffraction (XRD) patterns of this phase often contain secondary impurity phases perhaps associated with partial melting and/or volatilization of NiO at high temperatures. Solid solutions of BZN and BNN prepared by the standard mixed oxide method are disordered; ordered phases were obtained using wet chemical methods and the processing dependency of the ordering was suggested to arise from chemical heterogeneities in the samples prepared by conventional techniques. For (1–$x$)BZN–($x$)BNN, a zero $\tau_f$ composition was identified at $x = 0.7$ where a dense ceramic sintered at 1500$^\circ\text{C}$ had $Q \sim 5700$ at 10 GHz.

Compared with the pure Zn-containing BZN or BZT systems, for Ni- or Co-containing perovskites such as BNN, BaNi$_{1/3}$Ta$_{2/3}$O$_5$ (BNT), BCoN, and their BZN–BNN and BZN–BCoN solid solutions, the kinetics of cation ordering are considerably slower. Consequently, a very slow-cooling process (e.g., 2$^\circ\text{C}/\text{h}$) or prolonged low-temperature annealing (e.g., 3 days) is required to maximize the $Q \times f$ value. The enhancement in $Q \times f$ coincides with an improvement in the cation order.

To accelerate the kinetics of the cation ordering, we focused on preparing zero $\tau_f$ ceramics in the BW-substituted BZN–BNN and BZN–BCoN systems. The B-site vacancies introduced by the substitution of BW were previously found to be very effective in permitting the formation of well-ordered, high $c/a$, high $Q$ ceramics of pure BZN. While similar enhancements were found for the Zn/Ni Zn/Co systems, for shorter anneal times, the kinetically limited diffusion of Ni and Co produced a gradient in the ordering throughout the dense pellets. The gradient was found to be responsible for a dependence of their $Q \times f$ on the size and the resonant frequency of the ceramic.

II. Experimental Procedure

Samples in the (1–$y$)[(1–$x$)Ba$_3$(ZnNb$_2$)O$_9$–$x$Ba$_3$(NiNb$_2$)O$_9$]–$y$ Ba$_3$W$_2$O$_9$ and (1–$y$)[(1–$x$)Ba$_3$(ZnNb$_2$)O$_9$–$x$Ba$_3$(CoNb$_2$)O$_9$]–$y$
Ba$_3$W$_2$O$_9$ systems were prepared by standard solid-state methods. The concentration of BW was fixed at $y = 0.01$. Stoichiometric amounts of BaCO$_3$ (Cerac, 99.9%), Nb$_2$O$_5$ (Cerac, 99.9%), Nb$_2$O$_3$ (Cerac, 99.95%), ZnO (Cerac, 99.95%), NiO (Cerac, 99.95%), CoO (Cerac, 99.95%), and WO$_3$ (Cerac, 99.9%) were mixed and calcined at 1000°C to expel CO$_2$. The products were then ball milled in ethanol with Y-stabilized zirconia media and annealed at 1100°C–1200°C for 10 h with an intermediate ball milling for 4 h to obtain single-phase powders. After a second ball mill for 6 h, the samples were dried and pressed isostatically into pellets at 80,000 psi. The pellets were burried in sacrificial powders of the same composition on a piece of platinum (Pt) foil. The Pt foil tray with the pellets and their muffling powders was placed in an alumina crucible together with a small amount of additional ZnO and NiO:CoO powder; the crucible was covered with a flat lid. The effectiveness of this method in maintaining the initial stoichiometry of the pellets was monitored through the weight of the ceramics after each heat treatment; no gains or loss in weight were observed. BZN–BW–BNN could be sintered to a high density ($>94\%$) at 1420°C–1440°C and BZN–BW–BCoN at 1380°C–1400°C. The subsequent low-temperature annealing was conducted at 1300°C for various times (6–48 h) to monitor the improvement in the ordering and the dielectric properties.

XRD patterns were collected with a Rigaku diffractometer (The Woodlands, TX) using a CuKα source operated at 45 kV and 30 mA. Equilibration of the samples was gauged by the absence of any changes in the relative intensities of the weak superlattice peaks associated with the cation ordering and splitting of any reflections in the XRD patterns after additional heat treatment. The unit cell parameters were refined by a least-squares procedure using data collected over the range of 10°–140° 20 with a slow scan speed (0.2°/min) and a step size of 0.02°. Scanning electron microscopy (SEM) study was performed on the thermally etched ceramic samples using a JEOL 6300F FEG HRSEM (Peabody, MA) operated at 10 kV.

