EFFECTS OF A COMBINED SUPERCRITICAL EXTRACTION/THERMAL CYCLE ON BINDER REMOVAL CYCLE TIME, YIELD, RESIDUAL CARBON, AND DEFECT FORMATION IN MULTILAYER CERAMIC CAPACITORS

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Brandon D. Abeln

Dr. Stephen J. Lombardo, Thesis Supervisor

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The undersigned, appointed by the Dean of the Graduate School, have examined the thesis entitled

EFFECTS OF A COMBINED SUPERCritical EXTRACTION/ThermAL CYCLE ON BINDER REMOVAL CYCLE TIME, YIELD, RESIDUAL CARBON, AND DEFECT FORMATION IN MULTILAYER CERAMIC CAPACITORS

Presented by

Brandon D. Abeln

A candidate for the degree of

Master of Science

And hereby certify that in their opinion, it is worthy of acceptance.

Dr. Stephen J. Lombardo _________________________________

Dr. William Jacoby _________________________________

Dr. Qingsong Yu _________________________________
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Brandon D. Abeln

Dr. Stephen J. Lombardo Thesis Supervisor

ABSTRACT

In the fabrication of multi-layer ceramic capacitors (MLCs), organic blends of binder and plasticizer are often utilized to aid in the forming and handling of ceramic green bodies. However, these organic constituents must be removed before the green body may be sintered into a dense component. Typically the binder is thermally decomposed in air, but as the size and binder loading of the MLC increases, the time required for binder removal may last up to several days in order to avoid introducing defects into the green body. Furthermore, thermal decomposition may leave a carbon residue within the dielectric which can modify the electrical properties and sintering behavior of the MLC.

In this work, a combined cycle consisting of a supercritical extraction cycle (SCE) followed by a thermal cycle (TC), was used to remove binder from green multilayer ceramic capacitors. The dielectric was barium titanate and the binder consisted of poly(vinyl butyral) plasticized with phthalates. Supercritical extraction of approximately
one-third of the binder leads to an increase in the porosity and gas permeability of samples and to a decrease in the adhesion strength between layers. The partial removal of the binder also resulted in a decrease in cycle time and defects following the combined SCE/TC. Samples subjected to the combined SCE/TC also had 25-30% less residual carbon as compared to samples subjected to a thermal cycle alone.

Supercritical extraction of binder in carbon dioxide has been shown to successfully remove large amounts of low molecular weight (MW) organic species but becomes less effective as the molecular weight of the organic components increases. A potential strategy to enhance the removal of the high MW components is to partially decompose the species during the extraction process. This work also presents the binder removal efficiencies from tape cast films fabricated with titanate-based dielectrics. The organic systems evaluated are plasticized acrylic, poly(propylene carbonate), and poly(vinyl butyral) binders. The effect of temperature on binder removal and defect formation is assessed.
1.0 INTRODUCTION

Multilayer ceramic capacitors (MLCs) are widely manufactured for use in a number of electronic devices such as TVs, cellular phones, and personal computers [1-3]. MLCs are extensively used in electronic circuitry because of their low cost and high volumetric efficiency (capacitance per unit volume) [1,4-6]. From 2009 to 2011, the number of MLCs manufactured is expected to increase from 1.5 trillion to over 2 trillion [1]. Figure 1.1 is an image of a typical multilayer ceramic capacitor manufactured in industry with ceramic dielectric and Pt/Pd/Au metal electrodes.

![Image of a multilayer ceramic capacitor containing ceramic dielectric and Pt/Pd/Au metal electrodes.](image)

**Figure 1.1:** Image of a multilayer ceramic capacitor containing ceramic dielectric and Pt/Pd/Au metal electrodes.

The manufacture of noble-metal and base-metal multilayer ceramic capacitors is usually comprised of the steps outlined in Fig. 1.2. First, organic additives such as binder, plasticizer, and dispersant are milled together with an appropriate proportion of solvent and dielectric powder to aid in the subsequent processing steps. The type and quantity of binder used has an influence on the specific properties of the final MLC such
as the green strength, the adhesion strength between layers, and the permeability to gas flow. After milling, the slurry is tape cast into a thin film, which forms into a green ceramic tape upon drying. The term “green” refers to the samples at any point prior to the sintering step. Once the ceramic tape is dried, cut into individual pieces, and screen printed with metal electrodes, the individual pieces are stacked and laminated into MLCs. The conditions of the lamination process via the lamination time, temperature, and pressure can also affect the aforementioned properties of the MLCs [7-9]. Following lamination, the organic fraction must ultimately be removed in the binder removal process before the ceramic green body can be sintered into a dense component [10,11]. In this work, the specific characterization techniques and machinery used to fabricate, analyze, and process ceramic green bodies are further described in Chapter 2.

**Figure 1.2:** Steps involved in the processing of multilayer ceramic capacitors. Both a thermal cycle alone and a combined supercritical extraction/thermal cycle are outlined for the binder removal step.
For the binder removal step, the organic fraction is traditionally eliminated in furnacing operations in which the organic constituents of the binder are either vaporized, oxidized, or pyrolyzed into gas phase species. The evolution of these species into the pore space of the green body thus leads to an increase in pressure, which in turn causes stress [12-14] within the component that ultimately may lead to defects such as fracture or delamination [15-20]. To avoid failure of green components during binder removal—and thus increase yield—several strategies are available. Most often, once a binder system has been developed, heating schedules are specified in which multiple ramp rates, hold temperatures, and hold times are selected so that the rate of binder decomposition is sufficiently retarded in order to minimize the pressure and hence stress within the green body. The heating schedules become longer and more complex as the permeability of the green body decreases, which occurs when (a) the size of the capacitor increases, (b) the binder loading within the green body increases, or (c) the particle size of the dielectric powder decreases. Thus, in practice, specification of an ideal heating schedule becomes a compromise between achieving a short cycle time and a high product yield.

To avoid the aforementioned drawbacks associated with the thermal removal of binder, supercritical extraction (SCE) has been proposed and demonstrated as an alternative processing route [21-31]. Under supercritical conditions, organic species dissolve into the supercritical fluid and then diffuse out of the green body, thus effecting binder removal without additionally increasing the pressure in the pore space. It is well known that supercritical carbon dioxide is most effective in removing organic species of lower molecular weight such as short chain waxes [17-19] or plasticizers [25-28,30,31].
Higher molecular weight organic species such as polystyrene (MW = 150,000) have been known to dissolve in supercritical hexane [32]; however by operating at the critical temperature of hexane (234.5 °C) the rate at which binder inside the MLC is decomposed will be so rapid, that damage will probably occur [25]. For the case of MLCs, the preferential extraction of plasticizers may lead to 40-60% removal of the organic fraction [25,26], depending on the components in the binder blend, and substantial partial extraction of binder can lead to increases in both the porosity and gas permeability of the green body. Even though SCE may not be capable of fully removing all of the organic constituents, a subsequent thermal cycle (TC) may be used to remove any remaining organic species. This thermal cycle may be realized as a separate furnace operation or as part of the sintering cycle, and may be conducted under oxidizing, reducing, or inert conditions.

In summary, although supercritical extraction can remove binder, either partially or nearly completely from green bodies, its efficacy as an alternative processing strategy in terms of benefits to cycle time and yield—the same issues which pertain to thermal debinding—have not been specifically addressed in the literature. It is thus one aim of Chapter 3 to demonstrate that a combined supercritical extraction/thermal cycle (SCE/TC) can be used to decrease overall processing time while avoiding defects but still maintaining high yield. This combined cycle may be rapid because supercritical extractions times can be short, often on the order of hours, and the aforementioned increases in porosity and gas permeability of the green body may consequently facilitate a subsequent rapid thermal cycle. In addition to this objective, we have also observed in
earlier work that changing the conditions of an upstream process step in the manufacture of MLCs, namely in the lamination conditions, can also influence the yield during binder removal [9], and thus we report and discuss on this aspect of processing for the combined SCE/TC as well.

In Chapter 4, we note that another shortcoming of thermal debinding is that carbonaceous residues may remain in the green body and these may either inhibit sintering or degrade the electrical properties of the final components. The role of carbon contamination on device performance is especially important in the processing of MLCs with base metal electrodes in which non-oxidative conditions during binder removal are typically used at temperatures above 270 °C in order to avoid oxidation of the base-metal electrode material [33-37]. We therefore additionally demonstrate that a combined SCE/TC can also impact the residual carbon present in base-metal MLCs, which is a current processing trend for MLCs.

Although supercritical extraction has been shown to effectively remove 40-60 wt% of the organic fraction [25,26], defects such as cracks and delamination have been occasionally observed in MLCs [38]. Such defects were observed predominantly in MLCs that exhibited little loss of binder during the extraction cycle, and therefore had low permeability to gas flow. It was then postulated that failure occurred during depressurization due to pressure gradients that arise because of the low gas permeability. A mathematical model consequently was developed to quantify the magnitude of the pressure gradients [38]. The occurrence of defects in samples exhibiting low weight loss suggested the exploration of more aggressive extraction conditions by operating at higher
temperatures. Under these circumstances, binder degradation may also be contributing to the observed weight loss, and possibly to the occurrence of defect formation in the samples as well.

In Chapter 5, the origin of defect formation in green ceramic bodies is thus examined for samples fabricated with three binder systems: poly(propylene carbonate)-based, acrylic-based, and poly(vinyl butyral)-based. Three potential mechanisms for defect formation are additionally considered: failure during depressurization, failure from enhanced binder degradation due to the presence of residual air in the vessel, and failure due to mechanical vibrations from gas compression when going to high pressure. Experiments were conducted on both individual tapes and MLCs.

Finally in Chapter 6, we summarize the effects of operating with a combined supercritical extraction and thermal cycle on cycle time, yield, carbon content, and defect formation in multilayer ceramic capacitors. The chapter then addresses potential areas for further investigation to improve the manufacture and yield of MLCs.
1.1 REFERENCES


CHAPTER 2

GENERAL EXPERIMENTAL
2.0 FABRICATION OF GREEN TAPES

In this work, ceramic tapes were prepared by first combining dielectric powder, solvent, binder, plasticizer, and dispersant in appropriate proportions to create a slurry. After the contents were milled together, the slurry was poured onto a sheet of mylar which rested on the bed of a tape caster. The tape caster, shown in Figure 2.1, contained a doctor blade which spanned the width of the bed and was slightly elevated above the mylar. The doctor blade was then used to pull the bulk of the slurry across the length of the bed leaving behind a thin film, which formed into a green ceramic tape after drying.

