

Bimodal Grain Sized Barium Titanate Dielectrics Enabled Under the Cold Sintering Process

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Abstract:

In barium titanate (BaTiO₃) the relative permittivity varies as a function of grain size due to the influence of various sizes and scaling effects. The cold sintering process (CSP) has been applied to sinter nanocrystalline BaTiO₃ (< 200 nm), however in conventionally sintered BaTiO₃ a maximum relative permittivity is achieved at average grain sizes around 0.8 μm. In this work the feasibility of cold sintering 1 μm BaTiO₃ inclusions in ratios of fine-grained BaTiO₃ matrixes from 50 - 200 nm is investigated. Occurrences of both conformal sintering of inclusions into the matrix and constrained sintering with residual porosity are observed. Subsequently, electrical resistivities increased from $1 \times 10^8 \Omega\cdot\text{cm}$ to approximately $1 \times 10^{12} \Omega\cdot\text{cm}$ by a post CSP heat treatment of 500°C. Relative permittivity of annealed samples increases systematically following a logarithmic mixing law as a function of matrix grain size and increasing the ratio of inclusions to matrix.

1. Introduction

Barium titanate (BaTiO₃) or BT, is a high permittivity ferroelectric perovskite ceramic with a wide range of technologically critical applications including Positive Temperature Coefficient Resistors (PTCR) and Multilayer Ceramic Capacitors (MLCCs) [1], [2], [3]. Control of the microstructure and of the final grain size distribution is a critical factor impacting the final electrical performance of barium titanate.

For capacitive applications, there is a balance that is required between the magnitude of the permittivity and the reliability. The reliability of the devices that are cofired into a multilayer ceramic capacitor (MLCC) is improved with the reduction of the grain size and increasing the number of grain boundary barriers to limit oxygen vacancy migration under an applied field [4]. However, below grain sizes of ~0.8 μm there are size effects from clamping of the domain walls which limit the dielectric response of a material [5]. Additionally, for grain sizes below 40 nm there are intrinsic size effects where the paraelectric-ferroelectric phase transition temperature renormalizes, and the depolarization fields suppress the onset of ferroelectricity [6].

A number of sintering methods have been applied to densify BT and other dielectric ceramics, such as flash sintering, spark plasma sintering, hot pressing, two-step sintering, rate-controlled sintering, atmosphere firing and liquid phase sintering [7], [8], [9], [10], [11], [12], [13], [14]. Despite advances in these sintering technologies, limitations imposed by the thermal processing parameters required to activate densification such as cost, sustainability, and microstructural control leave several outstanding challenges unaddressed [15]. With the

36 additional needs for greater microstructural control and reducing processing temperatures, The Cold Sintering
37 Process (CSP) becomes an attractive sintering technique for the fabrication of future electronic devices.

38 CSP is a relatively new sintering technique enabled by a non-equilibrium, pressure solution creep mechanism
39 which permits the densification of ceramic materials at significantly reduced temperatures [16]. Densification by
40 pressure solution creep is enabled when a powder compact is pressurized in the presence of a transient fluid phase
41 that the material is soluble in. This enhances the interfacial dissolution of matter at high pressure contacts between
42 grains, and drives diffusion through chemical potential gradients, to lower pressure porous regions, resulting in
43 precipitation of solute in the pores [17], [18]. With densification of a diverse array of materials possible at
44 temperatures below 300°C, the incorporation of organic polymers and non-noble metals into composite
45 microstructures becomes feasible [19]. CSP can aid in enhancing the performance of dielectrics by adding low
46 volume fractions of a high impedance polymer at the grain boundary which improves resistivity, nonlinear
47 permittivity, reliability, and high field performance due to reduction of oxygen vacancy migration across the grain
48 boundary [20].

49 Thus far, the CSP has been successfully employed to sinter fine grained barium titanate ranging in particle size
50 from 50 to 200 nm with finite but minimal grain growth [21], [22], [23]. In these size ranges, the relative
51 permittivity of BT is decreased because of the size and scaling effects associated with the ferroelectric phenomena
52 [24]. However, the microstructural control enabled by cold sintering enables the potential to incorporate a bimodal
53 mixture of larger grain size inclusions to these smaller grain sized matrixes, to increase their dielectric
54 performance while still preserving reliability. With traditional sintering the relative permittivity approaches a
55 maximum at a particle size of ~ 800 nm. The objective of this paper is to cold sinter barium titanate with a
56 controlled bimodal mixture of fine grain barium titanate of 50, 100, and 200 nm with 1 μm inclusions, compare
57 this to corresponding unimodal distributions, and investigate the consequences on the microstructure and electrical
58 properties.

59 **2. Experimental Methodology**

60 **2.1. Precursor chemicals**

61 BaTiO₃ powder (US Research Nanomaterials, Inc., 99.9% reported purity) synthesized by a combustion method
62 with particle sizes ranging from 50 nm to 1 μm were sintered in this study.

63 Powders were calcined at 800°C under an oxygen atmosphere to remove organics and carbonates from the
64 precursor powder surfaces. The precursor powder was mixed and ground in a mortar and pestle for 10 minutes.
65 Precursor powder and the final bulk samples were stored at 80°C under vacuum. Ba(OH)₂·8 H₂O (Alfa Aesar, 99.7%
66 reported purity) was used as the transient chemical phase and stored under vacuum at room temperature to limit
67 carbonate formation.

