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Long- and short-range structure studies of KBT-KBZ solid-solutions using synchrotron radiation

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The relaxor ferroelectric $K_{0.5}Bi_{0.5}TiO_3$ has been synthesised in a solid-solution series with $K_{0.5}Bi_{0.5}ZrO_3$, as $K_{0.5}Bi_{0.5}Ti_{1-x}Zr_xO_3$. High-resolution synchrotron X-ray powder diffraction and X-ray absorption near edge structure spectroscopy were used to characterise the long-range average and local structural behaviour. Rietveld refinements against diffraction data show that a pseudocubic tetragonal region exists across the whole solid-solution series, with truly cubic symmetry only observed at $x = 1$ (pure KBZ). Variable-temperature diffraction data for $x = 0$ (pure KBT) showed a broad ferroelectric transition from tetragonal to cubic symmetry at approximately 683 K with a coexistence of both phases close to that temperature, accompanied by a marked volume contraction. Ti K-edge data showed that Zr doping has a minimal effect on Ti off-centering, and revealed no evidence for local clustering. Metal L-edges showed that Ti^{4+} cations remain off-centered with increasing Zr content, while Zr^{4+} cations approach a higher-symmetry coordination environment, most likely due to the increased size of the Zr atoms. Although there is a minimal effect on actual Ti-offsets, an effective dilution of these environments by Zr doping leads to a reduction in polar domains and a diminished ferroelectric response.

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1. Introduction

A thorough understanding of structure–property relationships underlies any rational search for new functional materials. In the case of ferroelectric materials, the most fundamental relationship is between the role of second-order Jahn–Teller distortions of metal cations with a d^0 electronic state,¹ and local (or long-range) ferroelectric order. In metal oxide perovskite (ABO_3) systems, the empty d orbitals of the B-site cation can hybridise with oxygen 2p orbitals causing a favourable offset of the B-site cation itself.² For a non-centrosymmetric crystal structure, this offset causes a net dipole.³ An increase in temperature can remove the B-site cation offset, rendering the material paraelectric at the Curie temperature T_C .

Ferroelectricity is widely found in perovskites and perovskite-related materials. The most important commercially is $PbZr_xTi_{1-x}O_3$ (PZT), which has a number of particular features, chief among which is the lead cation (Pb^{2+}) which enhances

the ferroelectric effect *via* additional distortions from its stereochemically active $6s^2$ electron lone pair. It has been found that the ferroelectric properties of PZT are further enhanced near a structural transition known as the morphotropic phase boundary (MPB).⁴ A great deal of research effort has been dedicated to finding lead-free alternatives to PZT in order to minimise the use of the toxic element, and encouraging the growth of MPBs appears to be a promising route.⁵ Bismuth-based materials have been shown to be feasible candidates, noting that Bi^{3+} is isoelectronic to Pb^{2+} and thus preserves the enhanced distortive effects. Charge balance upon replacing Pb^{2+} with Bi^{3+} can be achieved by an alkali metal cation of similar size, such as Na^+ or K^+ . There has been extensive research into bismuth titanates, including $Na_{0.5}Bi_{0.5}TiO_3$ (NBT) and $K_{0.5}Bi_{0.5}TiO_3$ (KBT), solid-solutions systems such as NBT-KBT,^{6,7} combinations with $SrTiO_3$,^{8,9} $BaTiO_3$,¹⁰ and the multiferroic $BiFeO_3$.¹¹

Relatively little research has been done into the effects of replacing the B-site cation, compared to the vast amounts of literature in A-site modifications. The $4d^0$ cation Zr^{4+} is only slightly larger than the $3d^0$ Ti^{4+} (ref. 12), and complete B-site substitution is possible in $BaTiO_3$ ¹³ and NBT¹⁴. The present study concerns the solid-solution between KBT and $K_{0.5}Bi_{0.5}ZrO_3$ (KBZ), first reported by Yamada *et al.*,¹⁵ and more recently the subject of a physical property study by Bengagi *et al.*¹⁶ A smooth structural transition was reported on the

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basis of sealed-tube X-ray powder diffraction (XRD) data, from the relaxor ferroelectric tetragonal KBT (Ti end member) to the paraelectric cubic KBZ (Zr end member). A pseudocubic region was reported for $x = 0.1$ to 0.4 , in which a dramatic decrease in the ferroelectric moment was observed.¹⁶

This work is specifically concerned with investigating chemical changes to the B-site and their effects on structural distortions in the KBT-KBZ system. Due to the presence of very heavy metals such as Bi, conventional XRD is inadequate to resolve the details of oxygen atomic displacements. Our approach is therefore to use a combination of high-resolution synchrotron XRD (SXRD) to investigate long-range average structure and X-ray absorption near edge structure (XANES) spectroscopy to investigate the local coordination environment of Ti^{4+} and Zr^{4+} cations. Our recent study using these techniques on the NBT-NBZ solid-solution concluded that Zr doping had little effect on Ti-offset,¹⁴ noting the slightly different sizes of Na^+ and K^+ , slightly different Zr doping effects can be expected in the KBT-KBZ series.

