Polarized Raman study of the phonon dynamics in Pb(Mg\textsubscript{1/3}Nb\textsubscript{2/3})O\textsubscript{3} crystal

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\textbf{I. INTRODUCTION}

Pb(Mg\textsubscript{1/3}Nb\textsubscript{2/3})O\textsubscript{3} (PMN) is one of the earliest members of the growing family of lead ferroelectric relaxors, which, due to its application potential, has been attracting attention of researchers for many years. However, despite significant efforts, many properties of lead compounds are still unclear and the very nature of the relaxor behavior has yet to be understood. The difficulties stem from the high complexity of these materials with a high degree of compositional, chemical and structural disorder.

Pb(Mg\textsubscript{1/3}Nb\textsubscript{2/3})O\textsubscript{3} has a perovskite structure with, on average, a cubic Pm3m symmetry, with Mg\textsuperscript{2+} and Nb\textsuperscript{5+} ions interchanging on B sites. Measurements of the dielectric constant show that the crystal does not undergo a sharp transition to a ferroelectric phase. Instead, the dielectric constant exhibits a broad maximum at $T_\text{max} \approx 270$ K. Below $T_a \approx 620$ K, it deviates from Curie-Weiss law\textsuperscript{2} and, at $T \lesssim 350$ K, exhibits strong dispersion.\textsuperscript{2} The structure of PMN is not homogeneous. If Mg and Nb occupation rates were uniformly and ideally equal 1:2 throughout the crystal, it would have a rhombohedral symmetry of R3m. But, a network of superstructure clusters with face centered cubic symmetry (Fm3m), destroys the picture. The size of these clusters is of the order of 2-3 nm; the distance between the centers of the neighboring clusters is near 2.5 nm.\textsuperscript{3,4}

The most widely accepted hypothesis, so-called “space-charge model”, associates these Fm3m clusters with antiferroelectrically ordered areas where Mg:Nb occupation ratio is equal to 1:1, i.e. with local composition Pb(Mg\textsubscript{1/2}Nb\textsubscript{1/2})O\textsubscript{3}.\textsuperscript{5,6} Such 1:1 clusters carry an excessive negative charge, which is compensated by the positively charged host matrix. Network of these charges acts as a source of random fields and may prevent the occurrence of the “normal” phase transition. Alternatively, the Fm3m clusters might be explained by the local order in the form of Pb[B\textsuperscript{2+}\textsubscript{1/2}B\textsuperscript{5+}\textsubscript{1/2}]\textsubscript{1/2}O\textsubscript{9/10}. This hypothesis avoids assumption of the considerable space charges and calls for other reasons to explain relaxor behavior, including the random fields due to the disorder-induced local anisotropy,\textsuperscript{7} effects related to different ionic radii,\textsuperscript{8} or both.\textsuperscript{9}

The presence of areas with special order causes crystalline structure to change in an intricate way. At the high temperature limit, the crystal has a uniform primitive cubic (Pm3m) structure. Lowering the temperature leads to distinguishability between Mg\textsuperscript{2+} and Nb\textsuperscript{5+} sites and to formation in the ordered regions of a face centered (Fm3m) superstructure. The temperature of the Fm3m clusters formation has yet to be determined. Husson’s data suggests they presence at $T \sim 850$ K.\textsuperscript{10} Our data (see below) raise the upper limit to a temperature higher than 1000 K.