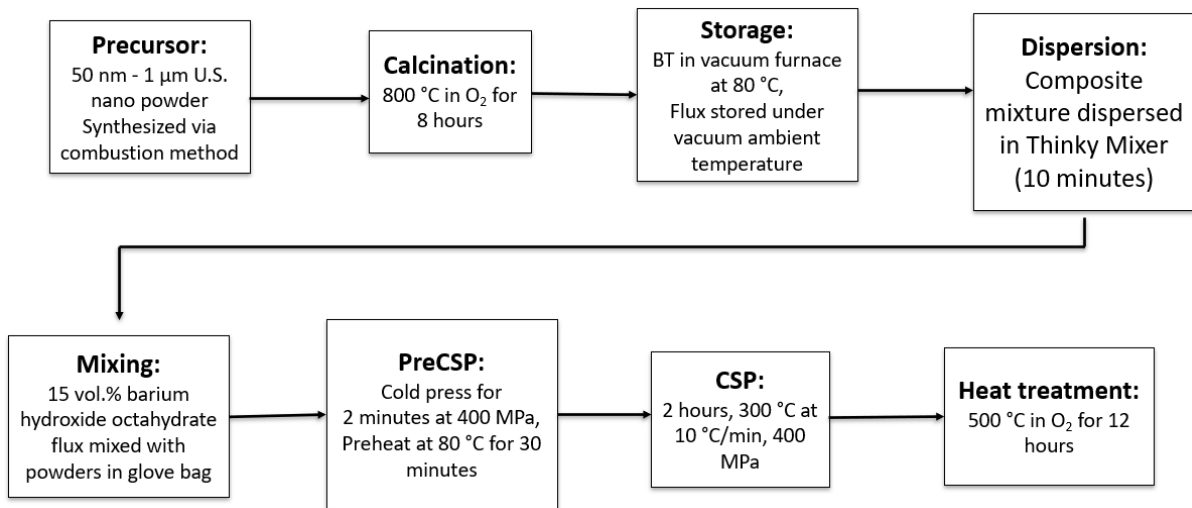
68 **2.2. Particle Preparation and Mixing**

69 In this study, both monodispersed and bimodal mixtures were sintered. The monodispersed samples were barium
70 titanate of 50, 100, and 200 nm. The bimodal mixtures were, 80 vol% of 50, 100, and 200 nm barium titanate
71 powders mixed with 20 vol% of 1 μm barium titanate. All samples had a total mass of 0.8 grams.

72 The bimodal mixtures were dispersed using a non-contact defoaming mixer (Thinky Corporation, model AR-250)
 73 for a total of 10 minutes, with intermittent stops every minute to scrape powder adhering to the container and to
 74 stir the powder. 15 wt% of the BaOH₂:8 H₂O flux was then mixed with the precursor powder in a mortar and
 75 pestle for a total of 10 minutes to disperse the flux.

76 **2.3. Cold Sintering Procedure & Post Annealing**

77 An Across International stainless-steel die with an inner diameter of 12.7 mm was used to carry out the CSP. Inner
 78 spacers were machine polished to provide a flat surface for the application of pressure. Four inner spacers were
 79 used inside the die: two at the top and two at the bottom. Round nickel foil was punched and inserted onto the
 80 surface of the pucks to provide an inert surface during sintering. A heater jacket was wrapped around the die with
 81 a thermocouple inserted in between the jacket and die to detect sintering temperature. The powder and flux
 82 mixtures were initially pressed at 400 MPa for 2 minutes before CSP to form a green body. Pressure is released,
 83 and then transferred to a Carver press with heated plates. The green body is then pressed to 400 MPa again. The
 84 heated plates, in tandem with the heater jacket (Tempco, band heater) are used to heat the system, with a thermal
 85 ramp rate of 10°C / min. The temperature is raised to 80°C (at which point the BaOH₂:8 H₂O melts) and held for
 86 30 minutes to allow for flux distribution, and then raised to 300°C and cold sintered for 2 hours. After cold
 87 sintering, the sample is brought down to room temperature and ejected. Fig.1 shows a summary of the basic cold
 88 sintering process. The experimental procedures are all conducted in a temperature and relative humidity stable
 89 environment with the temperature being 20°C and a relative humidity of 70%. Following the cold sintering, the
 90 samples are then loaded into a furnace (Carbolite Gero, cube furnace), and heat treated at 500°C under oxygen
 91 atmosphere to increase resistivities.



92
 93 Fig.1. Flow diagram of the cold sintering process adopted for this study.

94 **2.4. Characterization**

95 2.4.1 Density

96 Prior to annealing, the samples are sanded to a thickness of approximately 0.8 mm, and geometric measurements
97 are performed to calculate density. The ratio between the measured and theoretical density of barium titanate
98 ($\rho = 6.02 \text{ g/cm}^3$) is used to calculate the relative density of each sample.

99 2.4.2 Electrical Measurements

100 Sputtering of 100 nm platinum electrodes is performed in a rotary pumped coater (Quorum Technologies, model
101 Q150R). Current-Voltage (I-V) measurements were carried out at voltages between 10 and 100 V, and the slope
102 of the curve, together with the geometry of the samples was used to calculate the resistivity of the dielectric.
103 Impedance measurements were carried out after heating the samples at 150°C for 5 minutes to limit the influence
104 of adsorbed moisture on dielectric properties. Capacitance and dielectric loss were measured at frequencies of 10
105 kHz and 100 kHz in fixtures using an LCR meter (Keysight Precision LCR Meter, LXI) and temperatures between
106 -20°C – 150°C using a delta oven. Samples were approximated as parallel plate capacitors, and electrode areas
107 and sample thicknesses were used to determine relative permittivity. PE loops were measured at 10 Hz and
108 voltages between 10-60 kV using a Sawyer-Tower circuit (Trek, model 609 A).