![Figure 2.1: Schematic of a ceramic tape cast.](image-url)
2.1 SCREEN PRINTING OF ELECTRODES

Nickel paste was prepared by first combining dielectric powder, nickel powder, solvent, binder, and dispersant in various proportions to create a slurry. After the slurry was milled, the resulting nickel paste was poured onto a custom designed 400 calendared SS mesh screen (UTZ LLC, Clifton, NJ). Nickel metal electrodes were then printed 2-3 μm thick onto 1×1 inch pieces of ceramic tape. For printing, the nickel electrodes were applied to the tapes with either a screen printer (Model 645, Affiliated Manufacturers Inc., North Branch, NJ), shown in Figure 2.2, or a hand squeegee (80 durometer, Saati, Milan, Italy). When using a screen printer, the ceramic tape, with the mylar backing, was first placed onto a perforated loading platform and then held stationary with vacuum. During printing, the mobilized loading platform was automatically positioned under the screen where a squeegee then transferred metal electrodes onto the ceramic tape as the nickel paste was pulled across the screen.

When screen printing with a hand squeegee, the mylar backing of the ceramic tape was first attached to a piece of overlapped adhesive tape (Duck, Henkel Consumer Adhesives Inc., Avon, OH) and then pressed to a flat surface. After the screen was manually aligned over the ceramic tape, the nickel paste was pulled across the screen with the hand squeegee, as shown in Figure 2.3a. Figure 2.3b is an image of a ceramic tape after screen printing nickel metal electrodes.
**Figure 2.2:** Image of a screen printer.

**Figure 2.3:** Image of (a) a screen printing apparatus and (b) a ceramic tape containing nickel metal electrodes.
2.2 LAMINATION OF TAPES

A 25 ton press (Model 2518, Carver Inc., Wabash, IN) shown in Figure 2.4, was used to laminate individual green tapes into MLC samples. Two pressure gauges, designed specifically for the Carver press, were calibrated to measure both the pressure inside the hydraulic chamber in units of pound-force per square inch, as well as the applied load, in units of pound-force and metric tons. One pressure gauge was designed to measure small loads that ranged from 0-5,000 lb$_f$, while the other pressure gauge measured larger loads up to 50,000 lb$_f$. The control valve separating the two pressure gauges was closed prior to pressing if the calculated load was near or exceeded 5,000 lb$_f$.

![Image of a 25 ton Carver press with labels A-J:]
- A- Temperature Controller for Platen 1
- B- Temperature Controller for Platen 2
- C- Power Switch
- D- 5,000 lb$_f$ Gauge
- E- 50,000 lb$_f$ Gauge
- F- Control Valve
- G- Pressure Release Valve
- H- Lever
- I- Platen 1
- J- Platen 2

Figure 2.4: Image of a 25 ton Carver press.
For lamination, the temperature of each platen was adjusted with corresponding electronic temperature controllers (Model 6100, West, The Hyde Business Park Brighton, United Kingdom) which displayed both the actual platen temperature and the set-point platen temperature in units of degrees Fahrenheit as shown in Figure 2.5. The set-point temperature on each controller was adjusted by pressing the function button, which opened a settings menu where the directional arrows were pressed to increase or decrease the set-point value. The controller was then returned to the main display where the set-point temperature reflected the changed value after the function button was pressed once again.

![Image of the digital temperature controllers on a 25 ton Carver press.]

**Figure 2.5:** Image of the digital temperature controllers on a 25 ton Carver press.

Prior to lamination, the ceramic tape was first cut into individual layers and the mylar backing was removed. The layers were then stacked in a “bottom to top”
orientation with the “bottom” representing the side which was initially attached to the mylar backing. Next, the non-stick side of a new piece of mylar was determined by pressing some adhesive tape to each side and physically feeling the obvious difference in resistance while pulling the tape off. The side with the least resistance contained the non-stick coating. Mylar was then placed between the stack and the platens with the “non-stick” side of the mylar in contact with the ceramic tape as shown in Figure 2.6.

![Layered Tapes](image)

**Figure 2.6:** Image of a stack of layered ceramic tapes prior to lamination.

After the stack of tapes, surrounded by mylar, was placed on the center of the lower platen, the load required for lamination was calculated by multiplying the desired lamination pressure by the surface area of an individual layer. Once the platens were at the set-point temperature, the stack was manually laminated with the lever at a constant load. After pressing, the resulting MLC was removed from the press and cooled at room temperature for at least five minutes before the mylar was removed. The distorted shape of the MLC following lamination, referred to as “flow,” was then evaluated based on the expansion of the stack relative to the original size, as depicted in Figure 2.7.
Samples were fabricated for permeability and porosity measurements by first cutting ceramic tape into individual layers of $3.0 \times 3.0 \times 0.015$ cm. Next, four layers were stacked in a “bottom to top” orientation and laminated at 5 MPa for 10 minutes at either 85 or 95 °C. After lamination, the samples were cut with a surgical blade (# 11, Feather Safety Razor Co., Osaka, Japan) into disk geometry with an approximate diameter of 2.5 cm.

Samples were fabricated for peel test measurements by first laminating two green tapes of $4.0$ cm length $\times 1.0$ cm width together at 5 MPa for 10 minutes at either 85 or 95 °C to form a substrate. After forming an additional substrate, the two substrates were layered and then partially separated by mylar (see Fig. 2.8). For lamination, two 0.3 cm thick aluminum plates were placed on Platen 2 and heated to the lamination temperature of the press. After the temperature stabilized, as measured by a type K thermocouple attached to a digital multi-meter (Omega Engineering Inc, Stamford, CT), part of the layered substrate, not separated by mylar, was placed between the aluminum plates, as
shown in Figure 2.8. The two substrates were then laminated together at the above conditions to form a T-shaped peel sample.

**Figure 2.8:** Schematic of how two green ceramic substrates are laminated together to form a T-shaped peel sample where the filled-in arrows indicate the direction of the applied load.
2.3 CUTTING AND PHOTOGRAPHING MULTILAYER CERAMIC CAPACITORS

After the mylar surrounding the laminated MLC was removed (see section 2.3), the MLC samples were cut. To avoid delamination while cutting the MLCs, a metal ruler (Model# R590-12, Alvin & Company Inc., Bloomfield, CT) was placed on top of the sample and then pressed by hand. Next, the MLC was cut by running a surgical blade (Feather Safety Razor Co.) along the edge of the metal ruler as shown in Figure 2.9. After cutting, each edge of the MLC was observed at 40× magnification under a light microscope (Edmund Scientific, Barrington, NJ) to check for delamination.

Following observation and/or experiments, the MLC samples were photographed using a 6 megapixel digital camera (Canon EOS 10D, Lake Success, New York) attached to a 100 mm macroscopic lens and a 1:1× matched macro adaptor (Promaster, Japan). Photographs were taken under fluorescent lighting only while using a tripod (Bogen Model 3011, Manfrotto Nord, Bassano del Grappa (VI) – Italy) with the camera set to manual mode having a shutter release and f/stop setting of 1.0 second and 13 respectively.
Figure 2.9: Schematic for cutting Multilayer Ceramic Capacitors (MLCs).
2.4 GAS PERMEABILITY MEASUREMENTS

Permeability measurements were performed by measuring the molar flux of N\textsubscript{2} across four laminated tapes of disc geometry after lamination, after SCE, and after the combined SCE/TC. Following lamination, the thickness of each sample was measured with a drop pin gauge accurate to within 0.001 inches, as shown in Figure 2.10; the thickness was recorded as an average of five measurements at different locations. Next, the sample was loaded into a holder (Millipore, Billerica, MA) which had an open area of 1.56 cm diameter. The holder was then mounted into the permeability apparatus shown in Figure 2.11, which consisted of a pressure regulator (Valve 1, not shown) attached to a nitrogen tank, a pressure gauge (P\textsubscript{1}), a sample holder, a second pressure gauge (P\textsubscript{2}), a control valve (Valve 2), and a flow meter (50 ml, Kimble Chase Life Science, Vineland, NJ) connected in sequential order.

![Image of a drop pin gauge. The crosses on the sample represent the location of each measurement.](image)

**Figure 2.10:** Image of a drop pin gauge. The crosses on the sample represent the location of each measurement.
Figure 2.11: Image which outlines the flow of N$_2$ through various components within the permeability apparatus.

After the sample holder was mounted into the permeability apparatus, Valve 2 was completely closed and Valve 1 was slowly opened so that nitrogen could flow into the system. If P$_2$ remained at atmospheric pressure as P$_1$ approached 80 psi, then Valve 1 was closed and permeability of the sample was considered negligible. If a pressure increase was observed in P$_2$, then Valves 1 and 2 were adjusted so that P$_1$ and P$_2$ initially measured 20 psi and 5 psi, respectively, as nitrogen flowed from the tank to the flow meter.
The volumetric flow rate of nitrogen through the sample holder was determined with a flow meter, which consisted of a 50 ml tube and a rubber squeeze bulb filled with a mixture of soap and water. After the rubber squeeze bulb was compressed, the time required for the N₂ gas to push the resulting bubble up the flow meter from 40 ml to 10 ml was measured, as shown in Figure 2.12. Several bubbles were formed prior to each measurement to reduce friction at the wall of the 50 ml tube. The volumetric flow rate, \( V \), was determined as an average of three measurements for a given \( P_1 \) and \( P_2 \) and then converted to a molar flow rate, \( n \), from

\[
n = \frac{P_f V_f}{R T_f}
\]  

(2.1)

where \( R \) is the gas constant, \( T \) is the temperature, \( P \) is the pressure, and the subscript \( f \) denotes the conditions at the flow meter. The molar flux was then determined by dividing the molar flow rate by the area, \( A \), of the substrate as given by:

\[
N_m = \frac{n}{A}
\]  

(2.2)

Once the volumetric flow rate and the change in pressure from \( P_1 \) to \( P_2 \) across a sample of thickness, \( L \), were known, the permeability, \( \kappa \), was calculated from the measured flux data as

\[
\kappa = \frac{-2RTN_m \mu L}{P_2^2 - P_1^2}
\]  

(2.3)

where \( \mu \) is the viscosity of the gas. The permeability of the sample was recorded as the average of four measurements from four different flow rates of N₂.
Figure 2.12: Image of a 50 ml flow meter used to measure the volumetric flow rate of flowing nitrogen.
2.5 POROSITY MEASUREMENTS

After determining the permeability, the dry, suspended, and saturated weights of the sample were measured with a scale (Model R160P, Sartorius Co., Bohemia, NY) and then the porosity, \( \varepsilon \), was calculated using Archimedes’ method. After recording the dry weight, a 600 ml beaker (Model 14000, Kimble Chase Life Science) was filled with deionized water and placed on an inverted zirconia setter which rested on a hotplate (Fisher Scientific, Pittsburgh, PA). Once the temperature of the water reached 55 °C, as measured with a mercury thermometer (Model 14-985B, Fisher Sci., Pittsburgh, PA), the sample was placed on a 1.5 inch diameter 304 SS perforated disk (P.O. AC-30654660, Purolator EFP, Huston TX) which was then suspended in the water for 24 hours.