Cooling the sample leads to the nucleation of polar nanoregions (PNR’s) characterized by the local distortions. These distortions are responsible for the deviation of the dielectric constant from Curie-Weiss law at $T_a$. Their presence also causes a non-linearity in the temperature dependence of the optical refraction index.\textsuperscript{11} Size of the PNR’s is equal to 30-80 Å depending on temperature.\textsuperscript{11,12} The appearance of polar distortions is a consequence of the ion off-centering that is a common property of many perovskites. By structural similarity, one could expect that the mechanism of the polar regions formation in PMN is analogous to the one in KTa\textsubscript{1−x}Nb\textsubscript{x}O\textsubscript{3} (KTN).\textsuperscript{13} However, unlike KTN where Nb\textsuperscript{5+} is the only off-centered positive ion, PMN represents a more difficult case. Here, not only both ions of B-type, Mg\textsuperscript{2+} and Nb\textsuperscript{5+}, are shifted by ~0.1-0.2 Å, but also Pb\textsuperscript{2+}, ion of A-type, is off-centered by ~0.25-0.35 Å.\textsuperscript{14-22} The B-ions tend to shift along a (111) direction, thus having eight equivalent positions. The shifts of the Pb\textsuperscript{2+}-ions exhibit a spherically symmetric distribution.\textsuperscript{14,15} It is often assumed\textsuperscript{14,15,16,17} that Nb\textsuperscript{5+},
TABLE I: Interpretations of Raman scattering spectrum from PMN offered by various research groups. Note, many of the entries in the table mutually exclude each other.

| Research Group | Frequency cm\(^{-1}\) | Mode | Interpretation |
|----------------|----------------------|------|----------------|
| Husson (1990)  | 45, 130, 260, 420, 500-600, 780 | Pb-O stretching modes, O-B-O bending, Mg-O-Mg stretching, Nb-O...Nb stretching | Coexistence of tetragonal and rhombohedral phases |
| Dimza (1992)   | Writeable text | | |
| Krainik (1993)  | TO\(_1\) | All spectrum is due to one phonon processes, originates from rhombohedral symmetry clusters and consists of 9 modes | |
| Idink (1994)    | TO\(_1\), TO\(_2\), TO\(_3\), TO\(_4\), TO\(_2\)+TO\(_4\) or LO\(_4\) (1996) | Due to positional disorder of lead, Due to Fm3m clusters and vibration of oxygen octahedra | |
| Marsz (1996-98) | All spectrum originates from Fm3m clusters, similarly to the case of PbSc\(_{1/2}\)Ta\(_{1/2}\)O\(_3\), VV-E\(_g\)??, TO\(_1\), TO\(_2\), TO\(_3\), TO\(_4\), TO\(_2\)+TO\(_4\) or LO\(_4\) (1996) | Disorder-induced scattering from silent modes, Breathing mode | |
| Jiang (1999)    | All spectrum originates from Fm3m clusters, similarly to the case of PbSc\(_{1/2}\)Ta\(_{1/2}\)O\(_3\), VV-E\(_g\)??, TO\(_1\), TO\(_2\), TO\(_3\), TO\(_4\), TO\(_2\)+TO\(_4\) or LO\(_4\) (1996) | Disorder-induced scattering from silent modes, Breathing mode | |
| Lushnikov (1999) | VH-2T\(_{2g}\) | Disorder-induced scattering from silent modes, Breathing mode | |
| Siny (1999)     | VV and one VH have fractal character another VH-T\(_{2g}\) | Disorder-induced scattering from silent modes, Breathing mode | |
| Dujovne (2002)  | TA+disorder | Disorder-induced scattering from silent modes, Breathing mode | |

Due to its position in the cell and small radius, acts as the main ferroelectric agent.

At high temperature, the off-centered ions are free to reorient among all allowed off-centered positions. Cooling down and the growing role of the electric interactions causes the appearance of correlated regions consisting of several neighboring cells. Each such region is characterized by a giant electric dipole moment and a local distortion from the cubic symmetry. The PNRs are capable of reorientational motion as whole units. In KTN, the development of these regions leads, at some critical temperature, to a phase transition. However, in PMN, the polar regions develop under constraints imposed by the local anisotropy. As a result, no ferroelectric phase transition, but a glasslike freezing of the polar regions is observed. A bias electric field of magnitude ~1.8 kV/cm, applied in the (111) direction, is sufficient to compensate for the influence of the frustrating effects and to induce, at \(T_{de} \sim 210\) K, a transition to the rhombohedral R3m ferroelectric phase.

