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Permalink
https://escholarship.org/uc/item/9k19q4bv

Journal
Physical Review B - Condensed Matter and Materials Physics, 92(21)

ISSN
1098-0121

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Publication Date
2015-12-28

DOI
10.1103/PhysRevB.92.214302

Peer reviewed
Phonon coupling to dynamic short-range polar order in a relaxor ferroelectric near the morphotropic phase boundary

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Received 18 September 2015; published 28 December 2015

We report neutron inelastic scattering experiments on single-crystal PbMg1/3Nb2/3O3 doped with 32% PbTiO3, a relaxor ferroelectric that lies close to the morphotropic phase boundary. When cooled under an electric field \( E \parallel [001] \) into tetragonal and monoclinic phases, the scattering cross section from transverse acoustic (TA) phonons polarized parallel to \( E \) weakens and shifts to higher energy relative to that under zero-field-cooled conditions. Likewise, the scattering cross section from transverse optic (TO) phonons polarized parallel to \( E \) weakens for energy transfers \( 4 \leq \hbar \omega \leq 9 \) meV. However, TA and TO phonons polarized perpendicular to \( E \) show no change. This anisotropic field response is similar to that of the diffuse scattering cross section, which, as previously reported, is suppressed when polarized parallel to \( E \) but not when polarized perpendicular to \( E \). Our findings suggest that the lattice dynamics and dynamic short-range polar correlations that give rise to the diffuse scattering are coupled.

DOI: 10.1103/PhysRevB.92.214302 PACS number(s): 63.20.K−, 77.80.Jk, 78.70.Nx

I. INTRODUCTION

Relaxor ferroelectrics have great potential for applications due to their piezoelectric and dielectric properties [1–3], but there is much that is not understood about how their properties arise on a microscopic level [4]. The composition of many relaxors is related to simple perovskites with the \( ABO_3 \) formula but with one cation site randomly filled with two or more cations of different valences, resulting in strong, disordered electric fields. For the relaxor PbMg\(_{1/3}\)Nb\(_{2/3}\)O\(_3\) (PMN), the \( B^{4+} \) site is occupied by Mg\(^{2+}\) and Nb\(^{5+}\). The variation in valence can be reduced by doping with an ion of intermediate valence, e.g., Ti\(^{4+}\) in the case of PMN doped with \( x \% \) PbTiO\(_3\) (PMN-x\%PT). The resulting PMN-x\%PT phase diagram shows four basic regions [5–7]: a cubic paraelectric phase at high temperature for all \( x \); a region with relaxor behavior for low \( x \) with either cubic or rhombohedral symmetry; a tetragonal, conventional ferroelectric region for high \( x \); and a morphotropic phase boundary (MPB) region between the relaxor and tetragonal regions. The piezoelectric coefficients \( d_{33} \) are very large in the MPB region and abruptly drop for higher \( x \) [1,8–10]; understanding this behavior and exploiting the large piezoelasticity provide much of the motivation for exploring relaxor ferroelectrics. These PMN-x\%PT solid solutions with small \( x \) exhibit clear relaxor behavior characterized by large dielectric constants which have a broad maximum with respect to temperature and are highly frequency dispersive within this range. These relaxor behaviors are widely believed to be associated with polar nanoregions (PNR) or other short-range polar order, as shown by numerous x-ray and neutron diffuse scattering studies [11–25].

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We have previously characterized two distinct components of the diffuse scattering in PMN-x\%PT, which we label T1 and T2 [24], as shown in Fig. 1(a). These labels are intended to refer to the related phonons where a T1 mode is a transversely polarized phonon propagating along \( ⟨100⟩ \) and a T2 mode is a transversely polarized phonon propagating along \( ⟨110⟩ \). For example, near \( ⟨100⟩ \), a TA1 phonon mode would refer to the transverse acoustic phonon mode propagating along the \( ⟨010⟩ \) or \( ⟨001⟩ \) direction with polarization along \( ⟨100⟩ \), while near \( ⟨110⟩ \), a TA2 phonon mode would refer to the transverse acoustic phonon mode propagating along \( ⟨110⟩ \) with \( ⟨110⟩ \) polarization.

