MICROSTRUCTURE-PROCESSING-PROPERTY RELATIONS

IN CHEMICAL SOLUTION DEPOSITED BARIUM TITANATE FILMS

A Thesis in
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by
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ABSTRACT

This thesis explored the microstructure-processing-property relationships in chemical solution deposited BaTiO$_3$ films on Ni foils as model thin film capacitors. Different techniques, including X-ray diffraction, transmission electron microscopy and spectroscopic ellipsometry were combined to provide better understanding of microstructure and interface quality of BaTiO$_3$ thin films on Ni foil.

It was found that high quality thin films could be prepared using rapid thermal annealing 750°C in N$_2$ to crystallize the film without building up significant levels of NiO on the substrate. After building up the desired dielectric thickness, the films were heat-treated at 1000°C and later re-oxidized under controlled oxygen partial pressure conditions. The resulting films have dielectric constants of 1000-1300 which are stable as a function of temperature with loss tangents less than 2%.

Furnace annealed barium titanate films on Ni foil were characterized by X-ray diffraction and transmission electron microscopy (TEM). X-ray diffraction shows a well-crystallized polycrystalline perovskite phase in furnace annealed films with a high intensity of the 100 and 200 peaks. The films show equiaxed grains with average grain size of 42 nm. There are 5-6 grains across a 200 nm thick film, suggesting that it should be possible to use grain boundaries in films in order to control the capacitor reliability, as is done with bulk capacitors. NiO was detected by X-ray diffraction, but not by transmission electron microscopy, suggesting that the oxide is not a continuous barrier layer, but is distributed inhomogeneously over the surface. Electron energy loss microscopy shows the existence of C in barium titanate grains. In addition, high
resolution transmission electron microscopy and electron energy loss spectroscopy showed that an interfacial Ni-Ba alloy develops at the interface between the BaTiO₃ film and the Ni foil. This would be consistent with very reducing partial pressures locally during the processing, probably as a result of retained organics.

Decomposition of both powders dried from barium titanate solutions and barium titanate films was studied by differential scanning calorimetry (DSC), thermal gravimetric analysis (TGA), Fourier transform infrared spectroscopy (FTIR) and X-diffraction. It was found that flowing N₂ delays decomposition of organics in the powders, leading to retained carbonate phases. Thus, crystallization of barium titanate occurs via decomposition of a barium carbonate phase. Retained organics, especially C, in BaTiO₃ films was thus found to be critical in processing of dense chemical solution deposited films in low oxygen partial pressures.

The optical properties, film density and film thickness were studied by spectroscopic ellipsometry as a function of processing conditions. It was shown that the refractive index of amorphous dried films increases as drying temperature increases. Similarly, the refractive index of the films increases as the RTA temperature increases, probably at least in part due to crystallization of BaTiO₃ and removal of some of the intermediate phases. On annealing at 1000°C, there is slightly increase in the refractive index of the film due to further crystallization. The final refractive index is comparable to that of 95% dense barium titanate ceramics. Re-oxidation did not change the refractive index of the film.

To facilitate studies of the dielectric/electrode interface, the optical properties of thermally grown NiO on Ni foil were extracted. It was found that the Ni foil begins to
oxidize at 300°C in air. The real part of the high frequency dielectric constant of NiO is similar to that of barium titanate, which complicated modeling of NiO in barium titanate films on Ni foil.
# TABLE OF CONTENTS

LIST OF FIGURES ......................................................................................................................................................... ix

LIST OF TABLES ........................................................................................................................................................ xiv

ACKNOWLEDGEMENTS ................................................................................................................................................. xv

Chapter 1 Introduction ....................................................................................................................................................... 1
  1.1 Base-metal electrode multilayer ceramic capacitors (BME MLCCs) ......................................................... 1
  1.2 Objectives ............................................................................................................................................................ 6
  References ............................................................................................................................................................... 8

Chapter 2 Literature Review .......................................................................................................................................... 13
  2.1 Dielectric properties ........................................................................................................................................ 13
  2.2 Barium titanate materials for capacitor applications ................................................................................. 19
  2.4 Chemical solution deposition of barium titanate thin films .................................................................... 22
  References ............................................................................................................................................................. 26

Chapter 3 Experimental Procedure ............................................................................................................................... 29
  3.1 Chemical solution deposition of BaTiO₃ Film ......................................................................................... 29
    3.1.1 BaTiO₃ solution ........................................................................................................................................ 29
    3.1.2 Film deposition and crystallization ..................................................................................................... 32
  3.2 Reducing furnace system ................................................................................................................................. 34
  3.3 Spectroscopic ellipsometry ............................................................................................................................. 38
    3.3.1 Introduction to ellipsometry ............................................................................................................... 38
    3.3.2 Data collection ....................................................................................................................................... 41
    3.3.3 SE measurement with a compensator .............................................................................................. 42
5.4 Spectroscopic ellipsometry studies ......................................................... 118
  5.4.1 Amorphous dried BaTiO$_3$ films ....................................................... 118
  5.4.2 BaTiO$_3$ films on sapphire substrates ............................................... 122
  5.4.3 Optical properties of nickel oxide ..................................................... 130
  5.4.4 BaTiO$_3$ films on Ni foil ............................................................... 135
  5.5 Electrical properties of BaTiO$_3$ films on Ni foil ................................ 141

References ........................................................................................................ 145

Chapter 6 Conclusions and Future Work ....................................................... 150
  6.1 Conclusions .............................................................................................. 150
  6.2 Future Work .............................................................................................. 152
    6.2.1 Organic removal and decomposition ................................................. 152
    6.2.2 Interface of barium titanate film and Ni foil ..................................... 153
    6.2.3 Spectroscopic ellipsometry for NiO on Ni foil investigation............... 154

References ........................................................................................................ 155
LIST OF FIGURES

Fig. 1.1 Schematic of a multilayer ceramic capacitor........................................... 2

Fig. 1.2 Trend of dielectrics and electrode thickness and active layer count of Kemet’s commercial (0603) Class 2 MLCCs................................................................. 2

Fig. 1.3 Dielectric properties of a) (Ba,Sr)TiO₃ films on Ni foil [31], b) BaTiO₃ films on Ni foil [28] and c) BaTiO₃ films on Cu foil [32]......................................................... 6

Fig. 2.1 a) Capacitor with vacuum between two parallel plates b) Polarization of charges and net surface charges at the faces of dielectric c) Capacitor with inserted dielectric showing bound charges at the dielectric faces and an increase in free charge on the metal plates................................................................. 14

Fig. 2.2 Phasor diagram for a) perfect capacitor and b) real capacitor (V is applied voltage, Iᵥ is charging current and Iᵥ is loss current)........................................... 16

Fig. 2.3 a) Polarization mechanisms [2] b) Frequency dependence of ε′ and ε′′ [1].....18

Fig. 2.4 Schematic of the cubic perovskite structure of the high temperature BaTiO₃ unit cell (Adapted from Kingery et al. 1991 [4])......................................................... 20

Fig. 2.5 Phase transition sequence in BaTiO₃ (Figure from Moulson et al. [5])........20

Fig. 2.6 Dielectric constant ε (equivalent to εᵣ) and its inverse as a function of temperature for a BaTiO₃ single crystal [7]......................................................... 22

Fig. 3.1 Schematic of methanol-based solution preparation method..........................31

Fig. 3.2 Schematic of 2-MOE-based solution preparation method..........................31

Fig. 3.3 Schematic of preparation of BaTiO₃ thin films........................................33

Fig. 3.4 Stability of Ni and various oxidation state of titanium (Circles, ○ and ●, shows typical setting values and measured values for the pO₂ for each heat treatment) (Adapted from Yang et al. [2])......................................................... 34

Fig. 3.5 Schematic of reducing furnace system....................................................37

Fig. 3.6 Reflection of light at an interface..........................................................38

Fig. 3.7 Schematic of rotating-analyzer spectroscopic ellipsometer [4]..................39
Fig. 3.8 Reflection of light from a thin film on a substrate………………………………………40
Fig. 3.9 Bright field TEM image showing BaTiO$_3$ film detached from the Ni foil due to conventional polishing preparation. This complicated studies of the interface quality……………………………………………………………………………………………………………………………………………………………………46

Fig. 3.10 Preparation of cross-sectional TEM samples by conventional polishing……..48

Fig. 3.11 Diagram of the FIB lift out process………………………………………………50

Fig. 3.12 (a – c) Series of top-view secondary ion images showing the thinning procedure for cross-sectional TEM sample by FIB a) a lift-out sample is attached to a Cu grid b) the sample is thinned by milling along the interface c) one-third of the sample is further thinned. (d-e) side view secondary electron images d) side view of c), e) final sample which is thinned until all of the W in a region disappears…..51

Fig. 4.1 TEM cross-sectional micrograph of 750°C annealed BaTiO$_3$ films on Ni foil showing a) partial film damage by ion milling at the top of the film and b) discrete crystallization layers in a region where the film is thick and large amount of W is still intact. BT= BaTiO$_3$…………………………………………………………………………59

Fig. 4.2 TEM bright-field micrograph of microstructure and interface of BaTiO$_3$ films on Ni foils annealed in RTA at a) 700°C and b) 750°C…………………………………………………………61

Fig. 4.3 Investigated area and corresponding electron diffraction patterns of BaTiO$_3$ films on Ni foils: a-b) are for a 700 RTA’d annealed film, c-d) are from a furnace annealed film (1000°C)……………………………………………………………………65

Fig. 4.4 AFM scan area 0.5 x 0.5 m of BaTiO$_3$ films annealed at a) 700°C (RTA), b) 750°C (RTA) and c) 1000°C………………………………………………………………………………65

Fig. 4.5 Bright-field micrographs of the microstructure and interface of BaTiO$_3$ films on Ni foils annealed in furnace 1000°C (Courtesy of Dr. Ian Reaney)………………………………………………………………………68

Fig. 4.6 Cross-sectional bright field image of BaTiO$_3$ film on Ni foil (Courtesy of Dr. Gaiying Yang). Positions for EELS analysis are marked…………………72

Fig. 4.7 EELS spectra of BaTiO$_3$ grain and Ni foil, showing existence of C in the BaTiO$_3$ (Data courtesy of Dr. Gaiying Yang)………………………………………………72
Fig. 4.8 EELS spectra collected from BaTiO$_3$ film as a function of distance from Ni foil a) broad range spectra b) resolved spectra of the fine structure of Ti L$_{2,3}$ and O K edges including that of PME BaTiO$_3$ (Data courtesy of Dr. Gaiying Yang).................................75

Fig. 4.9 HRTEM image of furnace annealed BaTiO$_3$ film/ Ni foil interface (Data courtesy of Dr. Gaiying Yang)..........................................................78

Fig. 4.10 EELS spectra collected from Ni region and interfacial layer between BaTiO$_3$ and Ni (Data courtesy of Dr. Gaiying Yang)........................................79

Fig. 4.11 View of a larger section of the BaTiO$_3$/Ni interface showing the interface layer and the lack of NiO (Data courtesy of Dr. Gaiying Yang)..............79

Fig. 4.12 Binary alloy phase diagrams of Ba-Ni (Adapted from Massalski [30])......80

Fig. 5.1 Surface morphology of low purity (99+ %) Ni foil........................................86

Fig. 5.2 SE data of Ni foil (99+ %) and Ni foil after annealing at 200-450°C for 2 min in air.................................................................89

Fig. 5.3 Schematic of SE models for Ni foils with and without NiO......................89

Fig. 5.4 Dielectric functions of Ni single crystals (SC) at 4 K and 293 K. The modified room temperature data set developed in this work is also shown...........90

Fig. 5.5 Experimental and modeled data with and without NiO for an as-received Ni foil........................................................................92

Fig. 5.6 Experimental and modeled data for Ni foils annealed at different temperatures ................................................................................93

Fig. 5.7 SE – determined depth profiles from as-received and annealed Ni foils.......96

Fig. 5.8 Plot of total thickness of NiO+air layer and effective thickness of NiO.......97

Fig. 5.9 SE data collected from BaTiO$_3$ films on 99% and 99.99% Ni foils...........100

Fig. 5.10 Microstructure and surface morphology of high purity Ni foils (99.99 %) a) as-received and b) after pre-annealing at 900°C. Both SEM and AFM data are shown.................................................................101

Fig. 5.11 XRD patterns of BaTiO$_3$ films on high purity Ni foils RTA’d at different temperatures..............................................................................104
Fig. 5.12 XRD patterns of furnace annealed BaTiO$_3$ films that were RTA’d at different temperatures……………………………………………………………………………104

Fig. 5.13 Slow x-ray diffraction scan of RTA annealed BaTiO$_3$ films on Ni foil……..105

Fig. 5.14 Slow scan x-ray diffraction pattern of furnace annealed BaTiO$_3$ films on Ni foil The films were RTA annealed at different temperatures (600-750°C) before the furnace anneal step……………………………………………………………………105

Fig. 5.15 Slow scan of x-ray diffraction showing NiO after re-oxidation of 700°C RTA’d BaTiO$_3$ film…………………………………………………………………… 106

Fig. 5.16 Microstructure of furnace annealed BaTiO$_3$ films (for different RTA temperatures) on Ni foil…………………………………………………………………107

Fig. 5.17 TGA and DSC curves of dried powder from methanol-based BaTiO$_3$ Solution………………………………………………………………………………109

Fig. 5.18 XRD patterns of powders calcined in air at different temperatures…………111

Fig. 5.19 TGA curves of dried powders from 2-methoxyethanol (2-MOE) and methanol based BaTiO$_3$ solutions (2-methoxyethanol data courtesy of Dr. Song Won Ko)……………………………………………………………………114

Fig. 5.20 FTIR spectra and thickness of deposited 2-methoxyethanol-based BaTiO$_3$ films dried at different temperature (Data courtesy of Dr. David Lu)………116

Fig. 5.21 FTIR spectra of 2-methoxyethanol-based BaTiO$_3$ films RTA’d at 750°C a) in air and b) in flowing N$_2$. The films were dried at different temperatures (125°C, 390°C or 500°C) (Data courtesy of Dr. David Lu)………………117

Fig. 5.22 Model fits compared to measured SE data for an amorphous BaTiO$_3$ film dried at 200°C on a silicon wafer (prepared from 0.2 M 2-methoxyethanol solution)…………………………………………………………………………..120

Fig. 5.23 Refractive index of dried amorphous BaTiO$_3$ films (prepared from a 0.2 M 2-methoxyethanol solution)……………………………………………………………………121

Fig. 5.24 Total thickness of amorphous BaTiO$_3$ films as a function of drying temperature (prepared from 0.2 M 2-methoxyethanol solutions)…………………..121

Fig. 5.25 Delta and psi as a function of wavelength at various points in the processing of a BaTiO$_3$ film on a (0001) Al$_2$O$_3$ substrate……………………………..125
Fig. 5.26 Model fits with 2 layer (Al₂O₃/Osc/Osc+air) and 3 layer (Al₂O₃/Osc+air/Osc/Osc+air) model of 750°C RTA’d BaTiO₃ film on (0001) Al₂O₃

Fig. 5.27 Model fits with 2 layer (Al₂O₃/Osc/Osc+air) and 3 layer (Al₂O₃/Osc+air/Osc/Osc+air) model of 1000°C furnace annealed BaTiO₃ film on (0001) Al₂O₃ (Note: Osc means oscillator.)

Fig. 5.28 Refractive index of RTA’d annealed, crystallized and re-oxidized BaTiO₃ films

Fig. 5.29 Depth profiles of BaTiO₃ films on (0001) Al₂O₃ after each annealing

Fig. 5.30 Total thickness of BaTiO₃ films on (0001) Al₂O₃ after each annealing

Fig. 5.31 Optical properties of a NiO single crystal (SC) and NiO films of different grain size (F1 and F2)

Fig. 5.32 Model fits of NiO thermally grown on high purity Ni foil at 600°C

Fig. 5.33 Depth profile and refractive index of NiO thermally grown at 600°C

Fig. 5.34 Dielectric function of the thermally grown NiO and a BaTiO₃ ceramic Sample

Fig. 5.35 Model fits of 750°C RTA’d BaTiO₃ film on high purity Ni foil

Fig. 5.36 Unreasonable film structure of RTA’d BaTiO₃ film on high purity Ni foil when modeled with NiO

Fig. 5.37 SE model details of BaTiO₃ films on Ni foil RTA’d at temperature from 600-750°C

Fig. 5.38 Refractive index of RTA’d BaTiO₃ films on high purity Ni foil

Fig. 5.39 Total thickness determined by SE for BaTiO₃ films on high purity Ni foil as a function of RTA temperature

Fig. 5.40 Frequency dependence of dielectric constant and loss tangent furnace annealed BaTiO₃ films on Ni foil

Fig. 5.41 Temperature dependence of dielectric constant and loss tangent of 750°C RTA’d and furnace annealed BaTiO₃ films on (Kodak) Ni foil
LIST OF TABLES

Table 3.1 Reducing setting condition and actual pO₂ measured by an oxygen sensor
........................................................................................................................................37

Table 4.1 Reported grain size and average grain size (in parentheses) of BaTiO₃ –
based films prepared by various groups under a variety of different conditions........64

Table 5.1 AFM and SE surface roughness of 99+% and pre-annealed 99.99% Ni foil
............................................................................................................................................99
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Chapter 1

Introduction

1.1 Base-metal electrode multilayer ceramic capacitors (BME MLCCs)

Multilayer ceramic capacitors (MLCCs) have been widely used in electronic circuits, including computers and cellular telephones, because of their small size and high volume efficiency. Basically, an MLCC consists of alternating layers of dielectrics and internal electrodes, as shown in Fig. 1.1. The total capacitance of a MLCC is the sum for all of the dielectric layers. Such a structure allows great flexibility in preparing a wide variety of capacitance values in a range of case sizes.