To measure the suspended weight, the sample was first removed from the 600 ml beaker and placed onto another perforated disk (P.O. AC-30654660, Purolator EFP) which was suspended from the bottom of the scale. Next, the disk was submerged into a different container which contained deionized water and approximately 3 ml of a wetting agent (Triton X-100, Ricca Chemical Co., Arlington, TX) at room temperature. After determining the suspended weight, excess water droplets on the surface of the sample were removed with a damp paper towel by lightly patting the sample so the saturated weight could be measured. Once the dry \( (D) \), suspended \( (S) \), and saturated \( (W) \) weights of the sample were known, the percent porosity, \( P \), was calculated with:

\[
P, \% = \frac{W - D}{W - S} \times 100
\]  (2.4)
2.6 ADHESION-STRENGTH MEASUREMENTS

The adhesion strength between tape layers was determined by performing peel tests [1-4] on the T-shaped peel sample described in section 2.3. Following lamination, the peel sample would either be subjected directly to the peel test or first to a particular cycle, e.g., SCE, and then to the peel test. For the peel test, the two unlaminated parts of the T shaped sample were placed in the grips of a tensile tester (TA TX Plus Texture Analyzer, Stable Micro Systems Ltd, Surrey, UK) and then pulled apart as shown in Figure 2.13. During testing, the load applied to the sample was measured as a function of displacement as the top crosshead moved vertically at a speed of 0.4 cm s$^{-1}$. Since the elongation of sample due to strain was considered negligible during testing, the adhesion strength, $\sigma$, was given by

$$\sigma = \frac{F_1}{b} \quad (2.5)$$

where $F_1$ is the force applied to the sample and $b$ is the sample width.
Figure 2.13: Schematic of a peel sample placed into the grips of a tensile tester which pulled the sample apart as the top crosshead moved vertically away from the base.
2.7 RESIDUAL CARBON MEASUREMENTS

Following thermal oxidation, the residual carbon (RC) content of samples was determined with a residual carbon analyzer (Model C-144, LECO, St. Joseph, MI) shown in Figure 2.14. For each style of MLC evaluated, 5 MLCs were prepared and the average percent residual carbon (%RC) was calculated. Because less than 100 measurements were taken at each condition, 90% confidence intervals (C.I.) were evaluated using statistical analysis software (SAS) according to:

\[ C.I. = \bar{a} \pm t^* \frac{\sigma}{\sqrt{n}} \]  

(2.6)

where \(\bar{a}\) is the sample mean, \(\sigma\) is the standard deviation, \(n\) is the number of samples, \(t^*\) is a value obtained from the t-distribution tables, and \([ t^* \frac{\sigma}{\sqrt{n}} ]\) is the standard error of the mean.

For RC analysis, the MLC was first chopped with a razor blade (PO# 11140, ACE Hardware Corp., Oak Brook, IL) into a powder yielding an average particle diameter of 1 mm. The weight of the MLC was then determined within 0.1 mg by placing the sample into a zirconium silicate boat (Part# 529-204, LECO, St. Joseph, MI) which rested on a scale (Model LA 120S, Sartorius Co., Bohemia, NY). Next, the chamber inside the RC analyzer was purged with flowing O\(_2\) for 5 seconds at 3.5 LPM after which the chamber door was opened and a rod was used to push the boat into the furnace against the heating element, which was maintained at 927 °C. During analysis, O\(_2\) flowed through the analyzer at a constant 2.5 LPM and the evolved carbon dioxide was measured with an infrared detector. If the volume of a single MLC exceeded the capacity of the boat, then
one third of the pulverized MLC was analyzed per run, and the %RC was recorded as an average of 3 runs.

![Figure 2.14: Image of a residual carbon analyzer and other components required for measuring the residual carbon content of MLCs.](image)

To determine the precision and accuracy of the RC measurements, three runs, each containing 250 to 400 mg of a 1% carbon standard, were evaluated prior to analyzing the first MLC. If the accuracy was less than 0.015%, then the RC analyzer was recalibrated with software which performed a linear regression over the previous three measurements. Next, three additional runs were evaluated and the calibration process continued until results within 0.015% were achieved.
2.8 SUPERCRITICAL EXTRACTION EXPERIMENTS

The supercritical extraction (SCE) experiments were conducted in a SS 316 6.25 cm diameter 500 ml high pressure non-stirred vessel (P.O. 1834HC2, Parr Inst. Co., Moline, IL). The lid of the extraction vessel, shown in Figure 2.15, was equipped with a thermocouple well (P.O. A1453HC, Parr Inst. Co.), a gauge block assembly (P.O. 4317PK, Parr Inst. Co.), two different needle valves (P.O. NA176VB, Parr Inst. Co.); (P.O. 30VM-4071, Autoclave Engineers, Erie, PA), and a gasket mount. The original rupture disk included with the gauge block assembly was discontinued, so a replacement disk (P.O. 526HCPH, Parr Inst. Co.) which ruptures at 5000 psi was used. Depending on whether the experiment was conducted in supercritical carbon dioxide or supercritical hexane, either a polyteflon (PTFE) (P.O. 457HC2, Parr Inst. Co.) or carbon fiber (grafoil) gasket (P.O. 457HC3KL, Parr Inst. Co.) was installed, respectively.

![Image of the supercritical extraction vessel lid](image)

**Figure 2.15:** Image of the supercritical extraction vessel lid.
The phase of the extraction fluid at ambient conditions determined how the samples were loaded into the extraction vessel. When utilizing carbon dioxide as the supercritical fluid, the sample was placed onto a perforated pedestal which was then lowered into the vessel as shown in Figure 2.16. When utilizing hexane as the supercritical fluid, the vessel was first charged with $400 \pm 4$ ml of liquid hexane (HPLC grade $\geq 95\%$ purity, Sigma-Aldrich) and then the MLC was placed in a sample holder made from a SS 304 perforated metal sheet. Next, the sample holder was wired to the top of the thermocouple well so the sample rested above the liquid hexane, as shown in Figure 2.17.

![Figure 2.16: Schematic of a MLC loaded into a vessel.](image)

The split compression ring (P.O. A246HC2, Parr Inst. Co.) shown in Figure 2.18 was used to seal the vessel cover to the vessel by tightening one compression bolt from each ring at a time with the second bolt located 180 degrees from the first. The amount of torque applied to each compression bolt varied with the type of gasket installed. For the PTFE and grafoil gasket, each compression bolt was tightened at 30 ft-lbs and
55 ft-lbs, respectively. After placing a SS sleeve (P.O. A247HC, Parr Inst. Co.) around the split compression ring, the assembled vessel was lifted by the inlet and outlet tubing stems and inserted into the furnace (“Heater Assembly” P.O. 4920, Parr Inst. Co.) of the supercritical extraction apparatus.

**Figure 2.17:** Schematic of a MLC loaded into a vessel containing 400 ml of hexane.

The supercritical extraction apparatus was comprised of a furnace, a controller (Model 4836, Parr Inst. Co.), a type J thermocouple (P.O. A472E2, Parr Inst. Co.), a flow meter (Model VFB, Dwyer Inst. Inc., Michigan City, IN), a compressor (Model AG-152, Haskel Eng. and Supply Co., Salisbury, Australia), a computer (Model M6300, Dell Inc., Round Rock, TX), and high pressure gas cylinders. Figure 2.19 is a diagram which outlines the direction of gas and electrical current flow through each component of the SCE apparatus used for all the experiments in this investigation. Because the flow meter was calibrated for air, the actual flow of gas was obtained with
\[ Q_2 = Q_1 \times \sqrt{\frac{1}{\text{S.G.}}} \]  

(2.4)

where \( Q_2 \) is the actual flow corrected for the specific gravity (S.G.) of the gas used, and \( Q_1 \) is the measured flow.

Figure 2.18: Image of the split compression ring used to attach the vessel lid to the vessel.

A unique procedure was required to heat the vessel when the set-point temperature (\( T_{\text{set}} \)) was below 200 °C. Table 2.1 lists values for \( T_{\text{set}} \) along with the corresponding parameters necessary to heat and maintain the vessel temperature in air. For heating, the integral and derivative settings on the controller were always set to 60 and 0, respectively, while the primary proportionality band (\( P_1 \)) and percent power level (PL.1) settings varied with \( T_{\text{set}} \). The remaining control parameters were left at the values
set by the manufacturer. For each experiment, the amount of power used to heat the vessel to $T_{set}$ ($PL_{1ramp}$) was different than the amount needed to maintain the temperature at $T_{set}$ ($PL_{1dwell}$). To heat the vessel, a constant amount of power at $PL_{1ramp}$ was supplied to the furnace for a length of time ($t_{ramp}$), which depended on both the value of $T_{set}$ and the atmosphere inside the vessel. After the temperature reached $T_{set}$, the value of $PL_{1ramp}$ was manually changed to $PL_{1dwell}$.