The two diffuse scattering components can be distinguished by their anisotropic responses when field cooled (FC), i.e., after applying electric field above the ferroelectric transition temperature \( T_c \) and then cooling below \( T_c \). With a field applied along \( ⟨111⟩ \), a redistribution of T2-diffuse scattering intensity between two differently oriented components polarized along \( ⟨110⟩ \) and \( ⟨110⟩ \) has been observed in the structurally similar perovskite PbZn\(_{1/3}\)Nb\(_{2/3}\)O\(_3\) doped with \( x \% \) PbTiO\(_3\) (PZN-x\%PT) [25–27] and in PMN [28]. This situation has been interpreted in terms of a domain effect, in which applying a \( ⟨111⟩ \) field creates a single [111]-polarized ferroelectric domain (as opposed to the eight possible [111]-polarized domains present in the zero-field-cooled state) that favors certain orientations of polar nanoregions (PNR), resulting in the redistribution of diffuse scattering intensities along certain \( ⟨110⟩ \) directions [27]. An electric field along \( ⟨001⟩ \), on the other hand, does not seem to significantly affect the T2-diffuse scattering in the \( H0L \) plane [29]. A \( ⟨001⟩ \) field does, however, affect the T1-diffuse scattering, but the T1-diffuse scattering does not show a redistribution of scattering intensity under an external field. Instead, a suppression of [001]-polarized T1-diffuse scattering occurs under [001]-field cooling, while...
the [100]-polarized T1-diffuse scattering remains unaffected, as has been shown in PZN-x%PT [24,30]. This behavior is not due to a domain effect; otherwise, we would expect the intensity and other characteristics of the T1-diffuse scattering near [001] under zero-field cooling (ZFC) to be in between those near [001] and [100] under FC conditions. Thus, the changes induced by a [001] field should be due to changes within domains rather than changes in the sizes of domains, in contrast to applying a [111] field.

Ultimately, we would like to know how the local structures that give rise to the diffuse scattering affect bulk properties. When these local structures are intrinsically altered by, say, application of electric field, changes in the bulk structure would reveal connections between bulk and local structures. Phonons provide important measures of the dynamic properties of the bulk lattice, and their couplings to the diffuse scattering are therefore one of the key issues to pursue in understanding relaxor properties. For example, transverse acoustic phonons propagating along (110) (T2 phonons) are expected to couple with T2-diffuse scattering modes [31]. Evidence for this diffuse-TA2 phonon coupling has been shown with the help of an external [111] electric field, which breaks the pseudocubic symmetry and reveals a clear difference between TA2 phonons measured near (220) and (220) [32]. It would be interesting to see if there is a similar coupling between phonons and diffuse scattering along the T1 directions under a [001] field, given that the effect of a [001] field is not a domain effect like that of a [111] field, but coupling between the T1-diffuse modes and the TA1 or TO1 phonon modes has not yet been thoroughly studied.

In this paper we report neutron scattering experiments on PMN-32%PT with a field applied along [001]. When cooled below \(T_c \approx 430 \text{ K} \) [29], in addition to the expected suppression of T1-diffuse scattering measured near (001), we also see a clear change in the intensities of the TA1 phonons near (001), whereas the TA2 phonons near (100) are unaffected. These changes illustrate that there may be a TA1-phonon/T1-diffuse mode coupling, evocative of the TA2-phonon/T2-diffuse mode coupling previously seen in PZN-4.5%PT [32]. This coupling appears to be limited to large-wavelength phonons. In addition, we observed a suppression of spectral weight for the transverse optic phonons in the T1 direction (TO1 phonons) within 4 to 9 meV near (002), but no change was seen near (200). The changes in the TA1 and TO1 phonons were present at 400 K but much less pronounced at 200 K.