The relative permittivity, $\varepsilon_r$, was measured from 100 Hz to 1 MHz using the parallel-plate method combined with an HP 4284 A precision LCR meter (Global Test Supply, Wilmington, NC) and a Delta 9920 environment chamber from −150° to 200°C. Measurements of the dielectric loss, $Q = 1/\tan \delta$, and the temperature coefficient of resonant frequency, $\tau_t$, at microwave frequencies were performed using cavity methods with an Agilent HP8720ES network analyzer (Agilent Technologies, Inc., Palo Alto, CA). The temperature coefficient of the resonant frequency ($\tau_t$) was calculated from data collected in the temperature range of 25°–75°C according to $\tau_t = \Delta f / f_0 \Delta T$, where $f_0$ is the reference frequency at 25°C.

### III. Results

**(1) BZN–BNN–BaW$_{2/3}$O$_3$ (BW) System**

In a previous study of BZN, low-level (≤ 2 mole %) substitutions of BW accelerated the kinetics of the cation ordering and reduced the sintering temperatures to enable the formation of well-ordered, high $c/a$, high $Q \times f$ ceramics after a short anneal (12 h) at 1300°C. For this investigation of BZN–BNN solid solutions, the concentration of BZN was fixed at 1 mole%, the ratio of Zn:Ni was varied from $x = 0$ to 1, and the compositions were formulated as $0.99[(1-x)\text{BZN}-(x)\text{BNN}]-0.01\text{BaW}_{2/3}\text{O}_3$ with $x = 0, 0.5, 0.7$, and 1.0 after annealing at 1300°C for 24 h.

Compared with BZN–BW ($x = 0$), where the highest degree of order was achieved after 12 h of annealing at 1300°C, the kinetics of the ordering in the BZN–BNN–BW solid solutions were slow and at least 20 h of annealing were required for the intensity of the $(100)_{\text{superlattice}}$ reflection to reach their maximum value. The slower kinetics in the Ni-containing solid solutions also resulted in a range of non-equilibrium states where the extent of the order varies throughout the depth of the ceramic samples. By slicing the pellets to different depths after each heat cycle, the evolution of the variation in the order was examined from the surface to the interior of the dense pellet. To avoid any preferred orientation, the pellet slices were crushed before characterization by XRD. Figure 2 shows the changes observed in the lower angle regions of the XRD patterns throughout a pellet of a sample with $x = 0.7$ (0.99(0.3BZN–0.7BNN)–0.01BW) after sintering at 1430°C for 5 h (Fig. 2a), and after a subsequent anneal at 1300°C for 12 h (Fig. 2b) and 20 h (Fig. 2c), respectively. The XRD patterns of the as-sintered sample contain weak and diffuse superlattice reflections; these become sharper and more intense as the annealing time is increased. In all three samples, the intensity and width (FWHM) of the ordering reflections vary throughout the depth of the pellet. The change in the relative intensity and width of the $(100)_{\text{superlattice}}$ reflection, the $c/a$ ratio, and the cell volume throughout each pellet is shown in Fig. 3. While every signature of the order is improved with prolonged annealing, it is evident that the extent of cation order, particularly in the pellets annealed at 1300°C for 12 h, is higher in the near surface region compared with the interior. The large difference between the order at the surface and interior in the 1300°C 12 h annealed ceramic is reduced in the poorly ordered as-sintered samples and also in the samples that approach an equilibrium well-ordered state after annealing at 1300°C for 20 h. The dispersion in the degree of order leads to significant changes in their dielectric response, which are described below.