109 2.4.3 Microstructural Characterization

110 Fractured surfaces were imaged via scanning electron microscopy (FESEM, Verios, Thermo Fisher Scientific)
111 and energy dispersive spectroscopy (EDS). Thin cross-sectional transmission electron microscope (TEM)
112 specimens were prepared by using focused ion beam (FIB, FEI Helios 660). The local microstructures of the
113 cross-sectional samples were observed by FEI Titan3 G2 double aberration-corrected microscope at 300kV.

114 2.4.4 X-Ray Diffraction

115 X-ray diffraction was performed (XRD, Empyrean, PANalytical) with a scan step of 0.07°/s using Cu K α ,
116 scanning between 20°C to 80°C. XRD was carried out on cold sintered samples ground into powder along with
117 a silicon internal standard. The precursor powder, and barium hydroxide octahydrate flux were also checked by
118 XRD.

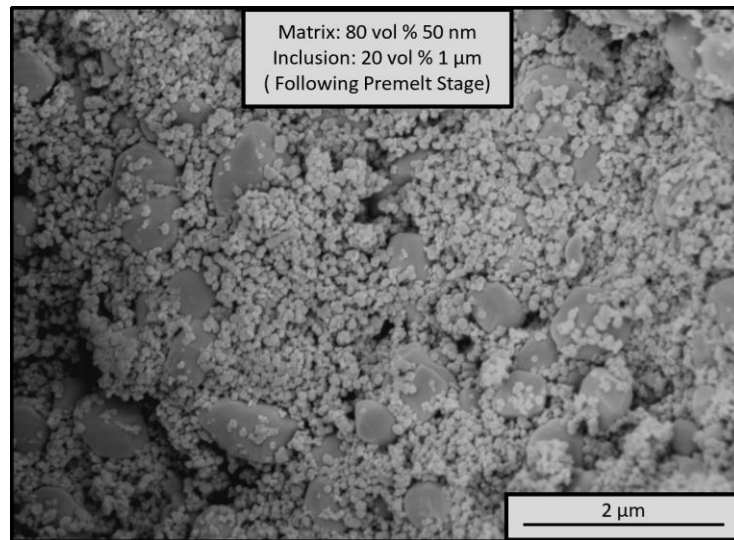
119 **3. Results and Discussion**

120 **3.1 Powder Processing to enable Cold Sintering of Bimodal Grain Sized BaTiO₃**

121 Earlier we described details of cold sintering of BT using both KOH-NaOH and BaOH₂·8H₂O transient phases
122 [20], [21], [22]. Here we emphasize two additional aspects of importance, the surface chemistry with respect to
123 BaCO₃, and maximizing the particle and transient chemistry prior to the cold sintering.

124 When sintering powders, it is important to understand surface chemistry, as this can be in the form of a BaCO₃
125 phase in the case of BT [25]. This is always important in high levels of industrial reproducibility of BT based
126 capacitors. Surface chemistry is also critical in the cold sintering process, as it limits the kinetics of the low
127 temperature diffusional processes that must be operational in the pressure solution mechanisms [26]. It is therefore
128 important to ensure that there is no Barium Carbonate (BaCO₃) present on the surface of the starting powder, as
129 this can negatively impact the sintering and permittivity of BT. This is ensured by initially exposing the BT
130 powders to an 800°C calcination.

131 Also crucial to densification, especially with a bimodal particle size mixture is the initial green body state of the
132 sample. There is evidence suggesting that poor mixing of particles or inhomogeneous distribution of the flux can
133 result in differential strains, causing macroscopic defects and poor densification [27]. The pre-pressing and pre-
134 melting step aids in the formation of a more compact geometry and ensures further breaking of particle aggregates
135 and improved densification [28]. In Fig.2, a SEM images shows an example of uniform mixing of the BT powders
136 after the mixing process, application of pressure, and pre-melting step at 80°C for an 80/20 vol% mix of 50 nm –
137 1 μm BT powder. Following the pre-melt step, there is also no surface impaction or faceting of the grains,
138 indicating that no cold sintering, or diffusional pressure solution creep occurred at this stage [29].