II. EXPERIMENTAL DETAILS

We purchased a PMN-32%PT single crystal from TRS Ceramics with dimensions \(10 \times 10 \times 2 \text{ mm}^3 \) and large [001] faces. The (001) surfaces were coated with gold to ensure a uniform equipotential surface during the application of a field \(E_{\parallel}[001] \). Another PMN-32%PT crystal from the same source was measured to have a cubic-tetragonal transition at \(T_c \approx 430 \text{ K} \) and a tetragonal-monoclinic transition near 355 K, both measured on cooling [29]. Neutron inelastic scattering experiments were performed on the HYSPEC (Hybrid Spectrometer) time-of-flight spectrometer at the Spallation Neutron Source at Oak Ridge National Laboratory [33]. The software package \textsc{mantid} was used in the processing of the data [34]. The incident energy \(E_i \) was set to 20 meV. The crystal symmetry was pseudocubic with lattice parameter \(a = 4.00 \text{ Å} \). All neutron scattering momentum transfers \(Q \) are reported in terms of reciprocal lattice units (r.l.u.), and energy transfers \(\hbar\omega\) are reported in meV. Measurements were performed in the \(HOL\) scattering plane. Fields of 0.5–8.0 kV/cm were used. The T1-diffuse scattering at (001) changed significantly with a field of 0.5 kV/cm and had almost no additional change with higher field, indicating that 0.5 kV/cm was sufficient to alter the T1-polarized short-range order. For pseudocolor plots, the data were smoothed. Error bars represent statistical error and correspond to one standard deviation from the observed value.

III. DATA AND ANALYSIS

In Fig. 1(b), the temperature dependence of the T1-diffuse scattering near (001) is shown with neutron scattering intensity measured at wave vector \(Q = (-0.125,0,1)\) for FC and ZFC conditions. The intensity was integrated within \(0.95 \leq L \leq 1.05 \text{ r.l.u.}, -0.1375 \leq H \leq -0.1125 \text{ r.l.u.}, \) and \(-0.5 \leq \hbar\omega \leq 0.5 \text{ meV} \). For temperatures up through 400 K a clear suppression of intensity is seen with applied field, but this difference disappears above the ferroelectric transition between 400 and 450 K. In Figs. 1(c) and 1(d) we show that the suppression of T1-diffuse scattering under FC conditions is direction dependent, being absent for [100]-polarized diffuse scattering measured near (100) [Fig. 1(c)] but present for [001]-polarized diffuse scattering near (001) [Fig. 1(d)]. These data were taken at 200 K as transverse scans across the Bragg peaks, with integration ranges of \(\pm 0.5 \text{ meV} \) for \(\hbar\omega\) and \(\pm 0.05 \text{ r.l.u.} \) for the \(H \) and \(L \) directions for Figs. 1(c) and 1(d), respectively. The direction dependence of the suppression of T1-diffuse scattering under [001]-field cooling is consistent with previous reports on the related PZN-x%PT system [24,30].
FIG. 2. (Color online) Pseudocolor plots of transverse acoustic phonons near (100) (left) and (001) (right) for ZFC (top) and FC (bottom) conditions. Neutron scattering intensity (indicated by color, in arbitrary units) is plotted against energy and momentum transfer. These data were taken at 400 K. White areas represent lack of data.

Figure 2 shows the dispersions of the TA1 phonons near (100) and (001) under FC and ZFC conditions, in which a change in intensity under field can be seen near (001) but not (100). These are pseudocolor plots of intensity vs energy and momentum transfer in slices across (100) and (001) at 400 K. The intensities were integrated within ±0.05 r.l.u. along H for (10L) and L for (H01). The TA1 phonons disperse out from the Bragg peaks and have maxima around roughly 5–6 meV. The width of the phonons with respect to energy is evident. The effect of field can be seen more clearly in Fig. 3, which shows the phonon intensities near (100) and (001) in a similar pseudocolor plot but with $\hbar \omega$ fixed at 2 meV and integrated within ±0.5 meV. There is a clear decrease in intensity with field near (001), but no clear change near (100).

