MULTILAYER CAPACITOR DIELECTRICS PRODUCED FROM
METALLO-ORGANIC PRECURSORS

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1. INTRODUCTION

The metallo-organic decomposition (MOD) process for preparing films or powders has a number of advantages over conventional processes. The metallo-organic compounds used in the present study had oxygen as the hetero atom to bond a metal atom to an organic ligand. These compounds were each dissolved in an appropriate solvent and the solutions mixed to achieve the desired stoichiometry. After adjusting the rheology, the formulation was deposited on an appropriate substrate to make a film or contained in a platinum dish to make a powder, and pyrolyzed to yield the inorganic product. Since the mixing of the starting materials is on the molecular level, the inorganic species exist as atoms or molecules in intimate contact immediately after decomposition. This leads to much more rapid formation of compounds and sintering of films, which translates to lower temperature processing of equilibrium phases and more dense films.

This report is divided into three parts. The first details studies of the formation of PbTiO$_3$ films by the MOD process, and demonstrates that defect free films 0.5 to 2.0 µm thick can be prepared over 2 cm X 2 cm areas. The second part of the report describes a study of grain size effects on the dielectric properties of BaTiO$_3$. The use of the MOD process gave 45 nm powders which were pressed and sintered to give ceramic bodies with controlled grain size. The third part of the report discusses preliminary studies of the formation of lead magnesium niobate by the MOD process.
2. PREPARATION AND PROPERTIES OF PbTiO$_3$ FILMS

2.1 General

Lead titanate (PbTiO$_3$) is one of the most polar ferroelectrics with a Curie point of 490°C [1]. However, very little is known about the intrinsic properties due to the difficulty in growing single crystals of appreciable size. PbTiO$_3$ single crystals grown by the aid of KF flux [2] showed a tetragonal lattice with $c = 0.4150$ nm and $a = 0.3904$ nm. The spontaneous polarization is high at room temperature; some authors suggest the value of $P_s$ to be greater than 50 µC/cm$^2$ [2,3]. The relative permittivity in an a.c. field of 10 kHz is about 10000 in the neighborhood of the Curie temperature [2]. Modified PbTiO$_3$ ceramics have been investigated extensively for high-temperature and high-frequency applications [4-6]. The excellent piezoelectric properties of PbTiO$_3$ ceramics, due to their very small aging rate of dielectric constant [7] and large anisotropy in piezoelectric constant between longitudinal and transverse modes, have been utilized in practical electronic device applications [8-10].

The preparation of PbTiO$_3$ thin films provides a unique opportunity to control either the composition or the structure in such a manner as to give useful non-bulk properties to the resultant film. The principal useful property is the dimension of the structure perpendicular to the substrate. Thin films of amorphous PbTiO$_3$ prepared by RF sputtering on cooled substrates [11] contained Pb-metal crystallites that showed a tendency to increase toward the surface and caused a high electrical
conductivity. Another problem with RF sputtering is that it was found to be difficult to control the composition of the film [12]. The surfaces of films produced by physical vapor deposition [13] were much smoother than those prepared by RF sputtering.

The metallo-organic decomposition (MOD) process used to grow PbTiO$_3$ thin films discussed in this section has a number of advantages over conventional processes. Different types of MOD films have been discussed in various papers and technical reports by the Vests [14-18]. The metallo-organic compounds used in the present study had oxygen as the hetero atom bridge between a metal atom and an organic ligand. These compounds were each dissolved in an appropriate solvent and the solutions mixed to achieve the desired stoichiometry. The homogeneous solutions were spun onto the substrates to form uniform wet films, which were then heated to produce the desired inorganic phase. Since the mixing of the starting materials is on the molecular level in the solution, the inorganic species exist as atoms or molecules in intimate contact immediately after decomposition, which leads to rapid formation of compounds in films and lower temperature processing of equilibrium phases. Thus, the MOD technique has great potential of overcoming the difficulties encountered with other techniques for film formation.

2.2 Experimental

2.2.1 Materials Selection and Synthesis
2.2.1.1 Selection of Compounds

Lead neodecanoate \([\text{Pb}(\text{C}_9\text{H}_{18})_2]\) and titanium di-methoxy-di-neodecanoate \([(\text{CH}_3\text{O})_2\text{Ti}(\text{C}_9\text{H}_{18}\text{COO})_2]\) were the precursors used for \(\text{PbTiO}_3\). Platinum amine 2-ethylhexanoate \([\text{Pt}(\text{C}_7\text{H}_{15}\text{N})_2(\text{C}_9\text{H}_{18}\text{COO})_2]\) was chosen to produce a thin Pt film on Si-wafer substrates. To increase the adhesion between the Pt film and the substrate, a small amount of bismuth was added to the platinum amine 2-ethylhexanoate solution in the form of bismuth 2-ethylhexanoate \([\text{Bi}(\text{C}_7\text{H}_{15}\text{COO})_2]\). The above choices of metallo-organic compounds for the MOC process were based on the following requirements: high metal content; high solubility in common organic solvents; thermally decompose without melting or evaporating; stable under ambient condition (e.g., do not gel); and compatible with other compounds in the formulation. The specific compounds selected were a compromise among these various requirements. All the metallo-organic compounds used in this study were synthesized in house for better control over experimental variables.

2.2.1.2 Titanium Di-Methoxy-Di-Neodecanoate

Eighty six grams (0.5 moles) of titanium methoxide was transferred to a 1000 ml flask under an inert atmosphere. Two hundred ml (1 mole) of 95% pure neodecanoate acid and about 20 ml of pure methanol were added to the flask without disturbing the inert atmosphere. The reaction mixture was refluxed under the inert atmosphere for about three hours at 65°C. A pale yellow viscous liquid was obtained. Methanol was
removed by vacuum distillation, and the liquid was dissolved in xylene.

Reaction:

\[(\text{CH}_3\text{O})_4\text{Ti} + 2\text{C}_9\text{H}_{19}\text{COOH} \rightarrow (\text{CH}_3\text{O})_2\text{Ti(C}_9\text{H}_{19}\text{COO})_2 + 2\text{CH}_3\text{OH}\]

2.2.1.3 Lead Neodecanoate

Two hundred and forty grams (0.73 moles) of \(\text{Pb(NO}_3\text{)}_2\) were dissolved in 700 ml of water. In a 2000 ml beaker, 280 ml (1.4 moles) of neodecanoic acid (95% purity) was neutralized with 87 ml of \(\text{NH}_4\text{OH}\) \((\text{NH}_3\) assay: 30%). After stirring the solution for 20-25 minutes, the \(\text{Pb(NO}_3\text{)}_2\) solution was added with constant stirring. A pale yellow, gummy lead neodecanoate soap settled to the bottom of the flask. It was separated from the aqueous solution and dissolved in xylene. Reactions:

\[\text{NH}_4\text{OH} + \text{C}_9\text{H}_{19}\text{COOH} \rightarrow \text{C}_9\text{H}_{19}\text{COONH}_4 + \text{H}_2\text{O}\]

\[2\text{C}_9\text{H}_{19}\text{COONH}_4 + \text{Pb(NO}_3\text{)}_2 \rightarrow 2\text{NH}_4\text{NO}_3 + \text{Pb(C}_9\text{H}_{19}\text{COO})_2\]

2.2.1.4 Platinum Amine 2-Ethylhexanoate

In a round bottomed flask, 1.54 g (0.0029 moles) of \(\text{H}_2\text{PtCl}_6 \cdot 6\text{H}_2\text{O}\) was dissolved in absolute methanol (8 ml) and cooled in an ice-bath (0-5°C). To this, 1.28 grams (0.0074 moles) of 1,2-diammino propane was added dropwise and stirred for 0.5 hours. The precipitate was washed with methanol and then dissolved in deionized water. An ammonium salt
of 2-ethylhexanoate acid was prepared by mixing 2.7 grams (0.019 moles) of 2-ethylhexanoic acid and 1.2 ml (0.019 moles) of 50% ammonium hydroxide in a 100 ml flask. With continuous stirring, the aqueous platinum amine solution was added dropwise at room temperature, and this mixture was stirred for 2 hours. A yellow oily portion at the bottom was separated and extracted in xylene (~10 ml). Reactions:

\[
\begin{align*}
H_2PtCl_6 \cdot 6H_2O + 4C_3H_7N_2 &\rightarrow Pt(C_3H_7N_2)_4Cl_4 + 2C_3H_7N_2 \cdot HCl + 6H_2O \\
C_7H_{15}COOH + NH_4OH &\rightarrow C_7H_{15}COONH_4 + H_2O \\
Pt(C_3H_7N_2)_2Cl_4 + 4C_7H_{15}COONH_4 + 4NH_4Cl + Pt(C_3H_7N_2)_2(C_7H_{15}COO)_4
\end{align*}
\]

2.2.1.5 Bismuth 2-Ethylhexanoate

A mixture of 6.4 ml (0.004 moles) of 2 ethylhexanoic acid and 4 ml (0.06 moles) of 58% ammonium hydroxide in 25 ml of water was stirred. Five grams (0.01 moles) of Bi(NO\(_3\))\(_3\) \cdot 5H\(_2\)O in 10% NH\(_4\)OH was added slowly to the solution. The reacted oily portion was separated and dissolved in xylene. Reactions:

\[
\begin{align*}
NH_4OH + C_7H_{15}COOH &\rightarrow C_7H_{15}COONH_4 + H_2O \\
Bi(NO_3)_3 \cdot 5H_2O + 3C_7H_{15}COONH_4 &\rightarrow Bi(C_7H_{15}COO)_3 + 3NH_4NO_3 + H_2O
\end{align*}
\]
2.2.2 Substrates

Two kinds of substrates were used in this study. Pure Pt foil was used mainly to study the crystal structure and phase transformations for the PbTiO$_3$ films for various experimental variables since it will limit the chemical impurity effects. However, the difference of thermal expansion between Pt and polycrystalline PbTiO$_3$ ($\alpha = 8.9 \times 10^{-6}/^\circ\text{C}$ and $-3.3 \times 10^{-6}/^\circ\text{C}$ for Pt and PbTiO$_3$, respectively [19]) made it difficult to form even a small area of crack-free film for studies of the electrical properties. Multilayer-Pt-coated Si wafers (p type, 111 orientation) were also used as substrates because the improved thermal expansion match between PbTiO$_3$ and Si ($\alpha = 2.6 \times 10^{-6}$ at room temperature [19]) made it possible to form nearly crack-free PbTiO$_3$ films 2 cm $\times$ 2 cm.

2.2.3 Control of Film Thickness

The thickness of single layer PbTiO$_3$ films on the substrates was found to have considerable effect on features such as crystal structure, grain size, relative amount of residual amorphous phase and the surface roughness of the fired films. In addition, the mismatch of thermal expansion between film and substrate also requires careful control of film thickness in order to get crack-free films. For this study, the conventional spinning technique was used to form a uniform wet film on the substrate. The spinning was carried out in a Class 100 hood in order to minimize dust particle streaks in the films. The film
thickness was established by controlling the spinning speed and time, and the viscosity of the solution of metallo-organic compounds. Figure 2.1 illustrates the effect of spinning speed, and Fig. 2.2 shows the effect of the concentration of the metallo-organic compounds in solution on both film thickness and viscosity of the solution.