The drive to increase the capacitance obtainable in a given case size has driven two factors continuously: a decrease in the dielectric thickness, and an increase in the number of layers. As an example, Fig. 1.2 shows the evolution of Kemet’s commercial (0603) Class 2 MLCCs [1]. The dielectric thickness decreased from 6 µm (1997) to <1 µm (2007) over the last decade. In order to continue this trend in miniaturization, Randall et al. projected that MLCC would have 1900 layers of 0.17 µm thick dielectrics in 2010 [1]. These layer thicknesses are below what can currently achieved by tape casting of the dielectric, and so alternative means of preparing the capacitors are being considered.

The most commonly used dielectrics for MLCC are barium titanate-based ceramics [2, 3]. The composition and microstructure of the barium titanate are modified to meet EIA (Electronic Industries Association) specifications for the temperature
coefficient of capacitance [2, 4]. For example, the core-shell structure in X7R capacitors is produced by introduction of dopants and use of a controlled sintering profile. The result is a more uniform dielectric constant over a wider temperature range. Base metals, i.e. Ni and Cu have been used as internal electrodes in MLCCs, because of instability in the price of Pd (and Pd-Ag alloys) [2]. The use of base metal electrodes requires cofiring in highly reducing atmosphere to prevent oxidation of the electrodes. Oxygen vacancies

![Fig. 1.1 Schematic of multilayer ceramic capacitor](image)

![Fig. 1.2 Trend of dielectrics and electrode thickness and active layer count of Kemet’s commercial (0603) Class 2 MLCCs](image)
introduced in the dielectric layers can be partially ameliorated by post-oxidation. In manufacturing of MLCCs, typical reducing atmospheres for cofiring and re-oxidation are 1300°C in 10⁻¹⁰ atm pO₂ and 800°C in 10⁻⁸ atm pO₂, respectively [5, 6].

Despite re-oxidation, some oxygen vacancies which cause insulation resistance degradation [7-11] remain in the dielectrics. It was shown that the reliability of the capacitor significantly depends on the Schottky barriers at the electrode-dielectric interface and the grain boundaries that limits migration of the oxygen vacancies toward the cathode [5, 6, 8]. Addition of some rare-earth ions i.e. Y³⁺, Dy³⁺ and Ho³⁺ was found to considerably improve the insulation resistance and lifetime of low pO₂ treated barium titanate [12-16].

There have been some reports concentrating on the microstructure of the dielectric layer and the interface between the Ni electrode and the dielectric in MLCCs. Opitz et al. showed that in the re-oxidation step, oxygen diffusion along the Ni electrodes is faster than the direction perpendicular to it [17]. Oxygen vacancies in low pO₂ - processed barium titanate dielectrics resulted in the appearance of dislocation loops observable in a transmission electron microscope [5, 18, 19]. Later work showed that clustering of oxygen vacancies in some degraded barium titanate grains results in a modulated structure with {111}planar defects [20, 21]. Yang et al. [5] showed that the oxygen content in cofired barium titanate is not uniform, and that a higher oxygen vacancy concentration is observed near the Ni electrode. The local variation of barium titanate reduction was attributed to the effect of residual carbon. However, the uniformity of the oxygen content in barium titanate is significantly improved after the re-oxidation step. Yang et al. [22] also revealed the existence of a discrete layer (4-15 nm
thick) of Ni-Ti-Ba metallic alloy between the Ni and barium titanate in many commercial MLCCs. The alloy layer is formed during high temperature cofiring and is stable through the re-oxidation step. On the other hand, Chen et al. [23] reported interdiffusion of Ni into the dielectric layer. It was found that the thickness of the diffused layer increased as the dielectric thickness decreased.

MLCCs production has traditionally employed tape casting for the barium titanate and screen printing of the Ni electrodes. With progressive reduction in dielectric thickness, there have been major advances in optimization of the tape cast slurries and processes to obtain thinner green sheets. These include controlling solution dispersion, developing of nanopowders, etc. [24-26].

Alternatively, thin film approaches to multilayer capacitors have also been explored recently. Sakabe et al. [27] demonstrated thin film multilayer capacitors prepared on Pt-coated MgO substrates by metalorganic chemical vapor deposition. With 12 layers of 0.26 µm thick (Ba,Sr)TiO$_3$ dielectrics, the capacitors show very promising capacitance per unit volume (33 µF/mm$^3$). Koutsaroff et al. [28] have fabricated double layer BST thin film capacitors with Pt electrodes on glazed alumina with a BST and TiO$_x$ buffer layer by metal-organic decomposition. The capacitance density achieved was 36.5 fF/µm$^2$. In addition, Nagata et al. have shown feasibility of making capacitors by microcontact printing [29] of thin films.

Development of thin film versions of MLCC requires an understanding of the factors that control the dielectric properties of single layers. In general, it is found that the permittivity of BaTiO$_3$ films rises substantially with the processing temperature [30, 31]. Typically, in order to achieve permittivities greater than >1000, coupled with low
loss tangents, process temperatures >800 °C should be employed. Among the substrates that have been employed for this purpose are Ni and Cu. For example, the thermal expansion coefficient of Ni foils (13x10^{-6}/°C) and barium titanate (cubic, 9.8x10^{-6}/°C) allows processing at temperature greater than 1000°C without film cracking. Dawley et al. (see Fig. 1.3a) have shown that a permittivity of 1500 can be obtained from highly oriented barium strontium titanate films deposited on Ni foils via chemical solution deposition with subsequent heat treatment at low oxygen pressure (900°C, 10^{-18} atm pO_2) [32]. Likewise, Nagata et al. (see Fig. 1.3b) has shown comparable permittivities in pure barium titanate films on high purity Ni foils [29]. Ihlefeld et al. (see Fig. 1.3c) reported much higher dielectric constants (ε_r ~ 2500) with high tunability from barium titanate films on Cu foil [33]. Following the procedures adopted in MLCCs production, Ihlefeld’s and Nagata’s work utilizes two step heat treatments: a sintering step and re-oxidation. Yuan et al. studied pulse laser deposited barium titanate thin film on Ni tape using NiO as a buffer layer [34]. The electrical data for these films appear to be characteristic of a lossy capacitor.
Fig. 1.3 Dielectric properties of a) (Ba,Sr)TiO$_3$ films on Ni foil [32], b) BaTiO$_3$ films on Ni foils [29] and c) BaTiO$_3$ films on Cu foil [33]

1.2 Objectives

Although barium titanate thin films on Ni foil have shown promising electrical properties, there are many unresolved questions that form the basis for the research presented in this thesis. Firstly, the science inherent in processing chemical solution derived films under low oxygen partial pressure conditions is not well understood. In particular, the role of pyrolysis conditions, rapid thermal annealing, the high temperature crystallization step, and the reoxidation in controlling the film thickness, residual carbon content, interface quality, and grain size are not at present, understood. Secondly, there are few methods available to non-destructively characterize the BaTiO$_3$/Ni interface. In
this work, X-ray diffraction, spectroscopic ellipsometry, and transmission electron microscopy are employed in an attempt to develop highly sensitive means for interrogating the interface in the model system of films on foils. Finally, the microstructure-processing-property relations in the films were explored.
References


[31] W. C. Yi, T. S. Kalkur, E. Philofsky, L. Kammerdiner and A. A. Rywak, "Dielectric Properties of Mg-doped Ba0.96Ca0.04Ti0.84Zr0.16O3 Thin Films Fabricated by Metalorganic Decomposition Method," *Applied Physics Letters*, 78 [22], 3517-3519 (2001).


2.1 Dielectric properties

The dielectric constant ($\varepsilon_r$) is a macroscopic measurement describing the ability of a material to polarize under an applied electric field, relative to that of vacuum. Fig. 2.1a shows a schematic of a parallel plate capacitor with vacuum as the dielectric. The charge stored in the capacitor when it is connected to a DC voltage is due to the permittivity of vacuum. When a dielectric is inserted between the metal plates, polarization in the dielectric creates bound surface charges (see Fig. 2.1b). The surface charges induce free charges on the metal plates, as shown in Fig. 2.1c. The capacitance (C) of the capacitor is proportional to the dielectric constant of the insulator and also depends on its geometry as described in Eq. 2.1.

$$C = \varepsilon_0 \varepsilon_r \frac{A}{d}$$  \hspace{1cm} \text{Eq. 2.1}

where $\varepsilon_0$ is the permittivity of vacuum ($8.85 \times 10^{-12}$ F/m). The thickness (m) and active area ($m^2$) of capacitor are given by $d$ and $A$, respectively.
Fig. 2.1 a) Capacitor with vacuum between two parallel plates

b) Polarization of charges and net surface charges at the faces of dielectric

c) Capacitor with inserted dielectric showing bound charges at the dielectric faces and an increase in free charge on the metal plates

Since the total charge developed on the metal plates is induced by the permittivity in vacuum and the polarization of the dielectric (P), the dielectric displacement (D) of the capacitor is described as [1]:

$$
\bar{D} = \varepsilon_0 \bar{E} + \bar{P}
$$

where $E$ is the applied electrical field. For a small AC signal, the polarization responds linearly to the electric field.

$$
\bar{P} = \varepsilon_0 \chi \bar{E}
$$

or

$$
\bar{D} = \varepsilon_0 \varepsilon_r \bar{E}
$$

where $\chi$ and $\varepsilon_r$ are the dielectric susceptibility and relative dielectric permittivity of the dielectric material, respectively and $\chi = \varepsilon_r - 1$. 
For an AC circuit, the polarization oscillates with the electric field. Because of inertia, part of the current is phase shifted with respect to the applied voltage (see Fig. 2.2a and b). As a result, there is current loss and energy dissipation in the ac circuit. Consequently, in an AC field, the permittivity of dielectrics is complex:

\[ \varepsilon_r = \varepsilon_r' + i\varepsilon_r'' \]

where \( \varepsilon_r' \) and \( \varepsilon_r'' \) are the real and imaginary parts of the complex dielectric constant (\( \varepsilon_r \)), respectively. \( i = \sqrt{-1} \). The dielectric loss is given by:

\[ \tan \delta = \frac{\varepsilon_r''}{\varepsilon_r'} \]

In reality, dielectrics are not perfect insulators, but contain some point defects, i.e. anion and cation interstitials or vacancies, as well as electronic carriers that lead to some conductivity. Therefore, practically, the measured \( \tan \delta \) consists of two types of loss: 1) dielectric loss due to inertia of charge oscillation and 2) leakage current due to flow of charge through dielectric or dc conductivity.
Fig. 2.2 Phasor diagram for a) perfect capacitor and b) real capacitor (V is applied voltage, I is the total current, I_C is the charging current and I_{loss} is the loss current)

The dielectric constant is attributed to various polarization mechanisms: electronic, ionic, dipolar (or orientational) and space charge polarization, as shown in Fig.2.3a [1]. The total polarization under an applied electric field can be described as

$$\vec{P}_{total} = \vec{P}_{elec} + \vec{P}_{ion} + \vec{P}_{dip} + \vec{P}_{sc}$$

Electronic polarization (\(\vec{P}_{elec}\)) occurs in all dielectrics. The process involves displacement of the nucleus with respect to the center of the electron cloud. Ionic polarization (\(\vec{P}_{ion}\)) is the relative displacement of anions and cations in ionic crystals. Dipolar (or orientational) polarization (\(\vec{P}_{dip}\)) results from aligning the permanent dipoles which exist in some dielectrics. Space charge polarization (\(\vec{P}_{sc}\)) is due to movement of charge carriers until they are blocked by insulating interfaces.

Each polarization mechanism has varying time response. A given polarization mechanism contributes to the dielectric only at frequencies that are low enough that the
charges can keep up with the oscillation. **Fig. 2.3b** shows a schematic of the dielectric constant and loss, showing the contributions of the polarization mechanism over a wide frequency range. The electronic and ionic polarization mechanisms respond to the applied field up to the optical ($\sim 10^{14}$ Hz) and infrared ($\sim 10^{11}$ Hz) regions, respectively. The polarization shows a resonance when the applied frequency matches the natural angular frequency of the electronic and ionic mechanisms. Dipolar and space charge polarization contribute to the dielectric constant at lower frequency. If the frequency is so high that dipoles cannot follow the alternating field, the polarization from the dipoles decreases and becomes zero. Relaxation processes are also associated with dielectric loss.
Fig. 2.3 a) Polarization mechanisms [2]

b) Frequency dependence of $\varepsilon'$ and $\varepsilon''$ [1]. Note that in ferroelectric materials, the dipolar polarizability is not lost until the GHz frequency range.
2.2 Barium titanate materials for capacitor applications

Barium titanate is the most widely utilized material for multilayer capacitors because of its high dielectric constant over a wide temperature range. The prototype structure of barium titanate is perovskite. The general formula of the perovskite structure is ABO₃, where A is a large cation with 1+, 2+, or 3+ ion charge (i.e. Ba²⁺) and B is a small and highly charged (3+, 4+ or 5+) cation (i.e. Ti⁴⁺). As shown in Fig. 2.4 for the cubic prototype, the Ba²⁺ ions (A site) are at the eight corners of the unit cell and the O²⁻ ions are at the face centers of the cube. Each Ba²⁺ ion is surrounded by 12 O²⁻ ions that are at the face centers of nearby cubes. The Ti⁴⁺ ion (B site) sits at the center of six oxygen ions, forming a TiO₆ octahedral cage. Another point of view (right hand side of the figure) shows a dodecahedral hole filled with Ba²⁺ and corner sharing of Ti octahedra in the perovskite framework.

In barium titanate, the cation sizes (Ba and Ti) do not perfectly fit the ideal perovskite structure, which is described using the tolerance factor [3]

\[
t = \frac{(R_A + R_O)}{\sqrt{2}(R_B + R_O)}
\]

where R is the radius of each ion. A, B and O are Ba, Ti and O, respectively. The tolerance factor for barium titanate (1.06, or >1), suggests that Ti⁴⁺ ion is too small for the octahedral site. At temperatures higher than Tc (130°C), the high symmetry cubic phase is stable. The cubic structure shows a high dielectric constant with paraelectric behavior, i.e. the net polarization returns to zero after removing an applied electric field. However, as temperature decreases, the TiO₆ cage is distorted by displacement of the Ti⁴⁺ ion off the center of the unit cell, and the cubic structure becomes unstable.
Fig. 2.4 Schematic of the cubic perovskite structure of the high temperature BaTiO₃ unit cell (Adapted from Kingery et al. 1991 [4])

Fig. 2.5 Phase transition sequence in BaTiO₃ (Figure from Moulson et al. [5])
Fig. 2.5 shows the structure changes in barium titanate with temperature. Below $T_c$, the structure is distorted by shifting the Ti$^{4+}$ ion closer to one O$^{2-}$ ion at a cube face (i.e. parallel to one of the <100> directions of the original cube). As a result, the cube is stretched along the edge parallel to the Ti$^{4+}$ displacement; the perpendicular edge contract, resulting in a tetragonal structure. Upon further cooling, the structure changes to orthorhombic and rhombohedral distortions at approximately 0°C and -90°C, respectively. The Ti$^{4+}$ ion is shifted along the pseudocubic <110> for the orthorhombic phase and <111> for the rhombohedral phase. Below $T_C$ (in the tetragonal, orthorhombic and rhombohedral structures), a spontaneous polarization is retained even in the absence of an electric field. These distorted structures possess ferroelectricity because the spontaneous polarization is reorientable.

The dielectric constant of a barium titanate single crystal as a function of temperature is shown in Fig. 2.6. The anomaly in the dielectric constant at $T_c$ is due to the high distortability of the structure at the phase transition. The dielectric constant above $T_c$ follows the Curie-Weiss law.

$$\varepsilon'_r = \varepsilon_o + \frac{C}{(T - T_o)}$$

where $C$ is the Curie constant and $T_o$ is the Curie-Weiss temperature. As barium titanate undergoes its first order transition, $T_o$ is slightly less than $T_c$ and a plot of the inverse dielectric constant ($1/\varepsilon$) has a finite value at $T_c$. Below $T_c$, the dielectric constant along the $c$ axis is smaller than that along the $a$ axis because the displaced Ti$^{4+}$ is tightly bound to the closest oxygen along the $c$ axis [6].
Fig. 2.6 Dielectric constant $\varepsilon$ (equivalent to $\varepsilon_r$) and its inverse as a function of temperature for a BaTiO$_3$ single crystal [7]

2.3 Chemical solution deposition of barium titanate thin films

BaTiO$_3$ thin films have been prepared by many techniques, i.e. rf-magnetron sputtering, pulsed laser deposition, molecular beam epitaxy, metalorganic chemical vapor deposition and chemical solution deposition [8-12] Chemical solution deposition (CSD) has some unique advantages over the others. In principle, a stoichiometric and homogeneous composition can be obtained due to reaction at the molecular level. Incorporation of dopants into the composition is conveniently done by adding the desired precursors to the solution. In addition, large film area and inexpensive deposition methods, i.e. spin coating and dip coating are very attractive for industrial production.
Chemical solution deposition involves fabricating thin films from a chemical solution, drying to remove the liquid in the gel network and performing appropriate heat treatment to convert the as-deposited film into the desired crystalline phase. The solution is prepared by dissolving precursors containing the necessary cation species in a solvent. The cation precursors are usually carboxylates i.e. metal acetates, M(OOCCH₃)ₓ and metalloorganic compounds i.e. alkoxides, M(OR)ₓ, where M is the metal and R is an alkyl group. The carboxylates are often used as precursors for lower-valent cations, i.e. Pb²⁺, Sr²⁺ and Ba²⁺ [13]. There are several chemical routes for making thin films. The two main routes: sol-gel or chelation are included here. Hybrid processes also exist. Other processes such as metallo-organic decomposition (MOD) [14, 15] and Pechini routes [16, 17] are also used, however they are not described here.

One of the solution synthesis routes is the sol-gel process. The standard reactions in the process are hydrolysis and condensation of the alkoxide reagents:

**Hydrolysis:**

\[ M(OR)_n + H_2O \rightarrow M(OR)_{n-1}(OH) + ROH \]

**Condensation (water elimination):**

\[ 2M(OR)_{n-1}(OH) \rightarrow M_2O(OR)_{2n-2}(OH) + H_2O \]

**Condensation (alcohol elimination):**

\[ 2M(OR)_{n-1}(OH) \rightarrow M_2O(OR)_{2n-3}(OH) + ROH \]

Hydrolysis is the reaction of a water molecule with the alkoxide, producing a hydroxyl metal alkoxide \( M(OR)_{n-1}(OH) \) and a released alcohol molecule \( (ROH) \).
Condensation is the further reaction of two hydroxyl metal alkoxides to form a polymeric species with metal-oxygen-metal (M-O-M) bonds.