For a clearer view of how the phonon dispersion is affected by field, in Fig. 4 we show constant-Q scans at $(-q,0,1)$ and $(1,0,q)$ for various $q$ values, taken at 400 K. The data were integrated over ±0.05 r.l.u. in both the H and L directions. Each data set was fitted to the sum of a Gaussian function for elastic scattering and Voigt functions for the acoustic phonons at ±$\hbar \omega$. We can see that there is little change near (100) for all $q$, but near (001) a clear change is seen, with both a suppression of intensity and an increase in energy transfer for $q = 0.1$ (and possibly also $q = 0.2$). These data suggest that the electric field effect is strongest for low $q$. We note that we have not seen a clear field effect on longitudinal acoustic (LA) phonons measured along [100] near (100) and along [001] near (001) or on TA2 phonons measured along [101] near (101), suggesting that the [001] field primarily affects T1 phonons. This situation is similar to how T1-diffuse scattering intensities respond to [001] fields, with T1-diffuse scattering suppressed near Bragg peaks with wave vector $G \parallel [001]$ but unaffected for
FIG. 5. (Color online) Pseudocolor plots illustrating the transverse optic modes near (200) (right) and (002) (left) for ZFC (top) and FC (bottom) conditions, with neutron scattering intensity plotted as color (in arbitrary units) as a function of energy and momentum transfer. White areas represent lack of data.

FIG. 6. (Color online) $H_{0L}$ slices at 400 K and $5.5 \leq \hbar \omega \leq 6.5$ meV, focusing on the TO phonons near (200) and (002). These are pseudocolor plots, with intensity plotted as color (in arbitrary units) and momentum transfer along (2,0,L) and (H,0,2) delimited on the axes. White areas represent lack of data.

FIG. 7. (Color online) Constant-energy cuts across the transverse optic modes along (20L) (right) and (H02) (left) for \( h\omega = 4, 6, 8, \) and 9 meV, integrated over \( 1.9 \leq H \leq 2.1 \) for (20L) and \( 1.9 \leq L \leq 2.1 \) for (H02), and within an energy range of \( \pm 0.5 \) meV. For each subplot, neutron scattering intensity is plotted against momentum transfer. All data were taken at 400 K.
FC conditions. Intensity was integrated within ±0.5 meV for energy transfer and ±0.1 r.l.u. for momentum transfer in the $H$ or $L$ direction transverse to the direction of the scan. We see that there is a consistent suppression of spectral weight near (002) but not near (200). This suppression can be seen from 4 to 9 meV; we note that the difference disappears outside of this range. As for TO1 phonon energy, it is difficult to observe changes in the TO1 dispersions due to their steepness and to the phonons becoming highly damped at small $q$. For comparison, we note that in conventional ferroelectrics there have been examples of optic modes being affected by field [43,44], and the effect is only predicted to be large for soft modes close to zero energy in the vicinity of a structural phase transition [45].

To illustrate the effect of temperature, in Fig. 8 we show representative data of the TA1 and TO1 modes at 200 K to contrast with the 400 K data in Figs. 4 and 7, respectively. In Fig. 8(a), we show a constant-$Q$ scan showing the TA1 mode at (−0.1,0,1), with intensity vs $h\omega$ plotted and intensity integrated within ±0.05 r.l.u. in the $H$ and $L$ directions. In Fig. 8(b), we show a constant-$h\omega$ scan showing the TO1 mode, integrated within 7.5 $h\omega$ ≤ 8.5 meV and 1.9 $L$ ≤ 2.1 r.l.u. In both plots, we see a similar suppression of intensity near (001) and (002). We also saw a similar lack of change near (200) (not plotted), but near (100) a spurious feature prevented us from determining if there was a change in TA1 phonon spectral weight. From these data, we can see that the field effect at 200 K seems to be less pronounced than that at 400 K, at least for the acoustic phonon mode.