Raman spectra of PMN have been measured and reported by several research groups. But, the high complexity of the processes caused difficulties in their interpretation, especially due to possible coexistence of clusters with Pm3m, Fm3m and R3m symmetry. It has been shown that the major spectral lines are of first-order character. In cubic crystals first-order Raman scattering is prohibited by symmetry. Explanation of the appearance of first-order scattering in principle is the one side of the problem. Another side is the assignment of particular lines. The question of the soft ferroelectric mode (TO\(_1\)) and the strong Raman line at \(~45\) cm\(^{-1}\), which is close to where the TO\(_1\) is expected to be found, has been like a stumbling block for investigators. Interpretations of the PMN Raman spectrum offered by various research groups are summarized in Table I. This table reflects the variety of mutually excluding opinions ranging from those that explain the light scattering by disorder in general (groups of Husson, Krainik, Marsz, Dujovne) to those that connect it to the presence of some particular type of disorder, the one that can lead to the appearance of clusters with symmetry allowing first-order scattering, like face centered cubic (Jiang, Lushnikov), rhombohedral (Dujovne) or tetragonal (Dimza). The assignment of particular lines is even more contradictory. Recently made first-principle calculations can confirm only few of them. We believe, the arguments might be resolved by taking into account the possibility of coexistence of different mechanisms, each playing a role in the scattering processes.
prohibits softening, at $T \sim T_d$ K, due to the drastic increase of damping, it disappears. The overdamping continues down to $T_{do} \sim 210$ K. It also causes significant, almost sixfold, broadening of the TA phonon branch. It seems reasonable to expect that the Raman line at 45 cm$^{-1}$ is also affected by the described phenomenon. However, none of the known Raman reports show any significant effects that could be attributed to this overdamping.

The formation of the lower-symmetry clusters in the host crystal is accompanied by relaxational and reorientational dynamics. This dynamics should be reflected in the light scattering spectrum either directly or through interaction with the phonon modes. Theoretical considerations show that internal relaxational motion leads to the appearance of the central peak (CP) in the Raman spectrum. Interactions of this motion with phonons can cause softening of them. These predictions have been tested on several types of crystals with internal degrees of freedom, like alkali-halide-cyanide (K(CN)$_x$-KBr$_{1-x}$, K(CN)$_x$-KCl$_{1-x}$) compounds. Recently, we have analyzed the reorientational motion in ferroelectric KTa$_{1-x}$Nb$_x$O$_3$ crystals. In this paper we discuss the role of the relaxational and reorientational dynamics in the scattering processes from PMN crystal.

In the paper presented we offer a complex analysis of temperature dependencies of polarized (VV) and depolarized (VH) Raman spectra from a single crystalline PMN sample. This analysis is based on multiple peak decomposition. An attempt to apply similar kind of analysis has earlier been made by Siny et al. But, it was restricted to the low frequency VV spectra in a limited temperature range (with maximum temperature $\sim 620$ K), concentrating on the central peak (CP) only. To obtain the complete picture, we consider the temperature evolution of the whole Raman scattering spectrum (up to 1000 cm$^{-1}$) in the 100-1000 K temperature range. We interpret our results in connection with recently published neutron scattering data and with the concept of interaction of phonons with internal relaxational motion. As a result of this work, we present our view on the picture of structural transformations in PMN crystal and on the origin of its Raman spectrum.

II. EXPERIMENTAL SETUP AND RESULTS

We have investigated the Raman scattering from (100) cut PMN single crystal. The crystal was grown by the high temperature solution technique using 30 wt% PbO as flux. The growth conditions were optimized based on the pseudo-binary phase diagram established for PMN and PbO compounds. The as-grown crystal exhibits a pseudo-cubic morphology with (100) cub growth steps. A (100) / (110) cub-oriented crystal of a volume of 53 mm$^3$ was cut from a large as-grown crystal and polished with fine diamond paste (down to 0.25 mm). It showed very high optical quality and satisfied the requirements for light scattering studies. The scattering was excited by propagating in (100) direction 514.5 nm light from a 200 mW Ar$^+$-ion laser, focused to a 0.1 mm spot. The scattered light was collected at an angle of 90° with respect to the incident beam (i.e., in 010 direction) by a double-grating ISA Jobin Yvon spectrometer equipped with a Hamamatsu photomultiplier R-649. For most of the measurements, the slits were opened to 1.7 cm$^{-1}$. However, in order to acquire more precise data in the central peak region, the temperatures close to the maximum of the dielectric constant ($100 < T < 350$ K), the slits were narrowed to 0.5 cm$^{-1}$. Each polarization of the scattered light, $\langle x|zz|y \rangle$ (VV) and $\langle x|zx|y \rangle$ (VH), was measured separately. In order to exclude differences in sensitivity of the monochromator to different polarizations of the light, a circular polarizer was used in front of the entrance slit.