2. Experimental

~5 g samples of polycrystalline $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{1-x}\text{Zr}_x\text{O}_3$, where $x = 0$ to 1 (in 0.1 steps), were prepared by first weighing stoichiometric amounts of Bi_2O_3 (Aithaca, 99.999%, dried at 1023 K for 20 h before use), K_2CO_3 , TiO_2 , and ZrO_2 . The reagents were finely ground with a mortar and pestle with the addition of acetone to improve homogeneity. The mixture was then reacted as a powder in alumina crucibles initially at 1023 K for 8 h , then reground and heated to 1223 K for 8 h twice. At high temperatures, significant potassium loss was noted by Bengagi *et al.*¹⁶ An additional $10\text{--}15\%$ K_2CO_3 was therefore added to each sample to help minimise impurities. Long heating times improve the stability of the tetragonal phase of KBT, therefore the $x = 0$ and 0.1 samples were additionally heated to 1223 K for 72 h .¹⁶ Progress was monitored by X-ray powder diffraction (XRD) using a PANalytical X'pert PRO diffractometer in Bragg-Brentano geometry with a sealed-tube source of unmonochromated $\text{Cu-K}\alpha$ radiation ($\lambda_{\alpha 1} = 1.5405\text{ \AA}$, $\lambda_{\alpha 2} = 1.5443\text{ \AA}$).

Synchrotron X-ray powder diffraction (SXRD) data were obtained at the Australian Synchrotron, Melbourne, on the Powder Diffraction beamline.¹⁷ Samples were ground into a very fine powder and placed in 0.3 mm quartz capillaries for high-temperature measurements of KBT, or 0.3 mm glass capillaries for room-temperature measurements. Data were collected in Debye-Sherrer geometry at $13\,500\text{ eV}$. The corresponding refined wavelength against a LaB_6 NIST standard was $\lambda = 0.953296(1)\text{ \AA}$. Data collection under non-ambient conditions were achieved using a hot air blower. The temperature range was controlled between room temperature and 973 K (10 K steps , 5 K min^{-1} ramp) with $2 \times 150\text{ s}$ (2 detector positions) acquisition time at each step from $10^\circ \leq 2\theta \leq 90^\circ$. Room-temperature samples were collected over $2 \times 120\text{ s}$ acquisition times.

Initial structural models were adapted from the already reported structure of KBT.¹⁸ Rietveld refinements against diffraction data were carried out using the GSAS¹⁹ program with the EXPGUI²⁰ interface and extracted atomic positions were visualised using the VESTA software package.²¹ Cell parameters, anisotropic atomic displacement parameters (ADPs) as well as instrumental parameters and a zero offset were refined.

XANES measurements at the Ti L-edge for $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{1-x}\text{Zr}_x\text{O}_3$, where $x = 0$ to 0.9 were collected on the Soft X-ray beamline at the Australian Synchrotron.²² Powder samples were thinly dusted on double-sided carbon tape (SPI Supplies) and inserted into the vacuum chamber *via* a load lock. The pressure inside the analysis chamber was maintained at better than 10^{-9} Torr. All spectra were taken simultaneously with a total electron yield (TEY) signal measured from a Ti metal reference foil in the beamline. Approximately 10% of the beam intensity is removed by the reference foil. The Ti L₃-edge of metallic Ti is well known, with a distinct feature located at 453.8 eV .²³ This allowed for a precise energy calibration of the spectra from the various samples. The Ti K- and Zr L₃-edge XANES spectra were collected on beamline 16A1 at the National Synchrotron Radiation Research Center (NSRRC) in Hsinchu, Taiwan.²⁴ Samples were finely ground and dispersed onto Kapton tape and placed at a 45° angle to the incident X-ray beam. Spectra were collected in total fluorescence yield (TFY) mode using a Lytle detector with a step size of 0.2 eV near the absorption edge. The Ti K-edge spectra were calibrated against metallic Ti foil with the maximum in the first derivative of the K-edge set to 4966.4 eV . The Zr L₃-edge spectra were calibrated against elemental Zr with the maximum in the first derivative of the L₃-edge set to 2222.3 eV .²⁵ All XANES spectra were normalised using the Athena²⁶ software.