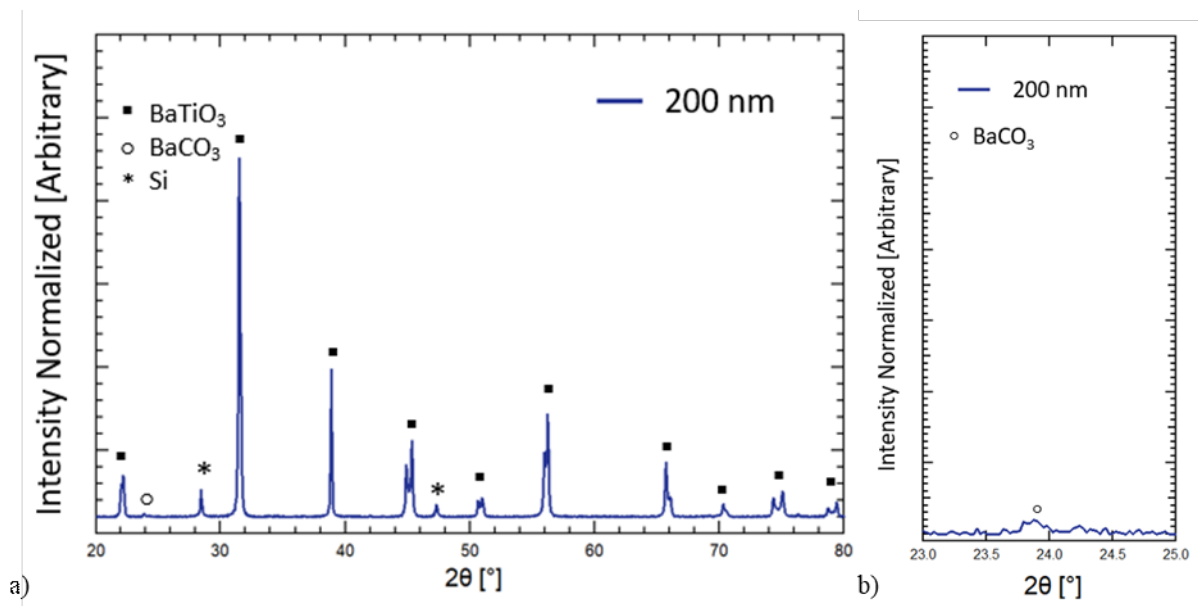
139

140 Fig.2. Green body after the pre-melt stage. There is a rounded morphology in both 50 nm and 1 μm BT powder
141 here in an 80/20 mix, indicative of just particle compaction without any sintering beyond the particle
142 rearrangement densification.

143 After cold sintering at 300°C, the monodispersed 100 nm and 200 nm BT samples achieved densities of > 93%
144 which is consistent with final stage sintering behavior and comparable to densities achieved in previous CSP
145 studies of BT powders in this size range [22]. However, the 50nm powder did not successfully densify because of
146 mechanical failure during the sintering process which could be attributed to agglomeration during the particle
147 mixing step resulting in inhomogeneous distribution of the flux. As a result, the relative densities in the 50nm case
148 cannot be reported. A lower relative density of 80% was obtained in the pure 1 um case, due to a lower driving
149 force for sintering [30]. In all bimodal mixtures, relative densities of ~93% were obtained. These values are found
150 in Table 2.

151 To assess the evolution of phases before and after cold sintering, X-ray diffraction was performed. Fig.3 (a)
152 presents a wide range 2θ XRD scan of a BT cold sintered sample of 200 nm precursor powder. All BT peaks are
153 present, but there is additionally a BaCO₃ peak around 2θ = 23.8° as seen in Fig.3 (b). The intensity of the peak
154 increases with decreasing BT particle sizes, 200nm, 100 nm and 50 nm and increases with an increase in volume
155 fraction of the nano particles relative to the 1.0 μm BT (Fig.S.3-5). It is inferred that there is a carbonate reaction
156 between the water phase from the decomposition of the BaOH₂·8H₂O and CO₂ from the air under the cold
157 sintering process. The increase in surface area from increasing amounts of smaller particle sizes in the bimodal

158 particle mixtures, increases the amount of the carbonate reaction. The BaCO₃ content being of lower permittivity
159 in BaTiO₃ will lower the overall relative permittivity.

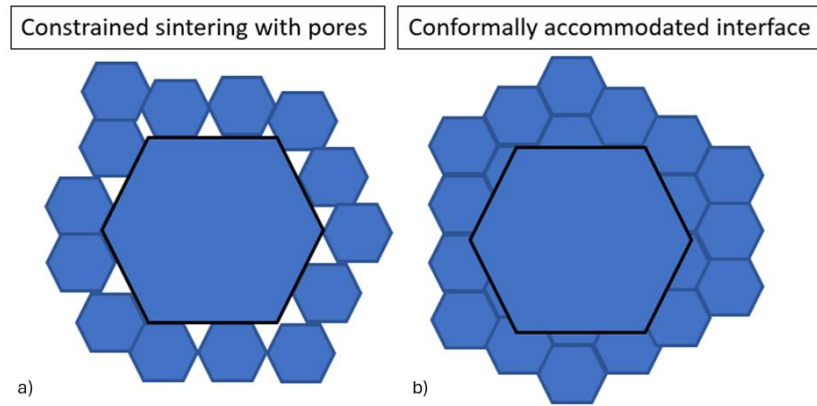


160
161 Fig.3. (a) 2 θ X-ray diffraction scan of a selected cold-sintered BaTiO₃ pellet (200 nm precursor powder). BaTiO₃ peaks are
162 marked with boxes, carbonate peaks are marked with open circles, Si peaks from a standard are marked with asterisks. (b) A
163 close-up of the barium carbonate peak taken between 2 θ = 23.0 – 25.0°.

164

165 3.2. Characterization & Microstructural Analysis of Cold Sintered Barium Titanate 166 Bimodal Ceramics

