LETTERS

Pressure-Induced Ferroelectricity in (NH₄)₃H(SO₄)₂

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(Received July 12, 1976)

Dielectric properties of single crystals of $(NH_4)_3H(SO_4)_2$ were measured at various hydrostatic pressures up to about 8 kbar below room temperature. The broad peak of dielectric constant along the *c**-direction which appears around -25 °C at 1 atm becomes more and more prominent as pressure increases, and then a ferroelectric phase (Phase VI) develops above 4.9 kbar. Another ferroelectric phase (Phase VII) is also found above about 5 kbar in a lower temperature region. The pressure-temperature phase diagram is given.

Triammonium hydrogen disulfate (NH₄)₃H $(SO_4)_2$ has five phases at atmospheric pressure.¹⁾ In order of descending temperature, they are denoted as I (stable above 140°C; trigonal), II (140°C~-8°C; monoclinic), III $(-8^{\circ}C \sim -132^{\circ}C)$, IV $(-132^{\circ}C \sim -140^{\circ}C)$ and V (below -140 °C). Dielectric and thermal anomalies have been observed for the transitions between them.^{1,2)} However, ferroelectric activity has not been observed for any of these atmospheric phases. In addition to these phase transitions, there is a broad peak of dielectric constant along the c^* -direction at around -25°C.¹⁾ The maximum value of the dielectric constant is about 60 at 1 atm. The peak is too broad to be attributed to a phase transition. The present study deals with the effect of hydrostatic pressure on this anomalous dielectric-constant peak.

Single crystals of $(NH_4)_3H(SO_4)_2$ were grown from an aqueous solution by slow evaporation at about 30 °C. Several *c*-plate specimens were polished with #1200 mesh carborundum powder, and then they were attached with silverpaste electrodes. A Cu-Be pressure bomb was served for the dielectric measurements at high pressures. The pressure-transmitting fluid used was 50-50 mixture of *iso*- and *n*-penthane. Dielectric constant was measured with a threeterminal capacitance bridge at a frequency of 100 kHz. Ferroelectric hysteresis loops were observed by using a conventional Sawyer-Tower circuit.

Figure 1 shows the temperature dependence of dielectric constant at various pressures. The broad dielectric-constant peak which is seen at about -25 °C at 1 atm becomes more and



Fig. 1. Temperature dependence of the dielectric constant of $(NH_4)_3H(SO_4)_2$ along the *c**-direction at various hydrostatic pressures. Frequency: 100 kHz.

more prominent as pressure increases. The reciprocal of the maximum dielectric constant linearly decreases with pressure as $1/\varepsilon_{max} =$ $C(p_0-p)$, where the parameters C and p_0 are estimated as $C = (3.20 \pm 0.03) \times 10^{-3} \text{ kbar}^{-1}$ and $p_0 = 5.1 \pm 0.1$ kbar. Slightly below the critical pressure of p_0 , the peak splits into two rather sharp maxima indicating that a high pressure phase develops between them. The phase is denoted as phase VI. The lowertemperature peak, however, is masked by the appearance of another pressure-induced phase (Phase VII) at pressures higher than about 5 kbar in a lower temperature region. The transition from VI to VII is of the first order being accompanied with a discontinuous change in dielectric constant and thermal hysteresis. Both the pressure-induced phases of VI and VII are shown to be ferroelectric from dielectric hysteresis measurements. Figures 2(a) and (b)



Fig. 2. Ferroelectric hysteresis loops of $(NH_4)_3$ H(SO₄)₂. (a) in Phase VI (-51.4°C, 5.7 kbar), (b) in Phase VII (-86.6°C, 5.6 kbar). Frequency: 50 Hz.





show the ferroelectric hysteresis loops at 50 Hz in Phase VI and in Phase VII, respectively. Figure 3 indicates the pressure-temperature phase diagram of $(NH_4)_3H(SO_4)_2$ given by the dielectric-constant measurements. In the figure, the contour lines of equal dielectric constant are indicated for non-ferroelectric phases of II and III. The maximum positions of the broad dielectric constant peak in constant-pressure measurements are indicated by solid circles. It is noted from Fig. 3 that a ridge of dielectric constant lies almost parallel to the II–III phase boundary.

The present results show that the broad dielectric peak around -25° C at 1 atm corresponds to the polarization-fluctuation which brings about the ferroelectric structure of Phase VI at high pressures. The pressure-dependence of the dielectric constant vs temperature curves shown in Fig. 1 quite resembles to the content-dependence of them in the ammonium Rochelle salt-Rochelle salt system.³⁾ Although in the present compound the appearance of Phase VII makes the problem complicated, the ferroelectric process in Phase VI will be analogous to that in Rochelle salt.

Acknowledgement

The present author devotes his hearty thanks to Dr. K. Ozawa for his cooperation in highpressure arrangements.

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