![Flow diagram of the supercritical extraction apparatus.](image)

**Figure 2.19:** Flow diagram of the supercritical extraction apparatus.

**Table 2.1.** Parameters used to heat a 500 ml vessel in air to various set-point temperatures ($T_{set}$).

<table>
<thead>
<tr>
<th>$T_{set}$ (°C)</th>
<th>$PL_{1ramp}$ (%)</th>
<th>$t_{ramp}$ (min)</th>
<th>$P_1$ (band)</th>
<th>$PL_{1dwell}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>40</td>
<td>23.0</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>120</td>
<td>70</td>
<td>19.0</td>
<td>5</td>
<td>15</td>
</tr>
<tr>
<td>150</td>
<td>70</td>
<td>25.5</td>
<td>5</td>
<td>15</td>
</tr>
</tbody>
</table>
For most of the SCE experiments, the SCE cycle was comprised of a purge step, a pressurization step, a dwelling step, and a depressurization step. When operating with carbon dioxide as the supercritical fluid, the purge consisted of a three minute exposure to flowing gas at approximately 0.8 LPM at room temperature and pressure to remove air from the extraction vessel. During the purge, the controller settings were set to the same values required to heat the vessel in air to a given $T_{\text{set}}$, as described earlier. Next, the vessel pressure was equilibrated with the pressure in the CO$_2$ cylinder, and then a compressor was used to increase the pressure to an initial value ($P_{\text{initial}}$) listed in Table 2.2. Upon closing the inlet valve to the vessel, however, $P_{\text{initial}}$ decreased by ~2 MPa. The final pressure of 30 MPa was attained by supplying power at PL.1$_{\text{ramp}}$ (see Table 2.1) to the furnace for the amount of time, $t_{\text{ramp}}$, listed in Table 2.2. If the pressure reached 30 MPa at a temperature below $T_{\text{set}}$, then the supercritical CO$_2$ was vented at the vessel outlet so that the pressure remained constant. After dwelling at $T_{\text{set}}$ for 1 hour, the vessel was depressurized isothermally to atmospheric conditions. After depressurization, the extraction efficiency was determined from the weight loss of the samples, normalized by the total amount of organic phase initially present.

Table 2.2. Parameters used to thermally pressurize a 500 ml vessel to 30 MPa with carbon dioxide at various set-point temperatures ($T_{\text{set}}$).

<table>
<thead>
<tr>
<th>$T_{\text{set}}$ ($^\circ$C)</th>
<th>$P_{\text{initial}}$ (MPa)</th>
<th>$t_{\text{ramp}}$ (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>12</td>
<td>25</td>
</tr>
<tr>
<td>120</td>
<td>10.5</td>
<td>21</td>
</tr>
<tr>
<td>150</td>
<td>9</td>
<td>27</td>
</tr>
</tbody>
</table>
When the dwell step consisted of multiple exposures of supercritical CO$_2$ instead of a single exposure, the temperature and pressure were first held constant for 1 hour after which the vessel was depressurized in a non-isothermal manner at a constant rate of 8.0 LPM to 10 MPa. Next, the compressor re-pressurized the vessel to 30 MPa at $T_{\text{set}}$ with fresh CO$_2$. After re-pressurization, if necessary, the proportional band was slightly modified from the values indicated in Table 2.1 so the temperature remained within $\pm 2^\circ\text{C}$ of $T_{\text{set}}$. This sequence of steps —pressurization, 1 h dwell, depressurization— was then repeated for 2 additional cycles before the vessel was depressurized isothermally to atmospheric conditions.

When operating with hexane as the supercritical fluid, the vessel was first purged with argon gas at 0.85 LPM for 5 minutes to completely remove oxygen. During the purge, insulation (P.O. 9328K42, McMaster-Carr, Elmhurst, IL) was placed on top of the heating assembly, and the $P_1$, $P_2$, $I$, $D$, and $PL_{\text{ramp}}$ settings on the controller were changed to 47, 47, 41, 158, and 100, respectively, with $P_2$ representing the secondary proportionality band. The remaining control parameters were left at the values set by the manufacturer. Next, the temperature of the vessel was increased from room temperature to 270 $^\circ\text{C}$, which pressurized the vessel from 0.1 MPa to 26 MPa as the phase of the hexane changed from liquid to supercritical. After dwelling for 2 hours, the vessel was depressurized thermally to $\sim 0.1$ MPa and 30 $^\circ\text{C}$, after which the MLC was removed and dried in an oven at 80 $^\circ\text{C}$ for 24 hours. These steps —charging, pressurization, dwell, depressurization, drying— were then repeated for 1 additional cycle.
2.9 REFERENCES


CHAPTER 3
EFFECTS OF A COMBINED SUPERCRITICAL EXTRACTION/THERMAL CYCLE ON BINDER REMOVAL CYCLE TIME AND YIELD IN MULTILAYER CERAMIC CAPACITORS
3.0 INTRODUCTION

In the fabrication of multilayer ceramic capacitors (MLCs), organic blends of binder and plasticizer are used to aid in the casting and lamination of ceramic tapes. Following lamination, the organic fraction must be removed before the ceramic green body can be sintered into a dense component. The most common method for binder removal is thermal decomposition of the organic constituents into gas phase species. The evolution of these species into the pore space of the green body, however, leads to an increase in pressure, which in turn causes stress [1-3] within the component that ultimately may lead to defects such as fracture or delamination [4-9]. To avoid failure of green components during thermal binder removal, the rate of binder decomposition is sufficiently retarded in order to minimize the pressure and hence stress within the green body. Thus, in practice, thermal cycles (TC) alone become a compromise between achieving a short cycle time and a high product yield.

To avoid the aforementioned drawback associated with the thermal removal of binder, supercritical extraction (SCE) has been proposed and demonstrated as an alternative processing route [10-20]. Even though SCE may not be capable of fully removing the entire organic fraction, the resulting increase in porosity within the green body may permit a rapid removal of the remaining organic species during a TC. Thus, the objective of this study is to demonstrate that a combined supercritical extraction/thermal cycle (SCE/TC) can reduce the overall processing time while avoiding defects and still maintaining high yield. In addition to this objective, we have also observed in previous work that altering the lamination conditions during the manufacture
of MLCs has an impact on the processing yield during binder removal [21], and thus we report and discuss on this aspect of processing for the combined SCE/TC as well.
3.1 EXPERIMENTAL

In this study, green tape samples were prepared which consisted of 86.6 wt% barium titanate powder (Ferro X7R422H, Ferro, Niagara Falls, NY), 6.7 wt% poly(vinyl butyral) (PVB) resin (Butvar B98, Richard E. Mistler, Inc., Yardley, PA), 5.1 wt% butyl benzyl phthalate (Santicizer 160, Richard E. Mistler, Inc.), and 1.7 wt% Menhaden fish oil (Richard E. Mistler, Inc.). A 25 ton press (Model 2518, Carver Inc., Wabash, IN) was used to laminate 17 individual green tapes, each approximately 150 µm thick, at 5 MPa for 10 min at either 85 °C or 95 °C into MLCs; the dimensions after lamination were 2.0×1.5×0.25 cm and 2.0×1.5×0.23 cm, respectively.

The porosity and permeability of four laminated tapes were measured after lamination, after SCE, and after the combined SCE/TC. The porosity, $\varepsilon$, was determined using Archimedes’ method. The permeability measurements, which have been described in more detail elsewhere [22,23], were performed by measuring the molar flux, $N_m$, of $N_2$ gas through tapes of disc shape. For a pressure drop from $P_1$ to $P_2$ across a sample of thickness, $L$, the permeability, $\kappa$, can be determined from the flux data with

$$\kappa = -\frac{2RTN_m\mu L}{P_2^2 - P_1^2}$$

(3.1)

where $R$ is the gas constant, $T$ is the temperature, and $\mu$ is the viscosity of the gas.

The adhesion strength between layers was determined by performing peel tests, which have been described in more detail elsewhere [21,24-26]. For the peel test, the two un laminated parts of the T shape were placed in the grips of a tensile tester (TA TX Plus Texture Analyzer, Stable Micro Systems Ltd, Surrey, UK). During testing, the load was measured as a function of displacement as the top crosshead moved at a speed of
0.4 cm s\(^{-1}\). When the extension due to strain is negligible, the adhesion strength, \(\sigma\), is given by

\[
\sigma = \frac{F_i}{b}
\]  

(3.2)

where \(F_i\) is the applied force and \(b\) is the sample width. The adhesion strength is reported as the average of three samples.

The supercritical extraction experiments were conducted in a SS 316 (6.25 cm diameter, 500 ml) Parr high pressure non-stirred vessel equipped with a thermocouple well located in the vessel cover. The vessel was placed in a controlled-temperature furnace which maintained the temperature within ±2 °C. Extraction experiments were conducted in a semi-continuous mode of operation using CO\(_2\) as the supercritical fluid. The SCE cycle was comprised of a purge step, a pressurization step, a dwelling step, and a depressurization step. The purge consisted of a three minute exposure to flowing CO\(_2\) at room temperature and pressure to remove air from the extraction vessel. The vessel was next pressurized with carbon dioxide from 0.1 MPa to 10 MPa with a compressor over 3-4 minutes at room temperature. The vessel was then thermally pressurized to 30±1 MPa as the temperature was increased to 90 °C. The dwell lasted for 1 h as pressure and temperature were held constant. The vessel was then depressurized isothermally over a 6 h period to atmospheric conditions. After depressurization, the extraction efficiency was determined from the weight loss of the MLCs, normalized by the total amount of organic phase initially present.

Thermal oxidation of the binder in the MLCs was conducted in air in a box furnace. The TC consisted of a ramp at either 7.5 or 10 K minute\(^{-1}\) to 600 °C, a 1 h soak
at 600 °C, followed by cooling to 40 °C. Figure 3.1 shows the segments of the SCE cycle (~8 h total duration), the TC (~6 h total duration), and also the combined SCE/TC (~14 h total duration).

