Nonmonotonic Composition Dependence of the Dielectric Response of Ba$_{1-x}$Ca$_x$ZrO$_3$

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We use first-principles density functional theory calculations to investigate the dielectric response of BaZrO$_3$ based perovskites. In this study we have substituted Ca for Ba over a range of compositions to understand what causes the recently reported (Levin et al., J. Solid State Chem. 2003, 175, 170) nonmonotonic dielectric response of Ba$_{1-x}$Ca$_x$ZrO$_3$ (BCZ). We show that in all compositions studied there appear two new sets of IR active phonon modes that increase the dielectric constant $\varepsilon$. One set occurs below the normal IR active A-O modes, due to the “rolling cation” motion of Ca about its O$_{12}$ cage. The second set of new phonon modes occurs between the normal A-O and B-O mode frequencies, caused by the displacement of Ca along one Cartesian direction. As more Ca is added to BCZ, Ca−Ca interactions increase, leading to Ca displacements in all three Cartesian directions, stronger Ca−O bonds and increased O$_6$ tilts. Such structural deformations decrease $\varepsilon$. These effects are responsible for the anomalous compositional dependence of the BCZ dielectric response.

I. Introduction

Dielectric materials are widely used in wireless communication devices. These require a high dielectric constant, $\varepsilon$, and low dielectric loss.\(^1\) Barium zirconate, BaZrO$_3$ (BZ), is an ABO$_3$ perovskite dielectric material that is both chemically and mechanically stable. Since dielectric properties are often optimized by varying material composition, there is a need to understand the impact of doping on the local structure and properties of a material. One study showed that the dielectric constant of BZ can be increased by up to 50% by doping with small amount of Ca\(^2\) as seen in Figure 1. This is interesting because CaZrO$_3$ (CZ) has a lower $\varepsilon$ than BZ. Such a nonlinear dependence of $\varepsilon$ on composition was also observed in Ba-based perovskites with other B-site compositions.\(^3\)−\(^6\) In this paper, we use first principles calculations to study the BCZ solid solution at a variety of compositions, including some that were not accessible experimentally.\(^2\) We show how the changes in static structural features influence the dynamical behavior of these materials, giving rise to a new vibrational mechanism and enhanced dielectric response.

II. Methodology

In this study, two first-principles codes are used. An in-house solid state DFT code, BH, used in previous studies\(^7\)\(^,\)\(^8\) and the

Figure 1. Directionally averaged ($T = 0$ K) dielectric constant $\varepsilon$ vs Ca content $x$ in BCZ. The $\varepsilon$ values are also averaged over cation orderings. Black circles are the averaged diagonal elements of the $\varepsilon$ tensor. Low $T$ experimental values\(^6\) are shown as red squares. Also shown is the fraction of isolated Ca atoms, $x(1-x)^6$, as a function of $x$.\(^9\)

(2) Levin, I.; Amos, T. G.; Bell, S. M.; Farber, L.; Vanderah, T. A.; Roth, R. S.; Toby, B. H. J. Solid State Chem. 2003, 175, 170.
ABINIT software package\(^9\) are used to relax the ionic positions and lattice constants. The local density approximation (LDA) of the exchange correlation functional and a \(4 \times 4 \times 4\) Monkhorst—Pack sampling of the Brillouin zone\(^10\) are used for all calculations. All atoms are represented by norm-conserving optimized\(^11\) designed nonlocal\(^12\) pseudopotentials. All pseudopotentials are generated using the OPIUM code.\(^13\) The calculations are performed with a plane wave cutoff of 50 Ry.

At low temperatures, the ground-state structure of BZ is an irreducible \(2 \times 2 \times 2\) pseudocubic unit cell,\(^14\) containing \(\text{O}_6\) tilt angles that range between 3.6 and 4.2°. The computed BZ \(\varepsilon\) is 50 (45 from the ionic contribution and 5 from the electronic) at 0 K, larger than the value reported at room temperature. This is caused by the frozen tilts present at low \(T\). Approximating the ground-state BZ by a 5-atom cell\(^15\) leads to a significant overestimation\(^14\) of \(\varepsilon\) at \(T = 0\). Therefore, the majority of the calculations in our study were performed on \(2 \times 2 \times 2\) supercells of BZ. The ground-state structure of CZ at low temperatures is orthorhombic, displaying \(\text{O}_6\) tilt angles that vary between 17.1 and 17.5°. The computed CZ \(\varepsilon\) is 30, about half that of BZ. In the ground-state CZ structure, antiferroelectric displacements of Ca atoms are mostly along one Cartesian direction, with an average magnitude of 0.4 Å. This is over an order of magnitude larger than the displacements of Ba seen in BZ. We also performed calculations for \(3 \times 3 \times 1\) supercells. In these cases, Ca was substituted for Ba at \(x = 1/9\), \(x = 2/9\), and \(x = 1/3\). For \(1 \times 1 \times n\) and \(2 \times 1 \times n\) supercells, where \(n\) was varied from 2 to 6, \(x = 1/n\) and \(x = 2/n\) are considered. This was performed in order to see the effects of Ca substitution on nearest neighbor unit cells. Several different sets of initial ionic coordinates were used in the relaxations for each composition to ensure that the complicated potential energy surface of the material was adequately explored. For all compositions, we found that the DFT-LDA optimized lattice constants were within 1.5% of the experimental values.