IV. DISCUSSION

The electric field effects observed in our measurements can be summarized as follows: (i) there is no [001] field effect on the longitudinal acoustic (LA) modes or on the transverse acoustic modes propagating along the ⟨110⟩ (TA2) directions; (ii) for TA1 phonons polarized along ⟨001⟩, we observed, after field cooling, a reduction of intensity and increase of phonon energy near Bragg peak wave vectors $G$ || [001], but no field effect was observed for TA1 modes near $G$ || [100]. For diffuse scattering, a similar pattern in response to field cooling along [001] has been seen, with intensity suppression for $G$ || [001] but not for $G$ || [100] [24,30]. A much smaller effect on low-energy TO modes is also observed following the same rule, i.e., a reduction of intensity near the bottom of the TO mode measured for $G$ || [001] but no effect for $G$ || [100]. These results imply that coupling exists between the diffuse scattering along ⟨001⟩ (the T1 diffuse) and the TA1 and/or TO1 phonon modes along the same directions.

Previous work has shown strong coupling between the diffuse scattering along the ⟨110⟩ directions (the T2 diffuse) and the TA2 phonon modes in these lead-based relaxor materials [32]. The diffuse-phonon coupling along the ⟨001⟩ (T1) and ⟨110⟩ (T2) directions shares some common features. For example, when the diffuse scattering is suppressed by the external field, we always see a hardening of the corresponding TA phonon mode. Evidently, interaction between the PNR and the phonons tends to drive the phonons softer for both the T1 and T2 modes. This tendency suggests that short-range polar orders are likely related to lattice instabilities in these relaxor compounds.

On the other hand, the T1 and T2 modes differ in many aspects too:

(i) For the case of the T2 modes, the field effect on the PNR (and thus on the T2-diffuse scattering) is indirect. An external field along [111] helps establish a single-domain ferroelectric phase with [111] polarization. The change in the population of each domain induces a redistribution of PNR with different polarizations, resulting in the redistribution of T2-diffuse scattering intensities in reciprocal space. For example, under a [111] field there is an enhancement of the T2-diffuse scattering near the (220) Bragg peak but a reduction of the T2-diffuse scattering near the (220) Bragg peak. The TA2 phonon near (220) softens, while the phonon near (220) hardens. On the other hand, in zero field, all [111] domains are present to an equal degree. Measurements in zero field of the TA2 phonons near (220) or (220) average over both hardened and softened phonons, and the role of the [111] field is merely to obtain results for a single domain.

However, the [001] field effects on the T1-diffuse scattering and T1 phonons are not domain related. This can be demonstrated by comparing the ZFC and FC results. Under FC conditions, one sees changes (relative to the ZFC results) only for $G$ || [001] but no change for $G$ || [100]. If these changes were due to domain effects, one would have expected the ZFC results to lie in between (in intensity, energy, etc.) the FC results measured for $G$ || [100] and [001]. This is not the case. Therefore, the effect of the [001] field is more intrinsic, with the field directly affecting the short-range order and, consequently, the related phonon modes.

(ii) While there is no evidence of diffuse-TO coupling for the T2 modes [32], in our sample there appears to be a weak diffuse-TO coupling for $G$||[001] (TO1 modes).

(iii) The diffuse-phonon coupling for the T2 modes is strong throughout the entire Brillouin zone, but the coupling for the T1 modes is present only for a small range of $q$ values (~0.1 to 0.2 r.l.u.) away from $G$. This anisotropy of the diffuse-TA coupling revealed by our electric field measurements is also consistent with previous reports [46].

In order to understand these results, we consider the origin of the diffuse scattering and short-range orders in relaxors. The local random electric field generated by the $B$-site cations
believed to play important roles in Pb-based relaxor systems [47–52]. A direct link between the diffuse scattering in these relaxors and the random field has been demonstrated by comparing two isostructures with and without random B-site valences [53]. The random field in the system prevents long-range order from developing and induces short-range orders that also grow with cooling.

