Elastic precursor effects during Ba_{1-x}Sr_xTiO₃ ferroelastic phase transitions

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Elastic softening in the paraelastic phases of $Ba_{1-x}Sr_xTiO_3$ is largest near the transition temperatures and decreases on heating smoothly over extended temperature ranges. Softening extends to the highest measured temperature (850 K) for Ba-rich compounds. The temperature evolution of the excess compliance of the precursor softening follows a power law $\delta S \propto |T - T_C|^{-\kappa}$ with a characteristic exponent κ ranging between 1.5 in SrTiO₃ and 0.2 in BaTiO₃. The latter value is below the estimated lower bounds of displacive systems with three orthogonal soft phonon branches (0.5). An alternative Vogel-Fulcher analysis shows that the softening is described by extremely low Vogel-Fulcher energies E_a , which increase from SrTiO₃ to BaTiO₃ indicating a change from a displacive to a weakly order-disorder character of the elastic precursor. Mixed crystals of $Ba_xSr_{1-x}TiO_3$ to BaTiO₃. Using power-law fittings reveals that the elastic softening is still 33% of the unsoftened Young's modulus at temperatures as high as 750 K in BaTiO₃ with $\kappa \simeq 0.2$. This proves that the high-temperature elastic properties of these materials are drastically affected by precursor softening.

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I. INTRODUCTION

Ferroelastic materials [1] commonly display large elastic anomalies during structural phase transitions [2]. Structural collapses can lead to a total reduction of the effective moduli in the case of proper ferroelastics such as in BiVO₄ and LaNbO₃ [3-5]. In the case of improper ferroelastics, such as ferroelectric BaTiO₃ and antiferrodistortive SrTiO₃, a direct coupling between the acoustic [6] soft mode and the elastic moduli is symmetry forbidden while typical elastic softening still reduces the moduli by some 20-50% [7]. In ferroelectrics, the intrinsic softening of the low-temperature phase with respect to the paraelectric phase is due to the combined direct and converse piezoelectric effects [8,9]. Additional softening in the low-temperature phases may be due to mobile twin boundaries just below but close to the transition point, which vanishes if the twin walls are strongly pinned [10-12]. Thick domain walls were shown to be less prone to such pinning effects [13-15], and many examples of highly mobile wall movements during the softening process have been reported [16-21]. We argue in this paper that significant precursor softening is commonly observed in the paraelastic phase which can be-in some cases-directly related to intrinsic disorder of the high-temperature phase and dynamic local nanostructures. Conceptually, this effect is best observed when materials are disordered by extrinsic forces such as radioactive bombardment. Consider a single crystal without any domain boundaries which is then disordered by the radioactive decay of radiogenic impurities. Such samples will massively reduce their elastic moduli due to the structural heterogeneity. This situation is often encountered in so-called metamict materials such as zircon [22] and titanite [23] where the reduction in bulk and shear modulus is greater than 50%, while their structure is still unchanged. Structural disorder of the paraelastic phase and significant short-range order are thus expected if the ferroelastic phase transition is of the orderdisorder type and structural variations occur in nominally cubic materials [24-28].

Displacive systems show similar effects although to a lesser extent [29]. Following the initial theoretical analysis of Pytte [30,31] and Axe and Shirane [32], fluctuation contributions to elastic softening have usually been considered in terms of coupling between different vibrational modes [33–38]. In both cases, local fluctuations with high local correlations can be expected. Typical examples for local nanostructures are tweed structures with interwoven, dynamical strain fields. They were found by molecular dynamics

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simulations and diffractions experiments [39–42]. Similar effects were postulated by Pelc *et al.* [43] for high-temperature superconductors.