The degree of the hydrolysis reaction can be controlled by many approaches. For a classical sol-gel process, the hydrolysis reaction occurs readily and precipitation of hydroxide species may occur. Therefore, dry chemicals and a dry atmosphere are required for solution synthesis to minimize the reaction. Alcohol exchange reactions also can slow down hydrolysis by forming a less reactive alkoxide (M(OR’)_x):

\[
M(OR)_n + xR'OH \rightarrow M(OR')_x + ROH
\]

where OR’ is a less reactive polar group of an alcohol i.e. 2-methoxyethanol, which is a widely used solvent for perovskite thin films.

Another synthesis approach, called the chelating process, involves modifying the alkoxide with chelating legands, i.e. acetic acid or 2,4-pentanedione (acetylacetone) to reduce the hydrolysis reaction of the alkoxide precursor. For acetic acid, the chelation reaction is [18]

\[
M(OR)_n + xCH_3COOH \rightarrow M(OR)_{n-x}(OOCC\text{H}_3)_x + xROH
\]

The term “hybrid” refers to the use of carboxylate precursors for the A site and alkoxides for the B site in perovskite thin films. The carboxylate acid, often acetic acid, serves both to dissolve the carboxylates and also as a chelating agent for the alkoxides.

In chemical solution deposition, thin films are usually deposited through spin coating or dip coating of the solution on a substrate. After deposition, the as-deposited film is amorphous and contains a large fraction of entrapped solvent and organics associated in the gel network. To obtain the crystalline phase, generally the film is first
heat treated at low temperature (200-400°C) for drying and organic pyrolysis and then subjected to higher temperature (500-700 °C) for crystallization and densification.
References


M. L. Liu and D. S. Wang, "Preparation of La$_{1.2}$Sr$_{0.2}$CO$_{1.7}$Fe$_y$O$_{3-x}$ Thin-films, Membranes, and Coatings on Dense and Porous Substrates," *Journal of Materials Research*, 10 [12], 3210-3221 (1995).

Chapter 3

Experimental Procedure

3.1 Chemical solution deposition of BaTiO₃ Film

In this work, BaTiO₃ thin films were prepared by chemical solution deposition, following the work of Nagata et al. [1]. Undoped BaTiO₃ with a 1:1 Ba:Ti ratio was chosen for the microstructure and interface studies since it provides an excellent model system.

3.1.1 BaTiO₃ solution

The starting materials in this experiment were barium acetate (99.999%, Sigma-Aldrich) and Ti-isopropoxide (99.99%, Sigma-Aldrich). Acetylacetone was used as a chelating agent for Ti-isopropoxide, which is known to be a moisture-sensitive reagent. The precursors and chemicals were batched in a glove box under an Ar atmosphere. Two types of solvent were used: methanol and 2-methoxyethanol. It was found that 2-methoxyethanol-based solutions provide better wetting of the first deposited layer on a Ni foil surface.

The process-flow for preparation of methanol-based BaTiO₃ solutions is shown in Fig. 3.1. First, 0.01 mole of barium acetate was stirred in 12 ml of glacial acetic acid at 105°C for 1 hour or until completely dissolved, using a rotary evaporator flushed with flowing Ar. The flask containing the barium precursor was immersed in a heated silicone oil bath and rotated at 120 rpm. In a separate flask, 2 ml of acetylacetone and 0.01 mole of titanium isopropoxide were combined. Then, the dissolved Ba precursor was poured
into the flask containing Ti. A small amount of acetic acid (6 ml) was used to rinse the remaining Ba solution from the flask. The mixture of Ba and Ti precursors was allowed to react at 105°C for 1 hour. An appropriate amount (80 ml) of methanol was added to obtain 0.1 M BaTiO₃ solution and then the solution was stirred at room temperature until homogenized. It is noted that the volume ratio of acetic acid to methanol was approximately 1:4.

Preliminary work has shown that for the same solution concentration, the 2-methoxyethanol-based film thickness is a factor of two less than that of the methanol-based films. Therefore, 0.2 M 2-methoxyethanol based solutions were made as shown in Fig. 3.2. Preparation proceeded exactly as described for the methanol-based solution, including the dissolution of the barium acetate and the mixing of the Ba and Ti precursors. However, the mixture was vacuum distilled at 105°C until the volume of the solution was approximately 10 ml, before re-diluting the solution to the desired molarity. It is expected that the majority of the volume loss was due to acetic acid since its boiling point (~ 118°C) is lower than that of acetylacetone (~ 140°C). Then, 40 ml of 2-methoxyethanol was added and stirred until homogenized.

Keeping the ratio of acetic acid to solvent to 1:4 served two purposes. First, the possibility of changing the solution characteristics (and so factors such as the degree of cross-linking, etc.) is avoided. Secondly, it was found that if the concentration of acetic acid in the solution was too high, it attacked and damaged the Ni foil surface. This complicated interpretation of the resulting spectroscopic ellipsometry data due to the roughened interface.
Fig. 3.1 Schematic of methanol-based solution preparation method

Fig. 3.2 Schematic of 2-MOE-based solution preparation method
3.1.2 Film deposition and crystallization

High purity Ni foils (99.99%, Alfa Aesar) that were 25 μm thick were used to prepare BaTiO$_3$ films on Ni substrates. The Ni foil was pre-annealed at 900°C in approximately 1x10$^{-16}$ atm pO$_2$, using a reducing furnace which will be described in section 3.2. The pre-annealing step was to smooth the Ni surface and remove native oxide and organic contamination, if present [1]. Since rolled Ni foils have a finite level of surface roughness, which will be presented in chapter 5, Al$_2$O$_3$ and Si wafers were also used as substrates in some cases to provide smoother surfaces.

**Fig. 3.3** shows a schematic of the preparation of BaTiO$_3$ thin films. A syringe with a 0.1 μm filter was used to dispense the solution onto the substrate. After covering the substrate surface with solution, the spinner was rotated at 3000 rpm for 30 seconds. After each deposition, the films were dried at 180°C for 3 min and then 280°C for 3 min on hotplates in air. After every two spinning/drying steps, the organic was further removed at 350°C for 1 min. Films were then heat-treated at temperatures between 600 and 750°C for 1 minute under a N$_2$ gas flow (23 SLPM) in a rapid thermal annealer (RTA 600, Modular Process Technology Corp., San Jose, CA). Spinning, drying and RTA annealing were repeated until the desired thickness was obtained. The films were then crystallized at 1000°C for 1 hour in 5x10$^{-17}$ atm pO$_2$ (setting value) in a reducing furnace and then re-oxidized at 600°C for 30 minutes in 1x10$^{-10}$ atm pO$_2$ (setting value). The heating rate of both steps in the reducing furnace was 5°C/min. The conditions employed in this work are comparable to those reported by Nagata et al. [1]. A schematic showing the oxygen pressures and temperatures employed for some of the steps is shown along with the Ellingham diagram in **Fig. 3.4**. It is clear that the conditions used for
reoxidation could lead to oxidation of the Ni foil, if the BaTiO$_3$ layer does not provide a sufficient kinetic barrier against oxygen diffusion.

![Fig. 3.3 Schematic of preparation of BaTiO$_3$ thin films](image)

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Fig. 3.3 Schematic of preparation of BaTiO$_3$ thin films
Fig. 3.4 Stability of Ni and various oxidation state of titanium (Circles, ○ and ●, shows typical setting values and measured values for the pO₂ for each heat treatment) (Adapted from Yang et al. [2])

3.2 Reducing furnace system

As mentioned in sections 3.1.1 and 3.1.2, high temperature annealing of either the Ni foil or BaTiO₃ films on Ni foils requires a reducing atmosphere to prevent oxidation of the Ni. In this experiment, high purity nitrogen and either hydrogen or forming gas (5% H₂ and 95% N₂) were flowed into a tube furnace to control the oxygen partial pressure. Part of the nitrogen gas was directly fed to the furnace (dry nitrogen) while an appropriate amount was passed through water to carry water vapor into the furnace (wet
nitrogen). The oxygen partial pressure within the furnace was controlled by the equilibrium of hydrogen and water vapor through the following chemical reaction.

\[
\text{H}_2(\text{g}) + \frac{1}{2}\text{O}_2(\text{g}) \leftrightarrow \text{H}_2\text{O}(\text{g}); \quad K_p = \frac{P_{\text{H}_2\text{O}}}{P_{\text{H}_2}P_{\text{O}_2}^{\frac{1}{2}}} \tag{3.1}
\]

and

\[
\ln K_p = -\frac{\Delta G^\circ}{RT} \tag{3.2}
\]

\(K_p\) is the equilibrium constant which is dependent on temperature (T in Kelvin) and \(\Delta G^\circ\) (J) is the standard Gibbs free energy of formation of water vapor [3].

\[
\Delta G^\circ = -246,740 + 54.81T \tag{3.3}
\]

For a desired temperature and oxygen partial pressure, \(K_p\) is calculated from equation 3.2 and 3.3, then the ratio of \(P_{\text{H}_2\text{O}}\) and \(P_{\text{H}_2}\) required in the furnace were obtained. The total pressure and total gas flow in the furnace were 1 atm and 500 cc/min, respectively. Once the flow rate of \(\text{H}_2\) is defined (usually in the range of 1 to 50 cc/min), the flow rate of wet \(\text{N}_2\) (which helps determine the water vapor pressure, \(P_{\text{H}_2\text{O}}\)) is calculated. The dry \(\text{N}_2\) is added to maintain the total gas flow (500 cc/min). Typical gas flows for different reducing conditions, i.e. crystallization and re-oxidation are shown in Table 3.1. For the re-oxidation condition (600°C), \(P_{\text{H}_2} \approx 2P_{\text{O}_2}\) is used in equation 3.1 because there is no \(\text{H}_2\) gas in the system.

**Fig. 3.5** is a schematic of the reducing furnace. The pressure of water vapor and hydrogen was controlled by mass flow controllers (MFC, MKS Type 1179A). The outgoing gas was passed through a flask containing silicone oil to insure flow by
checking the bubble rate. An extra empty flask was put between the end of the furnace and the oil container to trap any oil from backstreaming into the furnace system. An oxygen probe with a zirconia-based sensor and a thermocouple was used to read the actual oxygen partial pressure and temperature at the center of the furnace. One sensor surface is exposed to the reducing atmosphere in the furnace while the other is in contact with an air reference containing 20% oxygen. The voltage developed due to the difference in oxygen content between the two surfaces was measured. Then, using an independent measurement of temperature, the oxygen pressure in the furnace was calculated from the Nernst equation.

\[
\text{pO}_2 \text{(atm)} = 0.209 \exp \left( -\frac{46.42E}{T} \right)
\]

where \( E \) is the oxygen signal (mV) and \( T \) is temperature (K). Since the operating temperature of the sensor is between 700°C and 1700°C, it is reasonable that the pO\(_2\) read during the 600°C re-oxidation step may differ significantly from the setting value, as shown in Table 3.1.
Fig. 3.5 Schematic of reducing furnace system

Table 3.1 Reducing setting condition and actual pO$_2$ measured by an oxygen sensor

<table>
<thead>
<tr>
<th>Reducing condition (°C, atm pO$_2$)</th>
<th>Gas flow rate (SLPM)</th>
<th>Actual pO$_2$ (atm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H$_2$</td>
<td>Wet N$_2$</td>
</tr>
<tr>
<td>900°C, 1x10$^{-16}$</td>
<td>10</td>
<td>136.5</td>
</tr>
<tr>
<td>1000°C, 5x10$^{-17}$</td>
<td>10</td>
<td>40.98</td>
</tr>
<tr>
<td>600°C, 1x10$^{-10}$</td>
<td>-</td>
<td>27.9</td>
</tr>
</tbody>
</table>
3.3 Spectroscopic ellipsometry

3.3.1 Introduction to ellipsometry

Spectroscopic ellipsometry (SE) was used to determine the depth profile, film thickness, microstructure and optical constants of the BaTiO$_3$ films on Ni foils and other substrates. Ellipsometry measures the change in the polarization state of a light beam after reflection from a sample. For the rotating analyzer spectroscopic ellipsometer which was used in this experiment, linearly polarized light is incident on a sample and elliptically polarized light is produced, as illustrated in Fig. 3.6. The basic configuration of the ellipsometer is shown in Fig. 3.7. White light from a Xe lamp is separated into different wavelengths by a monochromator. The monochromatic light is passed through a polarizer before hitting the sample surface. At each wavelength, the intensity of the reflected light is measured as a function of analyzer position by a photomultiplier tube (PMT), thus providing a measurement of the polarization state of the reflected light.

![Reflection of light at an interface](image)

**Fig. 3.6** Reflection of light at an interface
Two parameters, the relative phase shift ($\Delta$) and amplitude change ($\tan \Psi$) of the parallel ($E_\parallel$) and perpendicular ($E_\perp$) components of the light wave, are measured [5].

$$\Delta = \delta_r - \delta_i$$ \hspace{1cm} (3.4)

and

$$\tan \Psi = \frac{|r_\parallel|}{|r_\perp|}$$ \hspace{1cm} (3.5)

where $\delta$ is the phase difference between $E_\parallel$ and $E_\perp$ of the reflected and incident light; the subscript r and i denote the reflected and incident waves. The parameters $r_\parallel$ and $r_\perp$ are the complex Fresnel reflection coefficient of $E_\parallel$ and $E_\perp$, respectively. For reflection of light from a homogeneous isotropic material, the Fresnel coefficients are [5]:
where $\phi_0$ and $\phi_1$ are angles of incidence and refraction, respectively and $N_1$ and $N_2$ are the complex refractive indices of air and the sample, respectively at the wavelength ($\lambda$) used.

For multiple interfaces, like a thin film on a substrate as shown in Fig. 3.8, the total reflection coefficients count reflections from all interfaces and are defined as

$$r_{//} = \frac{r_{//,01} + r_{//,12}e^{-2i\beta}}{1 + r_{//,01}r_{//,12}e^{-2i\beta}}$$

Fig. 3.8 Reflection of light from a thin film on a substrate
and

\[ r_\perp = \frac{r_{\perp,01} + r_{\perp,12}e^{-2i\beta}}{1 + r_{\perp,01}r_{\perp,12}e^{-2i\beta}} \]  \hspace{1cm} (3.9)

where \( r_{\parallel,ij} \) and \( r_{\perp,ij} \) are the Fresnel reflection coefficients of the parallel and perpendicular waves traveling from medium \( i \) to \( j \), respectively.

\[ \beta = 2\pi \left( \frac{L}{\lambda} \right) N_1 \cos \phi_1 \]  \hspace{1cm} (3.10)

When there are multiple films, the measured delta and psi are influenced by the optical depth profile of the entire stack. \( \Delta \) and \( \tan \Psi \) can be expressed as:

\[ \tan \Psi e^{i\Delta} = \frac{r_{\parallel}}{r_\perp} = \rho(N_1, L_1, \lambda) \]  \hspace{1cm} (3.11)

Therefore, the measured \( \Delta \) and \( \Psi \) contain information on the optical properties and thickness of the film(s). Because it is relative measurement, rather than an absolute measurement of intensities, ellipsometry is a very accurate and reproducible. Data at different wavelengths obtained from spectroscopic measurements enable analysis of samples with complicated depth profiles.

### 3.3.2 Data collection

System alignment and calibrations of the rotating-analyzer spectroscopic ellipsometer were performed as described by Chindaudom [6]. Since the Ni foil is flexible, BaTiO\(_3\) films on Ni foils were flattened and fixed onto a glass slide using double
sided tape. The sample was put on the SE stage and aligned. The SE parameters, delta ($\Delta$) and psi ($\Psi$), were collected at wavelengths from 250 to 750 nm with an increment of 5 nm at an incident angle of 70 degrees. For samples in which there was at least one absorber in the film stack, the incident light was linearly-polarized at 30°. At each wavelength, data from 50 turns of the analyzer were averaged. Background spectra (obtained with the PMT voltage set to the value during the measurement, but with the light beam blocked by a shutter) were also collected and subtracted from the measured data before calculating $\Delta$ and $\Psi$.

### 3.3.3 SE measurement with a compensator

It has been reported that the error in $\Delta$ for a rotating analyzer ellipsometry measurement is proportional to $1/\sin\Delta$ [7]. Therefore, for a transparent film on a transparent substrate, the measured $\Delta$ is inaccurate because the value is near 0° or 180° for the whole measured spectrum [6, 7]. To resolve the problem, an achromatic compensator, or a quarter-wave plate, is used to add a 90° phase shift to the incident beam. As a result, accurate measurements of $\Delta$ near 90° can be obtained.

In this thesis, the compensator was used to measure $\Delta$ and $\Psi$ of BaTiO$_3$ films on Al$_2$O$_3$ substrates. The polarizer was fixed at 45° and the compensator was inserted between the polarizer and the sample to obtain a 90° phase shift or circularly-polarized incident light. Two SE measurements were required. First, the true phase shift induced by the compensator was measured without any sample in the straight-through configuration (i.e. with a 90° incident angle) for the entire wavelength range. Then, the
sample was aligned at a 70° angle of incidence and another set of data was measured. The Δ and Ψ of the sample was calculated by [6]

\[
\tan \Psi e^{i\Delta} = \frac{\tan \Psi_2 e^{i\Delta_2}}{\tan \Psi_1 e^{i\Delta_1}}
\]

where the subscripts 1 and 2 refer to the straight-through measurement and the sample measurement, respectively. It is noted that the back surface of the Al₂O₃ was roughened using a diamond file and then painted with a carbon adhesive to remove reflection from the back surface of the substrate.

### 3.3.4 SE data modeling

Spectroscopic ellipsometry is an indirect characterization method because the measured data need to be modeled. First, a model is constructed assuming discrete layers of optically homogeneous materials parallel to the sample surface. Then, the thickness and optical properties of each layer were used to calculate Δ and Ψ as defined in equation 3.11. To minimize the number of parameters that must be fit, it is important to be able to input as much information as is known for the optical properties of the various layers in the sample.