2.2.4 Firing

All samples were fired in a fused quartz muffle furnace with an air atmosphere. A programmable control unit was used to establish the time-temperature profile. Some of the samples were fired in a D.C. electrical field from 2.5 kV/cm to 8.5 kV/cm, which was established by two parallel stainless steel plates. Generally, the heating rate and the maximum firing temperature were found to effect the relative amount of amorphous PbTiO$_3$ and the formation of undesired pyrochlore phase. The thermal analysis of Pb(C$_9$H$_{19}$OO)$_2$ and Ti(CH$_3$O)$_2$(C$_9$H$_{19}$COO)$_2$ at a heating rate of 10°C/minute showed that the reactions

$$\text{Pb(C}_9\text{H}_{19}\text{COO)}_2 \rightarrow \text{PbO}$$

and

$$\text{Ti(CH}_3\text{O)}_2(\text{C}_9\text{H}_{19}\text{COO)}_2 \rightarrow \text{TiO}_2$$

were both complete at almost the same point: 370°C ± 2°C. Therefore, the majority of the studies of the effects of different time-temperature profiles were carried out below 370°C.
Figure 2.1 The dependence of single layer thickness on spinning speed.

Figure 2.2 The dependence of single layer thickness and viscosity of the solution of metallo-organic compounds on the concentration of the solution.
2.2.5 Electrodes

The electrode between the \( \text{PbTiO}_3 \) film and the Si wafer had to be a continuous film with good electrically conductive, low surface roughness and good adhesion to the substrate. To achieve these requirements, a multi-layer MOD technique was used to form four-layer Pt films on the Si wafers. It was determined that the desired surface finish could only be achieved if each layer produced a fired thickness of \( \approx 12.5 \) nm. Platinum amine 2-ethylhexanoate \( \text{Pt}(\text{C}_3\text{H}_{10}\text{N}_2)_2 \left( \text{C}_7\text{H}_{15}\text{COO} \right)_4 \) was dissolved in tetrahydrofuran \( \text{CH}_2(\text{CH}_2)_2\text{CH}_2\text{O} \) to form a solution containing 0.8 wt% Pt. The solution used to form the first layer on the Si wafer also contained Bi 2-ethylhexanoate in an amount such that the composition of the fired films was 5 wt% \( \text{Bi}_2\text{O}_3 \) and 95 wt% Pt. The geometry of a four-layer Pt film with total thickness 50 nm on a Si wafer is illustrated in Fig. 2.3(a). The sheet resistance measured by the Van Der Pauw method was 12 \( \Omega/\text{sq} \). To study the electrical properties of the \( \text{PbTiO}_3 \) films, planar capacitors were formed by vacuum evaporation of gold through a mask having holes of 1 mm in diameter. The planar capacitor structure is shown in Fig. 2.3(b).

2.2.6 Crystal Structure and Microstructure

X-ray diffraction analyses using CuK\( _x \) radiation were used to study phase formation and transformations, grain size, lattice parameters, and the preferred orientation of crystal growth. The microstructure of the films was studied using SEM and TEM.
Figure 2.3 (a) The structure of the multilayer Pt film. (b) The planar capacitor structure.
2.2.7 Electrical Property Measurements

The temperature dependences of dielectric constant, dissipation factor and D.C. resistivity for PbTiO$_3$ films were studied from room temperature to 500°C. The dielectric constant and dissipation factor were measured using an LCR meter (HP 4262) at a frequency of 1 kHz and field strength 5 kV/cm to 12 kV/cm depending on the film thickness. The D.C. resistivity was measured using a Keithley 173 A multimeter, which has a useful range to 300 MΩ. At temperatures above 500°C, the high conductivity of the samples made it difficult to measure dielectric constant and dissipation factor. The ferroelectric hysteresis loop was displayed on an oscilloscope using a modified Sawyer and Tower circuit [20].

2.3 Results and Discussion

2.3.1 Morphology of Films

The surface roughness of films formed using the same time-temperature profile was found to be directly related to the single layer thickness. Figure 2.4 shows the surface profiles for two single layer samples with thickness of 123 and 365 nm, respectively. The surface roughness is ~4 nm for the thinner sample and ~8 nm for the thicker sample (the Si wafer substrate had ~1 nm roughness). These surface finishes are much improved compared to films prepared by RF sputtering [11,12]. Figure 2.5 shows SEM pictures for the same samples. The film 123 nm thick (Fig. 2.5(a)) is a crack-free uniform film. In contrast,
Figure 2.4 The surface profile for single layer PbTiO$_3$ film with identical thermal processing with thickness (a) 123 nm and (b) 365 nm.
there are severe cracks on the surface of the film 365 nm thick (Fig. 2.5(b)). The cracking is the result of the large volume change in going from the wet film to the fired film. This volume change may be as much as a factor of 30 depending on the concentration of the metallo-organic compounds in solution. Thicker films require a much slower heating rate through the critical temperature ranges. For a given time-temperature profile, the film must be below some critical thickness in order to sustain the volume change without cracking. Figure 2.6 shows the cross-section of a typical structure of five layer PbTiO₃ deposited on a four layer Pt coated Si wafer. The very dense PbTiO₃ film produced by the MOD technique is apparent.

2.3.2 Metastable Phase in the PbO-TiO₂ System

A metastable phase was observed in PbTiO₃ films (1 μm thick) deposited on pure Pt foil. Figure 2.7 shows the x-ray diffraction patterns for the films at various firing temperatures. For films fired at 494°C or lower, the x-ray diffraction patterns showed that the tetragonal phase was the only crystal structure formed (of course there existed a certain amount of amorphous PbTiO₃). As the firing temperature was raised to 514°C, a new crystal structure with 2θ = 30.40°, 35.20° and 50.64° was formed. The relative intensities of those peaks compared to the tetragonal peaks reached a maximum when the film was fired at 540°C - 550°C. Further increasing the firing temperature decreased the relative intensity of the peaks of this metastable phase. Samples fired at 630°C or higher showed no metastable phase. The lattice parameter and the (h,
Figure 2.6 Micrograph (TEM) of the cross-section of substrate, Pt and PtTiO$_3$ films. (90,000 x).
Firing Temperature

631

601

583

569

536

514

494

475

435

$\rightarrow 2\theta$

Figure 2.7 X-ray diffraction patterns (with CuK$_{\alpha}$) for films (1 μm) fired on Pt foil at various temperatures. The indicated peaks are not characteristic of tetragonal PbTiO$_3$. 
1) values for \( \theta = 20.40^\circ \), \( 35.20^\circ \) and \( 50.64^\circ \) were calculated assuming that the metastable phase had the crystal symmetry \( \text{Fd}3\text{m} \) as did the pyrochlore structure observed in the \( \text{PbO} \)-\( \text{TiO}_2 \)-\( \text{SiO}_2 \) system [21]. Each unit cell of such a structure contains eight formulae of \( \text{Pb}_2\text{Ti}_2\text{O}_6 \). The position of Pb ions are fixed in the lattice of \( \text{d} \) subgroup and Ti ions in \( \text{c} \) subgroup [22]. For x-ray diffraction, the smallest value of \( h^2+k^2+l^2 \) is when \( h=k=l=2 \). Thus, it is confirmed that the smallest value of \( \theta \) (30.40\(^\circ\)) is the reflection from the \((2,2,2)\) plane. Thus, the lattice parameter \( a \) can be calculated as:

\[
d_{2,2,2} = \frac{\lambda}{2 \sin \theta} = 0.24902 \text{ nm}
\]

\[
a = \frac{d_{2,2,2}}{\sqrt{h^2+k^2+l^2}} = 1.019 \text{ nm}
\]

The \( h,k,l \) values for \( \theta = 35.20^\circ \) and \( 50.64^\circ \) were calculated by

\[
h^2+k^2+l^2 = \left( \frac{a}{d_{h,k,l}} \right)^2
\]

and it was found that \( h^2+k^2+l^2 = 16 \) and \( 32 \) for \( \theta = 35.20^\circ \) and \( 50.64^\circ \), respectively. Considering the \( \text{c,d} \) subgroup's limitations, the peaks at \( \theta = 35.20^\circ \) and \( 50.64^\circ \) must be reflections from \((4,0,0)\) and \((4,2,2)\) planes, respectively.

By the proper choice of firing conditions, the amount of amorphous phase formed in the films could be minimized. X-ray diffraction patterns for samples fired so as to minimize the formation of amorphous phase indicated no metastable phase peaks, as shown in Fig. 2.8. The pyrochlore phase was also never observed if the films were fully
Figure 2.8  X-ray diffraction patterns (with CuKα) for films fired at 560°C with the thermal processing (a) to minimize the formation of amorphous phase, (b) to maximize the formation of amorphous phase. The indicated peaks are not characteristic of tetragonal PbTiO₃.
crystallized from the amorphous phase below 500°C. The metastable phro-
chlore phase of Pb₂TiO₆ can be formed only from amorphous PbTiO₃, and
the tetragonal phase can not be transformed into the cubic phase at any
temperature. The phase transformation diagram for the PbTiO₃ system is
shown in Fig. 2.9.

2.3.3 Structure of Tetragonal PbTiO₃ Films

The axial ratio (c/a) for bulk tetragonal PbTiO₃ varies from 1.065
at 25°C to 1.02 at the Curie temperature (490°C) [23]. It was found
that the c/a values for films fired at the same temperature changed with
single layer thickness. Figure 2.10 shows that this ratio increases as
the layer thickness increases, but in all cases was smaller than that of
bulk tetragonal PbTiO₃.

For the same thickness films fired at the same temperature, it was
found the c/a values for those films fired in a D.C. electric field at
temperatures below T were smaller compared to those fired without a
field, and the effect increased with increasing field strength. Figure
2.11 shows the experimental results. The field strength dependence of
c/a value can be explained if it is assumed that the dipole moment of
the tetragonal PbTiO₃ unit cell is proportional to its axial ratio c/a.
In an electrical field E, the energy density stored in a dielectric can
be expressed as 1/2 PE, where P is the polarization or the dipole moment
per unit volume. The smaller c/a will reduce the energy stored in the
system thus giving a more stable state at thermal equilibrium. The
phenomenon observed presents an opportunity to control the value of c/a,
Figure 2.9 Phase transformation diagram for the PbTiO$_3$ system prepared by the MOD process.
Firing Temperature 552 °C
Firing Time: 8 hrs.

Figure 2.10 The dependence of c/a on the single layer thickness.
Figure 2.11 The dependence of \( c/a \) on C.D. electrical field strength during firing for the film with 2.42 kÅ in single layer thickness.
Figure 2.12 The grain size dependence of PbTiO₃ films on firing temperature.
Firing temperature: 552°C

Figure 2.13 The grain size changes with single layer thickness for the films fired at 552°C.
In general, the polycrystalline PbTiO$_3$ films formed by the MOO technique showed a random orientation of grains. However, as the firing temperature was increased, the films indicated a preferred growth direction with the c axis (001) perpendicular to the substrate surface. As shown in Fig. 2.7, the peak intensity ratio of (001) to (100) is about 1:2 for the films fired at 500°C, which means that the c axes of the grains in the film were randomly distributed. This ratio increased as the firing temperature increased and reached 1.2:1 for the films fired at 630°C. The increase in this ratio means more grains direct their c axis perpendicular to the substrate. Further efforts to grow films in which all of the grains direct their c axis perpendicular to substrate are currently in process.