After a structure was fully relaxed, response function\(^16\),\(^17\) calculations were performed using ABINIT to generate the mass-weighted dynamical matrix which was then used to calculate the directionally averaged dielectric constant \(\varepsilon\) at \(\Gamma = 0\). Contributions to \(\varepsilon\) come only from IR active \(\Gamma\) point phonon modes.\(^18\) Details of this method are presented in a previous study.\(^14\) A full listing of the diagonal elements of the \(\varepsilon\) tensor is provided in the Supporting Information, Tables I and II. A comparison of the experimental and computed \(\varepsilon\) for the entire compositional range of BCZ is presented in Figure 1. The magnitude of the experimental\(^2\) and theoretical dielectric constants show reasonable agreement, with both showing the trend of an initial sharp increase up to 10% Ca and subsequent gradual decrease in \(\varepsilon\) with greater Ca fraction.

III. Results

The unit cell volume is the most basic characteristic of the solid-state structure. The volume of BZ is 16% larger than that of CZ, and one might expect a simple linear change in the volume as Ca fraction is increased. Somewhat surprisingly, we find that as Ca is added to BZ, the volume of BCZ does not monotonically decrease (Figure 2). In compositions \(x = 0\) through \(x = 0.50\) for \(2 \times 2 \times 2\) supercells, the volume remains nearly constant. Then, when Ca becomes the majority A-site, the volume drastically decreases. This is supported by the change in volume from components to solid solution for \(x = 0.125\) through \(x = 0.50\) and then subsequent decrease at \(x = 0.875\). A trend that is different from this is the near monotonic increase in formation energy for these supercells. Both are presented in the Supporting Information. Comparison of our results with available experimental data shows close agreement between DFT and experimental lattice constants. The relaxed volumes are slightly underestimated by DFT-LDA, as expected. The constant volume for \(x < 0.50\) implies that the changes in \(\varepsilon\) for these compositions are not due to volume effects; rather they are caused solely by the changes in structural distortions at the same volume.

Several important structural changes are induced by Ca doping into BZ. While the Ba atom in BZ is located at the high-symmetry position in the center of the \(\text{O}_ {12}\) cage, Ca atoms in BCZ displace significantly toward a low-symmetry position. The small Ca forms shorter Ca—O bonds along (110), to satisfy bonding requirements, by displacing in (100). Ca displacements increase with increased Ca doping, from 0.3 Å at \(x = 0.125\) to 0.5 Å at \(x = 0.5\). This is somewhat larger than the 0.4 Å displacements of Ca in CZ and is due to the larger volume of BCZ.

The direction of Ca displacements shows a dependence on Ca concentration. At low doping levels, Ca displacements are parallel to each other along (100). As more Ca is added, Ca—Ca interactions increase and \(\text{O}_6\) octahedra rotate to accommodate the displaced Ca. This locally reduces the available volume and gives rise to Ca displacement components in other directions, such as (110). The total magnitude of Ca displacement continues to increase with increasing \(x\), although the displacement component along the (100) direction is nearly unchanged. Such distortions are also seen in CZ. Thus, as the Ca fraction rises, the Ca local environ-
The presence of Ca displacements induces Ba and Zr atoms to displace as well, opposing the Ca. These displacements are small (∼0.1 Å) and are localized on the Ba and Zr atoms adjacent to Ca. The structural trends are shown in Figure 3. Finally, the presence of small Ca atoms increases the average O6 tilt angle in BCZ with increasing x.