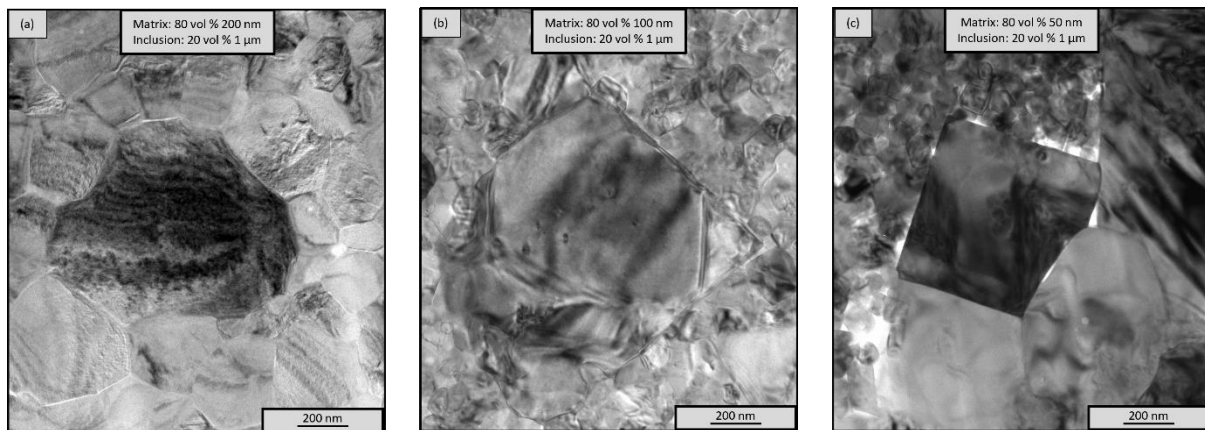
167 Before considering the reasons for microstructural development of the cold sintered bimodal mixtures, it is
168 important to consider the risk of constrained sintering that could limit densification locally at the interfaces
169 between the micron inclusions and nanoparticles within the matrix. Fig.4 shows the basic concept of constrained
170 sintering in bimodal particle size mixtures. Each of the larger particles potentially act as a constraining substrate
171 to the surrounding small particles. Such constrained interfaces could limit the densification along these surfaces,
172 as major densification must be normal to the interface. Earlier, a comprehensive review of constrained sintering
173 was made by Green *et al* [31]. From that review and subsequent studies [32], it is expected that samples sintered
174 under constraints will display anisotropic macroscopic shrinkage and porosity evolution, thereby limiting the
175 overall densification, and leaving residual porosity. However, with the pressure solution mechanism acting normal
176 to the stressed constrained surfaces, densification along these surfaces is expected to be permitted. This provides
177 microstructures with bimodal grains with limited porosity, high densities and is promising for the design of future
178 cold sintered particle size mixtures and composites with constrained surfaces.



179

180 Fig.4. A schematic illustrating sintering of large inclusions into a matrix phase. (a) Depiction of a large grain limiting shrinkage
 181 of the surrounding finer particles owing to constrained sintering at the interface and residual pores trapped at the interface. (b)
 182 Depiction of a case where creep diffusional processes dominate at the interface between the large and fine-grained particles
 183 resulting in a conformally accommodated interface between matrix and inclusion.

184 TEM imaging provides an excellent way to consider the interfaces between large and small grained BaTiO₃
 185 particles undergoing cold sintering. With the 80 vol% of 100 nm and 200 nm – 1 μm BT particle mixtures, the
 186 interfaces are conformal because of the stress transfer around these interfaces via the pressure solution creep
 187 processes (Fig.5. (a) and (b)). The interfaces are desirable as they remove a major porosity source. The 80 vol%
 188 50 nm - 1 μm interfaces however have a greater number of regions with porosity at their interfaces (Fig.5 (c)).
 189 This is most likely because of poor distribution of the flux in the 50 nm BT particles due to agglomeration.

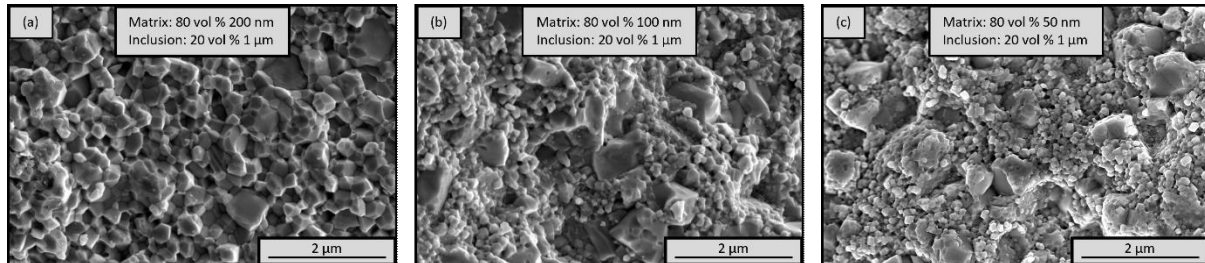


190

191 Fig. 5. TEM bright field images of bimodal mixtures with varying matrix size. (a) 200 nm matrix with a conformally sintered
 192 interface with impaction onto an inclusion. (b) 100 nm matrix with conformally sintered interface without substantial impact
 193 on the inclusion. A morphological change of the inclusion into a cubic geometry is observed. (c) Constrained sintering with
 194 substantial porosity around an inclusion. A morphological change of the inclusion into a cubic geometry is observed. Other
 195 TEM micrographs are shown in Fig.S. (8) and (9).

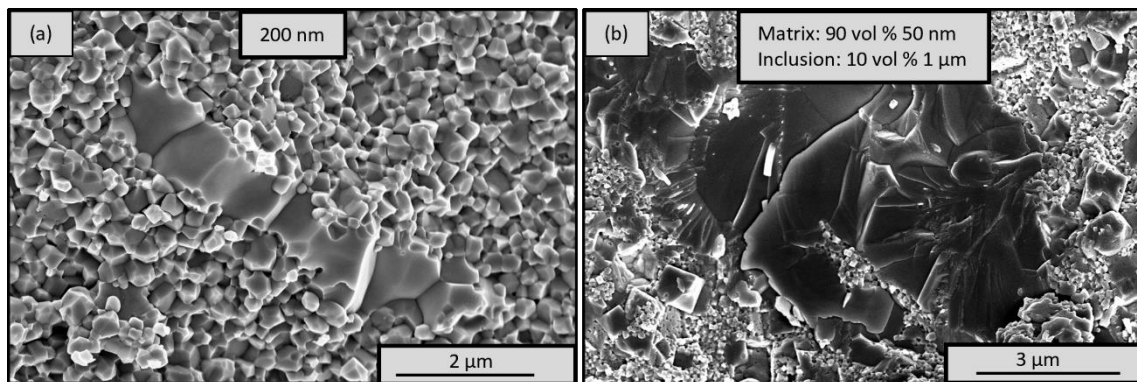
196 The BT grain structures of the bimodal mixtures from SEM images are also presented. Fig.6 compares the fracture
 197 surface of the 80 volume % of 200, 100, and 50 nm – 1 μm BT particle size mixtures. The highest density structure
 198 is observed in the case of the 200 nm – 1 μm mixture case. These interfaces are additionally very well faceted,
 199 and conformal with minimal pitting. However, as seen in the TEM images, as the matrix particle size is decreased
 200 the surfaces become less conformal, and in the case of the 50 nm – 1 μm mixture, there is an increase in porosity