For known materials, reference data of the optical properties were used if available. When the optical properties of the materials were not known, it was necessary to generate the data. Several approaches were utilized here. First, if the SE measurement is performed on an ideally smooth surface of a homogeneous material (i.e. a substrate),
the complex refractive index of the material can be directly inverted from the measured $\Delta$ and $\Psi$ by [6]

$$N_1 = \tan \phi_0 \left[ 1 - \frac{4\rho \sin^2 \phi_0}{(1+\rho)^2} \right]^{\frac{1}{2}}$$ \hspace{1cm} 3.13

where $N_1$ is the complex refractive index. However, generally, the sample surface is not perfectly smooth. Therefore, the complex refractive index obtained from the direct inversion without correcting for the real structure of the sample is not accurate. Since the surfaces of the Ni foils are very rough, this approach could not be employed. Instead, two sets of reference data of Ni were used (see section 5.1) Secondly, for unknown dielectric materials, a damped Sellmeier dispersion equation was used to describe the optical properties of the unknown phase [8].

$$N^2(\lambda) = A_1 + \frac{A_2 \lambda^2}{(\lambda^2 - A_3^2 - 2i\lambda A_4)}$$ \hspace{1cm} 3.14

where $A_1$-$A_4$ are fitting constants. In all cases, $A_1$ was assigned to 1.

If any layer in the model had two constituents, consisting of $f_A$ volume percent of phase A and $f_B$ volume percent of phase B, Bruggeman effective medium theory was applied to calculate the optical properties of the layer [5].

$$f_A \frac{\varepsilon_A - \varepsilon}{\varepsilon_A + 2\varepsilon} + f_B \frac{\varepsilon_B - \varepsilon}{\varepsilon_B + 2\varepsilon} = 0$$ \hspace{1cm} 3.15

where $\varepsilon$ is the complex dielectric function of the composite. This was used, for example, to calculate the net optical properties of low density layers as a mixture of the material and air.
Using one set of values for the optical properties and thickness of each layer, it is possible to calculate the Fresnel coefficients and the $\Delta$ and $\Psi$ values for each wavelength. Once this is done, an unbiased estimator of the error in the fit was then calculated [8]:

$$\sigma = \frac{1}{m - p - 1} \left[ \sum_{i=1}^{m} \left\{ (\cos \Delta_{i,\text{exp}} - \cos \Delta_{i,\text{cal}})^2 + (\tan \Psi_{i,\text{exp}} - \tan \Psi_{i,\text{cal}})^2 \right\} \right]^{1/2}$$

where $\Delta_{i,\text{exp}}, \Psi_{i,\text{exp}}$ are the measured data; $\Delta_{i,\text{cal}}, \Psi_{i,\text{cal}}$ are the calculated values from the optical model at the $i^{th}$ wavelength; $p$ is the number of unknown variables in the model and $m$ is the number of data points. A grid search in parameter space was then performed to obtain reasonable initial model parameters, where the values chosen were those that led to a minimum value for $\sigma$. Then, the program iterated around the best initial parameters, using a linear regression analysis. Once a best fit was achieved for a given model, a new model was attempted, and the process was repeated. For modeling of a BaTiO$_3$ film on a transparent substrate such as sapphire, the cosine of delta is very small, which underweights the importance of the $\Delta$ data. Thus, in this case, a difference of delta values was used to calculate sigma, rather than a difference in $\cos \Delta$ [6].

The best fit model was chosen based on the following criteria [8]:

1) A physically reasonable model

2) A small value of $\sigma$

3) Good agreement between experimental and calculated data

4) Reasonable values for the 90% confidence limits

5) Low values for the cross-correlation coefficients
3.4 Microstructural characterization

The microstructure and interface quality of BaTiO$_3$ films on Ni foils before and after a 1000°C furnace anneal were observed in a transmission electron microscope (Philips 420) with a 120 keV tungsten source. This enabled observation of changes in grain size, grain morphology, and the presence of second phases in the films.

Cross-section TEM samples were prepared either by conventional polishing and ion milling or a focused ion beam technique. It was found that conventional polishing is useful for BaTiO$_3$/Ni foil samples after furnace annealing. However, RTA annealed films readily detached from the Ni foil during polishing, as shown in Fig. 3.9. Therefore, focused ion beam milling was utilized for such samples as it enables preparation of cross-section TEM samples without a mechanical polishing step [9, 10].

![Fig. 3.9 Bright field TEM image showing BaTiO$_3$ film detached from the Ni foil due to conventional polishing preparation. This complicated studies of the interface quality.](image)

3.4.1 Conventional polishing and ion milling

Preparation of cross-section TEM specimens from Si single crystal substrates was described by Bravman et al. [11]. In this work, Prof. Ian Reaney of Sheffield University was extremely helpful in suggesting appropriate ways of preparing TEM samples using polishing and milling. Since the Ni foils employed here are quite thin (25 µm) and hard to handle, buffers were used to support the foils for polishing. In this experiment, Si wafers were chosen as the mechanical supports due to their ready availability. Two pieces of approximately 3 mm square were cut from a BaTiO₃ film on Ni foil. Ni foils are easily bent; therefore, the pieces were flattened by gently pressing them between two glass slides. Two Si pieces of equivalent size were also cut and cleaned in acetone. To make a cross-section stack, the Si wafers and the samples were glued in the following order: Si/Ni-BT/BT-Ni/Si, using M-bond 610 adhesive (Vishay Micro-Measurements, Raleigh, NC 27611). Both the BaTiO₃ films and the polished side of the Si pieces faced the center of the stack. The stack was heated on a hotplate at 150°C for 30 min to cure the glue. Pressure was applied to the stack to minimize the thickness of the glue layer and to avoid trapping air bubbles. Minimizing the thickness of the glue between the two BaTiO₃ surface also provided better uniformity during ion milling.

For the cross-section samples, the stack was mounted to a polishing stub using crystalbond. The sample was then gently polished with the polishing direction parallel to the interfaces. Polishing normal to the interface tended to break the stack and worsen the smear of the ductile Ni on top of the BaTiO₃. Both sides of the cross-section were polished using SiC paper #600 (16 µm grain) finished by #1200/4000 (5 µm grain) with a water rinse. To increase the mechanical stability of the stack during thinning, the first
completely polished side was glued onto a copper grid using M-bond adhesive before polishing the other side, as shown in Fig. 3.10. After thinning down to 40 µm thick, the sample on copper grid (3 mm in diameter) was cleaned in acetone and isopropanol. Then, it was further thinned using an EA Fishione Model 3000 ion mill operating at 5 kV and 5 mA. The inclination angle was set at 12 degrees and then 10 degrees to finish the ion milling.

**Fig. 3.10** Preparation of cross-sectional TEM samples by conventional polishing
3.4.2 Focused ion beam (FIB) TEM specimen preparation

Cross-sectional TEM samples of BaTiO$_3$ films on Ni foils were also prepared using an FEI Quanta 200 3D dual beam FIB (with the assistance of Trevor Clark). The system consists of an ion column (Ga), an electron column, a gas delivery system and a micromanipulator station. The system provides capability for ion milling, real-time monitoring of the sample and material (i.e. W) deposition. W deposition was performed using ion beam assisted chemical vapor deposition of chemical precursors.

In this experiment, the lift-out technique was used to prepare cross-sectional samples. The technique entails directly cutting a cross-sectional sample from the substrate and transferring it to a TEM grid where further thinning is performed. In this case, the BaTiO$_3$/Ni foil sample was coated with conductive carbon before mounting in the FIB to prevent charging as a result of ion/electron bombardment.

A 20 µm thick tungsten strip was first deposited on the desired region (20x2 µm$^2$), as shown in Fig. 3.11. This layer protects the BaTiO$_3$ surface underneath during subsequent ion milling. Then, the material surrounding the area of interest was wedge-cut free by ion milling. Then the piece of sample was lifted out and attached to a Cu grid (see Fig. 3.12a) using an omni probe. It is noted that an omni probe was attached to the sample by W deposition. On the Cu grid, the material at both sides along the interface of the sample was milled away until the width of the W strip was approximately 0.7-1 µm, as shown in Fig. 3.12b. The milling depth was just about half of the sample thickness, not all the way through the thickness. The material at either end and the bottom of the sample was retained to keep the sample mechanically stable. Then, the sample was further milled using an ion beam current from 300 to 100 pA, with alternate sample
tilting at 1 degree from the normal ion beam to enable milling on both sides. The final milling (100 pA) was finished when there was no W left in at least one region, as shown in Fig. 3.12e. It is noted that, as shown in Fig. 3.12c-e, one-third of the sample was thinned at a time.

Fig. 3.11 Diagram of the FIB lift out process
Fig. 3.12 (a – c) Series of top-view secondary ion images showing the thinning procedure for cross-sectional TEM sample by FIB a) a lift-out sample is attached to a Cu grid b) the sample is thinned by milling along the interface c) one-third of the sample is further thinned. (d-e) side view secondary electron images d) side view of c), e) final sample which is thinned until all of the W in a region disappears
3.4.3 Chemical analysis by electron energy loss spectroscopy (EELS)

A JEOL (2010F) transmission electron microscope with a field emission electron source combined with a Gatan Enfina parallel electron energy-loss spectrometer (EELS) was used for chemical analysis of BaTiO$_3$ films on Ni foils (with the assistance of Dr. Gaiying Yang). The microscope was operated at an accelerating voltage of 200 kV and the EELS spectra were acquired using diffraction mode. The resolution of the EELS spectra was 1.1 eV at the zero loss peak and the electron collection angle was 14 mrad.

Cross-sectional samples of furnace annealed BaTiO$_3$ film were prepared by conventional polishing, as described in section 3.4.1. Initial attempts to prepare the sample as described above resulted in very long milling times. To reduce the milling damage for the EELS specimens, the final thinning was performed on a cooling stage (using liquid N$_2$) in the EA Fishione Model 3000 ion mill. Nonetheless, it is possible that the long milling time may have influenced the oxidation state of the BaTiO$_3$ film.

3.5 Other characterization

3.5.1 Structural characterization

The crystallinity of the BaTiO$_3$ films on Ni foils was monitored routinely using a Scintag V X-ray diffractometer (Scintag Inc., Cupertino CA) with Cu K$_\alpha$ radiation. Data were collected from 20 to 60° 2θ with a step size of 0.02° and a 0.5 second acquisition time per step. A slow scan with a count time of 5 second was also used when a small amount of some phase, i.e. NiO, was expected.

3.5.2 Surface roughness

The surface roughness of Ni foil and BaTiO$_3$ films on Ni foil was determined by atomic force microscopy (Multimode IIIa, Digital Instruments), using tapping mode over
a scanning area of 5x5 µm or 3x3 µm, depending on feature size of the samples. The Si cantilever of 160 µm long was operated at its resonance frequency of 300 kHz.

3.5.3 Electrical measurement

The room temperature dielectric constant and loss tangent of the BaTiO$_3$ films were measured using a Hewlett-Packard 4192A impedance analyzer (Agilent Technology, Palo Alto, CA). The frequency dependence (0.1 kHz-1 MHz) was measured using an oscillating voltage of 0.03 V, using Ni and Pt as bottom and top electrodes, respectively. Capacitance as a function of voltage was measured by superimposing a small AC voltage (0.03 V) on DC Bias. Typical electrode size for these measurements is ∼50 µm in diameter.

The temperature dependence of dielectric constant was measured using a Hewlett-Packard 4284A LCR meter. The measurement was performed at 0.03 V AC at different frequencies (0.1 kHz-1 MHz). The capacitance was measured over a temperature range of 25-200°C with a heating rate of 2°C/min. A typical electrode size for this measurement is ∼150 µm in diameter.

The top electrode pattern was prepared by lift-off photolithography and Pt sputtering. First, LOR5A, used for better undercut control, was spin coated on the film at 4000 rpm for 45 s and then baked at 180°C for 2 min. After the sample was cooled down, photoresist was spin coated on the top of film using the same spinning conditions and baked at 100°C for 1 min. Then, the top of sample was exposed to UV (∼300 W) through a photomask for 30 s. The photoresist and LOR pattern were developed using CD-26 (Shipley, Marlboro MA) and rinsed with deionized water. A 1000 Å thick Pt film was deposited using the W.M. Keck Smart Materials Integration Laboratory (Lesker CMS-18)
sputtering system. The substrate was held near room temperature and sputtering was done using an Ar pressure of 2.5 mTorr. The sample was dipped in acetone and then CD-26 to lift-off the unnecessary Pt, leaving only the Pt electrode pattern on the film surface.
References


Chapter 4
Cross-sectional Transmission Electron Microscopy Studies
of BaTiO$_3$ Films on Ni Foils

This chapter details the use of transmission electron microscopy (TEM) to observe the microstructure and interface quality of BaTiO$_3$ films on Ni foil after heat treatment in either the RTA or the reducing furnace. The BaTiO$_3$ films used in these experiments were obtained from 0.1 M methanol-based BaTiO$_3$ solutions. The deposition method was described in section 3.1.2. Two layers were deposited on the pre-annealed Ni foil; each layer was dried and pyrolyzed. After deposition, the film was annealed in an RTA at temperatures of 700-750°C. Another two layers were deposited and the drying, pyrolysis and RTA annealing were repeated until 8 layers were processed. Then, the 8 layer film was annealed in the reducing furnace at 1000°C and re-oxidized at 600°C. Either focused ion beam (FIB) milling or a conventional polishing/milling method was used to prepare cross-sectional TEM samples, as described in section 3.4. A discussion of damage due to FIB milling of an RTA annealed BaTiO$_3$ films is in section 4.1. The microstructures of RTA’d films and furnace annealed films are presented in sections 4.2.1 and 4.2.2, respectively.

4.1 Damage Artifacts Produced by FIB Milling

As described in section 3.4, cross-sectional TEM samples of low temperature annealed (RTA) BaTiO$_3$ films could be prepared without the films detaching from the Ni
foil by FIB milling. It is well-known that the rastered Ga\(^+\) ion beam used in FIB milling creates several forms of damage (i.e. ion implantation, amorphization and re-deposition of milled materials) on the machined surface that can complicate TEM studies, including imaging, structural and chemical analysis [1-4]. Therefore, it is necessary to understand the nature and cause of the damage both to minimize damage issues in a specific sample and to avoid inaccurate TEM study and analysis.

Fig. 4.1a shows a large, uniform, electron-transparent area in a cross-section of a BaTiO\(_3\) film on Ni foil prepared by FIB. The cross-section shows the Ni foil, BaTiO\(_3\), conductive C and a W layer (which was deposited in the FIB before cutting and thinning the TEM sample). In the figure, it is noticeable that the BaTiO\(_3\) film exhibits discrete layers of crystallized grains. The layers are visible through the film thickness in areas where the thick W protective layer is intact. Fig. 4.1b shows an undamaged film cross-section. In contrast, where the W layer is thinner, the BaTiO\(_3\) grain structure at the top of the film is difficult to resolve. The disappearance of this feature is believed to be a result of damage induced by ion beam milling. These results suggest that the damage occurs mainly during final thinning, rather than during deposition of W layer. For these images, the final thinning conditions used a 30 keV, 100 pA beam. Although this current is already low, it is possible that the ion energy could be reduced to 5 – 10 keV to further reduce the damage. Alternatively, it might be possible to minimize the impact of ion-bombardment induced damage by finishing sample preparation using an Ar\(^+\) ion beam.

There are no reports available on the mechanism for damage in similar samples. Nonetheless, the loss of the grain structure suggests that perhaps amorphization of the surface is responsible [5, 6].
Fig. 4.1 TEM cross-sectional micrograph of 750°C annealed BaTiO$_3$ films on Ni foil showing a) partial film damage by ion milling at the top of the film and b) discrete crystallization layers in a region where the film is thick and a large amount of W is still intact. BT = BaTiO$_3$
4.2 Microstructure

4.2.1 RTA annealed BaTiO$_3$ films

Discrete crystallization layers are apparent in the BaTiO$_3$ films annealed at 700°C and 750°C. There is no significant difference in grain size across the film thickness, as clearly shown in the undamaged area in Fig. 4.1b. The high magnification bright-field TEM micrographs in Fig. 4.2a and b show the microstructure of the BaTiO$_3$ films on Ni foil after RTA annealing at 700°C and 750°C, respectively. It is noted that an additional Au layer was deposited between the BaTiO$_3$ and C coat in order to provide better contrast of the film surface. (Although Au is conductive, it is easily milled by Ga$^+$ ions. Therefore, the C layer is necessary for charging protection.) A BaTiO$_3$ thickness of approximately 200 nm was achieved with 8 coatings and 4 RTA anneals, yielding 25 nm for each layer of deposition. Although the TEM samples were partially damaged, it is shown that both RTA’d BaTiO$_3$ films have equiaxed polycrystalline grain structures. The grain size of the BaTiO$_3$ film annealed at 700°C varied from 5 to 20 nm with an average grain size of 12 nm. On increasing the RTA temperature to 750°C, the average grain size slightly increases to 18 nm and the grain size is between 10 and 30 nm. Both films contain a few nano-size (<10 nm) pores and no cracking was observed. The BaTiO$_3$ / Ni foil interfaces in samples annealed at 700° and 750°C show no evidence of any interfacial layer such as NiO.
Fig. 4.2 TEM bright-field micrograph of microstructure and interface of BaTiO$_3$ films on Ni foils annealed on RTA at a) 700°C and b) 750°C
The grain size of the chemical solution deposited based-BaTiO$_3$ films in this work is compared to previous reports in Table 4.1. In the table, the substrate type and heat treatment conditions of the films is included. The morphology of most of the films was reported to be equiaxed; it is noted if the authors stated otherwise. The grain size range of 5-30 nm of the RTA’ed films at 700$^\circ$C and 750$^\circ$C in this experiment was small. Neglecting the difference in precursor chemistry and substrate materials, this small grain size is reasonable, compared to the reported grain size of 15-20 nm of (Ba,Sr)TiO$_3$ films on Pt/Si substrates prepared using similar conditions [7]. A larger grain size (20-60 nm) was reported for films that were annealed at 700$^\circ$ or 750$^\circ$C for longer times (0.5-1 hr) [8-10].