For control purposes, we also took measurements without polarization analysis. Finally, to protect the photomultiplier from the strong Raleigh scattering, the spectral region from -4 to +4 cm$^{-1}$ was excluded from the scans. The data were collected in the temperature range from

![FIG. 1: Examples of Raman scattering spectra measured at various temperatures without polarization analysis and corrected by the population factor.](image-url)
1000 to 100 K. The cooling rate was 0.5-1 K/min. Every 50-20 K the temperature was stabilized and the spectrum recorded. We agree with Ref. that the reproducibility of the spectra in the temperature range approximately between 200 and 350 K was not very good. In this range, the spectra exhibited a strong dependence on the cooling rate and other experimental conditions.

Fig. 1 presents examples of light scattering spectra measured at different temperatures without polarization analysis. To facilitate comparison, we corrected the spectra by the Bose population factor:

\[ F(f, T) = \begin{cases} 
  n(f) + 1, & \text{for Stokes part} \\
  n(f), & \text{for anti-Stokes part}
\end{cases} \]

where

\[ n(f) = (\exp(hf/kT) - 1)^{-1}. \]

Figure 2 demonstrates scattering spectra measured in VV (a) and VH (b) geometries at different temperatures. These spectra we present in uncorrected, ”as-measured”, form.

As can be seen from Figs. 1 and 2 our spectra are consistent with those from Refs. In the high temperature region, a typical spectrum consists of two strong lines centered approximately at 45 cm\(^{-1}\) and 780 cm\(^{-1}\) (labelled A and B) and of three broad bands (C, D, E). Line A exhibits fine structure, which is explicitly seen on the spectra measured with polarization analysis (see Figs. 2 and 3). Lowering temperature leads to the splitting of the broad bands C, D and E into a number of narrower lines (labelled by indices) and to appearance of new line F.

As seen from Fig. 1 starting from 1000 K, the reduced intensity of the scattering, first grows until the temperature of \(\sim 700\) K then sharply decreases to a minimum located close to the Burns temperature \(T_d \approx 620\) K, after which it increases again. At \(\sim 550\) K its strength gets restored. Below this temperature, the intensities of the major lines (especially A and B) are determined primarily by the Bose factor. I.e. being corrected by it, they do not show significant temperature changes, demonstrat-
FIG. 3: Examples of multiple peak decomposition of Raman spectra measured at 1000 and 110 K both, in VV and VH geometry. For convenience, major peaks are labeled with letters. Inserts show examples of fits for the line A using the model of the two coupled harmonic oscillators, as described in Discussion.

The characteristic first-order behavior. The shape similarity of the low- and high-temperature spectra shows that even at 1000 K the scattering has first-order character. In the perfectly cubic crystal, first-order scattering is prohibited. Therefore, one should assume the presence of distortions in the form of lower symmetry clusters, like Fm3m or R3m. At high temperature, these clusters might be present in the dynamic form, having short lifetime. Lowering the temperature, their lifetime increases and they become progressively more static. The observed around T_d decrease of Raman intensity is also clearly seen in Fig. However, there are no reports on this effect in the literature; it might have been overlooked. This intensity drop indicates worsening of the optical quality of the crystal. In our opinion, it is consistent with the formation of Fm3m clusters in their evolution from dynamic to static form. The splitting of peaks is associated with the development of R3m clusters. The first signs of the splitting appear at T_d. Starting from T_f ~ 350 K (which is also close to where the dispersion of the dielectric constant appears), this effect becomes even more definite and the intensities of the components begin to grow. Below T_d ~ 210 K, which is the temperature of electric field-induced ferroelectric phase transition, the shape of the spectra is relatively stable.