The underlying physical picture for displacive precursor softening is that, associated with a soft mode at some specific point in reciprocal space, there will be a set of branches which also soften to some extent. Along with the soft mode itself, when the frequencies of modes along the soft branches decrease, their amplitudes become larger, and they can combine to produce stress fluctuations and associated strain fluctuations. The summation of all such combinations will yield a net softening of some specific acoustic modes depending on the dimensionality of the elastic softening [2]. The total effect increases as the amplitudes of the modes increase, reaching a maximum at the transition point. The detailed temperature dependence is usually uncertain because it requires an exact knowledge of mode mixing properties [44], including potential local modes which are not captured by conventional spectroscopy. Nevertheless, it was advocated that the resulting temperature dependence of the elastic softening can be described conveniently by a power law [2,29,45]:

$$\Delta C_{ik} = A_{ik} (T - T_{\rm C})^{\kappa}. \tag{1}$$

 A_{ik} and κ are properties of the material of interest. The temperature $T_{\rm C}$ is below the transition temperature $T_{\rm tr}$, when the free energies of the low- and high-temperature phases become equal, because the transitions are slightly first order. Such power laws are inspired by simple soft-mode mechanisms, although κ is sensitive to the details of mode-mode coupling, to the degree of anisotropy of dispersion curves about the reciprocal lattice vector of the soft mode, and to the extent of softening along each branch. This effect mimics the bilinear coupling between the order parameter and strain. In the most common case, where direct interactions are symmetry forbidden, there are still mechanisms which lead to softening in the displacive limit. For example, optical phonons with opposite wave vectors \mathbf{q} and $-\mathbf{q}$ can combine to produce a fluctuating strain field. The symmetry-allowed coupling is $e_{\text{local}}\langle Q^2 \rangle$, where e_{local} is the local strain field and $\langle Q^2 \rangle$ is the average two-phonon amplitude.

Alternatively, when local disorder leads to thermally activated dynamics with a broad distribution of activation energies and a threshold related to the phase transition temperature, the elastic softening may follow a Vogel-Fulcher statistics [46,47] with

$$\Delta C_{ik} = B_{ik} \exp\left(\frac{E_a/k_{\rm B}}{T - T_{\rm VF}}\right),\tag{2}$$

where B_{ik} is a materials parameter, E_a is the activation energy, and T_{VF} is the Vogel-Fulcher energy [47–49]. Even though Eq. (2) is not the result of a formal theory, such behavior is typical for glasses and for local clusters in order-disorder phase transitions. In Ba_{1-x}Sr_xTiO₃ the two archetypal softening mechanisms, namely, the displacive power law and a mixed order-disorder Vogel-Fulcher mechanism, can be expected to be related to the antiferrodistortive displacive transition in SrTiO₃ at 105.6 K [50] and the ferroelectricferroelastic transition in BaTiO₃ at 400 K, which contains aspects of disorder from Ti off-centering in the paraelectric phase [51–53].

TABLE I. Density of ceramic samples $Ba_{1-x}Sr_xTiO_3$, assuming a linear theoretical density that changes linearly between 5.13 g/cm³ of SrTiO₃ and 6.02 g/cm³ of BaTiO₃.

x	Density (g/cm ³)	Relative density (%)
0	5.64	93.7
0.03	5.85	97.5
0.1	5.83	97.9
0.3	5.67	97.8
0.5	5.35	95.9
1	5.02	97.9

In materials investigated in this paper the elastic precursor softening [2,54,55] is significant, including complex intermediate cases [56,57]. The power-law softening and, to a lesser extent, the Vogel-Fulcher softening describe the experimental observations very well over a wide temperature range. The two limiting cases, SrTiO₃ and BaTiO₃, behave very differently, and a smooth variation of the model parameters A, κ , B, and E_a is indeed observed in mixed crystals of Ba_{1-x}Sr_xTiO₃ as a function of the chemical composition.