The appearance of consecutive layers of BaTiO$_3$ grains confirmed that crystallization occurred during each RTA annealing at both 700$^\circ$C and 750$^\circ$C. Perovskite BaTiO$_3$ was also apparent in electron diffraction patterns of the RTA’ed films. Fig. 4.3 shows sample areas and the corresponding diffraction patterns of BaTiO$_3$ films on Ni foils. Fig. 4.3a-b shows a film crystallized using an RTA temperature of 700$^\circ$C. A 0.4 $\mu$m aperture covering only the Ni and BaTiO$_3$ was used to avoid diffraction from Au or C. Only BaTiO$_3$ rings appear, implying the observed area did not have NiO at the interface. Only two diffraction spots were seen for the Ni substrate. Fig. 4.3c-d shows the results for a furnace annealed film. A larger number of more continuous rings were observed because an aperture of 0.8 $\mu$m was used to cover more film area. Similar to the RTA’ed film, only perovskite BaTiO$_3$ rings appear and there are two Ni spots.

AFM was used to confirm the surface morphology of the films, as shown in Fig. 4.4. In AFM scans over a 0.5 x 0.5 $\mu$m area for BaTiO$_3$ films annealed at 700$^\circ$C and
750°C (See Fig. 4.4a), a somewhat larger feature size is observed, compared to that obtained from the cross-sectional TEM sample. This discrepancy is believed to be due to detection of clusters of BaTiO$_3$ grains, rather than primary grains, by AFM.

The equiaxed, polycrystalline structure of the BaTiO$_3$ films suggests bulk nucleation in this work. However, it has been shown elsewhere that the substrate materials and preparation conditions can also affect the microstructure of the films. Sedlar et al. showed isolated large grains in a Ce-doped BST film at the film/Pt-Si interface. In that case, the grain size decreased from the interface to the film surface because of multiple coating and annealing at 750°C [7]. Gust et al. observed that heterogeneous nucleation at the film/Pt-Si interface competes with major bulk nucleation due to the small lattice mismatch between the BaTiO$_3$ film and Pt (~3%) [9]. Using Pt-Si substrates, chemical solution deposited BaTiO$_3$ (or SrTiO$_3$) films with columnar structure can be achieved by controlling film thickness and performing multiple coating and annealing. It was shown that depositing very thin layers (10-20 nm) and annealing after each layer can induce nucleation at the bottom surface and local epitaxial growth within each columnar grain. This effectively suppresses bulk nucleation [11-13].
Table 4.1 Reported grain size and average grain size (in parentheses) of BaTiO$_3$ – based films prepared by various groups under a variety of different conditions

<table>
<thead>
<tr>
<th>Film/substrate</th>
<th>Annealing condition</th>
<th>Grain size and (avg grain size)</th>
<th>Total Thickness (nm)</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T(°C)/Time</td>
<td>Atmospheric</td>
<td></td>
<td></td>
</tr>
<tr>
<td>This work:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BaTiO$_3$/Ni foil</td>
<td>700°C, 1 min*</td>
<td>N$_2$</td>
<td>5-20 (12)</td>
<td>200</td>
</tr>
<tr>
<td></td>
<td>750°C, 1 min*</td>
<td>N$_2$</td>
<td>10-30 (18)</td>
<td>200</td>
</tr>
<tr>
<td></td>
<td>1000°C, 1 hr</td>
<td>10$^{-16}$</td>
<td>24-75 (42)</td>
<td>200</td>
</tr>
<tr>
<td>BaTiO$_3$/Pt-Si</td>
<td>700°C, 30 min</td>
<td>O$_2$</td>
<td>20-60 (40)</td>
<td>250</td>
</tr>
<tr>
<td>(Ba,Sr)TiO$_3$/Pt-Si</td>
<td>700, 1 h</td>
<td>air</td>
<td>50</td>
<td>400</td>
</tr>
<tr>
<td>Ce-(Ba,Sr)TiO$_3$/Pt-Si</td>
<td>750°C, 2 min*</td>
<td>n/a</td>
<td>(15-20)</td>
<td>200</td>
</tr>
<tr>
<td>BaTiO$_3$/Si</td>
<td>750°C, 1 hr</td>
<td>O$_2$</td>
<td>20-60</td>
<td>300</td>
</tr>
<tr>
<td>(Ba,Sr)TiO$_3$/Ni foil</td>
<td>900°C, 1 hr</td>
<td>10$^{-18}$</td>
<td>30-35**</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td>900°C, 1 hr*</td>
<td>10$^{-18}$</td>
<td>40-50**</td>
<td>400</td>
</tr>
<tr>
<td>BaTiO$_3$/Cu foil</td>
<td>900°C, 30 min</td>
<td>10$^{-9}$</td>
<td>100-300 (200)</td>
<td>600</td>
</tr>
</tbody>
</table>

* Multiple annealing
** No evidence of grain morphology reported
Fig. 4.3 Investigated area and corresponding electron diffraction patterns of BaTiO₃ films on Ni foils: a-b) are for a 700 RTA\'d annealed film, c-d) are from a furnace annealed film (1000°C)

Fig. 4.4 AFM scan area 0.5 x 0.5 um of BaTiO₃ films annealed at a) 700°C (RTA), b) 750°C (RTA) and c) 1000°C
In this experiment, the equiaxed morphology in films annealed at 700°C and 750°C suggested that the nucleation occurs throughout the films. The grain size was uniformly distributed across the film thickness, implying that repetition of the RTA annealing induced crystallization of newly deposited layers, but did not cause considerable grain growth of the previously-crystallized BaTiO$_3$. It is also possible that the large lattice mismatch of BaTiO$_3$ and Ni (~13% [16]) may not cause preferential nucleation at the film/substrate interface.

No clear interfacial layer between the BaTiO$_3$ films and Ni foil was observed for films annealed at 700°C or 750°C, indicating that covering the Ni foil with at least ~ 50 nm of amorphous BaTiO$_3$ film (first 2 layers) and using N$_2$ flow during both film pyrolysis at 350°C and RTA annealing 700°C or 750°C prohibited oxidation of the underlying Ni foil. It is possible that pyrolysis in the nitrogen ambient did not allow complete removal of carbon, so that the local oxygen partial pressure during the RTA step was lower than that of the ambient. This will be discussed further in section 5.3.2. Finally, it is noted that the observed cross-sectional BaTiO$_3$ films are on a single grain of Ni. Therefore, no comment can be made about the presence of an oxide layer (i.e. of contaminants in the Ni foil, or due to NiO formation) at the junction of the BaTiO$_3$ film and a grain boundary in the Ni substrate.

### 4.2.2 Furnace annealed BaTiO$_3$ films

The BaTiO$_3$ film, which was RTA annealed at 700°C, was crystallized in a tube furnace at 1000°C at 10$^{-16}$ atm pO$_2$. This pO$_2$ is below that for the Ni/NiO equilibrium,
so NiO is not expected to be thermodynamically stable. Subsequently, the sample was re-oxidized at 600°C at 10⁻⁶ atm pO₂.

The cross-section of the furnace annealed film was prepared by mechanical polishing and Ar⁺ ion milling. **Fig. 4.5a** shows a TEM cross-section showing two pieces of the same film. It is noticed that there is some variation in grain size and thickness between the two pieces of films. The film surface is rough due to grain growth at high temperature. As shown in **Fig. 4.5a** and **b**, the films exhibit polycrystalline, equiaxed grains. The average grain size of the furnace annealed film is 42 nm, and the grain size ranged from 24 to 75 nm (from the piece with the larger overall grain size). The grain size distribution is similar to that in the film before furnace annealing, implying that no abnormal grain growth occurs. The film contains 5-6 grains across a film thickness of ~200 nm. There are a few isolated voids (with a size on the order of the grain size) distributed in the film. The voids may be from either from porosity or damage due to (Ar⁺) ion milling. There is no interfacial layer between BaTiO₃ film and Ni foil, indicating that neither the heat-treatment, not the re-oxidation caused extensive Ni oxidation.
Fig. 4.5 Bright-field micrographs of the microstructure and interface of BaTiO$_3$ films on Ni foils annealed in furnace 1000°C. (High magnification figure is courtesy of Dr. Ian Reaney)
The change in microstructure of the furnace annealed BaTiO₃ film was expected to be dominated by the 1000°C step since the re-oxidation was performed at much lower temperature (600°C), where the cation mobility is low [17, 18]. However, considering the high pO₂ used at 600°C, the re-oxidation step can affect the interface quality and oxygen content in BaTiO₃ lattice, which will be discussed in section 4.3.

Compared to the RTA annealed film in Fig. 4.2a, the furnace annealed BaTiO₃ film has a much larger average grain size (42 nm) due to grain growth at higher temperature. Again, the AFM image in Fig. 4.3c showed a much larger cluster size.

There are few reports of preparing BaTiO₃ films by CSD method at temperatures higher than 800°C, as shown in Table 4.1. The grain size of the furnace annealed BaTiO₃ film in this experiment is comparable to that reported in Dawley et al.’s work where BST films on Ni foil were prepared at 900°C in a similar reducing atmosphere [14]. In contrast, the grain size of the furnace annealed film here is much smaller than that of BaTiO₃ films on Cu foil, which consists of two equiaxed grains of 300 nm size across the film thickness, as shown in Ihlefeld et al.’s work [15]. Since the solution chemistry of the two films was similar, the differences may be due to the annealing conditions. In this experiment, the BaTiO₃ film was pre-crystallized in an RTA at 700°C before being subjected to the furnace anneal (1000°C), while in Ihlefeld’s work, the BaTiO₃ was crystallized once at 900°C. In addition, the pO₂ used for annealing BaTiO₃ films on Ni foils (10⁻¹⁶ atm) is much lower than that of Cu foil (10⁻⁹ atm). It has been reported that prior H₂ heat treatments of BaTiO₃ ceramics can freeze abnormal grain growth in subsequent air sintering [19, 20]. Also, Polotai et al. showed that average grain size of a BaTiO₃ ceramic strongly decreased from 296 to 137 nm as the pO₂ during sintering
decreased from $10^{-1}$ to $10^{-19}$ atm [21]. Thus, it is speculated that the difference in grain size for films on Cu and Ni can be attributed to a combination of the number of crystallized grains before the high temperature furnace annealing and the effect of the pO$_2$ atmosphere. Since the RTA’d BaTiO$_3$ film contains a high concentration of small grains, grain impingement during growth may limit the final grain size during furnace annealing.

Although a columnar structure was reported to improve the dielectric properties of BaTiO$_3$ - based films [12, 22, 23], equiaxed grains are necessary to limit oxygen vacancy mobility in base-metal MLCCS. Many studies have shown that the dc insulation resistance degradation originates from electromigration of oxygen vacancies in the perovskite lattice [11, 24-27]. In equiaxed structures, the grain boundaries parallel to the interface provide Schottky barriers for migration of the oxygen vacancies created by the reducing atmosphere processing. Thus, it is encouraging that there are multiple grains across the thickness of a quarter micron thick CSD-derived layers.

4.3 Chemical analysis and oxygen nonstoichiometry

Oxygen nonstoichiometry is very important to the electrical properties and reliability of BaTiO$_3$ capacitors. In an attempt to investigate this parameter, furnace annealed (re-oxidized) BaTiO$_3$ films on Ni foils were investigated by electron energy loss spectroscopy (EELS) in a transmission electron microscope. The data were obtained with the help of Dr. Gaiying Yang. The energy loss peaks for Ba, Ti and O were observed using the Ba M$_{4,5}$ (781 eV), Ti L$_{2,3}$ (455.5 eV) and O K edges (532 eV), respectively. Since the films were processed at low pO$_2$, it is also of interest whether
there is any residual C in the film. The C K edge is at 283.8 eV. In this experiment, all EELS spectra were calibrated (to shift the peaks to the correct position) by using the onset of the O K edge. High resolution TEM imaging was used to confirm any contamination on the surface of the thin specimen.

Fig. 4.6 shows a cross-section of a bright field image of BaTiO$_3$ on Ni foil. EELS analysis with an electron beam size of ~ 10 nm was randomly performed on 4-5 grains of the furnace annealed BaTiO$_3$ film on Ni foil. All grains show similar EELS spectra, which consist of a strong C K edge peak associated with the existence of Ba M$_{4,5}$, Ti L$_{2,3}$ and O K edges, as shown in Fig. 4.7. EELS spectra of the Ni foil show only the Ni L$_{2,3}$ edge (854 eV) without the C K edge. The result suggested that C really existed in the BaTiO$_3$ film and was not solely surface contamination.
Fig. 4.6 Cross-sectional bright field image of BaTiO$_3$ film on Ni foil (Courtesy of Dr. Gaiying Yang). Positions for EELS analysis are marked.

Fig. 4.7 EELS spectra of BaTiO$_3$ grain and Ni foil, showing existence of C in the BaTiO$_3$ (Data courtesy of Dr. Gaiying Yang)
Oxygen non-stoichiometry is usually determined using the Ti L2,3 and O K edges. A reference EELS spectrum of an air sintered, precious metal electrode (PME) MLCC (Fig. 4.8b) shows the fine structure in the Ti L2,3 and O K edges for oxygen-stoichiometric dielectrics. The splitting of the Ti L3 edge (lower energy) into e_g and t2g features and the high intensity of the O K edge peak series are characteristic of a low concentration of oxygen vacancies in the PME dielectrics. Of interest for this discussion is the clarity of the splitting in the Ti L2,3 edge.

EELS analysis on a furnace annealed BaTiO3 film on Ni foil was performed at 3-4 areas as a function of distance from Ni electrode. Fig. 4.6 is marked with 4 spots where the EELS spectra were acquired. EELS spectra for the C K, Ti L2,3, O K and Ba M4,5 edges were first collected from each spot using an energy dispersion of 0.5 eV/channel to cover the broad range of C, Ti, O and Ba edge positions. Finer resolution of the Ti L2,3 and O K edges was obtained by re-measuring each spot with an energy dispersion of 0.1 eV/channel.

Fig. 4.8a shows broad range EELS spectra from the 4 spots in Fig. 4.6. It was found that, in this area of the BaTiO3 film, every spot contains at least some C. The lowest carbon concentrations were observed near the electrode and a grain boundary. Oxygen diffusion is faster along grain boundaries in BaTiO3, which would facilitate carbon oxidation from those regions. The EELS spectrum near the Ni foil (Point 1) show no splitting of e_g and t2g peaks and a lower intensity of the O K edge, implying a partial reduction of the Ti oxidation state from 4+ to 3+ which is usually associated with low oxygen content. The spectrum at Point 4 shows almost fully oxidized Ti, similar to the PME dielectric. The fact that near-stoichiometric BaTiO3 was observed in some regions
of the sample suggests that the reduced regions observed elsewhere were not simply the product of ion milling.

Two more areas were also examined. A lower amount of C was found near the electrode in both areas. In one of these, nearly stoichiometric BaTiO₃ was found near the Ni foil and the oxygen content slightly decreased as a function of distance from Ni foil. In another area, nearly stoichiometric BaTiO₃ was found at all depths. The existence of C is probably a consequence of incomplete removal of C from the organic precursors in the film. This is reasonable given the low pO₂ conditions used. The variation in oxygen content as a function of distance from Ni foil was thus variable in this re-oxidized film. It is suggested that this parameter be investigated more thoroughly by examining films after the high temperature heat treatment, but prior to re-oxidation.
Fig. 4.8 EELS spectra collected from BaTiO$_3$ film as a function of distance from Ni foil
a) broad range spectra b) resolved spectra of the fine structure of Ti L$_{2,3}$ and O K edges
including that of PME BaTiO$_3$ (Data courtesy of Dr. Gaiying Yang)
HRTEM was performed to investigate the BaTiO$_3$ / Ni interface, as shown in Fig. 4.9. The clear atomic arrangement in the BaTiO$_3$ confirms that there is no contamination on the sample surface or damage due to sample preparation. Atomic steps were observed along the Ni foil surface. The Moiré fringes in the Ni region are not fully understood, however it is suspected that the fringes are due to absorbed oxygen on the surface of Ni foil [28].

The Ni foil surface shows some image contrast at the top ~ 5-8 nm. Fig. 4.10 shows representative EELS spectra obtained from deeper in the Ni region and this interfacial layer. The Ni L$_{2,3}$ edge confirms that in both cases Ni is in a metallic state, not an oxide [29]. Thus, the associated O K edge is due to O absorbed on the Ni surface. The interfacial layer shows primarily metallic Ni L$_{2,3}$ with a small Ba M$_{4,5}$ edge and no O K edge. The result suggested the interfacial layer is a Ni-Ba alloy. EELS spectra of the interfacial layer and Ni region were confirmed at 3-5 spots and the same results were obtained. A wider view of the interface showing this is given in Fig. 4.11. It is noted that no NiO is observed.

Oxygen non-stoichiometry and the internal dielectric / electrode interfaces in BaTiO$_3$ MLCC with Ni electrodes were studied by Yang et al. [24, 25, 28]. It was shown that the oxygen content in co-fired BaTiO$_3$ increases as a function of distance from the Ni electrode after the sintering step, but becomes more spatially homogeneous after re-oxidation. They also discovered the presence of a Ni-Ti-Ba alloy layer (4-15 nm thick ) between the BaTiO$_3$ and the Ni in commercial MLCCs [28]. It was found that the metallic alloy layer develops during the co-firing step (1300$^\circ$C, $10^{-10}$ atm pO$_2$) and survives the re-oxidation step (800$^\circ$C, $10^{-8}$ atm pO$_2$). Carbon residue was proved to be
the cause of the metallic alloy interface layer and the local variation of oxygen content in the co-fired BaTiO$_3$. Belated decomposition of the organics, especially C, from the Ni paste may rob oxygen from the BaTiO$_3$ lattice, resulting in oxygen vacancies near the electrode [24].