A comprehensive analysis of our data and their comparison with results of other experiments allows to understand the meaning of the mentioned above temperatures T_d ~ 620 K, T_f ~ 350 K and T_d ~ 210 K in the temperature evolution of the Raman spectra and the lattice structure.

III. MULTIPLE-PEAK DECOMPOSITION OF SPECTRA PROCEDURE.

To be able to analyze the data, we decomposed the measured spectra using multiple peak fitting procedure. It was found that satisfactory fit could be achieved with the assumption that the central peak has a Lorentzian shape and that each of the other peaks is described by the spectral response function, i.e. damped harmonic oscillator, modified by the population factor φ:

\[ \Phi_i \sim \frac{\Gamma_i f_i^2}{(f^2 - f_0^2)^2 + \Gamma_i^2 f_i^2} F(f, T), \]  

(2)
where $\Gamma_i$ and $f_{0i}$ are the damping constant and the mode frequency.

As all of the peaks are much better resolved at low temperatures, we started our fitting procedure at the low-temperature end of the data set (at 110 K) and, then, observed the evolution of the peaks with increasing temperature (i.e., in order, opposite to the order of measuring). At the same time, we tried to minimize the number of peaks necessary to achieve a reasonably good fit. The control data set (measured without polarization analysis) has been used to calibrate the positions and widths of the weak and poorly resolved peaks from the VV and VH data sets. Since a large number of parameters is involved, results of a particular fit may depend on their initial values. To stabilize the results, the best-fit values of parameters obtained at the previous temperature were used as initial values for the next one. In this manner, several sets of fits were obtained and analyzed. It is remarkable, that in all of them the major parameters showed the same trends of behavior, confirming the importance of the mentioned above temperatures: $T_d \approx 620 \pm 50$ K, $T_f \approx 350 \pm 25$ K and $T_{do} \approx 210 \pm 25$ K. These trends are summarized in Table II. One of the sets have been selected to report the most interesting results in detail. Examples of the fits at the temperatures of 1000 and 110 K in VV and VH geometries are shown in Fig. 3. For clarity, we describe the observed phenomena from high to low temperatures, following the same order as in measurements (unless the opposite stated).

IV. MOST INTERESTING RESULTS OF DECOMPOSITION AND THEIR ANALYSIS.

A. Central peak.

The temperature dependences of the fitting parameters for the central peak (CP) are presented in Fig. 4. Circles correspond to the VV and triangles to the VH component of the peak. The existence of the CP is a direct consequence of the lattice relaxations, which are very sensitive to the restrictions imposed by the low-symmetry clusters. If relaxations are fast, the CP is low-intense and broad, whereas their slowing causes growth and narrowing of the peak.

We would like to point out a striking similarity of the temperature behavior of the CP in PbMg$_{1/3}$Nb$_{2/3}$O$_3$ (Fig. 1) and in KTa$_{0.85}$Nb$_{0.15}$O$_3$ (Fig. 3 in Ref. 3) crystals. We have shown that the temperature behavior of the CP in KTN can be explained by the model involving the relaxational motion of off-centered Nb ions and its progressive restriction with temperature decrease. In the cubic phase, Nb ions are allowed to reorient amongst eight equivalent <111> directions. The appearance of the PNR’s followed by a sequence of phase transitions down to a rhombohedral R3m phase, limits the ion motion to four, two and, finally, locks it in only one site. This model is in agreement with the diffuse neutron scattering studies of similar systems. The similarity of the CP behavior in these two very different materials, suggests that the temperature evolution of the polar clusters in PbMg$_{1/3}$Nb$_{2/3}$O$_3$ passes through the similar stages as in KTa$_{0.85}$Nb$_{0.15}$O$_3$ (KTN), however, it is not accompanied by appearance of the long-range order.