Similar to the results for pure BZN, the substitution of BW was effective in lowering the sintering temperature of the BZN–BNN ceramics, which reached a high density (95%) at 1420°C–1440°C, approximately 100°C lower than those reported in previous investigations of the system. After sintering at 1430°C for 5 h, thermally etched samples with $x = 0.5$ and 0.7 revealed a well-densified microstructure comprised of large homogenous grains with sizes in the range 5–8 μm; see Figs. 4(a) and (b). The ceramics of BZN–BNN–BW sintered at 1420°C–1440°C show no evidence for the involvement of a liquid phase; however, the SEM images collected from samples sintered at a higher temperature (e.g., 1480°C for 4 h) reveal features consistent with liquid phase sintering (Fig. 4c), which were also observed in the
As expected, the substitution of BNN reduces the temperature coefficient of resonant frequency of BZN–BW ($\tau_z = \pm 21\text{ ppm/°C}$); the variation of $\tau_z$ with $x$ is shown in Fig. 5. A $\tau_z = 0$ was obtained for the composition close to $x = 0.7$ (Zn:Ni = 3:7). The $\tau_z$ of all the samples annealed at 1300°C for 20 h were slightly lower than those of the as-sintered ceramic. This is probably related to the changes in the cation–anion bond strengths and bond lengths during the ordering process.\textsuperscript{13} The $Q \times f$ values and relative dielectric constants ($\epsilon_r$) after the 20 h annealing are shown in Fig. 6. To confirm their reproducibility, the data were collected from specimens prepared in identical ways from several different batches of the starting powders. The zero $\tau_z$ composition with $x = 0.7$ has a mean $Q \times f = 71\,000$ at 8.2–9.2 GHz, and a dielectric constant = 32.2.

Besides their variation with composition, the dielectric loss properties of BZN–BNN–BW also show an interesting dependence on the sample dimensions. In a standard cavity measurement, the approximate resonant frequency of an isolated dielectric resonator can be estimated from $f_{GR} = \frac{34}{\sqrt{\epsilon_r}} [a + 3.45]$,\textsuperscript{20} where $a$ and $r$ are the radius and thickness of the resonator, respectively. For pellets with the same composition and a similar relative density, diameter ($d = 2a \approx 6.8$ mm), and relative dielectric constant (Table I), $f$ is primarily determined by the aspect ratio ($t/d$ or $t/2a$) and should be inversely proportional to $t/d$. Therefore, it is generally observed that thicker pellets resonate at lower frequencies (see Fig. 7). For impurity-free dielectric resonators, where the losses are primarily because of phonon scattering, the loss typically increases linearly with $f$ and the $Q \times f$ product should be independent of $f$. In that case, the $Q \times f$ product should be independent of $t/d$ and higher $Q$'s should be observed in thicker pellets where the resonance occurs at a lower frequency. It should be noted that refinements to the above equation are necessary for analysis of $Q$ and $f$ of pellets investigated over a wide range of frequency and aspect ratio, and for ring resonators.\textsuperscript{20,21} However, for the range of frequency and aspect ratio of the samples explored in this investigation, the effect of these higher order corrections on the $Q$'s that we observed is less than 0.1%.

$Q \times f$ data for ceramics of BZN–BNN–BW with different aspect ratios are shown in Fig. 7; a reproducible and significant dependence of $Q \times f$ on the aspect ratio, in particular for those samples annealed at 1300°C for 12 h, is clearly evident. The dependence of the $Q \times f$ on the aspect ratio of the 1300°C pellets is further highlighted by the plot in Fig. 8(a) where the $Q \times f$ product of the $x = 0.5$ and 0.7 samples is seen to increase significantly with frequency, while the data for the corresponding 12 h annealed Zn (Ni-free) ceramic are essentially independent of $f$ over the range investigated. After a prolonged annealing (e.g., 20 h), the $Q \times f$ values of the BZN–BNN–BW increase at all frequencies and reach values 50%–80% higher than the as-sintered samples. The extended annealing also reduces the dependence of $Q \times f$ on the aspect ratio (Fig. 7), which remains almost constant ($Q \times f \approx 80\,000$ for $x = 0.5$; $\approx 70\,000$ for $x = 0.7$) over the 7.9–9.5 GHz range. The increase in the $Q$ and $Q \times f$ values is closely related to the increase in the cation order observed for each specimen (Fig. 3). Their dependence on the aspect ratio, and therefore frequency, after a 12 h anneal at 1300°C coincides with the formation of pellets with the largest gradient in the degree of order and ordered domain size (Fig. 3).

