Raman spectroscopy study of ferroelectric modes in [001]-oriented 0.67Pb(Mg$_{1/3}$Nb$_{2/3}$)O$_3$–0.33PbTiO$_3$ single crystals

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This work investigates the evolution of micro-Raman spectra of a [001]-oriented 0.67Pb(Mg$_{1/3}$Nb$_{2/3}$)O$_3$–0.33PbTiO$_3$ single crystal in the temperature range from −120 to 200 °C. For a poled sample, the $T_{1u}$ mode splits into an $E$ mode (remaining at 51 cm$^{-1}$) and a new $A_1$ mode (shifting to 95 cm$^{-1}$ at −120 °C) at the tetragonal to rhombohedral phase transition ($T_C$ ~ 70 °C), while other modes remain the same. This $A_1$ mode belongs to the rhombohedral phase. If the sample is depoled, this $A_1$ mode persists for temperatures much above $T_C$ on heating, implying that local regions with rhombohedral symmetry still exist in the tetragonal phase. The underdevelopment of long-range order in the depoled crystal causes the sample to have typical characteristics of a relaxor while poling will enhance the long-range order to make it into a regular ferroelectric. © 2005 American Institute of Physics. [DOI: 10.1063/1.1948514]

Lead magnesium niobate, Pb(Mg$_{1/3}$Nb$_{2/3}$)O$_3$ (PMN), is a typical relaxor ferroelectric which exhibits a diffuse phase transition characterized by a strong frequency dispersion of the dielectric susceptibility at temperatures near and below the $T_{\text{max}}$ (265 K at 1 kHz). PMN has quenched chemical disorder on the perovskite B site, which is occupied by either Mg$^{2+}$ or Nb$^{5+}$ cations. When doped with PbTiO$_3$ (PT), the compounds (1−$x$)PMN−$x$PT (PMN–PT) exhibit a drastic increase in piezoelectric properties. The crystals with compositions near the morphotropic phase boundary (MPB) ($x$ =0.33−0.34) have piezoelectric constant $d_{33}>2400$ pC/N, and the electromechanical coupling coefficient $k_{33}>90\%$, when they are poled along [001] of cubic coordinates. The origin of this giant piezoelectricity was suggested as due to orientation effect.

Since the PMN–PT system is a solid solution of relaxor and ferroelectric PT, there are many interesting physical phenomena in this system owing to the competition of the two. At the MPB composition, the relaxor and ferroelectric characteristics are somewhat balanced, hence, the conversion between nano-domains (relaxor nature) and macro-domains (ferroelectric nature) can be more conveniently studied in the MPB system.

Because its vibrational spectrum has a shorter characteristic length scale, Raman microprobe is a good microscope to study the nanodomain structures that are the key for the interpretation of the relaxor properties. Previous results on both relaxor ferroelectrics and relaxor-based ferroelectrics showed that, even in the high-temperature cubic phase, the Raman spectra do not differ qualitatively from those of low symmetry phases except intensity variations of some modes. Unique ferroelectric modes have not been observed in the low symmetry phases up to date. There are already some studies on Raman spectra of PMN–PT crystals in the literature. Here we will try to answer the question of whether the tetragonal to rhombohedral structural change has any corresponding changes in the vibrational spectrum. In this work, micro-Raman measurements were performed on a [001] oriented 0.67PMN–0.33PT single crystal, which could provide interesting information about the phase transition from tetragonal to rhombohedral phase since the composition is near the MPB. Most importantly, we have found a unique mode splitting that can signify this phase transition.

The single crystal used in this study is transparent with light-yellow color. It is [001] oriented with the composition of 0.67Pb(Mg$_{1/3}$Nb$_{2/3}$)O$_3$–0.33PbTiO$_3$. The crystal was cut into a plate and optically polished. The dimensions of the plate are 3 mm on each side and 0.7 mm in thickness. Before the Raman measurements, the crystal was annealed at 300 °C for half an hour in air to remove possible residual stresses and polarization. Micro-Raman scattering spectra were recorded in back scattering geometry using a JY T64000 micro-Raman spectrometer with a coherent Ar ion laser. The excitation line is at 514.5 nm and the light spot size is about 1 µm using the ×100 objective lens. The spectra were presented without polarization analysis in order to displaying all observable features.