3. Results and discussion

The solid-solution $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{1-x}\text{Zr}_x\text{O}_3$ ($x = 0$ to 1 with 0.1 steps) was successfully synthesised and confirmed by Rietveld refinement against laboratory XRD. Refined occupancies showed that the Ti/Zr ratio were within their nominal composition ranges and hence fixed for refinement models. Interestingly, the refinement for KBT ($x = 0$) by Bengagi *et al.* shows the (111) reflection to be significantly undercalculated.¹⁶ A better fit to the peak profile was obtained here by refining for strain factors. The improvement is shown in Fig. 1 with the relevant statistics. This suggests that closer to the Ti-rich end member of the solid-solution series, there is a significant contribution from microstrain in the material, which explains the increased synthesis time required to stabilise this phase. For $x = 0.1\text{--}1$, the strain parameter did not significantly improve the fit and was therefore not used. This suggests that the introduction of a slightly larger cation rapidly removes the microstrain by granting additional flexibility to the crystal lattice. The best statistics in Rietveld refinements using laboratory XRD data were achieved using a non-centrosymmetric tetragonal $P4mm$ (#99) model from $x = 0\text{--}0.8$ and a cubic $Pm\bar{3}m$ (#221) model for



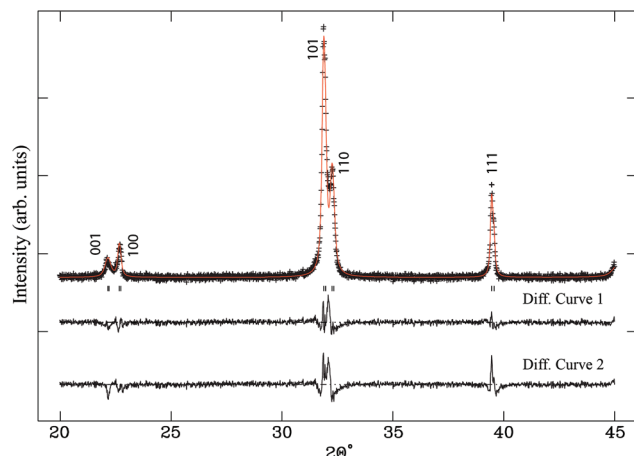


Fig. 1 The room-temperature laboratory XRD data of KBT. Black crosses represent data points with the calculated pattern shown as a red line. Bragg reflection markers are shown in black below the observed peaks. Diff. Curve 1 refers to the difference curve of the Rietveld refinement including strain parameters shown above ($\chi^2 = 1.673$, $R_p = 0.0264$). Diff. Curve 2 refers to a Rietveld refinement not including strain ($\chi^2 = 1.948$, $R_p = 0.0279$).

$x = 0.9$ – 1 . This preliminary result suggested that the tetragonal pseudocubic region extends to $x > 0.4$, above the previously reported limit.¹⁶

3.1. Synchrotron XRD

Higher-resolution SXRDX data were used to refine models to greater precision. The room-temperature data confirmed that the samples were pure apart from a very small (<2%) fraction of ZrO_2 in some of the samples, which could not be entirely eliminated despite the use of excess K to compensate for loss by evaporation.

In the conventional XRD data of Bengagi *et al.*¹⁶ diffraction peaks were generally observed to broaden with increasing Ti content through the pseudocubic region ($x = 0.1$ – 0.4). Visually, the same behaviour was observed in our SXRDX data for $x = 0.1$ – 0.4 , with only the pure KBT compound found to possess a

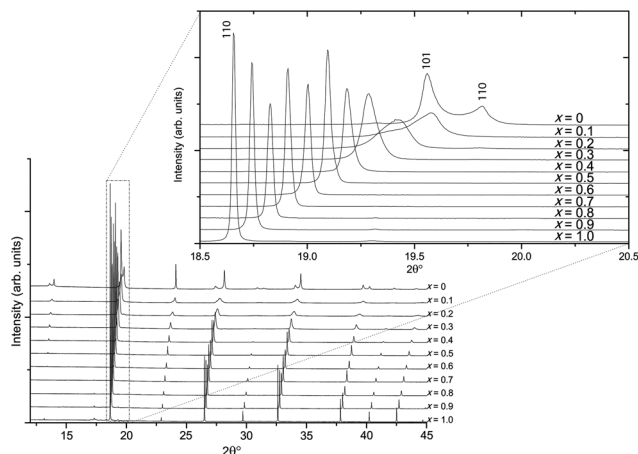


Fig. 2 Room-temperature SXRDX patterns of KBT-KBZ solid-solution.

clear tetragonal distortion (*i.e.*, split peaks) (Fig. 2). In order to test this more rigorously, Rietveld refinements were carried out against SXRDX data for all samples using both the non-centrosymmetric tetragonal $P4mm$ (#99) and cubic $Pm\bar{3}m$ (#221) space groups. A comparison of the fits for $x = 0.8$ is shown in Fig. 3. Refining equivalent parameters, the tetragonal model provides a significantly improved fit. Refined crystal parameters also were more plausible using a tetragonal model, as shown in Table 1, where the ADPs for the B-site cations and oxygen atoms are unreasonably large using the cubic model. Using the tetragonal model, large ADPs are only found for the K/Bi mixed occupancy A-site, as expected due to static disorder. The tetragonal model was superior for all samples except pure KBZ, where the best fit was obtained using the cubic model. The evolution of unit cell parameters and cell volume across the KBT-KBZ solid-solution is shown in Fig. 4. The unit cell volume is consistent with Vegard's law. The difference between the a and c parameters drops rapidly as x (Zr content) increases, but remains statistically significant until $x = 1$.