2.3.5 Electrical Properties

2.3.5.1 Dielectric Constant and Dissipation Factor

The room temperature dielectric constant of the planar capacitor structures (Fig. 2.3(b)) was found to be a strong function of relative humidity, as shown in Fig. 2.14. The origin of this effect is unknown, but its magnitude is impressive. Subsequent data were collected above 50°C were the effect of relative humidity is negligible. Figure 2.15 shows the temperature dependence of the dielectric constant and dissipation factor in the range 50°C to 500°C for a tetragonal PbTiO$_3$ film 1.83 μm thick (six layers each 305 nm thick). A dielectric constant was 100 - 110 at temperatures below 300°C, and reached its maximum value of 3 X 10$^4$ at the Curie point. The measured Curie temperature was 493 to
Figure 2.14 The effect of relative humidity on the room temperature dielectric constant of PbTiO$_3$ films.
Figure 2.15 The temperature dependence of dielectric constant and dissipation factor for a tetragonal PbTiO$_3$ film 1.33 μm thick.
495°C. The dissipation factor remained almost constant until 250°C and then increased rapidly with further increase in temperature. There was a relative minimum value of dissipation factor around the Curie temperature. The rapid increase of conductivity a few degrees above the Curie temperature made it difficult to measure the dielectric constant and dissipation factor at higher temperatures. Figure 2.16 shows the dielectric constant and dissipation factor changes with temperature for a glass-ceramic (tetragonal plus amorphous PbTiO₃) film with six layers, each 305 nm thick. A pseudo-Curie point was observed at 435°C, with K value equal to 2 X 10⁴. Similar to the results shown in Fig. 2.15, the dissipation factor had its relative minimum value at the pseudo-Curie temperature.

Figure 2.17 shows the temperature dependence of dielectric constant and dissipation factor for two samples both having the same thickness and crystal structure (fired with exactly the same temperature profile), but with one of them fired at an applied D.C. field of 8.5 kV/cm. As discussed in Section 2.3.3, the c/a value for the sample fired in the D.C. field will be smaller than that of the one fired without the field. Figure 2.17 indicates that the c/a value has no considerable effect on the dissipation factor in the range 50°C - 450°C, but it does effect the temperature dependence of the dielectric constant. The c/a values of the samples whose temperature dependence of dielectric constant are shown in Fig. 2.17 are 1.057 and 1.055, and it can be seen from Fig. 2.18 [23] that the film with c/a value of 1.055 corresponds to a temperature 30 to 40°C higher than the one with c/a value of 1.057. If the dielectric constant-temperature curve for the sample with c/a = 1.055 in
Figure 2.16 The temperature dependence of dielectric constant and dissipation factor for the film composed of tetraconal polycrystalline and amorphous phases.
Figure 2.17 The temperature dependence of dielectric constant and dissipation factor for films having some crystal structure but with different values of c/a.
Figure 2.18  Axial ratio of PbTiO₃ as a function.
Figure 2.17 is moved to right about 40°C, the two curves will match very well over the whole temperature range.

2.3.5.2 D.C. Resistivity

Figure 2.19 shows the temperature dependence of D.C. resistivity for one of polycrystalline films. No effect of film thickness on D.C. resistivity was found. The $\rho - 1/T$ relation is a combination of two almost-straight lines. The change of slope near 250°C indicates a change in conductive mechanism. The room temperature resistivity was $1.0 - 1.2 \times 10^{10} \ \Omega \ cm$, which is comparable to that given in Ref. [25].

2.3.5.3 Spontaneous Polarization

The P-E hysteresis loops were observed for the PbTiO$_3$ films deposited on Pt coating Si wafers, and an example is shown in Fig. 2.20. The spontaneous polarization and remanent polarization under an applied field of 100 kV/cm were about 5.1 $\mu$C/cm$^2$ and 3.82 $\mu$C/cm$^2$, respectively. In general, these values were found to be independent of thickness for films with the same phase composition. The coercive field was 33.2 kV/cm. Increasing the applied field beyond 100 kV/cm resulted in the destruction of the top electrodes (vacuum deposited gold films about 100 nm thick). The dielectric strength for the films grown from the metallo-organic precursors was definitely larger than 100 kV/cm.
Figure 2.19: The temperature dependence of D.C. resistivity.
Figure 2.20 Hysteresis loop for PbTiO$_3$ films under an applied field of 100 KV/cm (x: 100 KV/cm/div., y: 1.20 µC/cm$^2$/div.).
2.3.6 Reliability Studies

One of the objectives of this study was to grow thin dielectric films from metallo-organic precursors that were reliable films over an appreciable area. The reliability for the specimens with thickness 0.5 μm to 2 μm was defined as the percent of the planer capacitors (see Fig. 2.1), which gave equivalent dielectric properties with an applied field of 100 kV/cm. The reliability of the specimens was found to depend on many factors such as: the choice of substrate; the purity of the metallo-organic precursors; the process cleanliness; and the time-temperature firing profile. For the films made by the multi-layer spinning technique, the reliability was also found to depend on the single layer thickness; Fig. 2.22 shows this relation. Currently, 2 cm X 2 cm dielectric films 0.5 μ to 2.0 μ thick can be made from metallo-organic precursors with 100% reliability using the multilayer technique.

2.4. Summary

Ferroelectric PbTiO$_3$ films with thickness 0.5 μ to 2.0 μ were prepared by the MOD process using a multilayer spinning technique. The processing can yield highly reliable films with easy control of crystal structure and composition at a low firing temperature. Dielectric constants and dissipation factors were measured from 50°C to 500°C, and the dielectric constant reached $3 \times 10^4$ at the Curie temperature of 493-495°C. The temperature dependence of dielectric constant was found to
Figure 2.21 An illustration of planer capacitors made by PbTiO$_3$ film.
Figure 2.22 The reliability dependence on single layer thickness of films fired at the same temperature profiles.

Firing Temperature: 552°C

Reliability (%) vs. Single Layer Thickness (KÅ)
be a function of the c/a value, which could be modified by either control of the single layer thickness or the strength of an applied D.C. field during film preparation. The P-E hysteresis loops were measured, and the spontaneous polarization and remanent polarization under an applied field of 100 kV/cm were found to be 5.1 μC/cm\(^2\) and 3.29 μC/cm\(^2\), respectively. The dielectric strength for 0.5 - 2.0 μ thick films was larger than 100 kV/cm.

The value of c/a for tetragonal polycrystalline films in the range of single layer thickness 75-425 nm was found to be smaller than that of bulk materials. A metastable pyrochlore structure, which formed from amorphous PbTiO\(_3\), was observed in the temperature range from 500 - 550°C. At lower firing temperatures (<550°C), the grain orientation in polycrystalline tetragonal films was random. For firing temperatures higher than 600°C, a strong tendency for preferred orientation with the c axis perpendicular to the substrate surface was observed.

2.5. References


3. GRAIN SIZE EFFECTS IN BaTiO₃

3.1 General

The room temperature relative permittivities of a single crystal of BaTiO₃ are 3400 along the a axis and 150 along the c axis [1]. However, Kniepkamp and Heywang [2] reported a relative permittivity value of 3500 for a polycrystalline, 1 μm grain size BaTiO₃. Many other researchers have also reported similar or higher values [3-9]. Keizer and Burggraaf [10] observed similar effects of grain size on the permittivity and the Curie temperature of lanthana substituted lead titanate.

Very high permittivity values observed in fine grained ceramic cannot be explained by an averaging procedure. Various explanations have been offered for the grain size dependence of dielectric constant [11-16]. However, none of these theories can explain the sharp decrease of dielectric constant below a certain grain size. The sharp decrease of dielectric constant with decrease in grain size, below 10 nm, was explained by V.P. Dudkevich et al. [17] and Srinivasan et al. [18] using
the Lyddane-Sachs-Teller relationship. However, this effect is insignificant for the grain sizes higher than 10 nm. The dielectric constant of powdered BaTiO$_3$ was observed to be very low by various researchers [19-21]. Goswami [20] attributed the lower dielectric constant to a surface defect layer.

3.2 Experimental

Barium titanate powder was obtained by decomposition of metallo-organic compounds using the procedure outlined elsewhere[22]. Approximately 0.15 g of BaTiO$_3$ powder was pressed at about 550 MPa to obtain disk shaped pellets (diameter ~ 6 mm, thickness ~ 1.25 mm). The pellets were placed on a bed of powder of the same composition in a platinum crucible and fired at temperatures ranging from 1050$^\circ$C to 1300$^\circ$C, for 5 to 30 minutes. The grain sizes of the thermally etched samples were determined using SEM. The grain sizes and densities of such samples are listed in Table 3.1. The sintered pellets were painted with a Pd-Ag thick film conductor composition $^+$, dried at 200$^\circ$C and fired at 850$^\circ$C for about 15 minutes. Thin gauge copper wires were soldered to both sides of the samples. To avoid the effects of condensed moisture, the samples were sprayed with a protective coating $^{++}$. Anomalous electrical properties were observed at the dew point if the coating was not used.

$^+$ JEM/200CX JOEL Inc., Peabody, MA
$^{++}$ No. 6120, DuPont Inc., Niagara Falls, N.Y.
$^{+++}$ Crystal Clear, No. 1302, Borden Inc., Columbus, OH.
Table 3.1 Density and grain size of BaTiO$_3$ ceramics.

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Temperature ($^\circ$C)</th>
<th>Grain size ($\mu$m)</th>
<th>Density (% theoretical)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>1050</td>
<td>0.3</td>
<td>58</td>
</tr>
<tr>
<td>30</td>
<td>1100</td>
<td>0.35</td>
<td>62</td>
</tr>
<tr>
<td>30</td>
<td>1150</td>
<td>0.4</td>
<td>64</td>
</tr>
<tr>
<td>30</td>
<td>1200</td>
<td>0.5</td>
<td>66</td>
</tr>
<tr>
<td>30</td>
<td>1225</td>
<td>2.8</td>
<td>78</td>
</tr>
<tr>
<td>05</td>
<td>1300</td>
<td>9.6</td>
<td>74</td>
</tr>
<tr>
<td>30</td>
<td>1300</td>
<td>15.5</td>
<td>79</td>
</tr>
</tbody>
</table>
The dielectric properties of the samples were measured as a function of temperature. An environmental chamber was used to vary the temperature of the sample in air from -70°C to +160°C. The temperature of the specimens was measured with an alumel-chromel thermocouple.

The dielectric constant and the dissipation factor of the specimens were measured using a digital LCR meter. All the dielectric properties were measured at 1 kHz test frequency, and using zero D.C. bias and 1 Vrms test signal. Spontaneous polarization measurements were carried out using the hysteresis loop bridge circuit suggested by Diamant et al. [23].