In a recent study, temperature-dependent dielectric behavior and related phase transitions of the [001] oriented 0.67PMN–0.33PT crystal were investigated by impedance measurements and high resolution x-ray diffraction. It was demonstrated that a phase transition from the rhombohedral symmetry to the tetragonal symmetry occurs near 70 °C on heating, which is more clearly shown in the poled sample than in depoled sample. In the present study, we performed Raman spectroscopy study on both poled and depoled samples. Figure 1 shows the light-scattering spectra for temperatures from 180 °C down to −120 °C for the poled sample. It can be seen that all bands are relatively broad, indicating band overlapping, which is typical for perovskite relaxor-based ferroelectrics. The Raman spectra at tempora-
tures above 30 °C are similar to those of the 0.81PMN–0.29PT, where four bands centered near 780, 580, 270, and 51 cm$^{-1}$ had been found. These four modes correspond mainly to Nb–O–Mg stretching mode, oxygen bending vibration, B-site ion against O stretching vibration inside the octahedral, and Pb against BO$_6$ octahedra translational vibration, respectively. On the one hand, the frequencies and bandwiths of the three bands centered near 780, 580, and 270 cm$^{-1}$ are essentially temperature independent, similar to the results found in 0.81PMN–0.29PT crystals. Interestingly, the Pb mode at 51 cm$^{-1}$ shows strong temperature dependence. This mode is in fact a composite mode, which starts to bulge at 30 °C then gradually splits into two peaks upon further cooling. The new mode shifts towards higher frequency with decreasing temperature. It is at 71 cm$^{-1}$ at 20 °C and keeps on shifting up until 95 cm$^{-1}$ at −120 °C.

The sample was depoled by heating above 200 °C then cooling back down to room temperature without applying electric field. Raman experiments were carried out during heating from −120 to 200 °C for the depoled sample and the spectrum was taken when the temperature was held at each selected temperature for 5 min. The results from −20 to 80 °C are shown in Fig. 2. The Raman spectra at temperatures lower than 0 °C are similar to those of the poled sample (Fig. 1). At −120 °C, the lowest frequency band consists of twin peaks at 51 and 96 cm$^{-1}$, respectively. The 96 cm$^{-1}$ peak has the tendency to merge into the 51 cm$^{-1}$ peak on heating, but never disappear completely like in the case of poled sample. It shifts to lower frequency on heating until 55 °C but comes back up again when the temperature rises above 60 °C. Its peak position is centered at 89 cm$^{-1}$ at 70 °C. At higher temperatures, these two peaks are difficult to identify owing to the increasing bandwidth. As a result, the shape of the peak becomes asymmetric.

The labeling scheme of Burns and Scott has been used here for clarifying the origin of the twin peaks. In the cubic perovskite paraelectric phase, there are 12 optic modes which belong to the $3T_{1u}+T_{2u}$ irreducible representation of the $O_h$ point group. The $T_{1u}$ modes are infrared active (IR) and the $T_{2u}$ mode is a “silent mode” since it is neither infrared nor Raman active. In the tetragonal ($C_{4u}$) and rhombohedral ($C_{3v}$) structures of the ferroelectric phase, each triply degenerate $T_{1u}$ mode splits into $A_1$ and $E$ modes, while the $T_{2u}$ silent mode splits into $B_1$ and $E$ modes. In addition, long-range electrostatic forces in the ferroelectric phase split all the $A_1+E$ Raman and IR active modes into LO and TO components. Thus, the $T_{1u}(TO)$ mode of the paraelectric cubic phase splits into $A_1(1TO)$ and $E(1TO)$ modes, while the $T_{2u}$ mode into $B_1$ and $E$ modes. The mode below 100 cm$^{-1}$ observed in other compounds has shown that, contrary to the group theory predictions, Raman-active vibrational modes are observable in the paraelectric phase due to the presence of low symmetry nanoclusters. This is consistent with our results. Most of the modes (such as modes near 780, 580, 270, and 51 cm$^{-1}$) do not obviously change on heating or cooling and have no soft mode behavior near any of the structural phase transitions. The persistence of Raman peaks in the cubic phase shows that the local symmetry deviates from the overall average.
modes occurs at the tetragonal-to-rhombohedral phase transition.

The presence of Raman modes in the paraelectric cubic phase is generally ascribed to the presence of nanoscale chemical/structural inhomogeneities in the relaxor and relaxor-based compounds. Such inhomogeneities result in local polar clusters, or nanodomains, which have lower symmetry. The short-range order nanodomains with off-axis shift of lead ions result in a break down of the translation symmetry, leading to the failure of the zone center selection rules and a broadening of the Raman lines. On the other hand, when long-range order is achieved, zone center processes are enhanced, hence soft modes may appear at the ferroelectric phase transitions.\(^\text{12}\) Our results show that the Pb related vibrational mode does have a noticeable change at the tetragonal to rhombohedral phase transition.

It was pointed out that the long-range order in relaxor-based ferroelectrics can be driven by either increasing the PT content or applying a dc bias.\(^\text{13}\) The change of dielectric constant with temperature at various frequencies is presented in Fig. 4 for both poled and depoled sample. It can be seen in the inset that the phase transition between the tetragonal and rhombohedral phases triggers a shoulder in the dielectric constant of the depoled 0.67PMN–0.33PT crystals.\(^\text{13}\) This transition near 70 °C becomes more evident for the poled sample as shown in Fig. 4(a), indicating that the poling process enhances the long-range order in the crystals below the Curie point. Based on this fact, we may understand our Raman data as the following.

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