It is believed that the Ni-Ba layer observed in the films occurred during the high temperature heat treatment and was retained through re-oxidation. It is important to note that for all of the firing conditions employed, BaO, rather than Ba is thermodynamically stable. This implies that there must have been a strongly reducing local environment to produce metallic Ba. It was suggested by Yang et al. [28] that such a highly reducing atmosphere could be induced by local CO/CO$_2$ reactions with Ni as a catalyst. The C in the films could be from incomplete organic removal at the low oxygen partial pressure employed during the RTA step. It is interesting that the alloy composition seen in the thin films does not apparently show any Ti. This could be a result of the lower process temperatures used for thin film firing, than for bulk ceramics.

The origin of the interfacial alloy is not well understood. One possibility is that a Ni-Ba liquid phase develops due to the existence of low melting temperature eutectics on the phase diagram (See Figure 4.12). Ba-containing crystallites could precipitate from the liquid phase on to the Ni (FCC) surface, producing a structure-matched FCC Ba-rich phase. This would explain the observation that the Ni-Ba alloy structure is coherent with that of the Ni.

It will be important in the future to determine the alloy composition more quantitatively, and to estimate its impact on the Schottky barrier layer at the film/foil interface. Further study of organic removal in low pO$_2$ will be discussed more in section
5.3.2. The observed spatial nonuniformity in the C content may be linked to the local variations in the Ti$^{3+}$/Ti$^{4+}$ ratio.

**Fig. 4.9** HRTEM image of furnace annealed BaTiO$_3$ film/ Ni foil interface (Data courtesy of Dr. Gaiying Yang)
Fig. 4.10 EELS spectra collected from Ni region and interfacial layer between BaTiO$_3$ and Ni (Data courtesy of Dr. Gaiying Yang)

Fig. 4.11 View of a larger section of the BaTiO$_3$/Ni interface showing the interface layer and the lack of NiO (Data courtesy of Dr. Gaiying Yang)
Fig. 4.12 Binary alloy phase diagrams of Ba-Ni (Adapted from Massalski [30])
References


Chapter 5
Processing Studies for BaTiO₃ Films on Ni Foils

This chapter focuses on the use of spectroscopic ellipsometry, X-ray diffraction, and FTIR to characterize the composition and depth profile of the Ni foils and BaTiO₃/Ni samples as a function of temperature. Of particular interest was developing an understanding of the effect of processing conditions on the BaTiO₃ microstructure, as well as evaluation of the conditions under which NiO forms. Spectroscopic ellipsometry is a useful complement to the transmission electron microscopy studies, because it is a large area technique with good depth resolution.

5.1 Oxidation of Ni foil

In order to develop appropriate processing conditions for BaTiO₃ films on Ni foils, it is useful to understand at what point in the processing oxidation of the foil surface would be expected. Thus, the oxidation of Ni foil surfaces was studied by spectroscopic ellipsometry (SE) as a function of process temperature. The Ni foil used for this study was 50 µm thick and 99+ % purity. The surface was visually rough. Atomic force microscopy (AFM) was used to determine the surface roughness of the as-received foil, as shown in Fig. 5.1. Over a 5x5 µm area, the average surface roughness was 27 nm, the root-mean-square (rms) roughness was 37 nm, and the peak to valley roughness is as large as 330 nm.
Ni foils (99+ %) were heated on a hot plate at 200-450°C at 50°C intervals for 2 min. in air. Although none of the XRD patterns (not shown here) had NiO peaks, the color of the Ni foil changed from silver (unannealed) to yellow gold after annealing at 450°C, suggesting the development of a transparent overlayer.

SE data from 250 to 700 nm collected from the Ni foils after each annealing step are shown in Fig. 5.2. It is clear that the SE data progressively change as the annealing temperature increases from 200°C to 350°C. A significant change was observed after the Ni foil was annealed at 450°C, implying either a considerably change in the depth profile of the sample or oxidation. It is noted that the intensity of the reflected light from the Ni foils is low due to their surface roughness. This caused difficulty in sample alignment on the ellipsometer.

SE data modeling was performed on these samples by using different possible model structures i.e. models with and without NiO, as shown schematically in Fig. 5.3. Surface roughness layers on the Ni foil were treated as a layer of Ni+air on Ni. Models
with NiO consist of either a layer of Ni+NiO or NiO+air on the Ni substrate. In most cases, it was necessary to model SE data only over the range from 300-650 nm since it significantly improves the fit. It is noted that reference data of optical properties for single crystal O-rich NiO [1] were used for the growing NiO layer. This point will be further discussed in section 5.4.1.

Two sets of reference data for Ni single crystals (measured at 4 K and room temperature) were used to describe the bulk Ni. The best available data are those measured at 4K, as a well-defined surface was prepared, and the optical properties are reported at a large number of wavelengths over the visible range. Unfortunately, the available data sets for Ni at room temperature are considerably more sparse, and it was not immediately clear what errors would result from using cryogenic data for the Ni. In an attempt to provide a better reference data set, the 4K data were scaled to overlap, as well as possible, with the available room temperature data, without changing the shape of $\varepsilon_1$ and $\varepsilon_2$ as a function of wavelength. Fig. 5.4 shows reference data for the dielectric function of Ni at 4 K, room temperature and the modified room temperature data. Both data sets, as well as mixtures of the two, were then used to model the data for the Ni foils.

It was found that mixing of the two Ni dielectric functions often decreases sigma and improves the fit to the experimental data on foils. In particular, the best-fit models for Ni foils annealed between 200 and 400ºC were obtained using mixed Ni reference data. The relative volume fraction of the 4K reference data was found to vary randomly from 0.12 up to 0.48 for the different annealing temperatures. However, the best fits for the untreated foils and 450ºC annealed foils were obtained using only reference data for a Ni single crystal at room temperature and 4 K as the substrate, respectively. At least
some of these discrepancies in the best Ni reference data may result from the fact that the peak-to-valley roughness on these foils, as determined by AFM, is approximately the same as the wavelength used to probe the samples, and is variable from point-to-point and sample-to-sample. Thus, some degree of diffuse scattering is likely to complicate the ellipsometry modeling. Since there is no clear trend for the variation in the optical properties of Ni substrate, only “Ni” will be used to represent the Ni phase in the following discussion.
**Fig. 5.2** SE data of Ni foil (99+ %) and Ni foil after annealing at 200-450°C for 2 min in air

**Fig. 5.3** Schematic of SE models for Ni foils with and without NiO
Fig. 5.4 Dielectric functions of Ni single crystals (SC) at 4 K and 293 K. The modified room temperature data set developed in this work is also shown.
To establish a baseline, data for an as-received Ni foil were first modeled. The best fit model for the sample included a layer of surface roughness, but no NiO. Compared to a model of Ni/Ni+NiO, the model involving only roughness showed an improved fit at short wavelengths, as shown in Fig. 5.5, and sigma decreased from 0.019 to 0.01. The roughness layer is 72.1± 4.1 Å thick with 21% volume air, as shown in Fig. 5.7. Thus the layer of surface roughness of the Ni foil detected by SE is thinner than that detected by AFM. This may be a result of differences in the sampling areas or problems associated with the inability to properly model asperities with a size scale on the order of the wavelength of light. It is noted that modeling the as-received sample with Ni/NiO+air gave an unreasonable model.

Best fit models of Ni foils annealed at 200°C or 250°C were also found to consist of rough Ni, the same as for the as-received Ni foil. For Ni foils annealed at 300°C and higher (up to 450°C), the best fit models were obtained from Ni/NiO+air. The results suggested that Ni oxidation begins when un-coated Ni foils are annealed at 300°C. Fits of experimental and modeled data of these annealed foils are shown in Fig. 5.6.
Fig. 5.5 Experimental and modeled data with and without NiO for an as-received Ni foil
Fig. 5.6 Experimental and modeled data for Ni foils annealed at different temperatures
Fig. 5.6 (cont.)
Fig. 5.6 (cont.)
### Fig. 5.7 SE – determined depth profiles from as-received and annealed Ni foils

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Layer Composition</th>
<th>Layer Thickness</th>
<th>σ</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Unannealed Ni foil</strong></td>
<td>Ni + air(0.21 ± 0.01)</td>
<td>72.1 ± 4.1 Å</td>
<td>0.010</td>
</tr>
<tr>
<td>200°C</td>
<td>Ni + air(0.44 ± 0.04)</td>
<td>48.7 ± 3.6 Å</td>
<td>0.012</td>
</tr>
<tr>
<td>250°C</td>
<td>Ni + air(0.48 ± 0.07)</td>
<td>36.7 ± 3.1 Å</td>
<td>0.014</td>
</tr>
<tr>
<td>300°C</td>
<td>NiO + air(0.71 ± 0.07)</td>
<td>54.8 ± 11.4 Å</td>
<td>0.011</td>
</tr>
<tr>
<td>350°C</td>
<td>NiO + air(0.54 ± 0.05)</td>
<td>48.8 ± 4.3 Å</td>
<td>0.012</td>
</tr>
<tr>
<td>400°C</td>
<td>NiO+ air(0.52 ± 0.02)</td>
<td>83.2 ± 3.6 Å</td>
<td>0.010</td>
</tr>
<tr>
<td>450°C</td>
<td>NiO+ air(0.36 ± 0.01)</td>
<td>204.6 ± 2.3 Å</td>
<td>0.013</td>
</tr>
</tbody>
</table>
Depth profiles of the best-fit models for as-received and annealed Ni foils are presented in Fig. 5.7. In addition to an increase in the thickness of the NiO+air layer, the volume fraction of air in the layer decreases from 0.71±0.07 to 0.36±0.01 as the annealing temperature increases from 300°C to 450°C. The total thickness of the NiO+air layer and the effective thickness of NiO for each sample are plotted in Fig. 5.8. The effective thicknesses of NiO were calculated by multiplying the volume fraction of NiO by the thickness of the layer. Thus, despite the difficulties associated with the rough Ni surface, the SE results of this experiment agree well with an *in situ* reflectance study of a heated Ni film [2], which showed oxidation occurs at 300°C.

**Fig. 5.8** Plot of total thickness of NiO+air layer and effective thickness of NiO
5.2 High purity Ni foils

In the previous section it was shown that, despite the rough surface, low purity Ni foil (99%) can be used in a SE study to observe oxidation and determine the NiO thickness at low temperatures. However, preliminary work found that these foils were less useful for SE study when severe oxidation occurred, i.e. the surface of the foil became seriously roughened and the amount of detected light was too low. In addition, it was found that the acetic acid used in BaTiO₃ solution attacked the Ni surface, presumably etching it, and further increasing the surface roughness. This made it impossible to obtain interference fringes in SE data from BaTiO₃ films on low purity Ni foils (See Fig. 5.9).

To minimize these difficulties, high purity cold-rolled Ni foil (99.99%) was chosen as the substrate for further study of the Ni-BaTiO₃ thin film interface by SE as its surface is smoother than the one of 99+% purity. Earlier work of Nagata et al. [3] reported that pre-annealing these Ni foils at 900°C results in a much smoother surface. Also, due to its higher purity, there were fewer oxidized contaminants on the surface after annealing, compared to the low purity Ni foil. In addition, the smooth and clean surface of the high purity Ni foil provides better dielectric properties for the resulting BaTiO₃ thin films [3].

In this experiment, a high purity Ni foil was pre-annealed at 900°C for 1 hour in 10⁻¹⁶ atm pO₂. Fig. 5.10 shows the surface microstructure and morphology of the as-received and pre-annealed Ni foil. The as-received Ni foil, which was cold-rolled and thus highly deformed, shows no clear grain features. The scratches and roughness on the surface are due to the rolling process. After pre-annealing, the Ni grain size varies
between 20 and 200 µm. The surface of Ni foil becomes much smoother due to diffusion at high temperature. In addition, grain boundary grooves develop at the surface in order to reduce the combined surface and grain boundary free energy \[4, 5\].

The surface roughness of the high purity Ni foils was investigated by AFM. The measured area was 5x5 µm², which is very small compared to the grain size. It was found that the average roughness \((R_a)\) decreases from 6 nm to 4 nm after annealing. Fig. 5.9 demonstrates that using high purity Ni foil as a substrate, interference fringes from the transparent BaTiO₃ film can be seen in the SE data. Also, the smoother surface of the underlying substrate results in good reflection and easier sample alignment. Models of the SE data for BaTiO₃ film on the Ni foil will be discussed in section 5.4.

As was the case for the lower purity foils, the surface roughness measured by ellipsometry was smaller than the AFM result (see Table 5.1). However, the correlation between the two was good. That is, foils that were measured to be smoother by AFM were also smoother by SE. Again, the differences may be due to some of the larger scale topology that complicates the SE modeling.

Table 5.1 AFM and SE surface roughness of 99+% and pre-annealed 99.99% Ni foil

<table>
<thead>
<tr>
<th>Sample</th>
<th>Surface roughness (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(d_{\text{AFM}}) (Å) (or (R_a))</td>
</tr>
<tr>
<td>Ni foil (99+)</td>
<td>270</td>
</tr>
<tr>
<td>Pre-annealed Ni foil (99.99%)</td>
<td>40</td>
</tr>
</tbody>
</table>
Fig. 5.9 SE data collected from BaTiO$_3$ films on 99% and 99.99% Ni foils
Fig. 5.10 Microstructure and surface morphology of high purity Ni foils (99.99 %) a) as-received and b) after pre-annealing at 900°C. Both SEM and AFM data are shown.
5.3 BaTiO$_3$ films on Ni foils

5.3.1 Structure and microstructure

Fig. 5.11 showed XRD patterns of a pre-annealed Ni foil and BaTiO$_3$ films RTA’d at different temperatures under flowing N$_2$. The pre-annealed Ni foil shows only a strong 200 diffraction peak, suggesting that the foil is at least partially oriented after re-crystallization. The film RTA’d at 600$^\circ$C shows an extremely low intensity of the BaTiO$_3$ 100 and 110 peaks, along with a trace amount of a second phase at a two-theta of 26.7$^\circ$. The second phase peak corresponded to the highest peak of an oxycarbonate (Ba$_2$Ti$_2$O$_5$CO$_3$), which was reported to be an intermediate phase in the crystallization of perovskite BaTiO$_3$ [6-9]. A recent paper suggests that this intermediate phase may, in fact, contain very little Ti [10]. For the films RTA’d at 650-750$^\circ$C, the “oxycarbonate” phase disappears and slightly higher intensity BaTiO$_3$ peaks are shown. However, the low intensity of the BaTiO$_3$ XRD peaks in RTA films suggested that the films are either not well-crystallized, or have small coherently diffracting region sizes even for RTA temperatures of 750$^\circ$C. This could be due to the small thermal budget and the low pO$_2$ (N$_2$ flow) used in the RTA step. Since the electron diffraction pattern showed no sign of an amorphous phase (diffuse ring), the low intensity of the RTA films is presumably due to the small crystallite size.

These same films were then annealed in a reducing atmosphere furnace at 1000$^\circ$C for 1 hour. The XRD patterns after furnace annealing are shown in Fig. 5.12. The patterns for all films correspond to a well-crystallized polycrystalline perovskite phase with a high intensity of the 100 and 200 peaks. There is no clear peak splitting for the 001/100, 101/110, 002/200 or 112/211 peak pairs. Thus, the BaTiO$_3$ films on Ni foil
have a pseudocubic perovskite structure with some degree of <100> orientation. The high intensity and narrow peaks suggest furnace annealed films exhibit larger grain size, compared to those of RTA’d films.

Since NiO peaks were not observed in XRD patterns measured with a 0.5 sec. collection time per point (Fig. 5.10 and Fig. 5.11), a slower scan with a 5 sec. collection time per point was performed on all films over the 2-theta range where the NiO 111 and 200 peaks are expected. Fig. 5.13 shows that slow x-ray diffraction scans on RTA films did not show NiO. For furnace annealed films (Fig. 5.14), there is a small NiO 200 peak, however, no NiO 111 peak is observed.

Thus, the slow scan x-ray diffraction showed that NiO appears after the furnace anneal. To isolate whether the NiO formed during the high temperature step, or during subsequent re-oxidation, slow scan diffraction was performed after each step. It was found that NiO formation occurs in the re-oxidation step (See Fig. 5.15). The re-oxidation was done at 600°C at a pO2 of 10^-6 atm, which is above the Ni/NiO equilibrium. Thus, it is possible that annealing for 1 hour could cause Ni oxidation. The very low intensity of the peak implies that there is only a small amount of NiO. There is no significant trend in the amount of NiO for the furnace annealed films as a function of the RTA temperature.
**Fig. 5.11** XRD patterns of BaTiO₃ films on high purity Ni foils RTA’d at different temperatures

**Fig. 5.12** XRD patterns of furnace annealed BaTiO₃ films that were RTA’d at different temperatures
The films were RTA annealed at different temperatures (600-750°C) before the furnace anneal step.
**Fig. 5.15** Slow scan of x-ray diffraction showing NiO after re-oxidation of 700°C RTA’d BaTiO$_3$ film

The surface microstructure of furnace annealed (1000°C) BaTiO$_3$ films on Ni foils was observed by SEM, as shown in **Fig. 5.16**. All the films are crack-free and consist of equiaxed grains with some porosity. The grain size of the furnace annealed films that were RTA’d at 600°C and 650°C appear to be about 80-100 nm. The 700°C RTA’d film shows a slightly smaller grain size of 50-80 nm while a slightly larger grain size of 80-150 nm is observed in the 750°C RTA’d film. It is noted that all the films also contain some grains of very small grain size, on the order of 25 nm or less, which are hard to resolve by SEM. It was also observed that there was some spatial non-uniformity in the grain size on a given film. It is recommended that in future work, the grain size be studied as a function of solution aging and/or variation of pO$_2$ during annealing.
The porosity tends to decrease as the RTA temperature increases. It is believed that in an RTA step with flowing N₂, organic removal is incomplete (See section 5.3.2) and that porosity is developed when the film is subsequently furnace annealed. As has been described by Scherrer, in some solution-deposited films, if crystallization is initiated before densification is complete, the resulting films can incorporate porosity [11].