(2) BZN–BCoN–BaW$_{25}$O$_{73}$ (BW) System

The formation of zero-$\tau_z$ compositions was also investigated in the BZN–BCoN–BW system where the concentration of BW was again fixed at 1 mole%. Figure 9 shows the XRD patterns collected from crushed BZN–BCoN–BW pellets annealed at 1300°C for $\geq 20$ h after sintering at 1380°C for 5 h. The pattern is free of peaks from impurity phases and all the solid solutions form a 1:2 ordered perovskite. Similar to BZN–BNN–BW, after annealing for intermediate times (12 h at 1300°C), the degree of order and domain size vary throughout the thickness of the pellet, with the interior showing a much lower degree of order.

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**Fig. 2.** Lower angle regions of the X-ray patterns of a sectioned pellet sample of 0.99(0.3BZN–0.7BNN)-0.01BW ($x = 0.7$): (a) after sintering at 1430°C for 5 h, and after subsequent anneals at 1300°C for 12 h (b) and 20 h (c). The depth of the sectioned sample is shown for each pattern.
order compared with the near surface regions. The XRD patterns and trends in the structural parameters collected from sliced pellets were almost identical to those shown in Fig. 2 for BNN and are not reiterated here. All of the BZN–BcoN–BW solid solutions could be sintered to a high density (\( \geq 95\% \)) after a 5 h heat at 1380°C, again, this is lower than the temperatures previously reported for the BW-free system.\(^7\)–\(^{11}\) The SEM images (Fig. 10) reveal a dense homogeneous microstructure with a grain size \( \approx 5–8 \) \( \mu \)m. Figure 11 shows the variation of \( \tau_f \) with composition; a zero \( \tau_f \) was obtained close to \( x = 0.7 \) (70% Co–30% Zn). The \( Q_{\text{f}}/C_{\text{f}}' \)s and relative dielectric constants of the samples after equilibration at 1300°C for 420 h are shown in Fig. 12. The zero-\( \tau_f \) composition has \( Q_{\text{f}}/C_{\text{f}} \approx 82000 \) at 7.7–8.8 GHz and \( \varepsilon = 34.4 \). The \( Q_{\text{f}}/C_{\text{f}} \) values of the pellets annealed for intermediate times also show an aspect ratio and frequency dependence, which again could be minimized by annealing at a low temperature for a longer time (typically 20 h at 1300°C).

The correlation between the changes in structure and \( Q \) is further illustrated by the data in Fig. 13, where the \( Q \times f \) of BZN–BCoN–BW and BZN–BNN–BW ceramics was monitored as a function of the extent of annealing at 1300°C. In each case, \( Q \times f \) increases systematically with the annealing time, consistent with the associated increase in the degree of ordering (Fig. 3). The changes in \( Q \times f \) reported in Endo \textit{et al.}\(^7\) for even longer heat treatments of BW-free BZN–BCoN are also plotted in Fig. 13. The rate of increase in \( Q \), which reflects the rate of increase in the ordering, is greater in the BW-containing solid solutions that reach a higher degree of order and higher \( Q \) after a shorter annealing time.