II. SAMPLE PREPARATION

Ceramic $Ba_{1-x}Sr_xTiO_3$ samples were prepared in two different laboratories with different solid-state sintering procedures. The BaTiO₃ sample is sample BT No. 1 of Ref. [58], prepared from commercial high-purity powder of BaTiO₃ (99.9%, Sigma-Aldrich). The powder was first heated at 800 K for 2 h to remove undesired organics and ball milled for 24 h in order to reduce and homogenize the distribution of particle sizes. It was then mixed with 3 wt % polyvinyl butyral (PVB) as a binder, uniaxially pressed at 150 MPa into a thick bar, isostatically pressed at 250 MPa, and sintered at 1350 °C for 2 h. The $Ba_{1-x}Sr_xTiO_3$ materials were synthesized starting from high-purity powders of BaCO₃ (99.8%, Alfa Aesar), SrCO₃ (98%, Merck), and TiO₂ (99.8%, Merck) mixed for 24 h and then calcined at 1150 °C for 5 h. The calcined $Ba_{1-x}Sr_xTiO_3$ powders were checked by x-ray diffraction (XRD) to be of perovskite tetragonal (P4mm) phase, milled for 24 h to obtain a homogeneous particle size distribution around 1 µm, and mixed with 3 wt % PVB as a binder. Bar ingots were uniaxially pressed at 190 MPa into metallic molds, isostatically pressed at 250 MPa, and conventionally sintered at 1350 °C for 4 h. All samples were checked by XRD, and their densities are listed in Table I. The resulting ceramics were cut into thin bars with lengths of 35-42 mm, widths of 4-6 mm, and thicknesses of 0.4-0.7 mm, mechanically polished, and annealed at 750 °C for 2 h to release stresses.

The SrTiO₃ sample was prepared by solid-state reaction of SrCO₃ (Aldrich, 99.9%) and TiO₂ (Aldrich, 99.9%) for 6 h at 1100 °C. The resulting powder was milled, sieved, pressed into a bar, and sintered in air at 1450 °C for 24 h. Thin bars 43 mm long and 0.5 mm thick were cut and polished for the anelastic measurements. The Ba_{0.5}Sr_{0.5}TiO₃ bar, 37 mm long and 0.44 mm thick, was prepared in a similar manner. The sample of SrTiO₃ had $T_{tr} \simeq 111$ K, which is 5.3 K above the value generally found. A possible explanation is a larger than



FIG. 1. Young's moduli of $\text{Ba}_{1-x}\text{Sr}_x\text{TiO}_3$ ceramics normalized to overlap at $T \gg T_{\text{C}}$. The dashed line is $1/S_{11}$ of the SrTiO₃ crystal. The curves are obtained plotting $(f/f_0)^2$, where f(T) is the resonance frequency of the fundamental flexural mode.

usual content of Ca impurities in the starting SrCO₃ powder. To avoid artifacts due to impurities, we also used a single crystal of SrTiO₃ with dimensions of $26.15 \times 3.4 \times 0.5 \text{ mm}^3$, cut from a wafer supplied by MTI with the edges parallel to the $\langle 100 \rangle$ directions.

III. EXPERIMENTAL METHODS

The dynamic Young's modulus E was measured by electrostatically exciting the free flexural modes of the bars suspended in vacuum on thin thermocouple wires, as described in Ref. [59]. The real part is deduced from the resonant frequency of the fundamental flexural mode [60]

$$f = 1.028 \frac{t}{l^2} \sqrt{\frac{E}{\rho}}.$$
(3)

For the $\langle 100 \rangle$ -oriented crystal of SrTiO₃ we measured $E = S_{11}^{-1}$. We do not consider the absolute values of the Young's moduli, but their normalized values, $E(T)/E_0 = (f(T)/f_0)^2$, where the temperature dependence of the density in Eq. (3) is ignored in comparison with the elastic moduli. Its linear dependence does not influence the fits, since it is absorbed in the fit of the background elastic modulus.

IV. RESULTS

Figure 1 presents the Young's moduli of $Ba_{1-x}Sr_xTiO_3$ normalized to overlap at high temperature. The purpose of this figure is to show that the anharmonic linear softening (i.e., the slope dE/dT) is similar for all compositions. The slope of the single crystal is different because the Young's modulus of a ceramic cubic material is

$$E^{-1} = S_{11} - \frac{2}{5} \left(S_{11} - S_{12} - \frac{1}{2} S_{44} \right), \tag{4}$$

while that of a $\langle 100 \rangle$ -oriented crystal is $E^{-1} = S_{11}$ [60], and the various S_{ij} may have different slopes.