201 because of the poorer distribution of the flux. With decreasing matrix particle size, there is also an increase in
202 pitting within the 1 μm inclusions. Which is believed to be a result of stress amplifications against those regions
203 (Fig.S.14.(c)), which accelerates dissolution in the direction normal to the stress (as noted in the conformally
204 sintered interfaces). These features are additionally consistent with geologically observed pressure solution creep,
205 in the pitting of pebbles within natural systems [17].



206
207 Fig. 6. SEM images of 200 nm 100 nm, and 50 nm matrix BT bimodal mixtures with volume fractions of 1 μm inclusions
208 (80/20). There is good mixing of the grain sizes, the grain morphologies are faceted, and in the 1 μm grains there are
209 indentations (surface holes) from the nanograin at the surfaces, a residual effect of pressure solution creep.

210 For completeness, we note that occasionally amongst several samples made, a highly unusual grain coarsening in
211 isolated regions with elongated morphology was observed and is shown in Fig.7 (a) and (b), Fig. S.16. At this
212 time, the origin of growth is not known, but it is suspected that enriched and trapped molten fluxes could drive
213 the local grain growth. Additionally, there is still abnormal grain growth in the mono-particle cases. The rare event
214 aspect of these observations means that we will not consider these in the following electrical analysis. However,
215 as this is a surprising observation, we report it here in the hope that future researchers may bring light to the
216 fundamentals of this mechanism.



217
218 Fig. 7. SEM images demonstrating various examples of the unusual abnormal grain structure observed in cold sintered BaTiO₃
219 samples. (a) Elongated abnormal grain growth in a pure 200 nm sample. (b) Coarsening is observed with growth encasing a
220 portion of the 50 nm matrix. Multiple growth processes are operating in this abnormal grain growth region.

221 3.3. Electrical Properties of Bimodal Mixtures and Dielectric Trends

222 In previous studies of cold sintered barium titanate, reported resistivities were orders of magnitude lower than
223 conventionally sintered barium titanate, as reported in Table 1. By applying a 500°C anneal in oxygen atmosphere
224 for 12 hours, the resistivities of the cold sintered BaTiO₃ were substantially raised to those similar in literature.

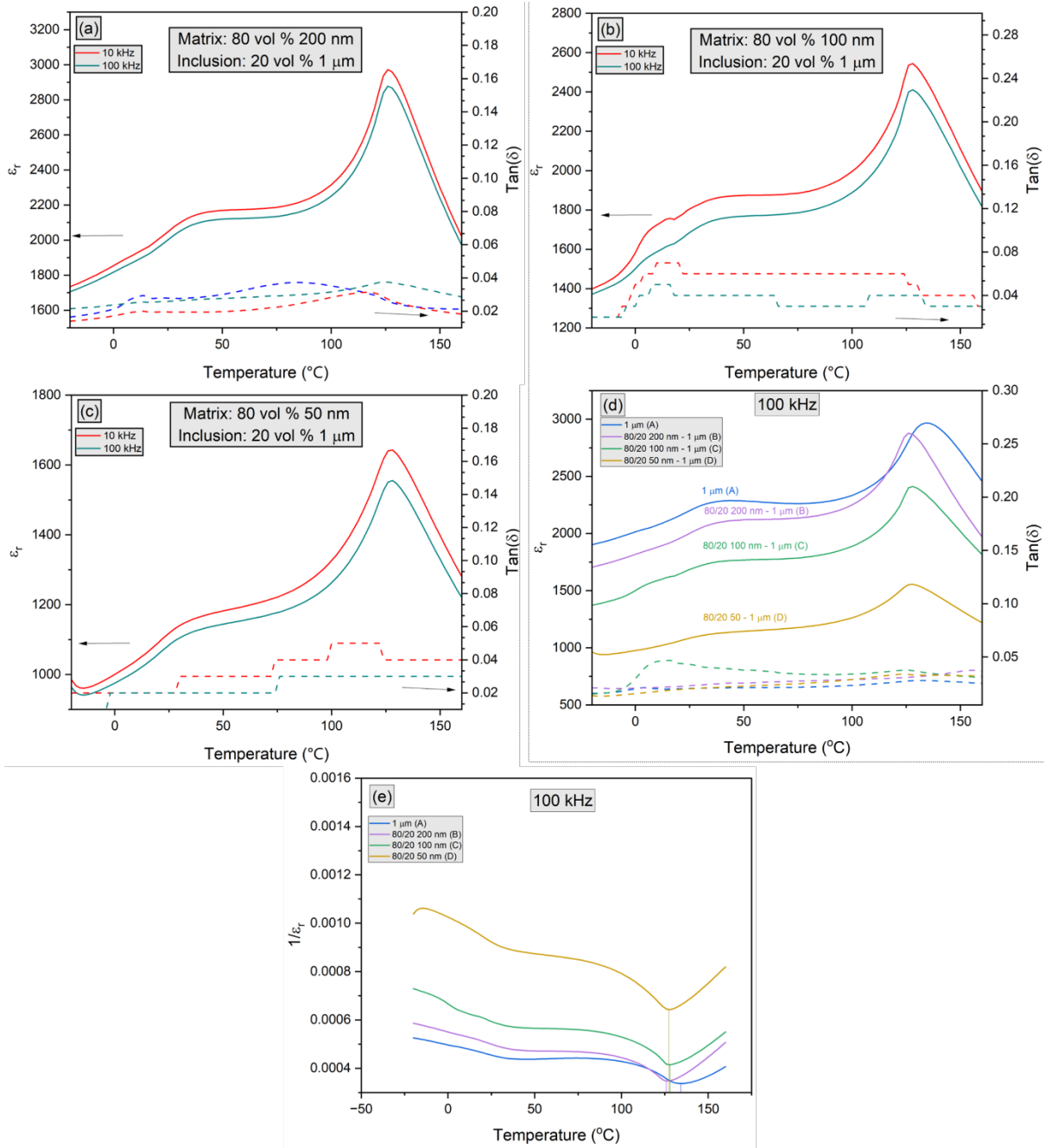
225 This annealing temperature is below that of which carbonates would decompose, and so the significant increase
 226 in resistivity could be due to enrichment of the grain boundaries with oxygen [33].