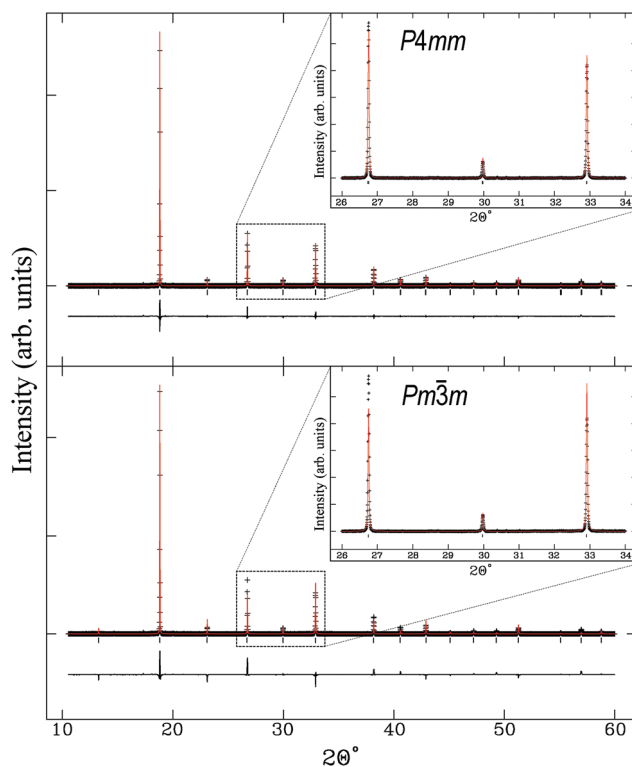


Fig. 3 $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{0.2}\text{Zr}_{0.8}\text{O}_3$ fit to tetragonal $P4mm$ (#99) $R_p = 0.0347$ and $\chi^2 = 9.294$ above and fit to cubic $Pm\bar{3}m$ (#221) $R_p = 0.0520$ and $\chi^2 = 29.98$ below.

Table 1 Atomic displacement parameters for $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{0.2}\text{Zr}_{0.8}\text{O}_3$

$100 \times U_{\text{iso}}$	$P4mm$ (#99)	$Pm\bar{3}m$ (#221)
A-site ($\text{\AA}^2$)	14.47(5)	3.48(4)
B-site ($\text{\AA}^2$)	0.89(2)	7.74(8)
Oxygen ($\text{\AA}^2$)	2.91(12)	21.7(4)

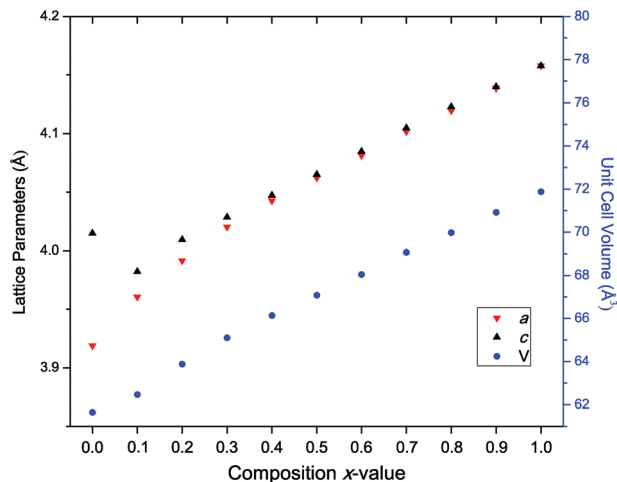


Fig. 4 The lattice parameters extracted from Rietveld refinement across the solid-solution with a (red ∇ triangles), c (black Δ triangles) and unit cell volume V (blue circles). For $x = 1$ only, the data were modelled in cubic $Pm\bar{3}m$ hence a and c are equal. Error bars are smaller than symbols.

The observation that the KBT-KBZ solid-solution retains tetragonal symmetry – *i.e.*, is pseudocubic rather than truly cubic – up to at least $x = 0.9$ is consistent with its relaxor ferroelectric behaviour for $x = 0$ – 0.9 (albeit very weak at higher x).¹⁶

Interestingly, the sodium analogue $\text{Na}_{0.5}\text{Bi}_{0.5}\text{ZrO}_3$ is reported to be rhombohedral/orthorhombic instead of cubic.¹⁴ This structural difference between K- and Na-versions can be explained by the tolerance factor:

$$t = \frac{\sqrt{2}r_{\text{A-O}}}{2r_{\text{B-O}}} \quad (1)$$

which predicts a cubic perovskite structure when $t = 1$, and distortions away from that value. Pure KBZ has a tolerance factor much closer to 1.0 (0.97) than NBZ (0.93), using bond lengths $r_{\text{A-O}}$ and $r_{\text{B-O}}$ calculated from Shannon's ionic radii.¹² For pure KBT, the tolerance factor is higher than the ideal value ($t = 1.03$); Zr doping across the KBT-KBZ solid-solution should reduce this and lead to an increase in symmetry. Our SXRD results bear this out, in agreement with previous work.^{15,16} This behaviour is different to the NBT-NBZ solid-solution, where the $t < 1$ for the whole range and lower symmetry structures (rhombohedral, monoclinic and orthorhombic) are always observed.¹⁴