**Figure 3.1:** The supercritical extraction cycle and the thermal cycle used for determining yield and cycle time of MLCs. For comparison, part of a typical cycle from industry is also shown.
3.2 RESULTS AND DISCUSSION

For the first experiment, an MLC, laminated at 85 °C and 5 MPa for 10 min, was subjected to a TC alone with a heating rate of 7.5 K minute⁻¹. As seen in the images in Fig. 3.2a, the top surface in the central portion of the substrate experienced a large defect during the TC; no damage was evident on the sides of the sample. Another MLC, laminated at the same conditions, was then subjected first to supercritical extraction; the sample lost 36 weight% during this step and displayed no defects afterwards. This sample was then subjected to the TC portion of the cycle, where the binder was completely removed, e.g., the total weight loss was 100%. Figure 3.2b shows that after the combined SCE/TC, no defects were evident in the sample.

Figure 3.2: Side and top images of MLCs laminated at 85 °C, 5 MPa for 10 minutes.
a) Sample after a TC conducted with a 7.5 K minute⁻¹ ramp to 600 °C with a 1 h hold.
b) Sample after a combined SCE/TC after exposure first to supercritical CO₂ at 90 °C, 29 MPa for 1 h followed by a 6 h depressurization. The MLC was then exposed to a thermal cycle identical to (a).
In the second experiment, the above procedure, for the same lamination conditions, was repeated, but this time during the TC the samples were heated at a higher rate of 10 K minute\(^{-1}\). Figure 3.3 shows that for the TC, the MLC once again experienced defects in the top surface and also large delaminations on the edge of the sample. For the MLC subjected to the combined SCE/TC, the top surface exhibited no defects but the edge now showed two small regions of delamination. Thus, an increase in the heating rate during the TC has lead to defects in samples subjected to both processes, although the severity is much worse for the TC alone. This suggests that the evolution of pressure, and hence stress, during the TC exceeded the strength of the green body in both cases.

![Image](image.jpg)

**Figure 3.3:** Side and top images of MLCs laminated at 85 °C, 5 MPa for 10 minutes.

a) Sample after a TC conducted with a 10 K minute\(^{-1}\) ramp to 600 °C with a 1 h hold.

b) Sample after a combined SCE/TC after exposure first to supercritical CO\(_2\) at 90 °C, 29 MPa for 1 h followed by a 6 h depressurization. The MLC was then exposed to a thermal cycle identical to (a). The arrows indicate delaminations.
Additional experiments were conducted, but now the lamination temperature was increased from 85 °C to 95 °C. Figure 3.4a shows that for the TC alone, in which the heating rate was 7.5 K minute⁻¹, defects occur in the green body; these defects, however, are not located on the surface but instead are restricted to the edges of the sample in the form of large regions of delamination. Figure 3.4b shows that after the combined SCE/TC, no defects were evident in the sample.

![Figure 3.4: Side and top mages of MLCs laminated at 95 °C, 5 MPa for 10 minutes.](image)

a) Sample after a TC with a 7.5 K minute⁻¹ ramp to 600 °C with a 1 h hold.
b) Sample after a combined SCE/TC after exposure first to supercritical CO₂ at 90 °C, 29 MPa for 1 h followed by a 6 h depressurization. The MLC was then exposed to a thermal cycle identical to (a).

In the final set of experiments, MLCs laminated at 95 °C were subjected to a TC alone, but now at the higher heating rate of 10 K minute⁻¹. For this heating rate, Fig. 3.5a
shows that the sample subjected to the TC alone exhibited defects on both the top and edges. The MLC subjected to the SCE/TC, however, exhibited no defects after the combined cycle, although defects were observed at this heating rate for a sample laminated at 85 °C.

Figure 3.5: Side and top images of MLCs laminated at 95 °C, 5 MPa for 10 minutes.

a) Sample after a TC with a 10 K minute\(^{-1}\) ramp to 600 °C with a 1 h hold.

b) Sample after a combined SCE/TC after exposure first to supercritical CO\(_2\) at 90 °C, 29 MPa for 1 h followed by a 6 h depressurization. The MLC was then exposed to a thermal cycle identical to (a).

The MLC yield relationships presented above between a TC alone versus a combined SCE/TC as a function of TC heating rate and lamination temperature are summarized in Table 3.1. All samples subjected to the TC alone, regardless of the lamination temperature and heating rate in the TC, experienced defects, although the
severity was worse for the higher heating rate. For samples subjected to the combined SCE/TC, the yield of defect-free parts was higher in three of the four sets of conditions as compared to the TC alone. An alternative way to view the yield results is that for a given type of heating cycle (TC versus SCE/TC) the propensity or severity of defect formation increased in three of the four cases as the heating rate was increased.

**Table 3.1.** Summary of defect formation in MLCs after exposure to a thermal cycle (TC) alone or a combined supercritical extraction cycle/thermal cycle (SCE/TC) versus lamination temperature and heating rate of the TC.

<table>
<thead>
<tr>
<th>Lamination Temperature (°C)</th>
<th>Heating Rate (K minute⁻¹)</th>
<th>TC Failure (Yes/No)</th>
<th>SCE/TC Failure (Yes/No)</th>
</tr>
</thead>
<tbody>
<tr>
<td>85</td>
<td>7.5</td>
<td>Yes*</td>
<td>No</td>
</tr>
<tr>
<td>85</td>
<td>10</td>
<td>Yes</td>
<td>Yes*</td>
</tr>
<tr>
<td>95</td>
<td>7.5</td>
<td>Yes*</td>
<td>No</td>
</tr>
<tr>
<td>95</td>
<td>10</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

* Sample experienced only slight damage

To clarify the trends in Table 3.1, the sample porosity, gas permeability, and adhesion strength were measured at different points in the process, and these data, along with sample weight loss, are summarized in Table 2. For lamination at 5 MPa for 10 min at 85 or 95 °C, each substrate experienced less than 0.5% weight loss, and the permeability of the laminated green tapes was too low to be measured. The fractional porosity decreased slightly from 0.19 to 0.18 as the lamination temperature was increased from 85 °C to 95 °C whereas the adhesion strength between layers increased from 48.5 to 59.5 N m⁻¹. This latter type of behavior for the effect of lamination on the adhesion
strength is consistent with previous work [21,24] whereby at higher lamination temperature, the viscosity of the sample, via the binder, decreases, and thus the interpenetration or flow of the tapes during lamination is enhanced, thereby increasing the adhesion strength.

Table 3.2. Porosity, permeability, adhesion strength, and weight loss for ceramic tapes as 4-layer substrates after lamination, after SCE, and after a combined SCE/TC.

<table>
<thead>
<tr>
<th>Lam. Temp. (°C)</th>
<th>After Lamination</th>
<th>After SCE</th>
<th>After SCE/TC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ε (-)</td>
<td>κ (m²)</td>
<td>σ (N/m)</td>
</tr>
<tr>
<td>85</td>
<td>0.19</td>
<td>~ 0</td>
<td>48.4</td>
</tr>
<tr>
<td>95</td>
<td>0.18</td>
<td>~ 0</td>
<td>59.5</td>
</tr>
</tbody>
</table>

* indicates samples were too fragile to measure

Following lamination, samples were subjected to SCE in carbon dioxide at conditions of 90 °C at 29 MPa for 1 h. For the substrate laminated at 85 °C, SCE resulted in a 36.5% weight loss of the organic content and an increase in the fractional porosity from 0.19 to 0.32. Supercritical extraction, likely via the weight loss and concomitant change in porosity, lead to an increase in the permeability, which was now measurable, and a decrease in the adhesion strength from 48.5 to 33.3 N m⁻¹. Similar trends in the porosity, permeability, and adhesion strength following SCE were observed for the samples laminated at 95 °C.

After the TC in air, the substrates experienced ~100% weight loss of the organic fraction, which corresponds to a porosity value of 0.48 for substrates laminated at 85 °C and 0.49 for substrates laminated at 95 °C. The complete loss of binder during the TC
rendered the samples too fragile for handling and testing; this behavior likely reflects a further decrease in the adhesion strength and masks an increase in the gas permeability.

To summarize this section, the combined SCE/TC leads to rapid binder removal cycles with high yield as compared to cycles practiced in industry for samples of similar dimensions, compositions, and physical properties (See Fig. 3.1). This likely arises because of the partial removal of binder during SCE, which increases the porosity and permeability of the samples, which in turn facilitates gas flow and thus mitigates the buildup of pressure and hence stress within the green body.
3.3 CONCLUSIONS

In this work, we have demonstrated that a combined supercritical extraction and thermal cycle can be used to remove binder from green MLCs, and this process affects both cycle time and yield. Supercritical extraction of a fraction of the binder results in an increase in the gas permeability and a decrease in the adhesion strength of the MLCs. Following the SCE segment, the MLCs then survive a rapid TC that they did not survive without prior SCE, which indicates that the trade-off in the increase in gas permeability following SCE has a more pronounced effect than the reduction in adhesion strength. This rapid cycle is much shorter than typical cycles of 50-150 h used in industry for samples of similar size and physical properties. The yield of the MLCs was also seen to depend on other upstream processing variables such as the conditions of lamination, whereby increasing the lamination temperature leads to a large increase in the adhesion strength, which persists even following partial binder removal via SCE.
3.4 REFERENCES


CHAPTER 4

EFFECTS OF A COMBINED SUPERCritical EXTRAction/Thermal CYCLE ON RESidual CARBON IN MULTILAYER CERAMIC CAPACITORS
4.0 INTRODUCTION

In the fabrication of multilayer ceramic capacitors (MLCs), organic additives such as a high molecular weight binder and a low molecular weight plasticizer are often used to impart strength to the ceramic green body. Before the MLCs can be sintered, however, all of the organic species must be removed from the green body. Traditionally, the organic fraction is removed by thermal degradation methods [1-3], but as the organic polymer becomes vaporized, oxidized, or pyrolyzed into carbonaceous gases, a carbon residue may remain inside the green body. These residues may either influence the sintering behavior or degrade the electrical properties of the final components. To avoid carbon contamination during processing of MLCs with base metal electrodes, the organic fraction is typically removed in non-oxidative atmospheres at temperatures above 270 °C in order to circumvent oxidation of the base-metal electrode material [4-8].