227 Table1. Resistivity Values for Barium Titanate Processed with Different Temperatures.

Sample	Highest Processing Temperature, T(°C)	Resistivity, $\rho(\Omega \cdot \text{cm})$	Reference
Cold Sintered BT-PTFE composite	225 °C, BHO flux	1×10^{12}	[34]
Cold Sintered pure BT	225 °C, BHO flux	1×10^8	[34]
Cold Sintered pure BT	500 °C, oxygen atmosphere anneal	$2 \times 10^{12} - 9 \times 10^{12}$	This work
Conventionally sintered BT	1190 - 1320 °C	$2 \times 10^{12} - 12 \times 10^{12}$	[35]

228

229 Fig.8, and Fig.S.10 show capacitance-temperature measurements that were then performed to determine the
 230 relative permittivity, dielectric loss, and Curie-Weiss behavior at 10 kHz and 100 kHz, over a temperature range
 231 from -20°C to 160°C.



232

233 Fig. 8. (a-c) Temperature and frequency dependence of relative permittivity and dielectric loss for various particle mixtures.
 234 (d) Changes in relative permittivity and dielectric loss as a function of matrix combinations. Highest permittivity of the cold
 235 sintered samples is 1 μm and is $\sim 90\%$ dense, then a systematic decrease with 80/20 bimodal mixtures 200 nm, 100 nm and 50
 236 nm that are $\sim 93\%$ dense. (e) Curie-Weiss plot for the data in (d).

237 Comparing the bimodal mixtures to the monodispersed powders, an increase in relative permittivity is
 238 observed due to the average grain size approaching 0.8 μm . In the case of the monodispersed powders, the 1 μm
 239 sample exhibited the highest permittivity out of all samples. However, this was still below values reported in
 240 literature due to the high volume of porosity. The 100 nm and 200 nm BT demonstrated permittivity values
 241 comparable to those reported in literature, with slightly depressed permittivity, likely due to carbonates, and
 242 intergranular voids (S. 8) [36], [37].

243 As seen in Table 2 and figure. 8e, curie constants were additionally calculated and compared to reported literature
 244 values using a modified Curie-Weiss law [38]:

$$245 \quad \frac{1}{\epsilon_r} - \frac{1}{\epsilon_m} = \frac{(T-T_c)^\gamma}{C} \quad (\text{eq.1})$$

246 where ϵ_r is the relative permittivity, ϵ_m is the maximum relative permittivity at the Curie temperature, C is the
 247 Curie-Weiss constant, and γ is the diffuseness degree. When γ is 1, a normal ferroelectric behavior is observed,
 248 while as γ approaches 2, relaxor ferroelectric behavior is measured. The γ value was calculated by linearizing the
 249 modified Curie-Weiss law and calculating the slope as reported in other work [38]. In this work, values of γ for
 250 bimodal mixtures were lower than that reported in other cold sintering studies, which may be due to experimental
 251 differences between these works [39]. In the mono sized particles, a substantially elevated γ is found with a
 252 depressed ϵ_r relative to expected values. It is likely that these deviations represent the presence of point defects
 253 from sources such as barium carbonate or residual flux after cold sintering which were unresolved by the current
 254 heat treatment. The residual space charge, the finite frequency dependence of the permittivity in the paraelectric
 255 regime and the small grains perturbs the magnitude of the Curie constant that would ideally be 2.5×10^5 C
 256 particularly in the case for the 1 μm BT particles.

257 Table2. Comparisons of the 500°C Annealed Cold Sintered Dielectrics of Mono and Bimodal BaTiO₃ Ceramics

Grain Size	ϵ_r (50 °C, 100 kHz) (Cold Sintered)	Tan(δ) 100 kHz	T _c	γ	C (x 10 ⁵ K)	Relative density
1 μm	2260	4.3%	134	1.75	39	80%
80/20 200 nm - 1 μm	2125	3.1%	126	1.06	2.2	93.9%
80/20 100 nm - 1 μm	1768	3.9%	128	1.15	0.45	93.8%
80/20 50 nm - 1 μm	1237	2.9%	128	1.07	0.45	92.9%
200 nm	800	2.8%	130	1.36	0.27	94.3%
100 nm	900	3%	128	1.27	0.22	92.2%

258

259

260 Finally, to then test whether the electrical measurements followed the initial bimodal size mixtures, a logarithmic
 261 mixing law (eq. 2) is used to track the increase in average relative permittivity from the 1 μm inclusions [40].