3.2. *In situ* synchrotron XRD

The temperature-dependent behaviour through the ferroelectric-paraelectric phase transition was studied by *in situ* variable-temperature SXRD. Data were collected for pure KBT in order to track the evolution of lattice parameters. This builds on previous work using conventional XRD.^{7,27} The evolution of the diffraction patterns (Fig. 5) clearly shows the transition from tetragonal to cubic symmetry at $T_C \sim 683$ K where the lower symmetry split peaks converge to higher symmetry single peaks. Slightly below this temperature, at approximately

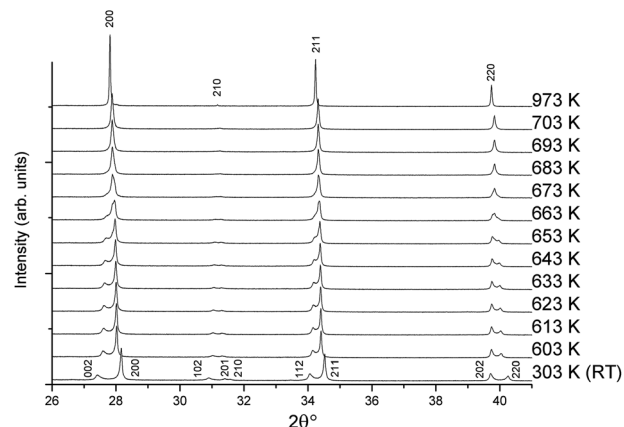


Fig. 5 Variable-temperature SXRD patterns from 303 to 973 K showing the tetragonal to cubic transition with indexed peaks.

663 K, there is a region of phase coexistence. This was not observed in earlier studies, but is a known behaviour of relaxor ferroelectric compounds, in which the presence of polar nano-regions creates a distribution of ferroelectric domains and hence a range of transition temperatures;^{27–29} and is in agreement with the broad peak in dielectric measurements reported by Bengagi *et al.*¹⁶ Around T_C , from 633 to 693 K, a two-phase model in Rietveld refinement was used to track the behaviour of the coexisting phases. This region is highlighted by the shaded area in Fig. 6.

Fig. 6 shows the Rietveld-refined lattice parameters and cell volume as the sample is heated from room temperature. The a and c parameters of the tetragonal phase converge to the common a parameter of the cubic phase in an essentially continuous second order manner, but with the unusual addition of a narrow coexistence region close to T_C , the origin of which is discussed above. Another unusual feature is the lattice volume contraction from the tetragonal to the cubic phase. We can rationalise this by considering the relationship between bond length and bond strength, which in solid-state chemistry is commonly considered in terms of effective bond valence sum (BVS) (eqn (2)):

$$\text{BVS} = \sum \left(\exp \left(\frac{R_0 - R_i}{b} \right) \right) \quad (2)$$

where R_0 is an empirically determined constant for a particular type of bond, R_i the actual bond length, and b a constant (normally 0.37 Å).³⁰ In the tetragonal phase, the A-site and B-site cations are offset from the centres of their 12-fold or 6-fold coordination environments respectively, bringing them closer to some O anions and further from others. Due to the exponential nature of eqn (2), this increases the total BVS compared to when those cations are at the centres of their coordination environments in the cubic phase. Thus, when the structure becomes cubic above T_C , the size of those coordination environments (hence the unit cell volume) must decrease in order to maintain the same BVS. The volume drop in the tetragonal phase within the shaded region is an