Fig. 5.16 Microstructure of furnace annealed BaTiO₃ films (for different RTA temperatures) on Ni foil
5.3.2 Organic decomposition at low temperature processing

*Methanol-based BaTiO$_3$ solution*

The previous section demonstrated that the residual porosity in BaTiO$_3$ films depends on the RTA annealing (600-750°C). It is important to note that the porosity is retained after the furnace anneal at 1000°C. To determine whether this could be related to retained carbon, thermal gravimetric analysis (TGA) and differential scanning calorimetry (DSC) on the dried powders were carried out simultaneously on a TA Instruments SDT 2960 to examine the decomposition behavior of a BaTiO$_3$ powder obtained from a solution. The BaTiO$_3$ solution was air dried at 150°C for 15 h and then ~20 mg of powder was heated to 900-1200°C at a heating rate of 5 or 10°C/min. The measurements were performed both in air and with the N$_2$ flow rate set at 100 cc/min. To complement this study, dried BaTiO$_3$ powders were also calcined at different temperatures (500-900°C) in air for 1 h and the phase content was identified by XRD.

*Fig. 5.17* shows a comparison of the thermal analysis results in air and N$_2$ for powder dried from a methanol-based BaTiO$_3$ solution. The total weight loss of 40% is the same for both air and N$_2$ flow. However, clearly the TGA curves measured in air and N$_2$ show complete decomposition at different temperatures (~750°C for air flow and ~1000°C for N$_2$ flow). In addition, the rate and amount of weight loss varies significantly at temperatures between ~400-600°C. Accordingly, the DSC curves reveal a sharp exothermic peak at ~450-500°C in air while a smaller and broader peak is shown in N$_2$. In addition, for measurements in N$_2$ flow, it is clear that the decomposition is completed at lower temperatures when slower heating rates (5°C/min) are used.
Fig. 5.17 TGA and DSC curves of dried powder from methanol-based BaTiO$_3$ solution
The data for the sample measured in flowing air will be described first. The weight loss (5.8%) from 25°C to 200°C is due to evaporation of the solvent and absorbed moisture, which agrees well with the endothermic peak shown at 55°C in the DSC curve. At 200-400°C, the weight loss (14.7%) and the associated DSC peaks can be attributed to pyrolysis of organics. At ~400-550°C, the powder heated in flowing air shows weight loss (12.6%) due to further removal of organics and a sharp exothermic peak at 490°C. Fig. 5.18 shows that powder calcined at 500°C for 1 h in air consists of amorphous material along with BaCO₃ and either Ba₂Ti₂O₅CO₃ or the calcite-like BaCO₃. These phases have previously been reported as intermediate phases in BaTiO₃ crystallization [6, 8, 9, 12-15]. The result suggested that the strong exothermic peak is due to formation of the intermediate phases. The exothermic peak at 605°C is indicative of crystallization of BaTiO₃, which is confirmed in Fig. 5.17. Upon heating, crystallization continues (as witnessed by the higher intensity of the BaTiO₃ peaks) and the amount of the second phases gradually decreases. The oxycarbonate phase and BaCO₃ completely disappeared when the powder was calcined at 700°C and 900°C, respectively. It is believed that the last weight loss (5.9%) is due to decomposition of the carbonate phase.

Similar TGA and DSC curves are observed when the BaTiO₃ powder was heated in flowing N₂ from room temperature to ~400°C at the same heating rate (10°C/min). Above 400°C, the TGA curve clearly shows a delay in the weight loss and the complete thermal decomposition is extended (from 750°C in air) to 1000°C. In contrast to DSC in air flow, the DSC in N₂ shows a small and broad exothermic peak at 450°C, suggesting carbonate phases are gradually formed in flowing N₂. The DSC curve in N₂ shows extra peaks at 740°C and 935°C, compared to that in air. Thus, it is believed that the delay in
weight loss is accompanied by slowing down other reactions, including carbonate decomposition and thus BaTiO₃ crystallization. As expected, the extra reaction peaks in DSC are shifted to lower temperature as heating rate decreases.

Fig. 5.18 XRD patterns of powders calcined in air at different temperatures
The results above show that organic burn-out is delayed when there is insufficient oxygen available at elevated temperatures. Therefore, it is possible that BaTiO$_3$ films RTA’d at $\leq 750^\circ\text{C}$ in N$_2$ might have remaining organics that are ultimately the origin of the film porosity on subsequent furnace annealing at 1000$^\circ\text{C}$ (see Fig. 5.16), especially as the heating rate in RTA ($10^\circ\text{C/s}$ ) is much faster than this thermal analysis experiment (5 and 10$^\circ\text{C/min}$). It would be important to further quantify, in future work, the amount of any residual carbon in the furnace annealed samples as well. Although a heating rate of 5$^\circ\text{C/min}$ in the furnace annealing will assist organic removal, the low pO$_2$ might retard the process. These results also suggest that it will be important to further optimize the pyrolysis steps to minimize the amount of carbon retained in the films.

Thermal analysis in air of dried powder from BaTiO$_3$ solutions has shown that the crystallization pathway from amorphous to perovskite BaTiO$_3$ occurred via a BaCO$_3$ intermediate. In contrast, 600$^\circ\text{C}$ RTA’d BaTiO$_3$ thin film on Ni foil shows only a small amount oxycarbonate as an intermediate phase (Fig. 5.11 in section 5.3.1). Future work should be done to determine whether this difference is a result of the fast heating rate and low pO$_2$ during the RTA step in flowing N$_2$.

**2-Methoxyethanol-based BaTiO$_3$ solution**

Thermal analysis was also performed on dried BaTiO$_3$ powders (with 1% Ca) from a 2-methoxyethanol-based solution to compare to the result of methanol-based powders. (The experiment was done by Dr. Song Won Ko.) A 2-methoxyethanol-based solution was prepared as described in section 3.1.1. The dried powders were analyzed in the TGA at a heating rate of 10$^\circ\text{C/min}$. 
**Fig. 5.19** shows the TGA results of the 2-methoxyethanol-based powders compared to those from methanol-based solutions. It can be seen that, in air, the weight losses from both powders were completed at approximately the same temperature (700-750°C for 2-methoxyethanol and 800°C for methanol). In N₂, the completed decomposition of the 2-methoxyethanol-based powders is delayed to 800°C, which is much lower than that of the methanol-based powders. Therefore, 2-methoxyethanol-based solutions may be more suitable for processing in low pO₂ conditions.
Fig. 5.19 TGA curves of dried powders from 2-methoxyethanol (2-MOE) and methanol based BaTiO$_3$ solutions (2-methoxyethanol data courtesy of Dr. Song Won Ko)
The organic removal from 2-methoxyethanol-based BaTiO$_3$ films was also investigated by a Nexus 670 (Thermo Nicolet) Fourier transform infrared spectroscopy (FTIR) technique. The experiment was performed by Dr. David Lu. A 0.3 M solution was spin coated onto SiO$_2$/Si substrates. Four layer films were dried at different temperatures (125$^\circ$C, 390$^\circ$C and 500$^\circ$C) in air for 5 min, where the drying temperatures were chosen as points above some weight loss event.

**Fig. 5.19** shows the FTIR spectra and thickness of the 2-methoxyethanol-based BaTiO$_3$ films dried at different temperatures. The as-deposited film shows peaks characteristic of carboxylate (at 1560 cm$^{-1}$) and carbonate groups (at 1425 cm$^{-1}$), OH (3300-3500 cm$^{-1}$) and C-H bonds (2880-3020 cm$^{-1}$) [16, 17]. The OH and C-H stretching vibrations are due to the presence of hydroxyls and organic molecules, respectively, in the film. These bands disappear after drying the film at 390$^\circ$C. The carboxylate group, which significantly decreases after 390$^\circ$C, almost disappears after 500$^\circ$C. Although the carbonate peak (which could arise, at least in part, from BaCO$_3$) intensity decreases as the drying temperature increases, some carbonate is clearly retained in the 500$^\circ$C dried film. The thicknesses of the films, measured by a profilometer, collapse to around 40% of the initial thickness after drying at 500$^\circ$C, as shown in **Fig. 5.20**.

**Fig. 5.21** shows FTIR spectra of 2-methoxyethanol-based BaTiO$_3$ films RTA’d at 750$^\circ$C in air or N$_2$ for 10 minutes. The films were previously dried at different temperatures (125$^\circ$C, 390$^\circ$C or 500$^\circ$C). The Ti-O and BaTiO$_3$ features confirm the crystallization of some BaTiO$_3$ after an RTA step in either air or N$_2$. However, in both cases (and for all drying conditions investigated), there are still features in the FTIR data characteristic of carboxylate and carbonate groups.
**Fig. 5.20** FTIR spectra and thickness of deposited 2-methoxyethanol-based BaTiO$_3$ films dried at different temperature (Data courtesy of Dr. David Lu)
Fig. 5.21 FTIR spectra of 2-methoxyethanol-based BaTiO$_3$ films RTA’d at 750°C a) in air and b) in flowing N$_2$. The films were dried at different temperatures (125°C, 390°C or 500°C) (Data courtesy of Dr. David Lu)
5.4 Spectroscopic ellipsometry studies

5.4.1 Amorphous dried BaTiO$_3$ films

Spectroscopic ellipsometry has previously been used to characterize the relative density, oxidation state and surface roughness of BaTiO$_3$ films and ceramics [18-24]. These results suggest that it is possible to use SE to follow changes in the BaTiO$_3$ films throughout the film processing. Thus, the optical properties and thickness of amorphous dried BaTiO$_3$ films were also studied by SE. In this experiment, eight layers of 2-methoxyethanol based BaTiO$_3$ solution (0.2M) were deposited on a freshly etched Si wafer. Si was chosen for this study due to its excellent surface smoothness and its stability over this temperature range. The SiO$_2$ was removed from the wafer surface by dipping the Si wafer in a small Teflon beaker with HF acid for 10 min. Then, the wafer was rinsed with deionized water and blown dry with N$_2$. The films were deposited on the wafer right after etching. After each layer deposition, the film was dried at different temperatures (200, 250, 300, 350°C) for 5 min.

SE data from 350 to 700 nm were used in modeling. The best fit model structure of these films consisted of a bulk film with a thin layer of surface roughness. The sigma value decreased from 0.089 to 0.038 with increasing drying temperature. It is also noted that the 350°C dried film shows no surface roughness. Fig. 5.22 shows the best fit to the experimental data for the amorphous BaTiO$_3$ film on Si dried at 200°C.

Fig. 5.23 shows the measured refractive index of each of the dried films. It can be seen that refractive index increases as the films are densified at higher drying temperatures. Accordingly, Fig. 5.24 shows that the film thickness collapses from 5332.0±64.1 Å to 3642.0±8.1 Å after drying at 200°C to 350°C, respectively. This is
consistent with the observed decrease in thickness with increasing drying temperature, as measured by profilometry (See Fig. 5.20). In section 5.3.2, it was shown that weight loss due to large amount of organics occurs at this temperature range.
Fig. 5.22 Model fits compared to measured SE data for an amorphous BaTiO$_3$ film dried at 200$^\circ$C on a silicon wafer (prepared from 0.2 M 2-methoxyethanol solution)
**Fig. 5.23** Refractive index of dried amorphous BaTiO$_3$ films (prepared from a 0.2 M 2-methoxyethanol solution)

**Fig. 5.24** Total thickness of amorphous BaTiO$_3$ films as a function of drying temperature (prepared from 0.2 M 2-methoxyethanol solutions)
5.4.2 BaTiO₃ films on sapphire substrates

To study the crystallization step, optical measurements were carried out on BaTiO₃ films on sapphire substrates. An Al₂O₃ (0001) single crystal was used as the substrate because it provides a stable surface which will not complicate SE modeling. In addition, the Al₂O₃ substrate has a small thermal expansion mismatch with BaTiO₃ up to at least 1000°C, giving crack-free films (7.5x10⁻⁶/°C for Al₂O₃ and 9.8x10⁻⁶/°C [25] for BaTiO₃).

Thin films of ~2000 Å were prepared as described in section 3.1.1. The film was RTA’d at 750°C in N₂ flow. A SE measurement was performed after the RTA annealing step (750°C in N₂ flow). Then, SE data from the same film area were measured after the high temperature step (1000°C, 10⁻¹⁵ atm pO₂) and again after re-oxidation (600°, 10⁻⁶ atm pO₂). As shown in Fig. 5.25, there are clear changes in the interference fringes after the 1000°C step; however, no significant change was observed after re-oxidation. In all cases, the data were collected using an achromatic compensator to minimize instrumental errors.

Modeling of the SE data was done using reference data for (0001) Al₂O₃ and a Sellmeier oscillator for the optical properties of the BaTiO₃ film. In all cases, good results were obtained from data modeled between 375 and 750 nm with the absorption part of the oscillator fixed to zero. Because this is an all-transparent system, the SE data were modeled using Δ and Ψ, rather than tanΨ, cosΔ for σ. This typically enables better fitting of the depth profiles, as discussed by Chindaudom [26].

Initially, modeling was performed with a 2 layer structure consisting of a bulk film with some surface roughness. The 2 layer model resulted in σ values that are unacceptably large (2.24°-3.44°). Much lower sigma (0.69°-0.81°) and improved fits
were achieved when a low density layer was added at the film/substrate interface (See. Fig. 5.26 and Fig. 5.27).

EELS analysis (section 4.3) showed retained C distributed in a furnace annealed film (on Ni foil). Thus, attempts were made to model with the optical property data using a mixture of BaTiO$_3$ and graphitic carbon. The resulting fit did not provide a reasonable volume fraction of C, and was discarded.

The optical properties and depth profiles of the films at the three different stages in the processing are shown in Fig. 5.28 and Fig. 5.29, respectively. Obviously, the results after 1000 °C and after re-oxidation show comparable refractive indices and film structures, suggesting that the re-oxidation step does not change the film substantially. The optical properties agree well with data for a ceramic sample with 95% density [27], meaning the film annealed at 1000°C is fully-crystallized and of reasonably high density. The RTA’d sample (750°C) shows a lower refractive index. Fig. 5.30 shows a plot of the film thickness after each annealing step. It can be seen that there is some densification after the furnace annealing, although the change in thickness is within the 90% confidence limits. This suggests that the major increase in refractive index after furnace annealing may be due to further crystallization, rather than densification.

There is also a slight change in the film structure between RTA’d and 1000°C annealed films (see Fig. 5.29). It was necessary to fix the volume fraction of air at 50% in the surface roughness layer to obtain a reasonable best fit model for the RTA’d film. In contrast, varying the % air in the layer improves the fit for the furnace annealed film. The thickness of the surface roughness layer increases from 83.3±1.8 Å (with fixed 50% air) for the RTA’d film to 206 ± 16 Å (with 32±3% air) for the 1000°C annealed film.
The larger surface roughness of furnace annealed films could be due to grain growth at high temperature. The result is consistent with the rough surface observed in the cross-sectional TEM sample in Fig. 4.5 (section 4.2.2).

These results suggest that BaTiO$_3$ films with good densities can be achieved by CSD processing, and that SE can be used to depth profile them. Thus, work was subsequently undertaken to study BaTiO$_3$ films on Ni foils.
Fig. 5.25 Delta and psi as a function of wavelength at various points in the processing of a BaTiO$_3$ film on a (0001) Al$_2$O$_3$ substrate.
Fig. 5.26 Model fits with 2 layer (Al$_2$O$_3$/Osc/Osc+air) and 3 layer (Al$_2$O$_3$/Osc+air/Osc/Osc+air) model of 750°C RTA’d BaTiO$_3$ film on (0001) Al$_2$O$_3$
Fig. 5.27 Model fits with 2 layer (Al₂O₃/Osc/Osc+air) and 3 layer (Al₂O₃/Osc+air/Osc/Osc+air) model of 1000°C furnace annealed BaTiO₃ film on (0001) Al₂O₃ (Note: Osc means Oscillator.)
Fig. 5.28 Refractive index of RTA’d annealed, crystallized and re-oxidized BaTiO₃ films
Fig. 5.29 Depth profiles of BaTiO₃ films on (0001) Al₂O₃ after each annealing.

<table>
<thead>
<tr>
<th>Annealing Step</th>
<th>Osc + air(Ω)</th>
<th>Osc</th>
<th>Osc + air(Ω ± Δ)</th>
<th>σ</th>
</tr>
</thead>
<tbody>
<tr>
<td>750°C RTA</td>
<td>83 ± 2 Å</td>
<td>1647 ± 25 Å</td>
<td>577 ± 16 Å</td>
<td>0.81°</td>
</tr>
<tr>
<td></td>
<td>Osc</td>
<td>1554 ± 57 Å</td>
<td>500 ± 16 Å</td>
<td>0.72°</td>
</tr>
<tr>
<td>Crystallization (1000°C)</td>
<td>206 ± 16 Å</td>
<td>1554 ± 57 Å</td>
<td>500 ± 16 Å</td>
<td>0.69°</td>
</tr>
<tr>
<td>Re-oxidation (600°C)</td>
<td>207 ± 15 Å</td>
<td>1557 ± 59 Å</td>
<td>495 ± 19 Å</td>
<td>0.69°</td>
</tr>
</tbody>
</table>

Fig. 5.30 Total thickness of BaTiO₃ films on (0001) Al₂O₃ after each annealing.
5.4.3 Optical properties of nickel oxide

In order to model the BaTiO$_3$/Ni interface, reference optical properties for the NiO must first be known. There have been many studies on the optical properties of nickel oxide [1, 2, 28-33]. The reported data in the visible range differ in both the refractive index and especially the extinction coefficient ($k$), as shown in Fig. 5.31. Powell et al. [28] showed that a flame fusion grown NiO single crystal (SC) is transparent for wavelengths above 400 nm. Later work by Franta et al. [33] for pulse laser deposited NiO films is in good agreement for the extinction coefficient, although a slightly lower refractive index was obtained (possibly because of a lower density). The near-zero extinction coefficient of both samples suggests they are stoichiometric NiO (which is normally green in color). Lopez et al. [2] studied the optical properties of thermally oxidized NiO films by spectroscopic ellipsometry. In their work, sputtered Ni films were oxidized in air at temperatures ranging from 389-580°C. The optical properties of the films were grouped according to grain size, as shown in Fig. 5.31. The difference in extinction coefficient of the films may also be due to a variation in the amount of O in the NiO film. Lu et al. showed a strong dependence of the transmittance on the O$_2$/Ar ratio used during RF reactive magnetron sputtering of NiO films [31].
In section 5.1, initial results on the oxidation of Ni foils were presented. To further clarify the optical properties of the NiO (to enable a more quantitative optical evaluation of the Ni/BaTiO$_3$ interface) additional work was performed. In this experiment, the optical properties of thermally grown NiO on high purity Ni foils were studied by spectroscopic ellipsometry. To obtain more accurate optical constants, films at least 800 Å thick were used. For this purpose, a NiO film was prepared by annealing

**Fig. 5.31** Optical properties of a NiO single crystal (SC) and NiO films of different grain size (F1 and F2)
the Ni foil at 600°C in 10⁻⁶ atm pO₂ for 1 hr. A Sellmeier oscillator (Osc) was used to describe the optical properties of any NiO present.