An alternative processing route to thermal degradation of the organic fraction is supercritical extraction [9-19]. Studies have shown that supercritical carbon dioxide can easily remove organic species of lower molecular weight such as short chain waxes [10-12] or plasticizers [10-12,15,16], but becomes less effective as the chain length increases. Supercritical hexane has been known to remove higher molecular weight organic species like polystyrene (MW = 150,000) [20]; but its application towards binder removal in MLCs is not well known in literature. For the case of MLCs, the preferential extraction of the low molecular weight plasticizers with supercritical carbon dioxide may lead to 40-60% removal of the organic fraction [13,14], depending on the components in the binder blend. By extracting a large fraction of the organic material, which does not lead to
carbonaceous products, less residual carbon may be left in the dielectric after the thermal binder removal step. In this work, we demonstrate the impact of a combined supercritical extraction and thermal cycle (SCE/TC) on the residual carbon present in base-metal and noble-metal electrode MLCs.
4.1 EXPERIMENTAL

Two tape compositions, identified by their binder type, were evaluated in this study. The first tape composition consisted of 87.7 wt% barium titanate powder (AD 342N, Ferro, Niagara Falls, NY), 8.8 wt% poly(vinyl butyral) (PVB) resin (BMS, Sekisui, Troy, MI), and 3.5 wt% dioctyl phthalate (Sigma-Aldrich, St. Louis, MO). The dried tapes were ~22 µm thick. To prepare the nickel metal electrode pastes, nickel powder at 75.8 wt% (JFE Ni Powder, JFE Mineral Co., Japan), 15.1 wt% barium titanate (BT 01, Sakai Chemical Industry Co., Japan), 4.6 wt% Witco Emphos (Witco Corp., Greenwich, CT), and 4.6 wt% ethyl cellulose (N-4, Ashland Aqualon, Wilmington, DE) were combined in a solvent and then screen printed onto the tapes. Next, 30 tapes were laminated in a press at ~21 MPa at 65 °C for 20 minutes, and the resulting dimensions of the MLCs were 2.54×2.54×0.067 cm. Samples were also prepared without the Ni electrode material.

The second tape composition consisted of 89.1 wt% of a titanate-based N2200 dielectric, 9.9 wt% acrylic-based resin (B72, Rohm & Haas, ON, Canada), and 2 wt% polyester adipate plasticizer (G50, C. P. Hall, Bedford Park, IL). The individual green tapes were laminated at 85 °C at 29 MPa for 10 min into MLCs that had 41 active layers and Pt/Pd/Au electrodes. The MLCs had dimensions of 2.2×2.0×1.0 cm following lamination.

For these samples, single or multiple SCE cycles were used, but individual cycles also varied with the type of supercritical fluid. When operating with carbon dioxide as the supercritical fluid, the SCE cycle, denoted as SCE_{CO_2}, was performed at conditions of
90 °C and 30 MPa for three 1-h exposures. Following SCE\textsubscript{CO\textsubscript{2}}, the TC varied with the type of MLC. For the PVB-based MLCs, the TC consisted of a ramp in air at 1 K minute\textsuperscript{-1} to 270 °C followed by a 12 h dwell period and then cooling to 40 °C. For the acrylic-based MLCs, the TC cycle consisted of a ramp in air at 2.0 K minute\textsuperscript{-1} to 800 °C, a 12 h soak at 800 °C, followed by cooling to 40 °C. Figure 4.1 shows the segments of the SCE\textsubscript{CO\textsubscript{2}}, the TC, and also the combined SCE\textsubscript{CO\textsubscript{2}}/TC for both the PVB-based and acrylic-based MLCs.

![Figure 4.1](image)

**Figure 4.1:** The extraction cycle with supercritical carbon dioxide, and the thermal cycles used for determining residual carbon content of MLCs.

When operating with hexane as the supercritical fluid, the vessel was first purged with argon gas for 5 minutes to completely remove oxygen. Next, the temperature of the
vessel was increased from room temperature to 270 °C, which pressurized the vessel from 0.1 MPa to 26 MPa as the phase of the hexane changed from liquid to supercritical. The SCE cycle, denoted as SCE$_{C_6H_{14}}$, was comprised of two 2 h cycles during which the pressure was held constant and the temperature was maintained within ±0.5 °C. After the vessel was thermally depressurized over a period of 12 hours to 0.1 MPa and 30 °C, the MLC was removed and dried in an oven for 24 hours at 80 °C. After SCE, the oxidation cycle consisted of a ramp in air at 1 K minute$^{-1}$ to 270 °C followed by a 12 h dwell period and then cooling to 40 °C.

The carbon content was measured with a residual carbon analyzer (Model C-144, LECO, St. Joseph, MI) by pulverizing the MLCs into a powder yielding an average particle diameter of approximately 1 mm. For each type of sample, the average percent residual carbon (%RC) and 90% confidence intervals were calculated from 5 samples at each condition. To determine the precision and accuracy of the RC measurements, a 1% carbon standard was evaluated prior to analyzing the samples.
4.2 RESULTS AND DISCUSSION

Table 4.1 summarizes the average percent weight loss for MLC samples subjected to various binder removal cycles. For the PVB-based MLC samples with and without nickel metal electrodes, both types of samples experienced 29-30% weight loss after being subjected to supercritical carbon dioxide at 90 °C, 30 MPa, for three 1 h cycles. For the acrylic-based MLCs, however, only a 3% weight loss occurred after extraction at the same conditions, which is consistent with the lower solubility of the acrylic/adipate binder [21]. The weight loss of the PVB-based MLC samples was further increased to 93% after exposure to supercritical hexane at 270 °C, 26 MPa for two 2 h cycles. After the following cycles (a) SCE\textsubscript{CO}_2/TC, (b) SCE\textsubscript{CO}_2/SCE\textsubscript{CH}_14/TC, and (c) TC alone, each sample lost up to 97-100% of the organic fraction, irrespective of sample type or prior processing conditions. The results above suggest that weight loss alone determined gravimetrically is insufficient to conclude which process is superior for binder removal.

To differentiate between the different processing routes, residual carbon analysis was conducted. Figure 4.2 shows the %RC for the PVB-based MLC samples with and without nickel electrodes. For both types of samples, the combined SCE\textsubscript{CO}_2/TC leads to 25-30% less carbon as compared to the TC alone. In addition, the use of a combined cycle leads to substantially less variability in the residual carbon content, as indicated by the 90% confidence intervals. Because of the overlap of confidence intervals in Figure 4.2, the sample means were compared for equality using a 90% confidence interval, one-sided test. This test indicates that for the MLC samples subjected to the SCE\textsubscript{CO}_2/TC and TC process, the differences in the means are statistically significant, which indicates the
SCE\textsubscript{CO\textsubscript{2}}/TC process is effective in lowering the \%RC as compared to the TC alone. For the MLC samples with nickel metal electrodes subjected to the SCE\textsubscript{CO\textsubscript{2}}/SCE\textsubscript{C\textsubscript{6}H\textsubscript{14}}/TC process, the mean \%RC was only reduced by 0.60\% as compared to the SCE\textsubscript{CO\textsubscript{2}}/TC results. However, the MLC samples without nickel metal electrodes showed an 8.65\% reduction in the mean \%RC and contained no overlap between the 90\% confidence intervals which means the SCE\textsubscript{CO\textsubscript{2}}/SCE\textsubscript{C\textsubscript{6}H\textsubscript{14}}/TC process was statistically a slight improvement over the SCE\textsubscript{CO\textsubscript{2}}/TC.

**Table 4.1.** Average percent weight loss for PVB-based and acrylic-based MLCs after exposure to different supercritical extraction and/or thermal cycles. The maximum operating temperature during the TC for the PVB-based and acrylic-based MLCs was 270 °C and 800 °C, respectively.

<table>
<thead>
<tr>
<th>Cycle Type</th>
<th>PVB-based With Ni Weight Loss (%)</th>
<th>PVB-based Without Ni Weight Loss (%)</th>
<th>Acrylic-based With Pt/Pd/Au Weight Loss (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCE\textsuperscript{*}</td>
<td>29</td>
<td>30</td>
<td>3.3</td>
</tr>
<tr>
<td>SCE\textsuperscript{*}/SCE\textsuperscript{++}</td>
<td>93</td>
<td>93</td>
<td>N/A</td>
</tr>
<tr>
<td>SCE\textsuperscript{*}/TC</td>
<td>97</td>
<td>~100</td>
<td>99</td>
</tr>
<tr>
<td>SCE\textsuperscript{++}/SCE\textsuperscript{++}/TC</td>
<td>98</td>
<td>98</td>
<td>N/A</td>
</tr>
<tr>
<td>TC</td>
<td>97</td>
<td>99</td>
<td>99</td>
</tr>
</tbody>
</table>

\textsuperscript{*} CO\textsubscript{2} was used as the supercritical fluid at conditions of 90 °C and 30 MPa. \textsuperscript{++} Hexane was used as the supercritical fluid at conditions of 270 °C and 26 MPa.

The origin of the lower \%RC for SCE\textsubscript{CO\textsubscript{2}}/TC as compared to the TC alone may be attributed to the fact that during the SCE\textsubscript{CO\textsubscript{2}} process, the binder, namely the low
molecular weight plasticizer component, is removed by dissolution into the supercritical carbon dioxide and then diffusion out of the body and not by thermal degradation. Thus, removing one-fourth to one-third of the binder by this process leads to less organic material being present which can subsequently crack to carbonaceous residue during the TC. Alternatively, the removal of the plasticizer, with the resultant creation of porosity, may allow for more facile oxidation of the remaining organic fraction during the TC without concomitant carbon deposition.

![Bar chart](image)

**Figure 4.2:** Percent residual carbon with 90% confidence intervals for MLC samples with and without nickel electrodes subjected to $\text{SCE}_{\text{CO}_2}/\text{TC}$ ($\text{SCE}/\text{TC}$); $\text{SCE}_{\text{CO}_2}/\text{SCE}_{\text{C}_6\text{H}_{14}}/\text{TC}$ ($\text{SCE}/\text{SCE}/\text{TC}$); and TC (TC) alone. Each subscript denotes the type of supercritical fluid used.
The logic mentioned above suggested that the addition of a SCE$_{C_6H_{14}}$ cycle to remove the high molecular weight binder component prior to the TC would further reduce the %RC in the MLC samples as compared to the SCE$_{CO_2}$/TC process. Though the supercritical hexane greatly increased the quantity of organic material removed, a less proportional effect was seen on the %RC remaining in the MLC samples following the TC. These results may be attributed to some residual hexane possibly remaining in the MLC samples following the TC.