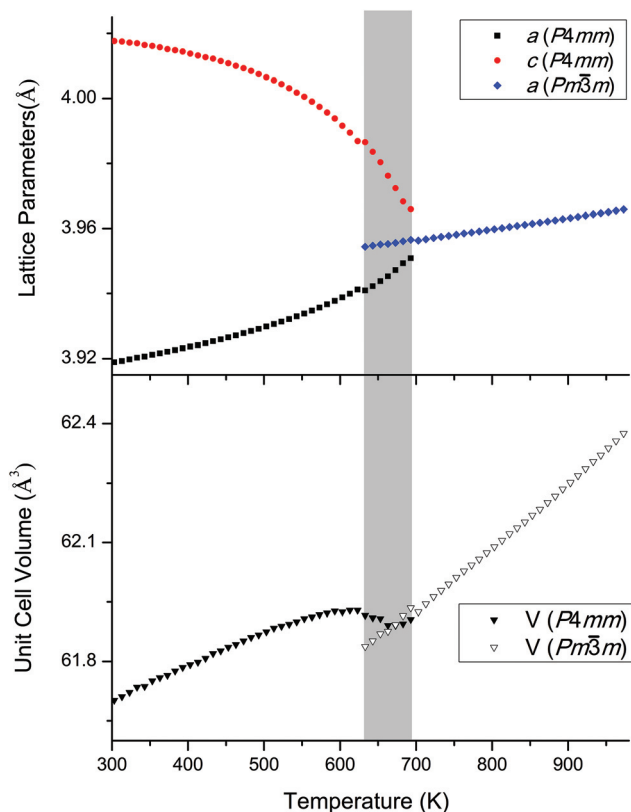


Fig. 6 Evolution of lattice parameters and cell volume for KBT obtained from Rietveld refinements against SXRD data. The grey region indicates the temperature range in which the tetragonal and cubic phases coexist.

unrelated artifact of the two-phase refinement, due to the imperfect deconvolution of peak widths, peak separation and phase fractions.

3.3. XANES

Relaxor ferroelectrics have complex behaviour due to their locally mixed compositions, sizes and charges across the crystal lattice.²⁹ While diffraction provides a good measure for the average crystal structure, knowledge of the local structure is crucial to understanding the behaviour of solid-solution ferroelectrics. We have used XANES as a probe of the local coordination environment. Changes in local cation environments cause changes in orbital hybridisation, which manifest as peak shifts and intensity changes at the metal absorption edges. The Ti K-edge is particularly sensitive to the oxidation state and coordination environment of Ti in systems such as these.¹⁴

Normalised Ti K-edge spectra across the KBT-KBZ solid-solution are shown in Fig. 7 (note that for $x = 0.9$, statistics were too poor to obtain usable data). There are two major features: the main edge feature (labelled A); and a pre-edge feature (labelled B1–B4). Feature A corresponds to the dipole-allowed transition of a 1s electron into unoccupied 4p states.¹⁴ The line shape of A implies minor changes to the hybridis-

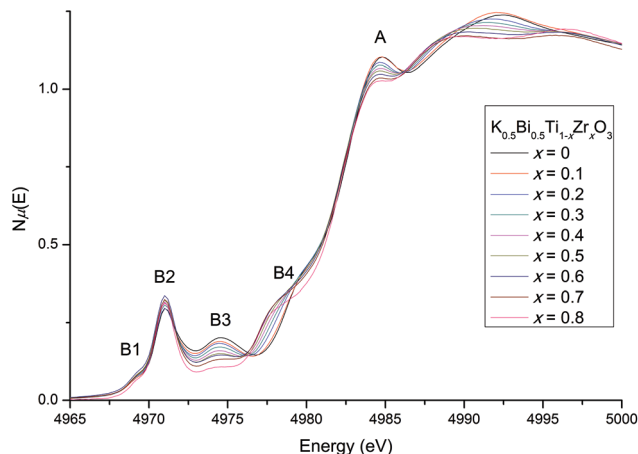


Fig. 7 The Ti K-edge (A) including the distinct pre-edge features (B1–B4). No energy shifts are observed at the main edge, but pre-edge changes indicate an increasingly centrosymmetric environment for the B-site with increasing Zr content, while the Ti^{4+} cation distortion remains largely unchanged.

ation of Ti 4p states with the bonding states of nearest- and next-nearest neighbouring atoms, largely due to the introduction of Zr content. Although small intensity changes are observed in the main peak A, likely due to replacing Ti 4p states with Zr 5p states, the lack of a shift in the main edge energy confirms that Ti remains tetravalent – *i.e.*, there is no evidence for oxygen vacancies.¹⁴

More subtle information about the coordination environment can be obtained by analysing the forbidden transitions in the pre-edge region. The pre-edge features arise from a dipole-forbidden transition of a 1s electron into unoccupied 3d states.³¹ The first pre-edge feature, B1, corresponds to the dipole-forbidden transition of a 1s electron into the 3d t_{2g} states for octahedrally coordinated Ti and remains largely the same with increasing Zr content. The higher-energy feature B2 arises from $1s \rightarrow 3d e_g$ transitions as well as non-local inter-site hybridisation, and B3 originates from the non-local $1s \rightarrow p$ transitions (where the empty p states of the absorbing Ti atom is hybridised with the empty d states of the Ti second neighbours *via* O 2p states).^{31,32} Similarly to previously reported Zr doping effects on NBT, a small increase is observed in B2 suggesting that the Ti-offset increases slightly with increasing Zr content.^{13,14} Together, B3 and B4 are sensitive to the local ratio of Ti/Zr around the absorbing atom, and therefore to local clustering.¹³ In the present case, B3 gradually decreases as Ti content decreases; while B4 increases as Zr content increases. These systematic trends suggest that Ti/Zr atoms are randomly distributed in the lattice,^{13,14} with no significant clustering. The local crystal structure from $x = 0.5$ to 0.9 (previously reported as cubic¹⁶) does not appear to ever become completely centrosymmetric, with Ti remaining off-centered. This is consistent with our high-resolution SXRD diffraction results, and the observation of weak ferroelectricity up to $x = 0.9$ by Bengali *et al.*¹⁶