The measurement of $BaTiO_3$ was already reported in Ref. [58] (Fig. 1 therein, sample BT No. 1). In addition to



FIG. 2. Young's moduli of $Ba_{1-x}Sr_xTiO_3$ corrected for porosity of the ceramic samples. The curves are obtained from those in Fig. 1 using Eqs. (3) and (5) with $p_c = 1/3$.

the step at the transition from the cubic paraelectric to the tetragonal ferroelectric phases, there are two additional minima due to the ferroelectric transitions to the orthorhombic and rhombohedral phases. In SrTiO₃ the structural transition is antiferrodistortive, with tilting of the TiO₆ octahedra around the c axis. A changeover of the character of the structural transitions in Ba_{1-x}Sr_xTiO₃ was projected at $x \approx 0.8-0.9$ [61]. The normalization of E in Fig. 1 only shows that the temperature slopes in the high-temperature limits of E(T) coincide even though the absolute values of the moduli are different. These absolute moduli are difficult to compare with each other due to irregularities in the sample shapes and differences in porosity, which are not included in Eq. (3) but reduce E in manners that depend on the amount and type of porosity [62-64]. We corrected for porosity effects using the linear decrease in the Young's modulus [62],

$$E = E^* (1 - p/p_c), (5)$$

where *E* is the measured modulus, E^* is the modulus of the dense material, and p_c is a critical porosity. We chose $p_c = 1/3$ because it yields 218 GPa for the maximum value reached by BaTiO₃ in the paraelectric phase, in good agreement with 214 GPa for the single crystal [64,65], and 275 GPa for the room temperature value of SrTiO₃, in agreement with 285 GPa given in Ref. [66]. The result is shown in Fig. 2.

The modulus of the paraelastic phase of $Ba_{1-x}Sr_xTiO_3$ increases with *x*, though the x = 0.3 and 0.5 compositions are well below a linear trend. This may be an artifact because the relative density measured by the Archimedes method underestimates the open porosity, and strong deviations from a simple formula like Eq. (5) may occur, as shown for $BaTiO_3$ [64].

We show in Fig. 3 the data normalized both in magnitude with respect to the maximum value of the modulus, and in temperature with respect to T_{tr} to emphasize the continuous change of the precursor softening above T_{tr} with composition, from a gradual decrease of BaTiO₃ to a sharp decay in SrTiO₃.

For numerical fits, the compliances S_{ij} are split into the background compliance S_{ij}^{bg} and the precursor softening ΔS due to the phase transition. The temperature dependence of



FIG. 3. Young's moduli of $Ba_{1-x}Sr_xTiO_3$ normalized to their maximum on a T/T_{tr} scale with T_{tr} listed in the inset; sc, single crystal (dashed line).

 S^{bg} is approximated to be linear with saturation below a temperature Θ , which can be described in terms of the quantum saturation expression [67]

$$S_{bg} = S_0 + S_1 \coth(\Theta/T).$$
(6)

The precursor softening is

$$\Delta S_{\text{power}} = A(T/T_{\text{C}} - 1)^{-\kappa}$$
(7)

for the power law and

$$\Delta S_{\rm VF} = B(1 - \exp\left[-E_a/(T - T_{\rm VF})\right]) \tag{8}$$

for the Vogel-Fulcher approach. If all the compliances S_{ij} have the same temperature dependences of background and precursor components, though with different amplitudes, the fitting expression

$$E/E_0 = 1/S = 1/(S_0 + S_1 \coth(\Theta/T) + \Delta S)$$
 (9)

is valid for both ceramics and a single crystal.

Figure 4 shows fits of the normalized moduli of ceramic BaTiO₃ and single-crystal SrTiO₃ with Eq. (9) and both Eqs. (7) and (8) and the parameters indicated in the legends. The dashed lines represent the background S_{bg} , which is obtained setting A or B = 0. For the case of the power law in BaTiO₃, S_{bg} is out of scale: from 1.18 at 760 K to 1.38 at 400 K. Similar fits are obtained for the intermediate compositions.