Starting from the high-temperature end (Fig. 4), the first important feature is the strong and narrow scattering in the VV geometry accompanied by a relatively weak scattering in the VH geometry. This indicates the presence of a symmetric slow relaxational motion, involving 180° reorientations of ions. Lowering the temperature, starting from ~ 900 K, a cessation of this motion causes an intensity decrease of the CP. Slow at first, this decrease becomes sharper and reaches minimum near $T_d \approx 620$ K. The prohibition of 180° reorientations in KTN, however, it is not accompanied by appearance of the long-range order.

FIG. 4: Temperature dependencies of the intensity (a) and half-width (b) of the Lorentzian approximation for the central peak in VV (circles) and VH (triangles) geometries of experiment.
TABLE II: Major peaks and their response to the temperature restrictions on ion motion. Only changing properties are shown. Tₜ ≈ 620 K is also characterized by the total loss of intensity of scattered light.

| Peak | Polarization | Region I. Unrestricted | Region II. Restricted | Region III. Formation of |
|------|--------------|-------------------------|-----------------------|-------------------------|
|      |              | T_d (T_d=620 K)         | (T_d=620 K)           | (T_d=620 K)            |
|      |              | T_d (T_d=620 K)         | T_p (T_p=350 K)       | T_p (T_p=350 K)        |
|      |              |                         |                       |                         |
| Central | VV | Intensity decreases | Intensity decreases | Intensity has a maximum at 300 K |
|        |     | broadens             | narrows               |                         |
|        | VH | Intensity decreases | Intensity increases | Intensity has a maximum at 300 K |
|        |     | width has a maximum at 450 K |
| Peak A  | VV | Softens, broadens    | Hardens and narrows  |                         |
|        |     | Broadens             | Broadens              |                         |
|        | VH-I| Broadens             | Hardens, intensity has minimum at 400 K |
|        |     | Softens              | Broadens and grows    |
| Peak C  | VV | Hardens and narrows  | Fine structure appears| Splits in two           |
|        |     |                     |                        |                         |
|        | VH | Hardens and narrows  | Fine structure appears| Splits in two           |
|        |     |                     |                        |                         |
|        | VH | Softens              | Splits in two         | Softens and narrows    |
| Peak D  | VV | Splits in two        | VV component appears  | VH component appears    |
|        |     |                     |                        |                         |
|        | VH | Softens              | VV component appears  | VH component appears    |
| Peak E  | VV | Splits in two        | Fine structure appears| Splits in two, VH appears|
| Peak B  | VV | Hardening            | Hardening             | VH appears             |

wavelength of light) dynamic fluctuations causing worsening of the optical quality of the sample.

With further decrease of the temperature, the optical quality of the crystal improves. From ~ 550 K, the four-cite reorientational motion of the PNR’s starts to slow down, which is marked by the narrowing of the VV component of the CP and increase of its VH intensities. The simultaneous broadening of the VV component up to the maximum at ~ 450 K, indicates new type of restrictions on the ion motion. Analogy with KTN suggests that below ~ 450 K the motion of R3m clusters becomes limited to two neighboring <111> orientations, averaging in monoclinic-like distortions. Such a rearrangement causes some decrease in intensity of the VV component (with minimum at ~ 400 K), while the VH intensity keeps growing. Below ~ 400 K, the slowing down of the two-site relaxational motion causes narrowing the CP and increase of intensities of both components. From T_f ≈ 350 K, these effects become especially dramatic. Further temperature decrease, starting from ~ 300 K, leads to the complete prohibition of intersite reorientational motion of R3m clusters, i.e. to appearance of static R3m clusters. This is marked by a sharp decrease in intensity of both components of the CP. At the temperature T_do ≈ 210 K, the process of conversion of PNR’s from dynamic to static form (freezing) is primarily finished. Below this temperature, the central peak is narrow and its intensity is small in both scattering geometries.