The Ti L-edge and Zr L-edge transitions correspond to the dipole-allowed transition of a Ti or Zr 2p electron into the unoccupied d states and hence probes the B-site distortion.^{23,33} Changes in line shape with composition for the L-edges of Ti and Zr are shown in Fig. 8 and 9 respectively. In

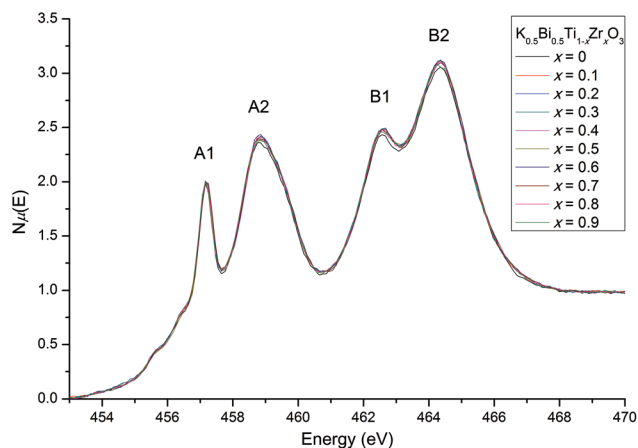


Fig. 8 Ti L₃- and L₂-edges. Asymmetry is observed in peak A2, with no line shape changes across the solid-solution series.

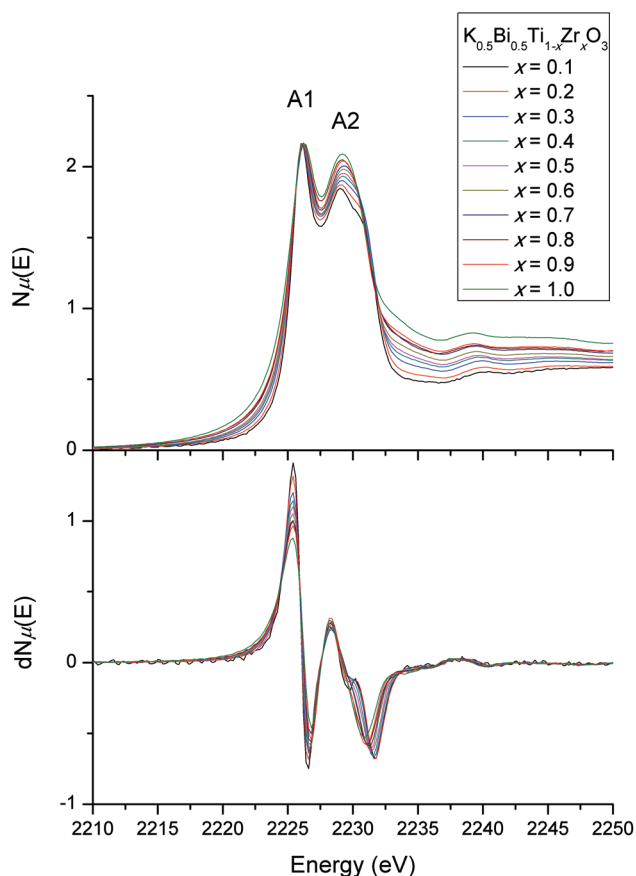


Fig. 9 Normalised Zr L₃-edge (upper) showing distinct line shape changes in A2 across the solid-solution series, indicative of increased symmetry with Zr doping. Line shape changes are emphasised by the plot of the first derivative (lower).

Fig. 8, the four strong features correspond to the Ti L₃-edge (two lower-energy transitions labelled A1 and A2) and the L₂-edge (two higher-energy transitions B1 and B2). The separation between the L₃- and L₂-edges comes from spin-orbit splitting of Ti 2p_{3/2} (L₃) and 2p_{1/2} (L₂). In an octahedral environment, the main edge splits into the 3d subbands, corresponding to excitations to the 3d t_{2g} (A1 and B1), and e_g (A2 and B2) orbitals.³⁴ The L₃-edge is usually more resolved due to the lifetime-related broadening of the L₂-edge. Consequently, after normalising the data for the Ti L₃-edge to A1, one can observe any line shape changes across the solid-solution series measured. Our interest lies in the A2 feature, the e_g states, which consist of d_{z²} and d_{x²-y²} orbitals. These orbitals are directed toward the ligand and are therefore sensitive to deviations in octahedral symmetry by Ti off-centering. In our normalised Ti L-edge data, peak asymmetry in A2 (less apparent in B2) is observed. This indicates a splitting of the e_g states due to a deviation from octahedral symmetry (*i.e.*, off-centered Ti⁴⁺). However, no obvious line shape changes are observed across the solid-solution series. This is consistent with the observed feature B2 in our normalised Ti K-edge data, where there were minimal changes in line shape with changes in composition, suggesting that Zr doping does not affect off-centering of the Ti⁴⁺ cations.