Figure 5 shows the dependence of the parameters κ , A and E_a , B on composition x. The error bars indicate the parameter regions where $\chi^2/\chi^2_{min} \leq 2$. The error bars for the Vogel-Fulcher (VF) parameters are quite large, because the VF expression tends to $B \times E_a/(T - T_{VF})$ for small $E_a/(T - T_{VF})$, and therefore the fit does not change reducing E_a below a certain value. This situation occurs at all compositions, where a decrease but no definite minimum of χ^2 is obtained decreasing E_a and simultaneously increasing B, except for SrTiO₃, where a clear minimum is found for $E_a = 0.50^{+0.05}_{-0.09}$ K.

The power-law dependence is well demonstrated with exponent $\kappa \simeq 0.2$ for BaTiO₃ increasing continuously to 1.5



FIG. 4. Fits of the normalized Young's moduli of BaTiO₃ and SrTiO₃ with Eqs. (6)–(9) for the power-law and Vogel-Fulcher analyses and the parameters indicated in the legends. The dashed lines represent S_{bg} , obtained setting *A* or B = 0. For the case of the power law in BaTiO₃, S_{bg} is out of scale: from 1.18 at 760 K to 1.38 at 400 K.

for SrTiO₃. The activation energy E_a decreases from approximately 3 to 0.50 K. Simultaneously, the amplitudes A and B decrease continuously. For both parameters a plateau for intermediate compositions near x = 0.4 appears to occur.

To better show the close adherence of the precursor softening of BaTiO₃ and SrTiO₃ to a power law, Fig. 6 presents double logarithmic plots versus reduced temperature of the precursor softenings, $\Delta S_{\text{power}} = (E/E_0)^{-1} - S_{bg}$, where S_{bg} is obtained from the fits setting A = 0. The exponents in the legends are obtained from the linear fits of the log-log plots and coincide with the κ parameters of the corresponding fits. The deviations from the power law at small reduced temperature depend largely on the first-order step of the modulus at the transition point between the cubic and tetragonal phases and the very large softening regime. Particularly impressive is the fact that BaTiO₃ closely follows a power law to $T_{\rm C}$ + 360 K, the maximum temperature measured.

We exclude that the measured temperature dependences of the Young's moduli are significantly affected by moving grain boundaries (GBs). Their relaxation operates at kilohertz frequencies above the temperatures where sintering and grain growth occurred. For BaTiO₃ and Ba-rich compositions the sintering temperatures are >1350 °C, about 800 K higher than



FIG. 5. Variation of power-law and Vogel-Fulcher fit parameters with composition *x*. The solid curves are guides for the eye.

the maximum temperature reached in our measurements, so that GBs can be considered immobile. The anelasticity due to GBs has been measured in SrTiO₃, having a sintering temperature lower than BaTiO₃, and the decrease in the elastic modulus due to GB motion occurs above 1300 K, accompanied by a rise in Q^{-1} above 0.01 [68]. In our measurements we always measure $Q^{-1} < 0.0005$ up to the highest temperature we reached. This background dissipation probably includes the tail of the GB peak, but at small levels of dissipation, so that it cannot influence the real part of the modulus. Regarding the possibility that the softening extended to very high temperature comes from near-surface regions of the grains, it does not seem to be the case for SrTiO₃, where there is no qualitative difference between the single-crystal and ceramic samples (Fig. 3). The different slopes of the linear dependences at high temperature are because in the crystal we measure S_{11} while in the ceramic sample we measure a mixture of all S_{ij} ; the differences close to T_{tr} should be rather ascribed to the impurities responsible for the shift in $T_{\rm tr}$. Notice also that the weaker softening of the ceramic sample with respect to the crystal in the tetragonal phase is presumably due to pinning of the domain walls from the impurities.



FIG. 6. Double logarithmic plots of the precursor softening vs reduced temperature of BaTiO₃ and SrTiO₃, $\Delta S_{\text{power}} = (E/E_0)^{-1} - S_{bg}$, where S_{bg} is obtained from the fits setting A = 0. The real temperatures are shown in the upper scales.