We should mention that the result of our analysis of the VV component of the CP exhibits similar tendencies to those in Ref.[54], except at the low temperature end. The discrepancy occurs due to the difference in experimental technique. In the central peak region, we did the measurements with much higher resolution (0.5 cm⁻¹, as compared to 2 cm⁻¹ from Ref.[54]). Consequently, we were able to provide a more correct separation of the slow relaxations from the elastic scattering, which was especially important at low temperatures.

B. Line A.

Figure[6] presents the temperature evolution of the fitting parameters of the peak A, showing its position (a), reduced intensity (b) and damping constant (c). This peak has a triplet structure, containing one component in VV (circles) and two components in VH (up and down triangles) geometries. The fitting parameters for this peak exhibit changes at mentioned above temperatures T_d, T_f and T_do, confirming their importance for the structural evolution of the crystal. However, the origin of this peak (see Table. [1]) requires clarification. From a comparison with the frequencies of the phonon modes determined from neutron scattering, it is clear that this line cannot be due to the zone center soft TO₁ mode (black stars in Fig. [5]). On the other hand, the lower frequency VH component, and possibly the VV component could be due to the disorder-induced zone boundary scattering on TA phonon (white stars in Fig. [5]). However, the higher frequency VH component would still not be accounted for.

In an attempt to account for both VH components simultaneously, we have tried to make use of a coupled oscillator model. In the approximation of linear coupling
between two harmonic oscillators\textsuperscript{57} this approach gains:

$$I(\omega) = \left[ n(\omega) + 1 \right] \left( \frac{CY - BZ}{B^2 + C^2} \right),$$

where

$$B = (\omega_1^2 - \omega_2^2)(\omega_1^2 - \omega_2^2) + \omega_2^2(\gamma_1^2 - \gamma_1 \gamma_2) - \omega_1 \omega_2 \Delta_{12}^2,$$

$$C = \omega_2^2(\gamma_1^2 - \gamma_2^2) + \gamma_2(\omega_1^2 - \omega_2^2) - 2(\omega_1 \omega_2)^{1/2} \Delta_{12} \gamma_{12},$$

$$Y = S_1^2(\omega_2^2 - \omega_1^2) + S_2^2(\omega_2^2 - \omega_1^2) - 2S_1 S_2(\omega_1 \omega_2)^{1/2} \Delta_{12},$$

$$Z = \omega(S_1^2 \gamma_2 + S_2^2 \gamma_1 - 2S_1 S_2 \gamma_{12}).$$

In these equations \(\omega_1\) and \(\omega_2\) are the resonant frequencies, \(\gamma_1\) and \(\gamma_2\) are the damping constants, \(S_1\) and \(S_2\) are the structure factors for two oscillators, \(\Delta_{12}\) and \(\gamma_{12}\) the real and imaginary parts of the coupling constant.

It was shown\textsuperscript{58} that a unitary transformation can be found, which would set either real or imaginary part of the coupling to be equal to zero. We have chosen to use imaginary coupling and set \(\Delta_{12} = 0\). We also set the structure factors \(S_1\) and \(S_2\) to be the same at all temperatures. The rest of the parameters were treated as being temperature dependent. In the result of fitting, we have found that the values of structure factors can be approximated as \(S_1 = 22.3\) cm\(^{-1}\) and \(S_2 = 20.6\) cm\(^{-1}\). The other parameters as a function of temperature are shown in Fig. 3: the best-fit frequencies (a), the damping constants (b) and the coupling coefficient (c).

The examples of the fits (inserts in Fig. 3) demonstrate that this model gives a good approximation to the shape of the peak A. However, a comparison of the best-fit frequencies (Fig. 6a) with the neutron scattering data\textsuperscript{43,44,45,46} rules out the possibility to explain the higher-frequency VH component by the interaction between ZC TO\(_1\) (black stars) and ZB TA (white stars) modes. The lower frequency oscillator can still be associated with disorder-induced scattering on ZB TA mode.