The changes in the structural properties described above give rise to a new dielectric response mechanism and increased value of ε. In BZ, ε is 50. In BCZ compositions below x = 0.15, computed ε is as high as 70. The breaking of symmetry induced by x = 0.125 Ca doping broadens the discrete Γ point phonon frequencies of BZ into a distribution of frequencies, as shown in Figure 4. Most importantly, new IR active phonon modes appear at frequencies that are not present in either BZ or CZ, as well as more modes centered around 400 cm⁻¹. All show IR active low frequency phonon modes that are small (≈0.1 Å) and are localized on the Ba and O bonds in the O12 cage, making the Ca-O bond weaker. Bond-valence sums support our DFT finding of weaker Ca-O bonds in BCZ. We find that Ca bond valence computed using relaxed DFT structures is 1.8 for Ca in BCZ and 2.05 for Ca in CZ. Similar conclusions about the weak bonding for Ca in BCZ were reached by Levin et al. using bond valence analysis of structures obtained from refinement of neutron diffraction data.²

Figure 3. Cation displacements for x = 0.125, 0.250, 0.375, and 0.500. Average displacement increases with the addition of Ca. Ba is shown as circles, Ca is shown as squares, and Zr is shown as diamonds. Average O6 tilt angle (in degrees) increases with the addition of Ca, shown here as triangles.

Figure 4. IR active phonon frequency comparisons of 2 × 2 × 2 BZ to BZ, shown in black, and CZ, shown in blue. Not shown are O6 modes above 400 cm⁻¹, which do not shift as significantly. (a) Baₐ₀.₈₇₅Ca₀.₁₂₅ZrO₃. (b) Ba₀.₆₅Ca₀.₃₅ZrO₃. All show IR active low frequency phonon modes that are not present in either BZ or CZ, as well as more modes centered around 130 cm⁻¹.

Figure 5. IR active phonon modes in Ba₀.₈₇₅Ca₀.₁₂₅ZrO₃. Ba atom is shown as blue, Ca is green, Zr is white, and O is red. Arrows depict relative atomic displacements. (a) The rolling cation motion associated with Ca. This mode occurs at 49 cm⁻¹, below (b) at 60 cm⁻¹ where Ca moves along the direction of its displacement from high symmetry. (c) All A-sites move, which occurs at 93 cm⁻¹.
The Ca motions in CZ and BCZ differ from the common rattling cation model of ionic vibrations in perovskites. In this model, it is assumed that the ionic vibrations are due to the oscillations of a cation from its central location within the unit cell and are similar in all three Cartesian directions. This is the case for Ba in BZ and for the Ba atoms in BCZ. However, in BCZ the oscillation of the Ca around its off-center, low-symmetry position is noticeably more pronounced in the two Cartesian directions perpendicular to the direction of Ca off-center displacement. This gives rise to an arc-like motion similar to that of a hindered rotor, with the Ca atom rolling on the surface of a sphere. The energy required to excite such a “rolling cation” motion in BCZ is less than that for the usual rattling cation mode, leading to a strong enhancement of the A-site dielectric response.

The introduction of Ca into BZ also transforms some of the modes that are Raman active in BZ into IR active modes in BCZ. This is caused by local dipoles created by Ca displacement. These modes occur around the frequency of the strongest Ca–O mode in CZ and are localized on Ca and its surrounding ions. At low doping levels, the local environment of displaced Ca in BCZ is similar to that of displaced Ca in CZ, so phonon modes at these frequencies are similar. However, as more Ca is added to BCZ, it causes the surrounding O₆ octahedra to increase their tilt angles to accommodate increased Ca displacement. This increased tilting causes not only Ba to displace but Zr as well, creating displacements which are further enhanced by an induced phonon mode. These modes are fairly high in frequency and therefore provide only a small increase in the dielectric response.

We now discuss the compositional dependence of the dielectric constant in BCZ. Figure 4a,b shows that as Ca content increases past \( x = 0.125 \), the A-O and B-O BZ-derived modes are further broadened over a wider range of frequency. The O₆ modes and electronic contributions, not shown, display the least compositional dependence. The total directionally averaged B–O contribution to the dielectric constant is in the range of 18–20, and the O₆ and electronic contributions are both \( \approx 5 \) for all compositions. The largest changes are observed for Ca–O rolling cation modes, where an increase in \( x \) causes stronger Ca–O bonds in all three Cartesian directions. This manifests as an increase in the phonon frequencies associated with these modes. Although at \( x = 0.25 \) the intensity of the low-frequency Ca–O modes is larger than at \( x = 0.125 \), their contribution to the dielectric constant is smaller due to the blueshift in the Ca–O mode frequencies and the \( 1/\omega^2 \) dependence of the dielectric constant. This leads to a decreased dielectric constant at \( x = 0.25 \) relative to \( x = 0.125 \). Further increase in Ca concentration introduces more Ca–Ca interactions and gives rise to further upshift of the Ca–O modes. Similar to the Ca–O modes, the frequency of the former Raman modes shifts up with increased Ca concentration so that the contribution to \( \varepsilon \) due to the former Raman modes first increases (at low \( x_{Ca} \)) and then decreases (for higher \( x_{Ca} \)) with increasing Ca content. For \( x > 0.5 \), the phonon modes become less broad and there is a general shift to higher \( \omega \) values, further decreasing the dielectric response.