V. DISCUSSION

The focus of the present analysis is the precursor softening in $Ba_xSr_{1-x}TiO_3$ beyond the classical critical regime, from a few kelvins above T_{tr} up to the highest temperature reached in our experiments. The major finding is that even in this huge temperature range it is possible to describe the temperature dependence of the elastic moduli in terms of a power law with exponent in line with predictions from models of anharmonic phonon softening or, with somewhat less success, with a Vogel-Fulcher expression. In both cases the fitting parameters evolve with the composition as expected from the increasing order-disorder character from SrTiO₃ to BaTiO₃.

The power-law and Vogel-Fulcher elastic softening theoretically converge for small Vogel-Fulcher energies with power-law exponents near unity. We can therefore define a rather strict "displacive limit" when the Vogel-Fulcher energy vanishes, which is nearly the case for $SrTiO_3$. In all other compounds we observe small but finite values of E_a which indicate some weak contributions of structural disorder. This component increases with increasing Ba content but always remains below the behavior of a typical order-disorder system. On the other hand, the power-law fits show that the anharmonic phonons coupled with the elastic moduli change their character, in particular their dimensionality, in the model in Ref. [2]. If a single branch flattens, the exponent κ becomes 1.5. If two orthogonal branches flatten while the third remains relatively steep, we expect $\kappa = 1$. Finally, if three orthogonal branches flatten, the expected value is $\kappa = 0.5$. The experimental observations are surprisingly close to these values for SrTiO₃ while somewhat exceeding the lower limit for BaTiO₃. Simultaneously, we observe that the precursor temperature interval is smallest for SrTiO₃, while it remains extremely large for Ba-rich materials with significant softening at least up to 800 K. Similar exponents, but smaller precursor temperature intervals, were previously observed in isostructural KMnF3 and KMn_xCa_{1-x}F₃ [29,69,70] with values of κ ranging between 0.4 and 1. In $PbSc_{0.5}Ta_{0.5}O_3$ (PST) the exponent is near 0.5 [71] with a smaller precursor interval, while $\sim 1/3$ has recently been found for structural fluctuations in cuprate superconductors [43].

An intriguing consequence of the precursor softening and the ensuing potential heterogeneity in SrTiO₃ was discussed in Ref. [72], where a strong coupling with the local carrier concentration was described. It was found that even very small amounts of dopants can stabilize the soft phonon branches. This is linked to the formation of polar nanoregions, which grow in size with decreasing temperature. Such nanoregions would modify the precursor exponent κ and would also lead to an increase in the VF activation energy. More generally [73], precursor effects are a key to understand the fundamental aspects of the structural phase transitions in perovskite structures which are unrelated to some traditional concepts of the structural tolerance factors and small-cell density functional theory (DFT) calculations. The mixing of transition mechanisms is most obvious in BaTiO₃ and related materials. Their phase transitions are accompanied by optic mode softening which remains incomplete near $T_{\rm tr}$ due to the first-order nature of the phase transition [74]. Simultaneously, local probe measurements show an order-disorder behavior [4,52,75]. The coexistence of both order-disorder and displacive mechanisms has been widely discussed [76,77]. Migoni et al. [78] pointed out that the most important ingredient of modeling of the soft mode activities is the directional anisotropic core-shell coupling at the oxygen ion lattice site, which is nonlinear with respect to the transition metal and harmonic with respect to the A-site cation. This mirrors our observation that dimensionality and hence the intrinsic anisotropy of the softening process play a major role in the precursor effects of $Ba_{1-x}Sr_xTiO_3$. We relate the difference between the end members for both model fits to the different phonon dispersions as calculated by Bussmann-Holder et al. [73], who found that the transverse acoustic zone boundary transverse acoustic (TA) mode couples strongly to the optic mode at finite momentum. The TA mode is presumed to be the origin of finite-size precursors and polar nanoregions. This mode is less temperature dependent than the soft transverse optic (TO) mode, which softens more strongly in BaTiO₃ and less in SrTiO₃. A crossing of the zone