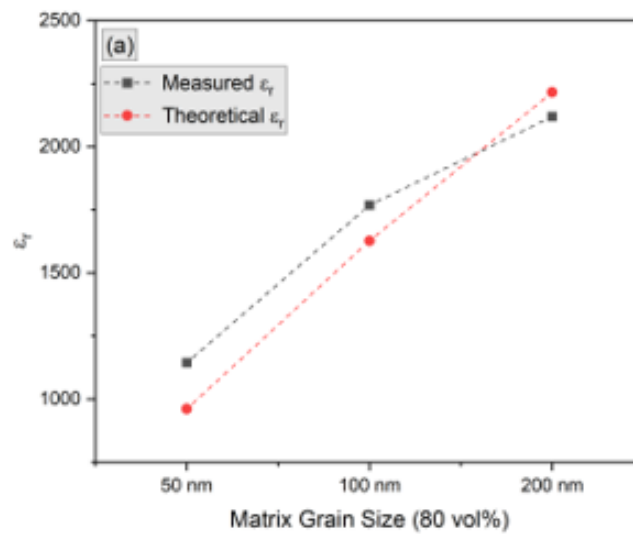
262 Here $\bar{\epsilon}$, is based on the individual relative permittivity of each constituent phase, porosity and the volume fraction
 263 and permittivity of each of the i^{th} phases given by f_i and $\epsilon_{r,i}$ respectively

264
$$\log \tilde{\epsilon} = \sum f_i \log \epsilon_{r,i} \quad (\text{eq.2})$$

265 The constituent phases of the bimodal mixtures will have a mixture of parallel and perpendicular connectivity
 266 between fine and large grain BaTiO₃ inclusions [41]. Additionally, there is assumed to be a random porosity
 267 associated with incomplete densification that also dilutes the permittivity and can be estimated from the relative
 268 density. As an approximation, the constituent phases are the large grained BaTiO₃, finer grained BaTiO₃, and the
 269 porosity with a relative permittivity of 1 (eq.3).

270
$$\log \tilde{\epsilon}_1 = f_1 \log \epsilon_{r1} + f_2 \log \epsilon_{r2} + f_3 \log 1 \quad (\text{eq.3})$$

271 In our analysis, we take the relative permittivity of samples at 100 kHz and 50°C. We assume the porosity volume
 272 fraction as 5 vol% to account for the variance in densities between the current work compared with those reported
 273 in the literature using various sintering techniques [24], [36], [37]. We take the trend of the average relative
 274 permittivity of the bimodal mixtures in 20/80 volume ratios with the matrix phase being the 80% phase for BaTiO₃
 275 with 50, 100 and 200 nm grain sizes, and the other 20% filler being the 1.0 μm particles, with 200 nm and 100 nm
 276 samples ~94% dense and 50 nm samples ~ 93% dense. This gives a relative permittivity change of a factor of 2
 277 between the extremes. The trend is consistent with eq.2, as shown in Fig.9. The theoretical curve uses ε_r values of
 278 1000 for 50 nm, 2000 for 100 nm, 3000 for 200 nm, and 5000 for 1 μm [24].



279
 280 Fig. 9. Improvements in relative permittivity as a function of matrix size at constant inclusion volume. A theoretical model
 281 using the logarithmic mixing law accounting for 5 volume percent porosity is used to predict the expected permittivity of a
 282 bimodal mixtures using published values of conventionally sintered 1.0 μm BT from literature [24].

283 The measured ε_r trends are generally in agreement with that predicted via the mixing law. In the theoretical analysis
 284 an expected increase in ε_r occurs as a function of matrix grain size due to the influence of the size and scaling
 285 effects. The measured ε_r shows slight deviations in shape from the mixing law, with the 200 nm matrix mixtures
 286 showing a depressed ε_r which could be accounted for by intragranular voids (S.8) and localized barium carbonate
 287 at the grain boundaries. A slightly higher measured ε_r compared to prediction occurs in 50 nm and 100 nm matrix
 288 mixtures, indicating an overestimation of porosity or average grain size in our model. Despite slight deviations

289 from the theoretical mixing law, this study demonstrates that controlled bimodal particle size mixtures of barium
290 titanate can be successfully designed by the cold sintering process.

291 **4. Conclusion and Future Work**

292 In this study, the degree of control on morphological evolution by the cold sintering process was assessed with
293 bimodal particle size mixtures. Through the pressure solution creep mechanism, densification of conformed
294 surfaces was made possible, and the bimodal particle size distribution in dense bimodal mixtures was mostly
295 preserved. It is important to note that rare events of abnormal grain growth were observed. Additionally, by
296 pairing this with an annealing step, an increase in resistivity, and decrease in dielectric losses are achieved and
297 compared to previous cold sintering studies. The dielectric properties in the 1 μm monodispersed BT have high
298 frequency dispersion with the cold sintering and this gives rise to an unrealistic Curie constant, C , due to the
299 additional space charge contribution. With the bimodal BT mixtures this is suppressed and shows a Curie Constant
300 that is more intrinsic and with small grain sizes being impacted by the size and interfacial effects. Electrical
301 performance of the relative permittivity was successfully enhanced by the incorporation of 1 μm grains, and
302 followed trends proposed by a theoretical mixing law accounting for porosity.

303 Future work should explore alternative techniques for improving mixedness of the barium hydroxide octahydrate
304 flux, as this made the sintering of finer grain BT more difficult. Additionally, while the abnormal grain growth
305 could be partially explained by those areas being enriched by the barium hydroxide octahydrate, its presence in
306 monodispersed cases questions this. While electrical properties demonstrated the preservation of a bimodal
307 particle size distribution, values of relative permittivity were lower than literature due to carbonate formation and
308 potential point defects. Alternative fluxes, sintering conditions including atmospheres, and post annealing
309 strategies could be explored as potential solutions.

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