3.3 Results and Discussion

3.3.1 Experimental Results

Since BaTiO$_3$ can be produced at considerably lower temperature by the MOD process, very fine particle size BaTiO$_3$ can be obtained. The crystallite size of BaTiO$_3$ powder obtained at 850°C was 0.045 μm. This powder was pressed into pellets and sintered under the conditions listed in Table 3.1. The microstructure of the sintered pellets was studied using the SEM. All the samples fired below 1300°C showed a relatively uniform grain size, and the average grain size of each was calculated by Hilliard's method [24]. The samples fired at 1300°C had a bimodal grain size distribution, and the average grain size was calculated using arithmetic average procedures. The porosity of the samples ranged from 0.045 to 0.05.

# No. 4262A Hewlett Packard Inc., Palo Alto, CA.

++++ Model SK 2101, Associated Testing Laboratories Inc., Wayne, NJ.
56 to 80% as determined by geometry. Since the samples were not 100% dense, the measured dielectric constants were corrected for porosity. Numerous formulae have been suggested for the dielectric constant of a heterogeneous system in terms of the dielectric constant and volume fraction of the individual phases [25]. In the present case, the dielectric constants were corrected for porosity using effective medium theory [26] for coordination number 6. Figure 3.1 shows the temperature dependence of the corrected dielectric constant of the various samples with differing grain sizes as a function of temperature. As the grain size decreased, the cubic-tetragonal transition temperature decreased by about 2-4°C and the tetragonal-orthorhombic transformation temperature increased considerably. To avoid the effect of transition temperature on the isothermal dielectric constant, the dielectric constant at 70°C was plotted as a function of grain size as shown in Fig. 3.2. As the grain size of BaTiO₃ increased, the dielectric constant increased, reached a maximum at about 0.4 μm and then decreased with further increase in grain size. A similar behavior was observed by Brandmayr [7], K. Kinoshita and A. Yamaji [9] and G. Arlt et al. [8]. However, Brandmayr and K. Kinoshita and A. Yamaji observed the maxima at about 1.0 μm and Arlt et al. at 0.7 μm.

3.3.2. Model

A maximum in the dielectric constant indicates that there are opposing factors which influence the grain size dependence of the dielectric constant. Grain boundaries are amorphous in nature, and hence have a lower dielectric constant than the grains. Since the
Figure 3.1 Variation of dielectric constant of BaTiO₃ ceramics of varying grain sizes with temperature.
Figure 3.2 A comparison of experimental behavior of dielectric constant with grain size and the one predicted by model.
fraction of grain boundary volume increases with decreasing grain size, the dielectric constant of polycrystalline BaTiO$_3$ should decrease with decreasing grain size. On the contrary, the contribution of internal stresses [11], internal electric field [12] and domain wall area [8], all of which increase the dielectric constant, increase with decreasing grain size.

The model used to calculate the dielectric constant of the system is shown in Fig. 3.3. Figure 3.3 shows an array of cubic grains surrounded by grain boundaries, which can be represented by a series and parallel combination of grain and grain boundary dielectrics. The total dielectric constant of a grain is made up of intrinsic dielectric constant which results from ferroelectric phenomenon and is affected by external factors such as electric field and stresses, and the extrinsic dielectric constant which results from domain wall motion etc. In this model, the following simplifying assumptions were made.

3.3.3 Assumptions

1. Grains are cubic.
2. Grain boundaries have a low dielectric constant.

The dielectric constants of the surface defect layers observed in the single crystals or polycrystalline powders are in the range of 5 to 200 [27-29].
3. Thickness of the grain boundaries follows a relation,
Figure 3.3 A schematic of the proposed model.
where, \( w = \text{thickness}, \ D_0 = \text{constant}, \ Q = \text{activation energy}, \ \text{gas constant} \)

\( t = \text{sintering time} \) and \( T = \text{sintering temperature} \)

This assumes that the boundary thickness is diffusion controlled and is determined by the sintering conditions.

4. The grains having grain size smaller than 0.4 \( \mu\text{m} \) are single domain grains. This assumption is made based on Arlt et al.'s [8] study.

5. The transformation stresses are experienced only by single domain grains. The stresses caused by the cubic-tetragonal transformation are completely relieved by formation of 90° domain in multidomain grains.

6. The transformation stresses are constant with respect to grain size, for grain size below 0.4 \( \mu\text{m} \).

7. There is no depolarizing electric field.

8. The dielectric constants of the porous bodies are corrected by effective medium theory to obtain the dielectric constants of the dense bodies.
3.3.4. Methodology

The following approach was used to calculate the dielectric constant of polycrystalline material; cgs units are used, as they are more convenient. The dielectric constants of the single crystal under a stress system was calculated using Devonshire's theory [30-31] as follows.

The elastic Gibb's free energy function can be written in terms of stresses and polarization [11] as follows.

\[
G_1 = -\frac{1}{2} s_{11} (X_x^2 + Y_y^2 + Z_z^2) - s_{12} (X_x Y_y + Y_y Z_z + Z_z X_x)
\]

\[
-\frac{1}{2} s_{44} (X_y^2 + Y_z^2 + Z_x^2) + (Q_{11} X_x + Q_{12} Y_y + Q_{12} Z_z) P_x^2
\]

\[
+ (Q_{12} X_x + Q_{11} Y_y + Q_{12} Z_z) P_y^2 + (Q_{12} X_x + Q_{12} Y_y + Q_{11} Z_z) P_z^2
\]

\[
+ Q_{44} (X_x P_x P_x + Y_y P_y P_y + Z_z P_z P_z) + A (P_x^2 + P_y^2 + P_z^2)
\]

\[
+ B (P_x^4 + P_y^4 + P_z^4) + C (P_x^6 + P_y^6 + P_z^6)
\]

\[
+ D (P_x^2 P_y^2 + P_y^2 P_z^2 + P_z^2 P_x^2)
\]

\[
+ G (P_x^4 P_y^2 + P_y^4 P_z^2 + P_z^4 P_x^2 + P_x^2 P_y^4 + P_y^2 P_z^4 + P_z^2 P_x^4)
\]

(2)

where \(X_x, Y_y, Z_z\) are the normal stress components,

\(Y_y, Z_z, X_x\) are the shear stress components,

\(S_{11}, S_{12}, S_{44}\) are the elastic moduli,
\( P_x, P_y, P_z \) are the components of polarization,
\( Q_{11}, Q_{12}, Q_{44} \) are the electrostrictive coefficients,
A, B, C, D and G are the coefficients of the energy functions.

The coefficients A, B, C, D and G are given by the following expressions [32].

\[
A = 3.7 \times 10^{-5} (T - \theta)
\]

where \( T \) is the experimental temperature and \( \theta \) is the temperature at which the inverse susceptibility is extrapolated to zero. The present inverse susceptibility studies on the coarse grained \( \text{BaTiO}_3 \) gave the value of \( \theta \) to be 118\( ^\circ \)C.

\[
B = - \left| \frac{\pi/\varepsilon_c + 2A}{2P_s^2} \right|
\]

where \( \varepsilon_c \) is the dielectric constant along the c axis and \( P_s \) is the spontaneous polarization.

\[
C = \frac{\pi/\varepsilon_c + A}{3P_s^4}
\]

\[
D = \frac{1}{P_0^2} \left| A(3\alpha^2 - 4) + B(3\alpha^2 P_T^2 - 2P_0^2) + C(3\alpha^2 P_T^4) \right|
\]

where \( P_0 \) is the polarization in the orthorhombic phase (at the second transformation temperature) = \( 8 \times 10^4 \) cgs units, \( P_T \) is the polarization in the tetragonal phase (at the second transformation temperature) = \( 6.5 \times 10^4 \) cgs units.
and \( \alpha = \frac{P_T}{P_0} = 1.23 \)

\[
G = -\frac{1}{P_0^4} \left| A(u^2 - 1) + B(a^2 P_T^2) + C(a^2 P_T^4 + P_0^4) \right| \quad (7)
\]

Previously [11,32-34], the coefficients A, B, C, D and G were calculated using dielectric constants of flux grown single crystals. Flux grown crystals have F as an impurity, and the Curie temperature of such crystals was 120°C. Melt grown crystals are more pure and have 130 \(+/-\) 2 as the Curie temperature. Since, in the present studies the observed Curie temperatures were in the neighborhood of 128°C, the reported [1] dielectric constants of melt grown crystals were used to calculate the values of the coefficients. Table 3.2 lists the values of the coefficients obtained along with the values of elastic and electrostrictive constants used in the free energy function.

Differentiating Eq.(2) w.r.t. \( P_s \), under the conditions that \( P_1 = P_2 = 0 \) and \( P_3 = P_s \), and under uniform compressive stress \( X \) along the c axis and equal tensile stresses \(-X\) along the a axes, the following expressions for dielectric constants are obtained. According to assumption 5, \( X = 0 \) for grain size \( > 0.4 \ \mu m \).

\[
\frac{4\pi}{\varepsilon_c} = 2(Q_{11}X - 2Q_{12}X) + 2A + 12BP_s^2 + 30CP_s^4 \quad (8)
\]

\[
\frac{4\pi}{\varepsilon_a} = -2Q_{11}X + 2A + 2DP_s^2 + 2GP_s^4 \quad (9)
\]

\[
E_c = 2P_sX(Q_{11} - 2Q_{12}) + 2AP_s + 4BP_s^3 + 6CP_s^5 = 0 \quad (10)
\]
Table 3.2 Values of the coefficients of Devonshire's free energy function at 70°C.

<table>
<thead>
<tr>
<th>Coefficient</th>
<th>Reference</th>
<th>Value in CGS units</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>[51]</td>
<td>$3.7 \times 10^{-5}(T-118)$</td>
</tr>
<tr>
<td>B</td>
<td>[51]</td>
<td>$-1.1972 \times 10^{-22}$</td>
</tr>
<tr>
<td>C</td>
<td>[51]</td>
<td>$1.5502 \times 10^{-22}$</td>
</tr>
<tr>
<td>D</td>
<td>[51]</td>
<td>$6.0 \times 10^{-13}$</td>
</tr>
<tr>
<td>G</td>
<td>[51]</td>
<td>$5.0 \times 10^{-23}$</td>
</tr>
<tr>
<td>Q_{11}</td>
<td>[118]</td>
<td>$1.23 \times 10^{-12}$</td>
</tr>
<tr>
<td>Q_{12}</td>
<td>[118]</td>
<td>$-0.48 \times 10^{-12}$</td>
</tr>
</tbody>
</table>

* Modified using melt grown single crystal data.
Equation (10) can be solved for $P_s$ to give

$$p_s^2 = \frac{-B + \sqrt{B^2 - 3C(X(-2q_{12} + q_{11})+A)}}{3C}$$

(11)

Using the value of $p_s^2$ obtained from Eq. (11) in Eq. (8) and Eq. (9), the values of $\varepsilon_c$ and $\varepsilon_a$ can be calculated at any stress level. The calculated values of dielectric constants $\varepsilon_c$ and $\varepsilon_a$ at stress, $X=0$, were 180 and 1850, respectively. The experimental values of dielectric constants $\varepsilon_c$ and $\varepsilon_a$ of stress free BaTiO$_3$ single crystal, reported by Wemple et al. [1], were 180 and 1850, respectively.