The diffuse scattering intensities are not entirely static and have a strong dynamic component [24,54,55], which is consistent with our results. Indeed, in our work, the coupling between the T1-diffuse scattering and the TA1 phonon appeared weaker than our results. Indeed, in our work, the coupling between the T1-diffuse scattering and the TA1 phonon appeared weaker at 200 K than at 400 K, where the dynamic component has also been shown to decrease with cooling [54,55] below TC. The existence of these dynamic/quasielastic components has been explained by theoretical work [56,57] and is essential for the coupling between the diffuse scattering and phonons.

The weak coupling of the T1-diffuse scattering to the TO1 mode can be understood from the fact that the short-range orders consist of a combination of acoustic (strain) and optic (polar) types of atomic shifts [24,26]. The polar component of the short-range order can couple to the TO mode and is also essential for the short-range order to respond to an external field. The coupling will likely diminish quickly when the TO phonon energy increases and moves farther away from the quasielastic component of the diffuse scattering, as is the case in our measurements. A more detailed understanding of the competing interactions on a microscopic level, such as the role of the Pb-O bond, whose covalent nature is said to be responsible for ferroelectricity in PbTiO3 [58], is desired but is beyond the scope of this paper.

The anisotropy of the diffuse-phonon coupling between the T1 and T2 directions is more intriguing. The T2-diffuse scattering is significantly stronger than the T1-diffuse scattering, extends to a larger q range in reciprocal space, and interacts with TA phonons almost throughout the entire branch. Overall, we could consider a picture where each PNR is actually a “core” of a region of short-range correlated (110) atomic shifts, contributing to the broader T2-diffuse scattering, whereas the polar/strain field surrounding the core extends to a much greater range and contributes to the narrower T1-diffuse scattering. (These core regions, being the PNRs, should not be confused with the virtually temperature-independent chemical disorder/short-range order that persists up to at least 900 K [59]; the PNRs are local polar structures that appear at the Burn’s temperature TD and grow with cooling.) The atomic shifts within the PNRs would be significantly larger than those in the surrounding region, leading to a much stronger T2-diffuse scattering than the weaker T1-diffuse scattering. The core of the short-range order, the PNRs, results from the local strong random field and cannot be directly suppressed by an external field [26,27,32]. However, the weaker polar/strain field around the core is less robust and can be partially modified by external field, showing the intrinsic field effect on the T1-diffuse scattering and its coupling to TA1 and TO1 phonon modes discussed in this paper.

An analog to this situation has recently been considered [60,61] where strong but dilute random fields are inserted into a system with a weak continuous random field. Theoretical work involving a magnetic system with a random field [61] suggested that a large correlation length or even a weak long-range order could be achieved. If we map the PNRs to the strong random field in the magnetic system, the large spin correlation length proposed by the theoretical work can be related to the weak polar/strain field surrounding the PNRs which gives the T1-diffuse scattering. Although not an exact analog, this picture does provide a crude description of the origin of the two types of diffuse scattering. For a better understanding of the source of these diffuse scattering components and their coupling to lattice dynamics, more detailed experimental work is required. Although numerous models have been proposed by various groups attempting to describe the diffuse scattering and short-range orders in these relaxor systems [15,22,62–68], our results simply suggest that there is a clear anisotropy in the diffuse scattering, their field dependence, and their coupling to the related phonon modes measured along the (100) and (101) directions.

We can compare elastic constants derived from the TA1 and TA2 phonon energies in our data with values reported for similar materials to get insight into the tendency for lattice instability in the T1 and T2 directions. Our values and those for related compounds in the literature are displayed in Table I. From our data, we obtained the elastic constant quantities C44 = 0.56(5) and (C11 − C12)/2 = 0.23(4) in units of 1011 N/m² based on the TA1 and TA2 phonons measured near (001) and (101) at T = 400 K. The value of C44 is slightly larger than in PMN but smaller than in PbTiO3 [46]. (C11 − C12)/2 = 0.23(4) shows a bigger change, being significantly reduced from its value in PMN, suggesting an increased lattice instability when the system approaches the morphotropic phase boundary. In fact, a similar value of (C11 − C12)/2 = 0.26(4) can be obtained from phonon data on PZN-4.5%PT [32].