The Zr L₃-edge (Fig. 9) corresponds to the transition of a 2p_{3/2} electron into 4d states, which are split into t_{2g} (A1) and e_g (A2) states. Unlike the Ti L-edge, the Zr L₃-edge line shape changes with Zr doping, particularly with respect to the e_g state. In general, the asymmetry of the e_g peak (A2), which is due to the splitting of the 4d e_g states into 4d_{x²-y²} and 4d_{z²} states, decreases with increasing Zr content. This indicates a decrease in Zr⁴⁺ cation off-centering with doping.^{35–37} The decrease in peak splitting is more obvious in the first derivative plot of the Zr L₃-edge shown in Fig. 9. Such changes in the Zr environment have been observed for BaTi_{1-x}Zr_xO₃ (BTO-BZO).¹⁴ The similar behaviour can be explained by similarities in the effective ionic radii (IR) of Ba²⁺ and K⁺/Bi³⁺ cations. In twelve-coordinate environments, IR(Ba²⁺) = 1.61 Å, mean IR(K⁺, Bi³⁺) = 1.51 Å and mean IR(Na⁺, Bi³⁺) = 1.385 Å. This gives ranges for *t*(BTO-BZO) = 1.06–1.00, *t*(KBT-KBZ) = 1.03–0.97 and *t*(NBT-NBZ) = 0.98–0.93. However, in practice, IR(K⁺, Bi³⁺) is larger due to the asymmetry of the Bi³⁺ cation, which brings *t*(KBT-KBZ) closer to *t*(BTO-BZO), but still different to *t*(NBT-NBZ).

In contrast, Zr cations did not approach higher symmetry environments in the NBT-NBZ series.¹⁴ This is probably due to the fact that the end-members were already stabilised in much lower-symmetry environments (rhombohedral, monoclinic and orthorhombic) as discussed above on the basis of calculated tolerance factors. For the KBT-KBZ series under investigation here, the combined results from the normalised metal L₃-edges suggest that B-site off-centering is reduced with Zr doping. However, it appears that this is due to the Zr⁴⁺ cations moving to more symmetric coordination environments, rather than a reduction in Ti off-centering (as shown by the lack of line shape changes in our observed Ti L₃-edge). This can be



rationalised by a cation size effect, where the larger Zr^{4+} cation environment “pushes” neighbouring Ti^{4+} environments at low Zr content compositions. At high Zr content, larger cations are more likely to remain centered in the octahedra rather than offset. The dominating presence of larger Zr^{4+} cations over fewer and smaller Ti^{4+} cations brings the average B-site closer to centrosymmetric, as shown by diffraction data. However, on a local scale, Ti^{4+} cations remain offset in the octahedra. This effect was also observed for BTO-BZO.^{13,14}

In summary, analysis of XANES spectra shows that Zr doping in this series of compounds effectively dilutes the number of non-centrosymmetric environments rather than affecting the Ti-offset directly. This in turn reduces the effective number of polar nano-regions and hence weakens the ferroelectric properties in the material.

4. Conclusions

Synchrotron X-ray techniques have provided new insights into the evolution of long- and short-range structure across the solid-solution series $\text{K}_{0.5}\text{Bi}_{0.5}\text{Ti}_{1-x}\text{Zr}_x\text{O}_3$, where $x = 0$ to 1 with 0.1 steps. Rietveld refinements against high-resolution SXRD data showed that the pseudocubic tetragonal region exists across the whole solid-solution, rather than $0 < x < 0.4$ as previously observed. Only the pure KBT end member was clearly tetragonal, and only the pure KBZ end member was clearly cubic. On heating through the tetragonal-cubic transition at T_c , a narrow region of tetragonal and cubic phase coexistence was observed despite the overall continuous second-order nature of the transition. This was explained by the presence of polar nano-regions in these relaxor ferroelectric perovskites. A marked volume contraction from the tetragonal to the cubic phase was explained in terms of the bonding (BVS) requirements of the cations as they move from an off-centered to a centered location. A future transmission electron microscopy study, such as has been carried out on the KBT end-member,²⁷ might be able to reveal direct evidence of these polar nano-regions.

Analysis of metal L-edge XANES data across the solid-solution series revealed that Ti^{4+} cations remain off-centered with increasing Zr content, while Zr^{4+} cations approach a higher symmetry coordination environment. Ti K-edge data showed a similar lack of change in the Ti off-centering. We can therefore conclude that Zr doping has a minimal effect on Ti off-centering, but rather, increases the centrosymmetry of the Zr local coordination environment with its size leading to a reduction in polar domains and hence diminished ferroelectric response.

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