G. Arlt [8] proposed a method of calculating the dielectric constant of a grain containing domains. The dielectric constant tensor for a grain containing domains is given by

$$\begin{vmatrix}
\varepsilon_{11} & 0 & 0 \\
0 & \varepsilon_{22} & 0 \\
0 & 0 & \varepsilon_{33}
\end{vmatrix}$$

(12)

where

$$\varepsilon_{11} = \frac{1}{3} \left[ -16\varepsilon_{c}^4 + 49\varepsilon_{a}^3\varepsilon_{c} + 526\varepsilon_{c}^2 + 405\varepsilon_{a}^2 + 49\varepsilon_{a} \right]$$

(13)

$$\varepsilon_{22} = \frac{1}{3} \left[ 2\varepsilon_{c}^2 + 11\varepsilon_{a} \varepsilon_{c} + 5\varepsilon_{a}^2 \right]$$

(14)
The dielectric constant of the ceramic was calculated using Bruggeman's [35] formula

\[
\varepsilon_{33} = \frac{1}{3} \frac{146\varepsilon_c^3 + 405\varepsilon_c^2\varepsilon_{a} + 324\varepsilon_c\varepsilon_{a}^2 + 81\varepsilon_{a}^3}{9(\varepsilon_c + \varepsilon_{a})(2\varepsilon_c^2 + 11\varepsilon_c\varepsilon_{a} + 5\varepsilon_{a}^2)} \tag{15}
\]

The dielectric constant of the ceramic was calculated using Bruggeman's [35] formula

\[
\varepsilon' = \frac{1}{4} \varepsilon_{22} + \left\{ \frac{2}{\varepsilon_{22} + 8\varepsilon_{11}\varepsilon_{22}} \right\} \tag{16}
\]

where \(\varepsilon_{22}\) in Eq. (16) is taken as a average of \(\varepsilon_{22}\) and \(\varepsilon_{33}\) given by Eq.(14) and (15) respectively. For a single domain grain \(\varepsilon_{11} = \varepsilon_c\) and \(\varepsilon_{22} = \varepsilon_{33} = \varepsilon_a\).

According to the assumptions made above, the total dielectric constant of the ceramic should be the result of the intrinsic dielectric constant, the grain boundary effects and the domain wall contribution for the grain sizes above 0.4 microns. For the grain sizes below 0.4 \(\mu\text{m}\), the grains are single domain grains; hence the domain wall contribution is absent. However, the intrinsic dielectric constant is affected by the residual transformation stresses below 0.4 \(\mu\text{m}\).

The domain wall contribution to the dielectric constant was calculated using G. Arlt's[08] method.

\[
\varepsilon_{dw} \propto \text{domain wall area}(A) \tag{17}
\]

\(\varepsilon_{dw}\) is domain wall contribution to the dielectric constant.

\[
A \propto \frac{1}{d} \tag{18}
\]

d is domain width. Thus,
G. Arlt [8] related the equilibrium domain size\(d\) to grain size\(a\) by the relation

\[
d = \frac{128\pi c a}{C_{11} S_s^2} \tag{20}
\]

where \(c_{11}, S_s\) and \(\sigma\) are the longitudinal elastic constant, spontaneous deformation and domain wall energy respectively. Therefore, the domain wall contribution to the dielectric constant can be expressed in terms of the grain size by:

\[
\varepsilon_{dw} = \frac{k}{d} \tag{21}
\]

Since, at the largest grain size, the internal stresses are completely relieved by formation of 90° domains and the volume fraction of grain boundaries is minimal, the dielectric constant of ceramics of 15.5 μm grain size \(\varepsilon_{15.5}\) was assumed to be the sum of intrinsic and domain wall contributions.

\[
\varepsilon_{15.5} = \varepsilon_r + \varepsilon_{dw} \tag{22}
\]

\(\varepsilon_r\) was calculated using Eq. (16) and subtracted from the measured value of \(\varepsilon_{15.5}\) to obtain the domain wall contribution. Using this, the value of \(K\) was calculated from Eq.(21) to be 2710.

The ceramics sketched in Fig. 3.3 can be represented by two capacitors of capacitance \(C_1\) and \(C_2\) in parallel as shown Fig. 3.3. The capa-
citances $C_1$ and $C_2$ are given by the following expressions

$$C_1 = \frac{\varepsilon_1 \varepsilon_2 (a - w)^2}{\varepsilon_1 (a - w) + \varepsilon_2 w}$$  \hspace{1cm} (23)$$

$$C_2 = \frac{w(2a - w)}{a} \varepsilon_1$$  \hspace{1cm} (24)$$

The total capacitance is the sum of $C_1$ and $C_2$. Thus, the effective dielectric constant is given by

$$\varepsilon = \frac{w(2a - w) \varepsilon_1}{a^2} + \frac{\varepsilon_1 \varepsilon_2 (a - w)^2}{\varepsilon_1 a (a - w) + \varepsilon_2 a w}$$  \hspace{1cm} (25)$$

where $\varepsilon_1$ is dielectric constant of the grain boundary and $\varepsilon_2$ is the dielectric constant of ceramic calculated using Eqs. (8) through (21).

The thickness of the defective layer $w$ is related to the processing conditions through Eq. (1). Using Eqs. (7) through (22) in Eq. (25) leaves the dielectric constant of the grain boundary, $D_0$, $Q$ and stress as the only unknowns in Eq. (25). The values of these parameters obtained by least square fit using all the data are listed in Table 3.3. The value of $Q$ obtained by the least square fit is in good agreement with the value of activation energy obtained in the kinetics of formation of barium titanate studies [22], and the value of the dielectric constant of the grain boundary is in good agreement with the previous studies [27-29]. Using these values, the dielectric constants were calculated at various grain sizes. Figure 3.2 shows the comparison between experimental behavior and the behavior predicted using this model. The experimental observations agree very well with the model.
Table 3.3 Values of the model parameters obtained by least square fit.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dielectric constant of grain boundary</td>
<td>90</td>
</tr>
<tr>
<td>Activation Energy (Q) (kJ/mole)</td>
<td>290</td>
</tr>
<tr>
<td>$D_0$ (cm$^{-2}$ hr$^{-1}$)</td>
<td>10$^{24}$</td>
</tr>
<tr>
<td>Stresses (dynes/cm$^2$)</td>
<td>1.26 X 10$^9$</td>
</tr>
</tbody>
</table>
The spontaneous polarization is linked to the ferroelectric phenomenon, and hence is related to the intrinsic dielectric constant. As grain size decreases, the crystallinity decreases and finally one reaches an amorphous state where ferroelectricity ceases. Thus, it was expected that the spontaneous polarization should decrease with decrease in grain size. Also, the random orientation of grains makes only a component of spontaneous polarization contribute to the measured value. The expected behavior of spontaneous polarization with respect to grain size was observed, as shown in Fig. 3.4. A linear porosity correction was made to the values of spontaneous polarization in Fig. 3.4.

G. Arlt and Sasko [36] studied the domain structure of polycrystalline BaTiO₃. They observed a spatial domain configuration as shown in Fig. 3.5 in unpolarized ceramic, where \( A = [011] \), \( B = [101] \) and \( C = [110] \) planes. The spontaneous polarization of such a configuration is \( P_a = \frac{\sqrt{2} P_s}{3} \), and the vector is directed along [110]. Since the grains are randomly oriented in space, the net polarization will be the spatial average of the \( P_a \) vector.

\[
P_{\text{ceramic}} = \int \int \int \frac{\pi \pi \cos \vartheta \cos \phi \, d\theta d\phi}{2 \pi^2}
\]

Integration yields \( P_{\text{ceramics}} = \frac{4\sqrt{2}}{3\pi} P_s = 0.19 P_s \). The value \( P_s \) for grain size 15.5 is 0.10 (Coul/m²) and that for the single crystal BaTiO₃ is 0.23 (Coul/m²). Thus, the value of the spontaneous polarization observed in the ceramic material is greater than the 0.2P_s for single crystals.
Figure 3.4 Variation of spontaneous polarization of BaTiO$_3$ ceramics of varying grain sizes with temperature.
Figure 3.5 A domain configuration observed in unpolarized BaTiO$_3$ ceramics.
Figure 3.6 Domain configuration observed in polarized BaTiO$_3$ ceramics.
This could be because in polarized ceramics the domain configuration is as shown in Fig. 3.6; this structure has a higher net polarization than the one shown in Fig. 3.5.

3.4 References


32. E. J. Huibregtse, W. H. Bessey and M. E. Dougard, "Electromechanical Behavior of Single Crystals of Barium Titanate for 25°C to
4. PREPARATION OF LEAD MAGNESIUM NIOBATE

4.1 General

Multilayer ceramic capacitors for electronic circuit miniaturization are attractive because of their potentially high capacitance per unit volume, together with their electrical stability, reliability, and low cost. The present material of choice for capacitors is mainly multilayer tape cast barium titanate bodies. The green sheets of modified barium titanates are cut to size, electroded, and then fired in air. The firing temperature is above 1350°C, and only expensive palladium or platinum metals can be used as internal electrodes. To reduce production
costs of terminating capacitors, new dielectric compositions with a low firing temperature (\(< 1000^\circ C\)) are being studied because at this low firing temperatures relatively cheaper Ag-Pd electrodes could be used. Complex solid solutions between ferroelectric compounds with a perovskite type structure and a general formula \(\text{Pb}(\text{Bi}^{\text{III}} \text{B}^{\text{IV}})_3\) are very attractive for this purpose. Besides a low sintering temperature, these compounds show a very high peak in the permittivity vs temperature curves \([1-6]\).

Smolenskii, et al. \([1]\) first reported a study on the compound \(\text{Pb}(\text{Mg}^{2+} \text{Nb}^{5+})_0\) hereafter referred to as PMN, with the perovskite crystal structure. Since then, this compound has been studied intensively by various workers. The investigations showed that the PMN system has a broad maximum of the dielectric constant just below room temperature. The magnitude of this maximum, (at 1 KHz, \(\sim 15000\) for ceramic PMN and \(>20000\) for single-crystal PMN), decreases and the temperature of this maximum increases with increasing frequency. This behavior is typical of a group of materials commonly referred to as relaxor ferroelectrics.

Recently, there has been much interest in PMN and PMN-based materials for electrostrictive applications. The electrostrictive strains generated in PMN-based ceramics are found to be an order of magnitude larger than those of \(\text{TiO}_2\)-based ceramics, due to the large dielectric constant of PMN. The induced electrostrictive strains are proportional to the square of the polarization.

The main limitation of the application of PMN in electronic devices
has been the lack of a simple, reproducible fabrication technique for pure PMN. The formation of pure perovskite PMN is difficult due to the formation of a pyrochlore type parasitic phase during fabrication. The main cause for this has been attributed to the volatilization of lead oxide during fabrication. Studies on the mechanism of formation and the ceramic processing of this system have shown that this parasite phase is very difficult to eliminate, even with an excess of PbO as regard to the stoichiometric composition [7-13].

Dielectric films produced by MOD processes are expected to have a finer grain size, more uniform surface and higher density compared to conventional tape cast films. Since the MOD films are relatively thinner it is also expected that the films produced using the MOD technique would give more capacitance per unit volume than the films produced using conventional routes.