Figure 4.3 shows the %RC for the acrylic-based MLC samples containing Pt/Pd/Au metal electrodes. For these samples, the %RC for both the TC and SCE$_{CO_2}$/TC is within 0.001% of each other. The 90% confidence intervals shown in Figure 4.3 indicate that a TC alone in air to 800 °C had less variability in the residual carbon content, as compared to the SCE$_{CO_2}$/TC process, which is inconsistent with the aforementioned PVB-based MLC samples. Because the mean %RC of the acrylic-based MLCs was so close to the sensitivity of the instrument, it may be considered inappropriate to unambiguously conclude the validity of these results. For this reason, no attempts were made to further reduce the %RC in the acrylic-based MLC samples with the addition of a SCE$_{C_6H_{14}}$ cycle.
Figure 4.3: Percent residual carbon with 90% confidence intervals for an acrylic-based MLC containing Pt/Pd/Au electrodes subjected to SCE\textsubscript{CO\textsubscript{2}}\textsubscript{2}/TC (SCE/TC) and TC (TC) alone. The subscript denotes the type of supercritical fluid used.
4.3 CONCLUSIONS

In this work, we have demonstrated that a combined supercritical extraction and thermal cycle has an effect on the residual carbon content in MLCs. For the PVB-based MLC samples, utilization of the SCE\textsubscript{CO\textsubscript{2}}/TC leads to 25-30\% less carbon as compared to a TC alone and to much less sample-to-sample variation. This reduction in carbon content and variation was observed in MLC samples prepared both with and without nickel electrodes. Exposing the PVB-based MLC samples to supercritical hexane effectively removed up to 93\% of the organic fraction, regardless of whether nickel was present. However, this cycle showed essentially no further reduction in the mean %RC for the samples with nickel and an 8.65\% reduction for the samples without nickel as compared to the SCE\textsubscript{CO\textsubscript{2}}/TC results. For the acrylic-based MLC samples, the %RC is nearly identical between the TC and SCE\textsubscript{CO\textsubscript{2}}/TC, and in fact was very close to the sensitivity of the instrument. This nearly complete removal of the organic fraction is attributed to a more complete oxidation of the MLC at higher temperature.
4.4 REFERENCES


CHAPTER 5

EFFECTS OF A COMBINED SUPERCRITICAL EXTRACTION AND THERMAL DECOMPOSITION OF BINDER ON DEFECT FORMATION IN GREEN CERAMIC BODIES
5.0 INTRODUCTION

In the fabrication of multilayer ceramic capacitors (MLCs), organic blends of binder and plasticizer are used to impart strength to the ceramic green body. However, all of the organic species must be removed from the green body before the MLCs can be sintered. Traditionally, thermal degradation methods are used for removal of the organic fraction [1-3], but as the binder loading of the MLC increases, the amount of time allocated to removing the organic fraction also increases to prevent defects from forming within the green body. Extraction of the organic material with supercritical fluids has been proposed and demonstrated as an alternative processing route [4-16].

In earlier work, defects such as cracks and delamination have been occasionally observed in MLCs subjected to supercritical extraction [15]. Such defects were mostly observed in MLCs that experienced little loss of binder during the extraction cycle which, in turn, resulted in a low permeability to gas flow. It was then postulated that the low gas permeability caused pressure gradients to form during depressurization from supercritical conditions which, in turn, led to the formation of defects. Defect formation in samples exhibiting low weight loss suggested the exploration of more aggressive extraction conditions by operating at higher temperatures. Under these conditions, degradation of the organic fraction may be contributing to the observed weight loss, and possibly to the occurrence of defects in the MLCs.

In this work, the origin of defects in samples is further explored for MLCs fabricated from a (a) poly(propylene carbonate)-based, (b) acrylic-based, and (c) poly(vinyl butyral)-based binder system. Three potential mechanisms contributing to the
formation of defects are additionally considered: failure during depressurization, failure from enhanced binder degradation due to the presence of residual air in the vessel, and failure due to mechanical vibrations from gas compression when going to high pressure.
5.1 EXPERIMENTAL

Three tape compositions, identified by their binder type, were evaluated in this work. The acrylic-based MLC samples nominally consisted of 89.1 wt% of a titanate-based N2200 dielectric, 9.9 wt% acrylic-based resin (B72, Rohm & Haas, ON, Canada), and 2 wt% polyester adipate plasticizer (G50, C. P. Hall, Bedford Park, IL). The individual green tapes were laminated at 29 MPa at 85 °C for 10 min into MLC samples that had 41 active layers and Pt/Pd/Au electrodes. The dimensions of the MLC after lamination were 2.2×2.0×1.0 cm. The acrylic-based MLCs were also cut using a vice and a razor blade (PO# 11140, ACE Hardware Corp., Oak Brook, IL) into 1/2, 1/4, and 1/8 fractions of the original thickness. After aligning the razor blade in parallel with the metal electrodes inside the MLC as depicted in Figure 5.1, the blade was pushed through the sample as the grips of the vice closed. The 1/8 MLC sample was then quartered perpendicular to the metal electrodes, producing a MLC that was 1/32 of the original size.

![Figure 5.1: Schematic for cutting acrylic-based multilayer ceramic capacitors.](image-url)
The poly(vinyl butyral) (PVB) based MLC samples nominally consisted of 87.2 wt% BaTiO$_3$ (X7R 422H, Ferro, Niagara Falls, NY), 6.5 wt% poly(vinyl butyral) resin, (PVB BL-1, Sekisui, Troy, MI) and 5.3 wt% dioctyl phthalate (DOP). The individual green tapes were laminated at 7 MPa at 85 °C for 10 min into a MLC sample which contained 45 active layers. The dimensions of the MLC sample after lamination were 2.1×1.5×0.17 cm.

The poly(propylene carbonate) (PPC) based tape nominally consisted of 86.6 wt% BaTiO$_3$ (X7R 422H, Ferro, Niagara Falls, NY), 9.4 wt% polypropylene carbonate resin, (PPC Q40-01, Empower Materials, Newark, DE), and 2.5 wt% DOP. The individual green tapes were too brittle for lamination into MLC samples. The size of each individual tape was ~2.1×1.5×0.013 cm.

Experiments were conducted with carbon dioxide in a SS 316 (6.25 cm diameter, 500 ml) Parr high pressure non-stirred vessel. Four cycles, described in Table 5.1, were evaluated which differ in the final pressure and whether air, and especially oxygen, was removed from the vessel with a purge cycle. Individual tapes were exposed to either CO$_2$ at low pressure (0.1 MPa) or to supercritical CO$_2$ at high pressure (30±1 MPa). For the high pressure cycle with a purge step, carbon dioxide was added to the vessel at an initial temperature of 22 °C until the pressure reached an initial value of 7-10 MPa. The final pressure of 30 MPa was attained by heating the vessel to the dwell temperature, causing the carbon dioxide to thermally pressurize. For the high pressure cycle with no purge step, the vessel containing air was pressurized with carbon dioxide at the dwell temperature. At the end of each cycle, the amount of organic removal was determined.
from the weight loss of the samples, normalized by the total amount of organic content initially present.

Table 5.1 Details of the four cycles evaluated.

<table>
<thead>
<tr>
<th>Purge (O₂ removed)</th>
<th>Low Pressure Cycle (0.1 MPa)</th>
<th>High Pressure Cycle (30 MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) 3 min CO₂ purge at 22 °C</td>
<td>a) 3 min CO₂ purge at 22 °C</td>
<td></td>
</tr>
<tr>
<td>b) Sample exposed to flowing CO₂ during heating ramp</td>
<td>b) Pressurization with CO₂ to 7-10 MPa at 22 °C followed by thermal pressurization during the heating ramp to 30 MPa</td>
<td></td>
</tr>
<tr>
<td>c) Dwell 1 h in flowing CO₂ at high temp.</td>
<td>c) Dwell 1 h at high temp.</td>
<td></td>
</tr>
<tr>
<td>d) Remove sample at dwell temp. in air</td>
<td>d) Depressurize at dwell temp. in 30±5 min, then remove sample in air</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>No Purge (O₂ present)</th>
<th>Low Pressure Cycle (0.1 MPa)</th>
<th>High Pressure Cycle (30 MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) No purge at 22 °C</td>
<td>a) No purge at 22 °C</td>
<td></td>
</tr>
<tr>
<td>b) Sample exposed to stagnant air during heating ramp</td>
<td>b) Sample exposed to stagnant air during heating ramp</td>
<td></td>
</tr>
<tr>
<td>c) Dwell 1 h in flowing CO₂ at high temp.</td>
<td>c) Pressurization with CO₂ to 30 MPa at dwell temp.</td>
<td></td>
</tr>
<tr>
<td>d) Remove sample at dwell temp. in air</td>
<td>d) Dwell 1 h at high temp.</td>
<td></td>
</tr>
<tr>
<td>e) Depressurize at dwell temp. in 30±5 min, then remove sample in air</td>
<td>e) Depressurize at dwell temp. in air</td>
<td></td>
</tr>
</tbody>
</table>
5.2 RESULTS AND DISCUSSION

Figure 5.2 shows the percentage of binder removed from the PPC-based samples after being subjected to each of the four cycles listed in Table 5.1. For each cycle, the degree of binder removal generally increased with increasing temperature. At low temperature, each sample experienced a higher percentage of weight loss when operating at 30 MPa instead of 0.1 MPa, regardless of whether a purge was used. When operating with a purge cycle, the maximum amount of binder removed was 74 wt% and occurred at 150 °C and 0.1 MPa. The weight loss of the samples may be attributed to an equilibrium that exists between propylene oxide (PO), carbon dioxide (CO₂), and PPC according to the reversible chemical reaction:

\[ \text{PPC} \rightleftharpoons \text{CO}_2 + \text{PO} \]  \hspace{1cm} (5.1)

From the data given in Figure 5.2, it is possible that the presence of carbon dioxide at high temperatures and low pressures may cause the chemical reaction equilibrium to shift to the right, which in turn leads to more weight loss. For these samples, no defects were observed at any temperature.