### IV. Discussion

High \( Q \) compositions with a zero \( \tau_f \) were identified in the BZN–BNN–BW and BZN–BCoN–BW solid solution systems. Consistent with the results obtained in previous work on pure BZN, BW was found to be an effective additive in accelerating the ordering kinetics and promoting the formation of a well-ordered structure with a high \( c/a \) distortion and a high \( Q \) in the BZN–BNN and BZN–BCoN systems. We believe that the enhancement in the order and the accelerated kinetics are related to the small concentrations of B-site vacancies introduced by BW and the formation of ordered structures with Ba\([\text{Zn}_{1-x}\text{Co}_{x}\text{Vac}_{0.01}]_{0.99}\text{WB}_{0.01}O_3\); these and the resultant changes in the local stability of the anions were identified and discussed previously for the BZN–BW system.\(^1\) Rather than re-iterate these arguments here, in this paper we focus on new features of the ordering discovered in the BZN–BNN–BW and BZN–BCoN–BW systems that lead to gradients in the degree of order in partially annealed dense pellets and produce an aspect ratio and frequency dependence of their \( Q_{\text{f}}/C_{\text{f}} \) values. In BW-substituted BZN, no obvious ordering gradient or aspect ratio dependence of the \( Q \times f \) is observed in either the as-sintered (1390°C, 5 h) or annealed (1300°C for 12 h) samples. The small concentrations of Zn vacancies introduced by BW are effective in removing any significant kinetic barriers to the formation of well-ordered ceramics, which form a homogenous structure comprised of large ordered domains with a high \( c/a \) and \( Q \) after a short anneal. The formation of ceramics with a gradient in the ordering in the dense pellets of BZN–BNN–BW and BZN–BCoN–BW is presumably related to a lower diffusivity of the Ni and Co cations compared with Zn. The increased kinetic limitations on the ordering process in the Ni- and Co-containing systems are consistent with the observations made on other perovskites such as Ba\(_3\)NiNb\(_2\)O\(_9\) (BNN), Ba\(_3\)NiTa\(_2\)O\(_9\) (BNT), BCoN, and their various solid solutions, which, compared with their Zn counterparts, require much slower cooling rates,\(^8\)\(^{11,13}\) longer sintering\(^9\) or longer low-temperature annealing\(^{16,17}\) to reach a high degree of order and a maximum \( Q \). The large gradients in the degree of order and ordered domain size (Fig. 3) and the associated dependence of the \( Q \times f \) on as-

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Fig. 3. Variation in the relative intensity (a), and width (b) of the (100) order reflection, the \( c/a \) ratio (c), and the cell volume (d), throughout the pellets of 0.99(0.3BZN–0.7BNN)–0.01BW.
pect ratio (Fig. 7) were only observed in dense pellets of BZN–BNN–BW and BZN–BCoN–BW with non-equilibrium, “intermediate” or “partial” order. The ordering gradient and aspect ratio dependence is present, but much smaller, in the poorly ordered as-sintered samples and is greatly reduced by prolonged annealing (20 h at 1300°C), although even in the longer annealed ceramics the ordered domain size and \(c/a\) of the 12 h annealed ceramics, which only approach those of the surface regions after a more extended annealing.

For ferroelastic systems,\textsuperscript{22} the formation of larger domains at the surface of a dense ceramic was shown to arise from the relaxation of the domain boundaries at a free surface. For our systems, thicker pellets with a lower surface to volume ratio show a larger ordering gradient and contain a higher volume percentage of a “constrained” ordered phase; we propose that this is the major factor in mediating their dielectric loss. As the aspect ratio is decreased, the thinner pellets have a smaller domain gradient, contain a lower volume percent of the constrained domains, and therefore have a higher \(Q \times f\). The change in the volume fraction of the constrained phase in turn is responsible for the dependence of \(Q \times f\) on the aspect ratio and frequency. When the “corrugating” effect is reduced and ordering becomes homogeneous throughout the pellet, the \(Q \times f\) increases and its dependence on frequency is removed.

The different degrees of constraint near the surface and in the interior of the pellets may explain our prior observation of a small gradient in the ordering in dense ceramics of pure BZN\textsuperscript{23} where the kinetics are limited by the slower diffusion of Zn in the BW-free stoichiometric system. It is possible that these types of constraint effects are also responsible for intra-grain domain gradients or “core-shell” structures that have been reported for 1:2 other ordered systems.\textsuperscript{24–27}

![Fig. 4. Scanning electron microscopic images of thermally etched BZN–BNN–BW with (a) \(x = 0.5\) (1430°C, 5 h sintering) and (b) \(x = 0.7\) (1480°C 4 h sintering).](image)

![Fig. 5. Variation of \(\tau_f\) with composition in 0.99\((1-x)\)BZN–(x)BNN–(0.01)BW.](image)