The SE data were modeled using two structures 1) Ni/Osc/Osc+air) and 2) Ni/Osc+Ni/Osc+air. **Fig. 5.32** shows that model with Ni/Osc+Ni/Osc+air gave a better fit in psi values. Room temperature Ni optical properties (See section 5.1) were used for the Ni foil as they provided a better fit and a lower sigma value. The ~7% Ni mixed in the bulk of the NiO layer implied two possibilities. The “Ni” either represents the surface roughness of the Ni foil or it is simply a means of describing the absorption of the NiO film. Given that the mixed layer is quite thick (717.6±26.2 Å) compared to the total film thickness (850.6±43.4 Å), the latter possibility is more likely. Single term Sellmeier oscillators do not always describe absorption well across a wide wavelength range. Thus, the “Ni” in the layer does not necessarily imply the existence of metallic Ni in the film.

The net optical properties of the NiO film were calculated using Bruggeman effective medium theory (Eq. 3.15) and are shown in **Fig. 5.33**. It can be seen that the optical properties determined here agree well with data for thermally oxidized NiO films from Lopez et al.’s work [2]. Consequently, the optical properties of the thermally grown NiO film (Osc+Ni) was used to described any NiO in BaTiO₃ films on high purity Ni foils. **Fig. 5.34** shows a comparison of the optical properties of the thermally grown NiO film on Ni foil from this experiment and a 95% dense BaTiO₃ ceramic. The primary difference between the two is in the imaginary part of the dielectric function.
Fig. 5.32 Model fits of NiO thermally grown on high purity Ni foil at 600°C
<table>
<thead>
<tr>
<th>Osc + air (0.50)</th>
<th>133.0 ± 17.2 Å</th>
<th>( \sigma = 0.055 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osc + Ni (0.07 ± 0.01)</td>
<td>717.6 ± 26.2 Å</td>
<td>Ni</td>
</tr>
</tbody>
</table>

Fig. 5.33 Depth profile and refractive index of NiO thermally grown at 600°C
Fig. 5.34 Dielectric function of the thermally grown NiO and a BaTiO₃ ceramic sample

5.4.4 BaTiO₃ films on Ni foil

The depth profiles and optical properties of BaTiO₃ films (on Ni foil) RTA’d at different temperatures (600-750°C) were studied by spectroscopic ellipsometry (SE). Modeling of SE data was performed using reference data for the thermally grown NiO (i.e. the data measured in this work for NiO on Ni foil, see section 5.4.3) and a Sellmeier oscillator to describe the optical properties of NiO and BaTiO₃, respectively. In most cases, the best fit was obtained when the absorption in the oscillator was zero. Room temperature Ni reference data (see section 5.1) were chosen to describe the optical properties of the Ni foil because it improves the fit overall. SE data were truncated to 350-650 nm to obtain reasonable models.
SE data of a BaTiO$_3$ film on Ni foil RTA’d at 750°C was first modeled with various possible film structures, which at least consist of a bulk layer and surface roughness. It was found that an interface layer of either (oscillator+Ni) or pure NiO near the substrate results in good fits to the experimental data (see Fig. 5.35) and similar sigma values were obtained (i.e. 0.081 for mix of oscillator and Ni and 0.078 for pure NiO). It is noted that the BaTiO$_3$ film refractive index extracted from both models is the same and comparable to that of a BaTiO$_3$ ceramic sample. The large sigma is again mainly due to error in the experimental data when delta is close to 0 and 180°.

Although the model with a NiO interface shows a slightly better fit for this specific film, the resulting depth profile is not realistic (See Fig. 5.36). The thickness of the BaTiO$_3$ film is 1026.6±65.3 Å, which is approximately half of the total thickness of the film with the same number of spin-coated layers (8 layer) on (0001) Al$_2$O$_3$. In addition, the NiO layer is quite thick (902.9±49 Å), which is in disagreement with the fact that XRD did not detect NiO peak on this same film (see Fig. 5.13). As was described in section 5.4.3, the refractive indices of BaTiO$_3$ and NiO are quite similar. Thus, it is believed that the “NiO” is simply an artifact created by the model’s attempts to create a slightly more optically lossy BaTiO$_3$ film. A similar situation also occurred on modeling other RTA’d films. Thus, it is believed that the model without NiO at the interface is correct for all RTA’d films. It might be useful in the future to conduct a comparable set of measurements on either an ellipsometer with a rotating compensator or a photoelastic modulator to remove the errors where $\Delta$ is near 0 or 180° to see if that enabled a clearer distinction on the presence of interfacial NiO.
Fig. 5.35 Model fits of 750°C RTA’d BaTiO₃ film on high purity Ni foil
Fig. 5.36 Unreasonable film structure of RTA’d BaTiO₃ film on high purity Ni foil when modeled with NiO

It was found that modeling without NiO results in a more sensible film structure. Fig. 5.37 shows that the total thickness of the 750°C RTA’d BaTiO₃ is 1984 ±242 Å. At the film substrate/interface film there is a 343 ±106 Å thick layer with 9±5 volume % of Ni mixed in the BaTiO₃ film. The thickness of this layer agrees well with the rms roughness (37 nm) of the pre-annealed Ni foil, suggesting that this layer is due, at least in part, to the surface roughness of the Ni foil. In contrast to bare Ni, SE is able to detect the surface roughness of the Ni foil better with a transparent (insulating) film on the surface. The best fit without a NiO interface agrees well with the XRD results and cross-sectional TEM studies on RTA’d annealed films, which showed no detectable NiO.

Depth profiles for the best fit models of BaTiO₃ films RTA’d at other temperatures modeled without NiO are shown in Fig. 5.37. All show a reasonable volume fraction of Ni and thickness of the mixed layer near the substrate. It is noted that the 600°C RTA’d film does not show surface roughness on the BaTiO₃, suggesting that the majority of the surface roughness develops at higher temperatures. Fig. 5.38 and Fig. 5.39 show the refractive index and total thickness of BaTiO₃ films RTA’d at different temperatures. It can be seen that as the RTA temperature increases, further densification
and film crystallization occur, resulting in a higher refractive index for the BaTiO$_3$ film. The film thickness decreases after annealing at 600°C or 650°C and becomes stable above 700°C.

**Fig. 5.37** SE model details of BaTiO$_3$ films on Ni foil RTA’d at temperature from 600-750°C

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Layer Configuration</th>
<th>Thickness (Å)</th>
<th>σ</th>
</tr>
</thead>
<tbody>
<tr>
<td>600°C RTA</td>
<td>Osc</td>
<td>1864 ± 141</td>
<td>0.101</td>
</tr>
<tr>
<td></td>
<td>Osc + Ni(0.08 ± 0.04)</td>
<td>399 ± 127</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ni</td>
<td></td>
<td></td>
</tr>
<tr>
<td>650°C RTA</td>
<td>Osc + air(0.50)</td>
<td>53 ± 23</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Osc</td>
<td>1666 ± 109</td>
<td>0.086</td>
</tr>
<tr>
<td></td>
<td>Osc + Ni(0.11 ± 0.05)</td>
<td>346 ± 83</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ni</td>
<td></td>
<td></td>
</tr>
<tr>
<td>700°C RTA</td>
<td>Osc + air(0.50)</td>
<td>51 ± 32</td>
<td>0.119</td>
</tr>
<tr>
<td></td>
<td>Osc</td>
<td>1537 ± 137</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Osc + Ni(0.07 ± 0.03)</td>
<td>381 ± 126</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ni</td>
<td></td>
<td></td>
</tr>
<tr>
<td>750°C RTA</td>
<td>Osc + air(0.50)</td>
<td>58 ± 22</td>
<td>0.081</td>
</tr>
<tr>
<td></td>
<td>Osc</td>
<td>1583 ± 114</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Osc + Ni(0.09 ± 0.05)</td>
<td>343 ± 106</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ni</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Fig. 5.38** Refractive index of RTA’d BaTiO$_3$ films on high purity Ni foil

**Fig. 5.39** Total thickness determined by SE for BaTiO$_3$ films on high purity Ni foil as a function of RTA temperature
5.5 Electrical properties of BaTiO$_3$ films on Ni foil

Dielectric property measurements as a function of frequency and temperature were performed on BaTiO$_3$ films on Ni foils. The dielectric properties and its relationship to structure and microstructure are also discussed.

Fig. 5.40 shows the room temperature dielectric constant as a function of frequency for furnace annealed BaTiO$_3$ films on Ni foils. The films were RTA’d at different temperatures (600-750°C) before crystallization (1000°C, $10^{-16}$ atm pO$_2$) and re-oxidation (600°C, $10^{-6}$ atm pO$_2$). The dielectric constants of each film were measured from 30 spots because there was some variation. It is suspected that the variation could be due to film thickness, variable surface roughness of the foil, porosity and grain size.

The films show very high loss at low frequency (<1 kHz) due to space charge polarization and DC conduction. The average dielectric constants at 10 kHz of 750 and 650°C’d RTA film are similar and approximately 1100 and 1160, respectively. The films RTA’d at 600°C and 700°C show relatively lower dielectric constants (880 and 950), respectively. This could be a result of differences in the grain size, porosity, and solution aging as discussed in section 5.3.1. The films RTA’d at 600°C and 650°C have similar grain size but the 600°C RTA film contains more porosity. Thus, it has a lower dielectric constant. The film RTA’d at 700°C shows a slightly higher porosity and smaller grain size, compared to the 750°C RTA film. The loss tangent at 10 kHz of all film is less than 2%. The apparent increase in the loss tangent at high frequencies is due to an RC resonance and is not characteristic of the film.

Overall, the dielectric constant of these films is somewhat lower than those reported previously for BaTiO$_3$ films on metal foil ($\varepsilon_r \sim 2500$ [34] and $\varepsilon_r \sim 1300$ [3]) and
highly oriented (Ba,Sr)TiO$_3$ film on Ni foil ($\varepsilon_r \sim 1500$ [35]). The lower dielectric constant could be due to microstructure. It was shown that these films contain some porosity due to incomplete organic removal in the RTA under low pO$_2$.

![Graph](image)

**Fig. 5.40** Frequency dependence of dielectric constant and loss tangent furnace annealed BaTiO$_3$ films on Ni foil

The temperature dependence of dielectric constant was measured for BaTiO$_3$ films on a smoother Kodak Ni foil that became available at the very end of this work. After an RTA at 750°C in N$_2$, the film was crystallized at 1000°C for 1 h in $10^{-12}$ atm pO$_2$ and then re-oxidized at 600°C for 30 min in $10^{-6}$ atm pO$_2$. The RTA’d film show a room temperature dielectric constant (at 10 kHz) of 375 with a loss tangent of 4%. The furnace annealed and reoxidized film shows a dielectric constant of 1300 with <2% loss tangent.
The temperature dependence of the permittivity for the RTA’d film in Fig. 5.41 is highly dispersive. This could be due to incomplete densification of residual organics in the film after the RTA step, as was discussed in section 5.3.2. The furnace annealed film is less dispersive. There is a shallow maximum in the permittivity above 100°C which suggests that tetragonal-cubic phase transition in the films.
Fig. 5.41 Temperature dependence of dielectric constant and loss tangent of 750°C

RTA’d and furnace annealed BaTiO$_3$ films on (Kodak) Ni foil
References


Chapter 6

Conclusions and Future Work

This thesis has shown the use of different characterization techniques, i.e. X-ray diffraction, transmission electron microscopy, and spectroscopic ellipsometry to characterize the phase purity, microstructure, and interface quality of barium titanate thin films on Ni foil. In addition, organic decomposition during low oxygen pressure processing was also investigated. The conclusions of this work are described in section 6.1. The recommended future work is included in section 6.2.

6.1 Conclusions

Barium titanate thin films were deposited on Ni foils by chemical solution deposition. Drying and pyrolysis of the deposited films were performed on a hot plate in air. The deposited films were pre-annealed in a RTA at 600-750°C with an intermediate pyrolysis at 350°C in flowing N2. Then, a high temperature heat-treatment (at 1000°C) and re-oxidation (600°C) were done in a reducing furnace. The resulting films have dielectric constants of 1000-1300 which are stable as a function of temperature, with loss tangents less than 2%. The average grain size of furnace annealed (1000°C) films is 42 nm. The low oxygen partial pressure during the heat treatment was suspected to be the reason for such a small grain size. There are 5-6 grains across the thickness of a 200 nm thick film, which will provide grain boundaries to help block the migration of oxygen vacancies through the barium titanate.
NiO was not found in the films until the re-oxidation step. While NiO was detected in X-ray diffraction scans, it was not observed in cross-sectional TEM. Thus, it is believed that the NiO is not uniform and is spatially distributed in the films. It was found that there is a 5-8 nm thick Ni-Ba alloy interfacial layer on the Ni foil surface in the furnace annealed films.

Since high temperature annealing of the films was performed at low oxygen partial pressures, organic decomposition for both powders dried from barium titanate solutions and barium titanate films was studied. It was shown that complete decomposition of organics in N$_2$ flow is extended to higher temperatures, compared to that in air. Thus, when fast heating rates are used, as in an RTA (10°C/s), residual carbon is expected. The retained organics can cause porosity development during the high temperature annealing. The presence of residual organics was confirmed in the cross-sectional TEM study.

Spectroscopic ellipsometry was used to track the evolution of film thickness, optical properties and film density as a function of the processing conditions. The decrease in thickness as a function of drying temperature is consistent with the weight loss observed in TGA results. The refractive index of the films increases as the RTA temperature increases, suggesting further crystallization of barium titanate and organic decomposition. The furnace annealed (1000°C) film show a refractive index comparable to that of a 95% dense ceramic.

Investigation of NiO in barium titanate films on Ni foils was attempted by spectroscopic ellipsometry. To facilitate studies of the dielectric/electrode interface, the optical properties of thermally grown NiO on a Ni foil were extracted. It was found that
the Ni foil begins to oxidize at 300°C in air. The real part of the high frequency dielectric constant of NiO is similar to that of barium titanate, which complicated modeling of NiO in barium titanate films on Ni foil.

6.2 Future work

6.2.1 Organic removal and decomposition

As shown in this work, organic removal and decomposition become a very significant issue in processing chemical solution deposited films in low oxygen partial pressures. The delay in decomposition results in remaining of organics especially C in the films. Different approaches are suggested to minimize the organics and C in the films in future work.

One approach for this would be to insure that the concentration of organics in the film is reduced as much as possible in air at temperature up to 500°C without Ni oxidation. It has been reported by Losego [1] that if more reducing conditions are employed during the pyrolysis of the first layer of PZT films on copper, then the perovskite layer acts as a kinetic barrier to future oxidation. Thus, it was possible to use more aggressive pyrolysis conditions when subsequent layers were added. It would be useful to find out if a comparable procedure can be employed for BaTiO₃ on Ni. This would require that the minimum thickness of barium titanate film that can protect against subsequent Ni oxidation at low temperatures should be determined.

Alternatively, investigations could be performed using organic removal and RTA steps in an air ambient, accepting the existence of Ni oxidation. The NiO could subsequently be removed in reducing atmospheres at high temperature. It will be
important in this case to minimize the amount of oxidation of the Ni because the volume change on reduction from NiO to Ni.

Another related issue is the intermediate phase. BaCO$_3$ and a small amount of oxycarbonate were found as intermediate phases in crystallization of barium titanate powders and films, respectively. In this experiment, the powders were calcined in air while the films were RTA annealed in N$_2$ flow. The origin of the difference in crystallization pathways should be explored. Thus, the processing might be adjusted to avoid or minimize the intermediate phase, especially in low pO$_2$. This, in turn would, enable a decrease in the crystallization temperature of the barium titanate films.

**6.2.2 Interface of barium titanate film and Ni foil**

This work showed the presence of C in the barium titanate film as well as the existence of a Ni-Ba alloy interface between barium titanate film and Ni foil. However, it is not clear whether, in thin films, the occurrence of the alloy layer (or reduced Ti) are associated with this local C or not. This point should be clarified. Towards this end, it may be useful to determine the oxygen nonstoichiometry and C in a crystallized film (without re-oxidation) to compare to the current results. In addition, the composition of the alloy phase should be measured quantitatively. This may enable the height of the Schottky barrier between the electrode and the film to be estimated. A lower barrier height is expected to be problematic for the insulation resistance of the thin film capacitors. Finally, it is interesting to quantitatively determine whether there is a composition change or gradient in the barium titanate near the interface due to the loss of some Ba to form the alloy phase.
6.2.3 Spectroscopic ellipsometry for NiO on Ni foil investigation

This work has shown that the Ni foil surface roughness and the similarity of optical properties between NiO and barium titanate complicated the spectroscopic ellipsometry study. In particular, the changes in delta and psi (especially near 0 or 180 °) due to surface roughness and to an interfacial layer are in many ways comparable. The differences between them, however, were masked by the degraded accuracy of these values in a rotating analyzer ellipsometry measurement. To improve the sensitivity of ellipsometry for this characterization, a much smoother Ni foil is required for a rotating analyzer instrument. Alternatively, it may be possible to remove the errors in Δ by using a rotating compensator ellipsometer.
References

VITA

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