FIG. 7. Temperature derivatives of the Young's moduli of Ba_xSr_{1-x} TiO₃. For clarity the curves are shifted by 10^{-4} K⁻¹ with respect to each other. The larger noise at the highest temperature in some curves is due to the fact that the measurements were very fast there (up to 7 K/min) in order to limit possible losses of O in high vacuum.

boundary and the zone center mode with decreasing temperature occurs in BaTiO₃ but not in SrTiO₃. The avoided mode crossing induces precursor dynamics at small momenta which are less developed in BaTiO₃, where the optic mode softens over the full momentum space. Bussmann-Holder *et al.* already argued that the softening of the optic mode in BaTiO₃ for all momenta is reminiscent of the breather-type excitation and a signature for the formation of polar nanoregions far above the actual phase transition, which mirrors our findings.

There appears no definite onset for the softening, because all compositions can be well fitted up to $4T_{\rm C}$ or the highest temperature reached experimentally with a power law or VF expression, without a characteristic temperature. This was expected for the displacive transition of nonpolar SrTiO₃, but not for Ba-rich compositions and especially BaTiO₃, where characteristic temperatures have been identified below which polar nanoclusters start forming, i.e., the Burns temperature $T_d \sim 550-590$ K and $T^* \sim 500$ K [46,79-83]. The experimental signatures of such onsets or crossover temperatures are deviations from linearity in the temperature dependence of the refraction index [79], of thermal expansion [81], or of the elastic constants at very high frequency [82], or a maximum in the elastic moduli at megahertz frequencies [46], bursts of acoustic emission [80], or a subtle change in the exponent of the power law describing the second harmonic generation [83]. Below T_d , nanoregions with static average displacements have been observed to develop with scanning transmission electron microscopy (STEM) and Raman spectroscopy [84].

Our data and their analyses do not show evidence of any characteristic temperature up to the highest temperature we reached. If different types of inhomogeneities, static and dynamic, existed in our samples, their evolution would be gradual, and the overall effect is a smooth progressive softening. This is better seen in the temperature derivatives of the moduli, shown in Fig. 7, where there is no evidence of inflections of the E(T) curves.

It appears possible that in $Ba_xSr_{1-x}TiO_3$ (away from the relaxor compositions $x \sim 0.15$) the variously estimated T_d should be considered as temperatures below which precursor phenomena are evident, without proper onsets, which do not appear in the elastic softening. Considering that there are many theoretical models and possibly concomitant mechanisms to account for elastic precursor softening, including the appearance of polar nanoclusters, local intrinsic correlations of precursor displacements [28], etc., it is surprising that the quantitative temperature evolution of the excess softening can be so well described by a simple power law or, to a lesser extent, a formula of the Vogel-Fulcher type. This observation also reveals that the bare elastic moduli (when the precursor softening is eliminated) are much bigger than the measured ones even at very high temperatures. As an example, with power-law fitting, at the highest temperature reached in BaTiO₃, 750 K, the bare elastic modulus is 34% larger than the experimentally observed value, and at 840 K the bare modulus of $Ba_{0.03}Sr_{097}TiO_3$ is 22% larger than the experimental value, so that BaTiO₃ would be even stiffer than SrTiO₃ in the absence of precursor softening.

VI. CONCLUSIONS

Elastic precursor effects are common in ferroelastic materials. An increasing number of publications report such effects although with little or no numerical analysis of the observed temperature dependencies. We proposed two frameworks to analyze such effects by power-law dependence and a Vogel-Fulcher dynamics. It is very likely that these approaches are indeed universal. We urge the community to analyze such effects in the paraelastic phase and/or paraelectric phase because several applications may be based not only on domain boundaries [85] but also on the dynamic effects in the highsymmetry phase. To demonstrate this behavior, we reported in this paper the precursor softening of two prototypical materials and their mixed crystals. We found a smooth crossover between a displacive and a (partially) order-disorder system so that the precursor dynamics can be tailored according to specific applications. We presume that other structural imperfections, such as oxygen defects, will contribute similarly to local structural disorder and enhance the BaTiO₃-type elastic softening with extremely wide temperature intervals and low exponents κ .

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