This project involved a preliminary study of the mechanism of formation of perovskite PMN in powder and film forms using the MOD technique of fabrication. The aim was to develop procedures to obtain pyrochlore-free perovskite phase. For this purpose, phase formation as a function of precursor composition and processing conditions was studied, and an attempt was made to understand the mechanism of perovskite PMN formation. Since the elimination of pyrochlore is a crucial initial step towards the fabrication of PMN ceramics with high dielectric constant (as is evident by the difference in the $\varepsilon$ values of single-crystal and polycrystalline ceramic PMN), attempts were made to study the crystal structure and composition of the pyrochlore phase formed during fabrication in order to gain an insight into the pyrochlore - perovskite
conversion reaction. In case of MOD films, the quality of the film being produced was also of considerable interest. The aim was to produce a film which was uniform, homogeneous and pin-hole free. For this purpose, preliminary studies were conducted on various substrates. For better quality control and consistency in the results, all the required compounds were synthesised in house.

4.2 Experimental

4.2.1 Selection of Compounds

The metallo-organic compounds selected were lead neodecanoate, magnesium neodecanoate and niobium tri-ethoxy-di-neodecanoate. The synthesis of lead neodecanoate was discussed in Section 2.2.1.3, and synthesis of the other two compounds is presented in the following sections. Individual xylene solutions of these MOD compounds were mixed in stoichiometric proportions to obtain a solution having a\( \text{Pb}_3\text{MgNb}_2\text{O}_9 \) composition. Xylene was chosen as the solvent because it had a low boiling point and was easily removed on heating. This 'PMN Solution' was used to study the decomposition characteristics, phase formation, and for structural studies. Some of the studies were conducted with the 'PMN Solution' having an additional amount of lead or magnesium MOD precursors to give excess PbO or MgO in the fired products.

4.2.2 Synthesis of Niobium Tri-Ethoxy-Di-Neodecanoate

Niobium tri-ethoxy-di-neodecanoate was prepared using niobium penta-ethoxide. Niobium penta-ethoxide is an extremely moisture and air
sensitive liquid, thus an inert gas atmosphere was required to transfer this substance. In a glove bag filled with nitrogen, a previously weighed 100 ml round-bottom flask with rubber septum, a bar magnet and the niobium compound in vials were placed. A vial was broken inside the glove bag and 25 grams of niobium penta-ethoxide was transferred into the flask under nitrogen. The flask was closed with the rubber stopper and weighed again to get the accurate weight of the niobium compound. The flask was then placed on a heating mantle supported over the magnetic stirrer and 84.31 ml of benzene and 30.9 ml of neodecanoic acid were added to the flask under nitrogen using a syringe. The solution was stirred and the rubber septum was quickly replaced by the condenser. The solution was heated and allowed to reflux for 4 hours. It was then filtered under a mild suction to remove suspended impurities. A pale yellow syrupy liquid of the niobium compound was obtained. The equation for the synthesis is given below.

\[
\text{Nb(O}_2\text{C}_2\text{H}_5\text{)}_5 + 2\text{C}_9\text{H}_{19}\text{COOH} = \text{Nb(O}_2\text{C}_2\text{H}_5\text{)}_5(\text{C}_9\text{H}_{19}\text{COO})_2 + 2\text{C}_2\text{H}_5\text{OH}
\]

4.2.3 Synthesis of Magnesium Neodecanoate

In order to prepare a MOD precursor of magnesium, attempts were made with various inorganic compounds of magnesium such as Mg(NO$_3$)$_2$, MgCO$_3$, MgO etc. However, a suitable metallo-organic compound of magnesium could not be obtained. Finally, Mg(OH)$_2$ was found to give the best results as far as the clarity of the solution, and the solubility in xylene was concerned.
Fifty eight grams of Mg(OH)$_2$, 198.0 ml. of purified neodecanoic acid and about 125 ml of xylene for refluxing purpose were mixed in a round bottom flask. After connecting it to a fractional distillation assembly, the content of the flask was heated on a heating mantel. The temperature of the flask was kept constant at 95°C. A magnetic stirrer was used during distillation to stir the content of the flask. Water vapors and some amount of xylene were collected in a beaker as the distillation product. After about one hour, heating was stopped and the whole assembly was allowed to cool. The content of the flask was poured in a china dish, and was reheated at 135°C in order to complete the reaction. The content of the china dish was dissolved in xylene and was filtered using a Buchner funnel to remove the unreacted magnesium hydroxide. A small amount of magnesium hydroxide, deposited at the bottom after 12 hours, was removed by filtering. The equation for the synthesis is given below.

\[
\text{Mg(OH)}_2 + 2\text{C}_9\text{H}_{19}\text{COOH} = (\text{C}_9\text{H}_{19}\text{COO})_2\text{Mg} + 2\text{H}_2\text{O}.
\]

4.2.4 X-ray Diffraction Studies

Compound identification and structural studies were performed on the fired products in bulk and film forms. The x-ray diffraction technique was used to determine the phase(s) present in the fired products as a function of precursor composition and processing conditions, and if appropriate, the crystal structures of the phases formed. An automated
x-ray diffractometer was used for this purpose. The phase analysis was performed using Cu Kα radiation and $2^0 (2\theta)$/min goniometer speed.

From crystal structure analysis, the high angle reflections of the respective phases were chosen for the lattice parameter determination. The accurate peak location was determined using the program "AUTOFI" adopted for the Siemens model. This program was designed to produce an intensity profile between specified two-theta angles and then to determine the peak position for the peaks in the range. Only peaks above a user-defined background level were located to prevent random fluctuation in the background being reported as peaks. The computer program fitted the Pearson Type VII distribution, using a least-squares method to all peaks in the 2θ range. Gaussian distributions are commonly too broad near the peak and too narrow at the tails. Functions proportional to a Cauchy distribution are usually unsatisfactory in the opposite way. Pearson Type VII distributions appear to closely represent the shape of a symmetric X-ray diffraction peak [14].

The determination of lattice constants of a phase was done by using a program which refines lattice constants for any symmetry by a modified least square method. With the help of this program, several types of extrapolation functions can be used either singly or in combination. In the present investigation, Nelson-Riley's extrapolation function was used. Knowing the (hkl) and determining the experimental value of 2θ and thus d, the lattice constant was calculated for each peak and plotted versus the function $1/2\left(\frac{\cos^2 \theta}{\sin \theta} + \frac{\cos^2 \theta}{\theta}\right)$. The extrapolated value of the lattice constant corresponding to $\theta = 90^0$ gave the minimum error. A complete error treatment was made with the help of the computer.
4.2.5 Characterization of MOD Compounds

The MOD compounds of lead, magnesium and niobium were dissolved in xylene and their decomposition behavior was studied using TGA. From the thermograms of individual compounds, information about the final firing temperature. The composition of final products was determined by comparing the weight loss during firing with the phases identified by the x-ray diffraction studies. The metallo-organic precursors were fired in a platinum crucible and the fired products were analyzed for the phases present. The weight percent concentrations of these phases were obtained by noting the difference in the weights of initial compounds and that of fired products. Results from the TGA and x-ray diffraction studies are given in Table 4.1.

4.2.6 TGA of 'PMN Solution'

The 'PMN Solution' prepared by using MOD compounds in xylene was studied for its decomposition behavior. The TGA was performed in the temperature range of 27-900°C using heating rates of 1°C/min, 5°C/min and 10°C/min. The results showed that the final decomposition temperature decreased as the heating rate decreased. The TGA results are summarized in Table 4.2.

4.2.7 PMN Powder Formation

The aim of PMN firing studies in the bulk form was to develop a procedure to obtain perovskite lead magnesium niobate with the minimum possible pyrochlore. Three 'MOD Solutions' were prepared, namely, a
Table 4.1 TGA and X-ray Diffraction Results of Metallo-Organic Compounds in Xylene.

<table>
<thead>
<tr>
<th>Metallo-Organic Compound</th>
<th>( T_d ) (^{\circ} \text{C} )</th>
<th>Wt. %</th>
<th>Fired Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead Neodecanoate</td>
<td>415</td>
<td>14.2</td>
<td>PbO</td>
</tr>
<tr>
<td>Magnesium Neodecanoate</td>
<td>400</td>
<td>2.23</td>
<td>MgO</td>
</tr>
<tr>
<td>Niobium Tri-ethoxy-di-neodecanoate</td>
<td>400</td>
<td>14.0</td>
<td>Nb_2O_5</td>
</tr>
</tbody>
</table>

*: \( T_d \) : Temperature at which decomposition is complete.

Table 4.2 TGA of 'MOD Solution' of Stoichiometry Pbn in Xylene.

<table>
<thead>
<tr>
<th>Heating Rate</th>
<th>( T_d ) (^{\circ} \text{C} )</th>
<th>Wt. %</th>
<th>Fired Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>1(^{\circ})C per min.</td>
<td>300</td>
<td>15.0</td>
<td></td>
</tr>
<tr>
<td>5(^{\circ})C per min.</td>
<td>325</td>
<td>15.0</td>
<td></td>
</tr>
<tr>
<td>10(^{\circ})C per min.</td>
<td>375</td>
<td>15.0</td>
<td></td>
</tr>
</tbody>
</table>

a: \( T_d \) : Temperature at which decomposition is complete.
stoichiometric lead magnesium niobate solution in xylene (referred to as "PMN Solution"), a "PMN Solution" with additional 2 wt % PbO, and a "PMN Solution" with additional 2 wt% MgO. The "PMN Solution" was prepared by mixing weighed amounts of the MOD compounds of lead, magnesium and niobium dissolved in xylene as per required for $\text{Pb}_2\text{MgNb}_2\text{O}_9$ composition. The "MOD Solutions" with additional 2 wt % PbO or MgO were prepared by adding the weighed amounts of the MOD compounds of lead or magnesium, as per the case, to the "PMN Solution". The fired products contained 100 parts by weight of PMN + 2 parts by weight of excess PbO or MgO if excess were added. A small amount of the "PMN Solution" was fired in a platinum crucible at the firing rate of 1°C/min to the decomposition temperature determined by TGA, in order to determine the wt % yield of the fired products.

The firing studies on the three above mentioned "MOD Solutions" were conducted by firing about 5 to 10 grams of these solutions in a platinum crucible, using a two stage process. The initial firing at the rate of 1°C per minute, for firing temperatures to ~420°C, was used for all the firing schedules. This slow heating rate at the early stages of the firing was desired because the vapor pressure of lead oxide can cause considerable lead losses during the removal of organic compounds, and because the final decomposition temperature to the PMN compound decreases with decreasing heating rates. The extent of lead loss in the form of lead oxide has a crucial effect on the formation of the perovskite phase as opposed to the pyrochlore phases during the final stages of firing.
The decomposition products from the initial firing were further heated from 600°C to 1100°C for times ranging from 30 min to 3 hours in order to complete the phase formation reactions. The phases in the fired samples were analyzed using x-ray diffraction with Cu Kα radiation.

4.2.8 PMN Film Processing

The second part of this project consisted of a preliminary study of the development of PMN films formed on various substrates by the decomposition of the 'MOD Solutions' prepared from metallo-organic precursors. The selection of metallo-organic decomposition technique to produce films is based on the fact that the MOD process produces a film which is denser, and more uniform in composition and microstructure. Also, the thickness of the films can be controlled easily by using this process. For the preliminary studies, the phase formation in PMN films was determined as a function of substrate materials, processing conditions and precursor compositions. The experiments that were conducted are described below.