The diffuse-phonon coupling discussed in this paper can also affect how one determines the elastic constants. We calculated the phonon velocities from our FC data rather than our ZFC data since we believe the FC velocities more closely resemble the velocities expected for q → 0. First, the diffuse-phonon coupling is expected to diminish as q → 0 [70]. Second, since the diffuse-phonon coupling tends to reduce the phonon energy (as discussed in this paper and Refs. [32,46]), at nonzero q the elastic constants should be calculated from data taken after this coupling effect is removed, such as after diffuse scattering is suppressed by field. For ZFC conditions, the value for C44 can be about 15% smaller than in FC conditions. This difference is, in fact, an artifact of calculating the phonon velocity using (reduced) phonon energies at nonzero q (we used data taken at q = 0.1 and q = 0.1 × √2 r.l.u. for the

<table>
<thead>
<tr>
<th>Material</th>
<th>C_{44}</th>
<th>(C_{11} − C_{12})/2</th>
</tr>
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<tbody>
<tr>
<td>PMN [46]</td>
<td>0.53(3)</td>
<td>0.48(6)</td>
</tr>
<tr>
<td>PMN-32%PT</td>
<td>0.56(5)</td>
<td>0.23(4)</td>
</tr>
<tr>
<td>PZN-4.5%PT [32]</td>
<td>0.72(2)</td>
<td>0.63(1)</td>
</tr>
<tr>
<td>PbTiO3 [69]</td>
<td>0.72(2)</td>
<td>0.63(1)</td>
</tr>
</tbody>
</table>
calculation in the T1 and T2 directions, respectively). If one could obtain the phonon velocity using smaller $q$ values near $q = 0$ where the diffuse-phonon coupling diminishes, the difference between ZFC and FC data should become negligible.

V. CONCLUSIONS

We have observed, when comparing field-cooling and zero-field-cooling conditions for different Brillouin zones with a field along [001], a change in the lattice dynamics of PMN-32\%PT that correlates with changes in diffuse scattering. Specifically, under field cooling we see a reduction of intensity and an increase of phonon energy for the TA$_1$ mode measured near (001) and propagating along [100] [(001) polarized] but no change for TA$_1$ phonons near (100) and propagating along [001] [(100) polarized]. This field effect is only clearly seen for wave vectors around 0.1 to 0.2 r.l.u. away from the Bragg peak. Meanwhile, the T1-diffuse scattering near (001) is suppressed under field cooling but is unaffected near (100). A similar effect is seen for the TO$_1$ mode, which is slightly suppressed near (002) from 4 to 9 meV but unaffected near (200). No clear field effect has been seen for the longitudinal modes near (001) or (100) or for the TA$_2$ mode near (101). The similarities in the effect of field on the T1-diffuse scattering near (001) and (100), the TA$_1$ phonons near (001) and (100), and the TO$_1$ phonons near (002) and (200) suggest the presence of diffuse-TO and diffuse-TO mode coupling which resembles the mode coupling observed in the T2 directions in related relaxor materials.

ACKNOWLEDGMENTS

This research at Oak Ridge National Laboratory’s Spallation Neutron Source was sponsored by the Scientific User Facilities Division, Office of Basic Energy Sciences, US Department of Energy. J.A.S. and G.Y.X. acknowledge support by the Office of Basic Energy Sciences, US Department of Energy, under Contract No. DE-SC00112704. Z.J.X. and R.J.B. are also supported by the Office of Basic Energy Sciences, US Department of Energy, through Contract No. DE-AC02-05CH11231. C.S. acknowledges the support of the Carnegie Trust for the Universities of Scotland and the Royal Society. The identification of any commercial product or trade name does not imply endorsement or recommendation by the National Institute of Standards and Technology.