![Fig. 6. \(Q \times f\) and \(\varepsilon\) data of (1–\(x\))BZN–(\(x\))BNN–BW after annealing at 1300°C for 20 h.](image)
Although the coring effects and associated $Q \times f$ dependence observed in the Ni/Co systems after a 12 h anneal were absent in similar-sized ceramics of BZN–BW annealed at the same temperature for the same time, it is possible, perhaps likely, that coring effects will be present in the faster-diffusing Zn systems after a shorter anneal time or in larger "puck-sized" ceramics where the constraints on the growth are even larger. For many years, it has been recognized that the $Q \times f$ factors of large pucks of "super $Q$" microwave perovskites such as Ba(Zn$_{3/3}$Ta$_{2/3}$)$_3$O$_3$ (BZT) prepared for 2 GHz applications are inferior to those of smaller ceramic pieces, where $Q \times f$'s $\geq$ 150 000 have been reported at frequencies close to 10 GHz. Indeed, the specifications of many BZT-based commercial ceramics are only quoted for a 10 GHz resonance, even though their main potential lies in 2 GHz base station applications. The deterioration in $Q \times f$ has often been thought to arise from variations in the degree of volatilization of ZnO in the different-sized parts or to other difficulties associated with the processing of large ceramic parts. We suggest that the reduction in $Q \times f$ at lower frequency is directly related to an increase in the coring of the order and a reduction in the volume percent of a well-ordered phase from the increased constraints on the growth of the 1:2 ordered structure within the interior of the large ceramic pucks. It is possible that this effect could be alleviated through very long anneal times; we propose that a more effective route would be to use chemistries where B-site vacancies are deliberately introduced.

### Table I. Dielectric Properties of BZN–BNN–BW with Different Aspect Ratios After Annealing at 1300 °C for 12 h

<table>
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<th>Aspect ratio</th>
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<td>Density (%)</td>
<td>f₀ (GHz)</td>
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<td>33.9 7.78</td>
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Fig. 7. Variation of $Q \times f$ with aspect ratio (t/d) for sintered and annealed samples of $(1-x)$BZN–$(x)$BNN–BW with (a) $x = 0.5$ and (b) $x = 0.7$.

Fig. 8. Resonant frequency dependence of $Q \times f$ for BZN–BW and BZN–BNN–BW ceramics annealed at 1300 °C for (a) 12 h and (b) 20 h.
into the starting powders, e.g., through the substitutions of BW investigated in this paper or the non-stoichiometric BZN powders described in our prior work, where the ordering kinetics and the stability of the ordered phase are greatly enhanced.

V. Conclusions

High $Q$, zero $\tau_f$ ceramics were obtained in the $Q(x) = 0.99[(1-x)Ba(Zn_{1/3}Nb_{2/3})O_3-(x)Ba(Co_{1/3}Nb_{2/3})O_3] - 0.001[BaW_{2/3}O_3]$ systems. The zero $\tau_f$ was obtained close to $x = 0.7$ in both systems.

Fig. 9. X-ray diffraction patterns of $0.99[(1-x)Ba(Zn_{1/3}Nb_{2/3})O_3-(x)Ba(Co_{1/3}Nb_{2/3})O_3] - 0.001[BaW_{2/3}O_3]$ with $x = 0, 0.4, 0.7$, and 1.0 after annealing at $1300^\circ C$ for $\geq 24$ h.

Fig. 10. Scanning electron microscopic images of thermally etched BZN–BCoN–BW with (a) $x = 0.4$ and (b) $x = 0.7$, after sintering at $1380^\circ C$ for 5 h.

Fig. 11. Variation of $\tau_f$ with composition in the $0.99[(1-x)BZN-(x)BCoN]-(0.01)BW$.

Fig. 12. $Q \times f$ and $\varepsilon$ data of BZN–BCoN–BW after annealing at $1300^\circ C$ for $\geq 20$ h.

Fig. 13. Change in $Q \times f$ with the time of annealing at $1300^\circ C$ for BZN–BW, BZN–BNN–BW, BZN–BCoN–BW. Data for BW-free BZN–BCoN were adapted from Endo et al.7

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with $Q \times f$ values of $\sim 71000$ for Zn–Ni and $Q \times f \approx 80000$ for Zn–Co. In agreement with the observations made on pure BZN, BW is ineffective in accelerating the ordering kinetics and promoting the formation of a well 1:2 ordered structure with a high $c/a$ distortion and a high $Q$ because of the small concentrations of B-site vacancies. Partially ordered dense ceramics show a large gradient in the ordered domain size, which results in an aspect ratio and frequency dependence of their $Q \times f$ values. The reduction in $Q \times f$ at lower frequency is due to an increase in the coring of the order and reduction in the volume percent of a well-ordered phase originating from the increased constraints on the growth of the 1:2 ordered structure within the interior of larger/thicker pellets.

References