I. In one set of experiments, xylene solutions that gave stoichiometric PMN, or PMN with excess PbO or MgO on thermal decomposition were spun on platinum foils and were fired at various temperatures. The phase formation studies were conducted using x-ray analysis, and the quality of the films were tested visually.

II. In another set of experiments, AlSiMag 838 substrates were used for film deposition. The uniformity of the films was tested by optical
microscopy and the phase formation studies were conducted using x-ray analysis.

4.2.9 Substrates

Selection of a suitable substrate was an important factor for obtaining a film with desired qualities. Several parameters were considered in the selection of a suitable substrate such as:

1) the chemistry of the substrate, because the development of film adhesion always involves some degree of chemical interaction between the film and the substrate;

2) the substrate should have a melting point well above the range of the desired working temperatures;

3) the substrate should not react with the metallo-organic compounds, producing phases that alter the properties of the film unfavorably;

4) the coefficient of thermal expansion of the substrate should be as close as possible to that of the products formed during the processing since this will affect the uniformity of the fired films;

5) the surface roughness of the substrate limits the minimum thickness of a film that can be continuous.

The preliminary studies were conducted on platinum foils (with cross-sectional area of 4 cm²) because they eliminate the possibility of any chemical interaction with the films and thus provide good samples for structural studies. However, due to frequent shape deformation of the platinum foils during handling and also due to the high cost of the foils, the platinum foils were replaced by alumina substrates.
Alumina (96 to 99.8% Al₂O₃) is the most common material used for substrates in hybrid microelectronics because it is quite good as far as all of the general requirements are concerned. They can withstand high temperatures and are available in a number of surface roughness specifications. The AlSiMag 838 (99.5% Al₂O₃) substrates were used for the later part of this study. The surface roughness of the AlSiMag 838 substrates, measured with a profilometer, was found to be ~0.15 μm.

4.2.10 Formation of Wet Films

The technique used for formation of wet films is an important initial stage whose selection has to be carefully considered. The thickness and the uniformity of the wet films affect all subsequent processes. The deposition of wet films on the substrate was done by spinning. This method gives a very uniform coating of a solution on a substrate. Also, layers of higher thickness can be obtained by repeated spinning operations.

The selection of proper spinning parameters, such as, the spinning speed and the spinning time, is very important. Low spinning speeds may lead to edge build up caused by the surface tension of the solution at the edge of the substrate, whereas very high speeds may cause streaking or thin coatings. Viscosity of the solution determines the spinning speed, and by proper selection of this parameter, coatings of various thickness can be obtained.

An LS-8000 lab spinner* was used during the course of this project.

*Integrated Tech. Inc.
The following procedure was used to deposit wet films. The 'MOB Solutions' prepared for the bulk firing studies were used as the starting solutions for film deposition. The viscosities of the starting solutions were adjusted for spinning operation by removing some of the xylene by bubbling air or nitrogen through these solutions. The initial solution had the viscosity of water and after bubbling gas through it the viscosity was similar to that of a syrup. The substrate was manually centered on the chuck; the chosen spinning parameters were set; the substrate was flooded with the solution until all parts of the substrate were wet and the solution just spilled over; the substrate was rotated for the preset spinning time and speed. The selection of spinning parameters was done by trial and error for each batch of solutions. In general, the spinning speed was in the range of 1500-2000 rpm and spinning time was between 15 to 25 seconds.

4.3 Results and Discussion

4.3.1 Formation of Lead-Magnesium Niobate in Bulk Form

The results from the firing studies of the stoichiometry 'PMN Solution' are reported in the Table 4.3. The wt % yield was obtained from the weight difference between the fired product and the starting 'Solution'. The third column in the Table 4.3 indicates the approximate percentage yield in terms of the relative contents of the perovskite phase vs. the pyrochlore phases obtained from x-ray data using the relationship given at the bottom of the Table 4.3.

The x-ray diffraction studies of the stoichiometric PMN samples
Table 4.3 Firing of Stoichiometry 'PMN Solution'.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Firing Schedule</th>
<th>Wt. %</th>
<th>Vol. %</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RT to 250°C/ 1°C/min + 250°C / 2 hrs.</td>
<td>14.74</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>RT to 400°C/ 1°C/min + 400°C / 2 hrs.</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>RT to 420°C/ 1°C/min</td>
<td>13.39</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>RT to 440°C/ 1°C/min</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>RT to 420°C/ 1°C/min + 500°C/ 2 hrs.</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>RT to 420°C/ 1°C/min + 550°C / 2 hrs.</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>RT to 420°C/ 1°C/min + 600°C/ 2 hrs.</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>RT to 420°C/ 1°C/min + 700°C/ 2 hrs.</td>
<td>-</td>
<td>82</td>
</tr>
<tr>
<td>9</td>
<td>RT to 420°C/ 1°C/min + 800°C/ 2 hrs.</td>
<td>13.40</td>
<td>90</td>
</tr>
<tr>
<td>10</td>
<td>RT to 420°C/ 1°C/min + 875°C/ 2 hrs.</td>
<td>12.81</td>
<td>95</td>
</tr>
<tr>
<td>11</td>
<td>RT to 420°C/ 1°C/min + 900°C/ 3 hrs.</td>
<td>11.86</td>
<td>81</td>
</tr>
<tr>
<td>12</td>
<td>RT to 420°C/ 1°C/min + 1000°C/ 2 hrs.</td>
<td>11.90</td>
<td>69</td>
</tr>
<tr>
<td>13</td>
<td>RT to 420°C/ 1°C/min + 1100°C/ 2 hrs.</td>
<td>11.60</td>
<td>65</td>
</tr>
<tr>
<td>14</td>
<td>RT to 350°C/ 1°C/min + 750°C/ 2 hrs.</td>
<td>-</td>
<td>82</td>
</tr>
</tbody>
</table>

* : Vol.% = \[ \frac{\text{Iperov.}}{\text{Iperov.} + \text{Ipyro.}} \] \times 100

Iperov. = Peak Height of the Most Intense Perovskite Peak (hkl 110).
Ipyro. = Peak Height of the Most Intense Pyrochlore Peak (hkl 222).
fired in platinum crucibles were conducted using Cu Kα radiation. The phases present in various samples were identified by matching the experimental x-ray patterns with the Powder Diffraction File. The correct identification of phases in the PMN system is very difficult because: 1) the intermediate phases, termed pyrochlores, have almost identical diffraction patterns; and 2) the diffraction patterns of the starting compounds, PbO, MgO and Nb₂O₅ have sets of lines which overlap with those of the pyrochlores and perovskite, making it difficult to determine the temperatures at which the various oxides started to react or a phase change took place. A list of possible compounds present in the PMN system at various temperatures and the location of their major peaks for Cu Kα radiation is given in Table 4.4.

Other factors which further complicate the phase identification, and which are partially responsible for the conflicting results available in the published literature are:

1) the excess PbO and/or MgO added during processing might be present at high temperatures due to incomplete reactions, or simply because they were added in excess of what was actually required;
2) depending upon the processing conditions, such as, firing time and temperatures, the amount of lead loss may vary, producing different PbO-Nb₂O₅ based phases for different sets of experiments;
3) the excess MgO, added to reduce the pyrochlore in the final samples, might be incorporated in the various product phases and thus affect their x-ray patterns.
Table 4.4 List of the Possible Compounds Formed in the PMN System and Their Major X-ray Peak Positions.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Crystal Structure</th>
<th>Powder Diffraction</th>
<th>File</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgO</td>
<td>Cubic</td>
<td>20</td>
<td>29.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td>d°A 3.05</td>
<td>2.05</td>
</tr>
<tr>
<td>MgO</td>
<td>Cubic</td>
<td>20</td>
<td>42.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td>d°A 2.106</td>
<td>1.689</td>
</tr>
<tr>
<td>PbO</td>
<td>Cubic</td>
<td>100</td>
<td>52</td>
</tr>
<tr>
<td>PbO</td>
<td>Orthorhombic</td>
<td>20</td>
<td>23.65</td>
</tr>
<tr>
<td>PbO</td>
<td>orthorhombic</td>
<td>20</td>
<td>29.11</td>
</tr>
<tr>
<td>(Nb₂O₅)ₓO</td>
<td>orthorhombic</td>
<td>20</td>
<td>28.35</td>
</tr>
<tr>
<td>Nb₂O₅</td>
<td>Monoclinic</td>
<td>20</td>
<td>22.50</td>
</tr>
<tr>
<td>PMN</td>
<td>Cubic</td>
<td>20</td>
<td>31.15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>d°A 3.15</td>
<td>3.93</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>31.15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>d°A 2.87</td>
<td>2.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>100</td>
</tr>
</tbody>
</table>
Table 4.4 Continued.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Crystal structure</th>
<th>Powder diffraction</th>
<th>File</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na$_3$BO$_4$</td>
<td>Cubic</td>
<td>20-403</td>
<td>24</td>
</tr>
<tr>
<td>4,8</td>
<td>6.09  3.05  1.86</td>
<td>1.87  1.16</td>
<td></td>
</tr>
<tr>
<td>100  100  50  25  14</td>
<td>100  200  40  40  30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Na$_3$PO$_4$</td>
<td>Trigonal</td>
<td>30-712</td>
<td>20</td>
</tr>
<tr>
<td>4</td>
<td>5.14  5.09  2.68</td>
<td>1.89  1.84</td>
<td></td>
</tr>
<tr>
<td>100  100  50  25  10</td>
<td>100  200  250  250  250</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Na$_3$SO$_4$</td>
<td>Cubic</td>
<td>20-750</td>
<td>20</td>
</tr>
<tr>
<td>4</td>
<td>6.12  5.06  2.68</td>
<td>1.87  1.60</td>
<td></td>
</tr>
<tr>
<td>100  100  55  70  85</td>
<td>100  200  400  400  620</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
The x-ray diffraction pattern of the sample 1 indicated that the decomposition of the "MOD Solution" was complete after slow 1\(^{\circ}\)C/min heating to 250\(^{\circ}\)C and holding for 2 hours, giving a mixture of the starting oxides. The broad peak observed in the 2\(\theta\) range of 20-35\(^{\circ}\) indicated the presence of very small particle size elemental oxides with their strongest peaks falling in this range, PbO (28.65\(^{\circ}\) for tetragonal phase and 29.11\(^{\circ}\) for the orthorhombic phase), MgO (29.95\(^{\circ}\)) and \(\text{Nb}_2\text{O}_5\) (28.35\(^{\circ}\) for orthorhombic phase). On further heating to 400\(^{\circ}\)C and holding for 2 hours, as in the case of sample 2, the x-ray peaks for a pyrochlore phase was observed along with those for starting oxides. Samples 2 to 7, fired in the temperature range 400-640\(^{\circ}\)C indicated the pyrochlore phase along with the PbO. Other oxides, MgO and \(\text{Nb}_2\text{O}_5\) could not be detected in the x-ray patterns, probably due to their very small particle size. The pyrochlore phase in this temperature range seems to be based on the composition Pb\(_3\)Nb\(_2\)O\(_8\), although a complete matching of the patterns was not possible.

The perovskite peaks were first observed in sample 7, fired at 640\(^{\circ}\). Further increases in the firing temperature continuously increased the perovskite content with the corresponding decrease in the pyrochlores. This trend was observed in the samples 8-10 and 14, fired between the temperature range 700-875\(^{\circ}\).