The possible explanation of the presence of two VH components in the peak A can be in the correspondence to TA modes with different polarizations in different crystalline planes of a distorted lattice. (A similar effect we have seen in monodomain KTa\(_1-x\)Li\(_3\)O\(_3\), where one or the other TA polarization was observed, depending on the orientation of the electric field.) These differently polarized modes are influenced by the polar nanoregions, which explains the increase of damping (Fig. 6b) and coupling (Fig. 6c) coefficients with lowering temperature.
It is also necessary to point out an observation concerning the temperature dependences of the frequencies of the components of the triplet A in approximation of independent oscillators (Fig. 5). These dependences seem to show anomalies at intersections with ZC TO$_1$ modes (black stars). For example, the VV component exhibits deeps at $\sim$ 700 and 350 K and VH-I component has a knee at $\sim$ 400 K.

![Graph](image)

FIG. 7: Amplitude of splitting between the components of the line E. The insert demonstrates that this splitting follows the critical dependence.

C. The most important parameters of other lines.

Among the changes occurring to the Raman spectrum with cooling the sample, the splitting of the phonon lines deserves special attention, since such a splitting is a direct indicator of the modifications of the local structure. As we already mentioned, the splitting occurs to the lines C, D, and E, which is especially well seen in the VV geometry (Fig. 2). At high temperatures, their widths are very big, so the lines almost merge in a single broad shoulder. Lowering the temperature, they become more discernible and the fine structure gradually develops. The first signs of it appear at $T_d \approx 620$ K. Below $T_{pr} \approx 350$ K, the fine structure becomes more obvious. These observations support the expressed above idea that the scattering originates from the distorted regions. Highly dynamic and disordered at high temperature, they become progressively more static with cooling down, and their motion becomes more correlated. This process is reflected by the temperature evolution of the lines C, D, and E.

Below the temperature of $T_{pr} \approx 350$ K, the splitting value of the line E exhibits especially interesting trend. As seen from Fig. 4 with temperature decrease below $T_{pr}$ the distance between the E-line components increases. Moreover, this increase exhibits the characteristic order parameter behavior, following the $(T_{pr}-T)^{-1/2}$ dependence, which is demonstrated in the insert to Fig.

![Graph](image)

FIG. 8: Temperature dependencies of the fitting parameters: position (a), reduced intensity (b) and damping constant (c) of the line B (located at $\sim$780 cm$^{-1}$.)

Finally, we would like to stop on the line B located at the upper end of the spectrum. Fig. 5 demonstrates the fitting parameters for this line: position (a), reduced intensity (b), and damping coefficient (c). This line is very strong in the VV geometry but it is rather weak in the VH one (i.e. characterized by A$_1g$ symmetry). It seems to be the least influenced by the ordering processes in the crystal. However, the temperature evolution of its parameters reflects development of the low-temperature phase, confirming the expressed above ideas about formation of dynamic polar clusters at $T_d \approx 620$ K, their slowing down, and appearance of the static order at $T_{pr} \approx 350$ K. As it is seen from the Table 1, the appearance of this line in the Raman spectrum has several contradictory explanations. However, according to the recent first-principle calculations, this mode most likely is a consequence of oscillations of oxygen ions.
V. CONCLUSION

This paper offers a comprehensive analysis of the Raman scattering spectra of PbMg1/3Nb2/3O3 in the temperature range between 100 and 1000 K. From our analysis, the structural evolution of the lattice occurs in several stages. Even at 1000 K, the PMN crystal is characterized by the presence of dynamic distortions of the lattice from cubic symmetry, which are responsible for existence of first-order Raman scattering at such a high temperature. At the Burns temperature $T_d \approx 620$ K, the first restrictions on the reorientational motion of the off-centered ions appear. These restrictions lead to slowing of the motion and growth of correlations between neighboring regions with R3m symmetry. They also introduce distinguishability between sites occupied by Mg and Nb, which causes onset of Fm3m symmetry in the 1:1 ordered areas. Further cooling leads to progressive growth of restrictions on ionic motion and, starting from $T_{pc} \approx 350$ K, to the appearance of the static polar nanoregions. At $T_{do} \approx 210$ K, the formation of the low-temperature phase is primarily finished. The shape of the phonon lines is determined by multiphonon processes involving phonons with different polarizations propagating in different directions, which interact with dynamic and static disorder.

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