The nonmonotonic dielectric constant dependence on composition is due to the local structural changes resulting from increased Ca content. For small Ca concentrations, the localized character of the low frequency modes means that Ca–Ca interactions will have a small impact on the Ca–O vibrational frequencies. This leads to a linear increase in the intensity of the low-frequency Ca modes and therefore \( \varepsilon \) as the number of Ca is increased.

The decrease of the dielectric constant with increasing Ca content is due to the structural changes induced by the presence of two Ca atoms in close proximity to one another, found frequently for nondilute solid solution compositions. The first Ca that is added to a BZ supercell displaces along (100). In 40-atom supercells with two Ca atoms (\( x = 0.25 \), Ca atoms coordinate their displacements mostly in (100), but with some components in (110) and (101). This is because the large O₆ tilts induced by the close proximity of two Ca atoms limits the available volume. At low doping levels, Ca displace parallel to each other along (100), but at higher doping levels, structural deformations limit their ability to do so. This effect becomes stronger with additional Ca, until large directional displacements are no longer feasible and smaller displacements are made along (111). Total Ca displacements increase, but the components in each direction are smaller than at lower dopant values. Off-center displacements in the directions perpendicular to the primary distortion direction strengthen the Ca–O bonds, increasing the Ca–O frequency and decreasing \( \varepsilon \).

The dielectric constant peaks at compositions for which Ca–Ca interactions are not strong. This suggests that the turnover should take place at \( x \) for which the population of Ca atoms with at least one Ca nearest neighbor is rising rapidly with composition. Simple statistics show that such a rise is present at small \( x \) (\( x < 0.15 \)) in BCZ, with the probability of Ca atom surrounded by only Ba nearest neighbors reaching its maximum value at \( x = 0.14 \), as shown in Figure 1. At this \( x \), the maximum amount of Ca having no nearest Ca neighbors is 6%. This explains why the turnover in \( \varepsilon \) of BCZ occurs at low \( x \), as observed experimentally and in our DFT calculations. For low \( x \), Ca–Ca interactions are rare and do not hinder the “rolling cation” motion. As \( x \) increases, nearest Ca–Ca neighbors become predominant, causing a decrease in the “rolling cation” motion. The mechanisms described above do not depend on the particular chemistry of Ba, Ca, and Zr ions but are due to a size mismatch on the perovskite A-site between the majority Ba cation and the minority Ca cations. Therefore we expect that the effects discussed above are relevant whenever doping a large cation perovskite A-site with a smaller cation.

IV. Conclusions

We performed a study of the nonmonotonic dielectric properties of the BCZ solid solution using first-principles calculations. We find that addition of Ca to BZ gives rise to two new sets of phonon modes. The first set are low frequency Ca–O modes that occur at frequencies below either Ca–O or Ba–O in their native structures. Their contribution to \( \varepsilon \) varied according to local structure. The
second set of frequencies were Raman active modes that had become IR active with the introduction of Ca. The presence of very low frequency Ca–O modes initially increases the dielectric constant of the material. The much lower frequencies of Ca–O vibrations in BCZ relative to CZ are due to placing Ca into a large O12 cage, giving rise to weak Ca–O bonds. Due to the large Ca displacement in its O12 cage, Ca oscillations are asymmetric and can be described by the “rolling cation” model, with Ca rolling on the surface of a sphere. At higher Ca content, Ca–Ca interactions lead to a blueshift of the Ca–O modes and a decrease of the dielectric constant. The effects described for BCZ should be general to all dielectric perovskite systems in which a smaller A-site cation has been substituted into a larger A-site perovskite for a range of compositions.

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Supporting Information Available: Formation energies, volumes of formation, and data for 2 × 2 × 2 and 1 × 1 × n, 2 × 1 × n, and 3 × 3 × 1 supercells (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.