The diffraction patterns of the samples 8 and 9 fired in the temperature range 700-800\(^{\circ}\) indicated an extra peak at 2\(\theta\) ~ 30.5\(^{\circ}\). This peak disappeared as the temperature was further increased (sample 10). Except for this peak, the rest of the peaks in the x-ray patterns could be separated into two sets, the perovskite phase and the pyrochlore.
phases. Samples 11-13 fired at temperatures above 875° showed a continuous decrease in the perovskite content and a corresponding increase in the pyrochlores with increasing firing temperature.

The pyrochlore in the samples fired in the temperature range 700-1100° matches closest to a mixture of Pb$_3$Nb$_2$O$_8$ and Pb$_2$Nb$_2$O$_7$ in terms of intensity, with the first type being the major pyrochlore phase. A complete matching of the experimental patterns with any of the listed pyrochlore phases was not obtained.

In summary, the x-ray diffraction and firing studies indicated that the perovskite was a major phase in the temperature range 700-1000°. The maximum in the perovskite content was obtained at 875°, with a further decrease in content as the temperature was increased due to the lead loss at high temperatures. It was also found that at low temperatures, pyrochlore was formed which converted into perovskite as the firing temperature was further increased. However, the complete transition from pyrochlore to perovskite was not observed.

The x-ray diffraction results for samples 15-18 having PMN with 2 percent excess PbO are given in Table 4.5. Sample 15, fired at 800° for 2 hours, showed an extra peak at ~30.5°, as in the case of the stoichiometric PMN sample fired at this temperature (sample 9). This peak also disappeared at higher temperatures (samples 16-18). The results of the experiments using "PMN Solution with 2 wt % excess PbO" showed that addition of extra PbO in the original solution (to compensate for the lead loss at high temperatures) increased the amount of the perovskite phase in the final product. Pyrochlore-free PMN was not
Table 4.5 Firing of 'PMN Solution with Excess 2 wt% PbO'.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Firing Schedule</th>
<th>wt.% Yield</th>
<th>Vol.% PMN</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>RT to 420°C/1°C/min + 800°C/2 hrs.</td>
<td>-</td>
<td>93</td>
</tr>
<tr>
<td>16</td>
<td>RT to 420°C/1°C/min + 875°C/2 hrs.</td>
<td>12.96</td>
<td>98</td>
</tr>
<tr>
<td>17</td>
<td>RT to 420°C/1°C/min + 1000°C/2 hrs.</td>
<td>12.34</td>
<td>76</td>
</tr>
<tr>
<td>18</td>
<td>RT to 420°C/1°C/min + 1100°C/2 hrs.</td>
<td>11.03</td>
<td>93</td>
</tr>
</tbody>
</table>

* Vol.% = \( \frac{\text{Iperov.}}{\text{Iperov.} + \text{Ipyro.}} \) *100%

Iperov. = Peak Height of the Most Intense Perovskite Peak (hkl 110).
Ipyro. = Peak Height of the Most Intense Pyrochlore Peak (hkl 222).
obtained in any of the samples, but the sample 16 showed 98 vol % PMN.

Phase formation in the "PMN solution with excess 2 wt % MgO" was studied as a function of firing temperatures and time. It was found that the addition of excess MgO gave significantly better results in terms of pyrochlore to perovskite conversion in the temperature range of 425–1000°C, as shown in Table 4.6. The perovskite peaks were first observed at temperatures as low as 420°C (sample 19). It was also observed that the reaction product of the "PMN Solution with 2 wt.% excess MgO" fired to 420°C at the rate of 1°C per min, followed by additional heating of 2 hours at 875°C (sample 21) did not indicate the presence of any pyrochlore phase in the x-ray diffraction pattern.

The perovskite PMN obtained from the PMN solution with excess of two percent MgO was used for determination of lattice constant. The lattice constant of this phase was found to be $a = 0.4038$ nm.

4.3.2 PMN Films

The x-ray diffraction results for films fired on platinum substrates are given in Table 4.7. The results of preliminary studies for deposition of PMN on platinum foils indicated that the removal of organic substances from the wet films was complete by 300°C producing an amorphous film, as was shown by the powder diffraction pattern of sample 24. A broad peak was observed at $2\theta \sim 30^\circ$ indicating the presence of lead, magnesium and niobium oxides in very fine particle size. On increasing the firing temperature to 520°C, as in the case of sample 25, the formation of the pyrochlore could be observed in the x-ray diffrac-
Table 4.6 Firing of 'PMN Solution with Excess 2 wt% MgO'.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Firing Schedule</th>
<th>Vol. 4</th>
<th>Figure No. **</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>PMN*</td>
<td></td>
</tr>
<tr>
<td>19</td>
<td>RT to 420°C/ 1°C/ min + 420°C/2 hrs.</td>
<td>18</td>
<td>4.19</td>
</tr>
<tr>
<td>20</td>
<td>RT to 420°C/ 1°C/ min + 800°C/2 hrs.</td>
<td>97</td>
<td>4.20</td>
</tr>
<tr>
<td>21</td>
<td>RT to 420°C/ 1°C/ min + 875°C/2 hrs.</td>
<td>100</td>
<td>4.21</td>
</tr>
<tr>
<td>22</td>
<td>RT to 420°C/ 1°C/ min + 1000°C/2 hrs.</td>
<td>69</td>
<td>4.22</td>
</tr>
<tr>
<td>23</td>
<td>RT to 420°C/ 1°C/ min + 1100°C/2 hrs.</td>
<td>65</td>
<td>4.23</td>
</tr>
</tbody>
</table>

* : Vol.% = \( \frac{\text{Iperov.}}{\text{Iperov.} + \text{Ipyro.}} \) *100%

Iperov. = Peak Height of the Most Intense Perovskite Peak (hkl 110).
Ipyro. = Peak Height of the Most Intense Pyrochlore Peak (hkl 222).

** : XRD Patterns are in Appendix D.
Table 4.7 Firing of Stoichiometry Film of PMN on Platinum Substrates.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Firing Schedule</th>
<th>Vol. % PMN*</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>RT to 300°C/ 10°C per min.</td>
<td>0</td>
</tr>
<tr>
<td>25</td>
<td>RT to 520°C/ 10°C per min.</td>
<td>0</td>
</tr>
<tr>
<td>26</td>
<td>RT to 620°C/ 1°C per min.</td>
<td>0</td>
</tr>
<tr>
<td>27</td>
<td>RT to 800°C/ 1°C per min.</td>
<td>94</td>
</tr>
</tbody>
</table>

* : Vol. % = \frac{\text{Iperov.}}{\text{Iperov. + Ipyro.}} \times 100$

Iperov. = Peak Height of the Most Intense Perovskite Peak \(\{hkl\,110\}\).

Ipyro. = Peak Height of the Most Intense Pyrochlore Peak \(\{hkl\,222\}\).
MULTILAYER CAPACITOR DIELECTRICS PRODUCED FROM METALLO-ORGANIC PRECURSORS (U) PURDUE UNIV LAFAYETTE IN TURNER LAB FOR ELECTROCERAMICS R W YEST ET AL.

UNCLASSIFIED 30 JUN 85 N00014-83-K-0321 F/G 9/1 NL
tion pattern. The x-ray pattern for pyrochlore in this case was found to be similar to that for powder samples. The perovskite peaks were first observed in the sample 26, fired to 620°C at the rate of 1°C per min. On further increasing the firing temperature, the pyrochlore content of the sample decreased with a corresponding increase in the perovskite, as shown for sample 27.

In summary, the best result was obtained with the film fired, first, continuously up to 800°C and then, for 1 hour at 800°C. However, the complete elimination of pyrochlore was not observed for any of the firing schedules. It was also noted that the films fired up to temperatures higher than 800°C (e.g. 875°C) showed relatively lesser amounts of perovskite than those fired up to 800°C. Also, as the firing time was increased (e.g. at 875°C), the amount of perovskite further decreased indicating that at higher temperatures the excessive lead loss took place.

4.4 Summary

1. The MOD precursors of lead, magnesium and niobium were prepared and characterized by thermogravimetric analysis. The lead neodecanoate, magnesium neodecanoate and niobium tri-ethoxy-di-neodecanoate were found to have almost the same temperature ranges of decomposition and peak decomposition temperatures of 415°C, 400°C and 400°C respectively. This is advantageous since the decomposition products of the three compounds would form simultaneously giving a uniform final fired product.
2. For the purpose of firing and x-ray diffraction studies, three ‘MOD Solutions’ in xylene having stoichiometric PMN, PMN with 2 wt % excess PbO, and PMN with 2 wt % excess MgO compositions were prepared by mixing the MOD precursors of the lead, magnesium and niobium in the required proportions. The TGA of the ‘MOD Solution’ having stoichiometric PMN composition indicated that the decomposition temperature of this solution was $375\,^\circ\text{C}$, which is lower than those for the elemental MOD precursors used for preparing this solution.

3. The firing studies of the ‘MOD Solutions’ indicated that the optimum firing schedule, to obtain perovskite as the major phase and to have low lead losses in the form of lead oxide evaporation, was an initial firing to $420\,^\circ\text{C}$ at the rate of $1\,^\circ\text{C}$ followed by a further heating in the temperature range $700-1000\,^\circ\text{C}$ for some constant interval of time ranging from 30 min. to 3 hours.

4. The firing studies of the stoichiometric ‘PMN Solution’ in bulk as well as film forms indicated that pyrochlore was the first product as the solid state reaction between the decomposed oxides of the lead, magnesium and niobium took place. The pyrochlore phases formed in the powder samples and in the films had similar diffraction patterns.

5. As the firing temperature of the ‘PMN Solution’ was increased, the low temperature pyrochlore phases converted into the perovskite lead magnesium niobate phase. This trend was continuous to approximately $800-875\,^\circ\text{C}$ for the bulk and film samples. Beyond this tem-
perature range, the pyrochlore content of the samples increased due
to the excessive lead loss.

6. Both the films and the powder samples of the stoichiometric 'PMN
Solution' did not produce a pyrochlore-free perovskite phase
irrespective of the firing time and temperatures. The maximum PMN
was 95 vol. % in the bulk and 94 vol. % in the films.

7. The addition of 2 wt % excess PbO increased the perovskite content
in the samples with the maximum at 875°C of 98%. However, the pyro-
chlore phases could not be eliminated completely.

8. In case of 'PMN solution with excess 2 wt % MgO', it was found that
the addition of excess MgO gave significantly better results in
terms of pyrochlore to perovskite conversion in the temperature
range of 425-1000°C. The perovskite peaks were first observed at
temperature as low as 420°C. The X-ray diffraction pattern of the
reaction product of 'PMN Solution with 2 wt.% excess MgO' fired
from RT to 420°C at the rate of 1°C per min followed by an addi-
tional heating of 2 hours at 875°C did not indicate the presence of
any pyrochlore phase. Traces of uncombined MgO were observed in
the powder diffraction pattern of this sample, indicating that even
less than 2 % excess MgO could be sufficient to eliminate most of
the pyrochlore phases.

4.5. References

1. G.A. Smolenskii and A. I. Agranovskaya, "Dielectric Polarization


END

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