When operating with no purge cycle, the presence of residual air showed a strong effect on the degree of binder removal at temperatures and pressures ranging from 120-150 °C and 0.1-30 MPa. When operating with no purge at 150 °C, regardless of pressure, the organic content of the samples was removed almost completely. This high degree of binder removal may be attributed to the oxidative degradation of the PPC binder. For the high pressure cycle, defect formation in the samples was observed at each temperature,
but for the low pressure cycle defects were only seen at 150 °C. These defects may be contributed to the fragile nature of the green body as a result of oxidative degradation.

**Figure 5.2:** Weight loss of PPC-based tapes subjected to different operating conditions. The asterisk (*) denotes that the sample exhibited defects.

The amount of binder removed from the PVB-based samples using each of the four cycles listed in Table 5.1 is shown in Figure 5.3. For these tapes, defect formation was never introduced into the green body under any set of conditions. Also, no significant advantage in the percent weight loss was evident between operating with or without a purge cycle. When operating at 90 °C, the degree of binder removal varied from 5 wt% to 41 wt% depending on whether a pressure of 0.1 MPa or 30 MPa was used, respectively. The high percentage of weight loss while operating at 30 MPa can be attributed to the extraction of the low molecular weight DOP constituent, which is known in the literature to be highly soluble in supercritical carbon dioxide [8-10,15]. For the
cycles conducted at 120-150 °C, the percent weight loss of each sample remained within 39-44%, regardless of the type of cycle used.

![Figure 5.3](image)

**Figure 5.3:** Weight loss of PVB-based tapes subjected to different operating conditions.

Figure 5.4 shows the percentage of binder removed from the acrylic-based samples after being subjected to each of the four cycles listed in Table 5.1. As compared to the results in Figures 5.2 and 5.3, the total amount of organic material extracted from each sample was substantially less. For this binder system, a higher percentage of weight loss was observed at each condition while operating at 30 MPa as opposed to 0.1 MPa. The maximum amount of binder removed was 8.6 wt% and occurred while operating at 90 °C and 30 MPa with a purge cycle. The low percentage of weight loss is consistent with the lower solubility of the acrylic/adipate binder in supercritical carbon dioxide [15]. Defect formation was observed in two of the acrylic-based samples while operating
without a purge step at a pressure of 30 MPa and at temperatures ranging from 120-150 °C. These defects may be due to mechanical vibrations of the sample which arise during rapid depressurization.

![Graph showing weight loss comparison for acrylic-based tapes subjected to different operating conditions](image)

**Figure 5.4:** Weight loss comparison for acrylic-based tapes subjected to different operating conditions. The asterisk (•) denotes that the sample exhibited defects.

The results in Figures 5.2-5.4 display some complex relationships between the conditions of extraction, the type of cycle used, the weight loss, and the formation of defects. The complexity may not provide conclusive evidence that a single mechanism underlies failure for all three binder systems, but some generalizations, however, are possible. First, the strong effect of thermal degradation on the strength of the green body may suggest that defect formation is more likely to occur at high temperature. This type
of defect was evident in the PPC-based and acrylic-based samples when operating with no purge, i.e., in the presence of oxygen.

Based on the results contained in Figures 5.2-5.4, additional experiments were conducted at the high pressure operating conditions under which defects were introduced into the green ceramic tapes. For these experiments, the vessel was depressurized from 30 MPa to 0.1 MPa over a period of 90-120 minutes instead of 30 minutes. Table 5.2 shows that the formation of defects in green ceramic tapes was avoided by using longer depressurization times.

Table 5.2. Effect of depressurization time on the formation of defects.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Pressure (MPa)</th>
<th>Depress Time (h)</th>
<th>Weight Loss (%)</th>
<th>Defects (yes/no)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPC-Based Single Tapes</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>90</td>
<td>30</td>
<td>0.5</td>
<td>24.7</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.5</td>
<td>24.4</td>
<td>no</td>
</tr>
<tr>
<td>120</td>
<td>30</td>
<td>0.5</td>
<td>65.0*</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>52.5</td>
<td>no</td>
</tr>
<tr>
<td>150</td>
<td>30</td>
<td>0.5</td>
<td>267.6*</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>97.5</td>
<td>no</td>
</tr>
<tr>
<td>Acrylic-Based Single Tapes</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>120</td>
<td>30</td>
<td>0.5</td>
<td>5.8</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>4.4</td>
<td>no</td>
</tr>
<tr>
<td>150</td>
<td>30</td>
<td>0.5</td>
<td>36.0</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>6.1</td>
<td>no</td>
</tr>
</tbody>
</table>

* Not all of the sample could be collected.
The results in Figures 5.2-5.4 and Table 5.2 indicate or suggest some best modes of operation to avoid defect formation within ceramic green bodies while extracting binder, i.e., when operating at high pressure. First, a purge cycle should be employed, which may avoid oxidative degradation of the sample. Also, longer depressurization times should be used, which may minimize both pressure gradients within the sample and the effects of vibration accompanied with depressurization. In addition, thermal expansion of the CO$_2$ during the ramp may be a preferable mode of operation to achieve high pressure, as compared to pressurizing the vessel at the dwell temperature.

The best modes of operation described above were then applied to a previous study, see Figure 5.5, which indicated that defects were always exhibited in acrylic-based MLC samples, regardless of the extraction conditions used. Table 5.3 lists all of the extraction conditions evaluated in the study as well as the appearance of the MLC after depressurization. In general, the occurrence of defects in the acrylic-based samples usually appeared as cracks or delamination located near the mid-plane of the MLC. The occurrences of these defects in the acrylic-based MLC were credited to the insufficient quantity of binder removed, which led to the formation of pressure gradients during depressurization from supercritical conditions, and then to failure of the green body. Because a much larger quantity of organic material was removed from the PVB-based MLC, defects were observed much less frequently.

Because the acrylic-based and PVB-based MLC samples depicted in Figure 5.5 differ in composition, porosity, permeability, and size, it is not straightforward to unambiguously determine the origin of defect formation. To assess the effect of size, the
acrylic-based MLC samples were cut into smaller pieces and then exposed to a cycle which was nearly consistent with the high pressure cycle containing a purge step listed in Table 5.1, except the vessel was depressurized over 7.5 hours instead of 30 minutes. Figure 5.6a shows images of an acrylic-based MLC sample before and after being subjected to supercritical carbon dioxide at 90 °C and 30 MPa for 1 hour; dimensions of the MLC sample were equivalent to 1/4 of the original size. After SCE, which removed 4.5 wt% of the organic content, a large defect along the edge of the sample was exhibited. This defect may be attributed to the formation of pressure gradients which arise during depressurization from supercritical conditions.

Figure 5.5: a) Image of an acrylic-based sample (2.2×2.0×1.0 cm) after depressurization over 19 h after SCE in CO₂ at 30 MPa at 90°C for 1 h [which resulted in defects]. b) Image of a PVB-based sample (2.1×1.5×0.17 cm) after depressurization over 6 h after SCE in CO₂ at 40 MPa at 90°C for 1 h [which exhibited no defects] [15].
Table 5.3. Summary of extraction results from acrylic-based MLC samples at various operating conditions.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Pressure (MPa)</th>
<th>Appearance</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>40</td>
<td>multiple cracks/delamination</td>
</tr>
<tr>
<td>55</td>
<td>10</td>
<td>vertical crack</td>
</tr>
<tr>
<td>90</td>
<td>30</td>
<td>multiple cracks/delamination</td>
</tr>
<tr>
<td>90</td>
<td>10</td>
<td>Delamination/other crack</td>
</tr>
</tbody>
</table>

In the next experiment, the above procedure was then repeated for the same type of sample, but this time, the MLC was cut to a dimension of 1/32 of the original size. Figure 5.6b shows images of the MLC before and after exposure to supercritical carbon dioxide. After SCE, which removed 4.8 wt% of the organic fraction, a small delamination was evident along the edge of the sample. Thus, even the smaller sized acrylic-based MLC samples experience failure upon depressurization from supercritical conditions. These results suggest that the formation of defects in the acrylic-based MLCs cannot be avoided using the best mode of operation. Furthermore, the enhanced porosity of the green body prior to depressurization from supercritical conditions remains the key factor for avoidance of defect formation in MLCs, as described previously [15].

While evaluating the effect of MLC size on defect formation, a noticeable change in weight loss became evident. Table 5.4 shows that the percent weight loss of the MLC tends to increase as the size of the sample decreases. The higher percentage of weight loss may be attributed to the increased surface area to volume ratio of the smaller sized MLCs which, in turn, may have led to shorter diffusion paths. Furthermore, by reducing
the MLC size, less organic material is initially present upon exposure to supercritical carbon dioxide.

**Figure 5.6:** Images of two different sized acrylic-based MLC samples before and after subjection to supercritical carbon dioxide at 90 °C and 30 MPa for 1 hour. The dimensions of each sample were (a) 2.2×2.0×0.24 cm and (b) 1.12×0.97×0.096 cm.

**Table 5.4.** Weight loss as a function of the size of acrylic-based MLCs after being subjected to supercritical CO₂ at conditions of 90 °C and 30 MPa for 1 h followed by a 7.5 h depressurization.

<table>
<thead>
<tr>
<th>Fractional Size of MLC</th>
<th>Weight Loss (%)</th>
<th>Damaged (yes/no)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.0</td>
<td>yes</td>
</tr>
<tr>
<td>1/4</td>
<td>4.5</td>
<td>yes</td>
</tr>
<tr>
<td>1/32</td>
<td>4.8</td>
<td>yes</td>
</tr>
</tbody>
</table>
5.3 CONCLUSIONS

In the presence of oxygen, more defects occur in green tapes, which were especially pronounced for the PPC-based and acrylic-based samples. For an enhanced extraction and/or minimization of defects in green structures, the best mode of operation employs (a) a purge cycle to remove and so as to avoid oxidization, (b) thermal expansion of the CO$_2$ during the ramp, and (c) longer depressurization times. The acrylic-based MLCs always exhibited defects following SCE, regardless of the samples size or operating conditions. This suggest that the formation of defects in the acrylic-based MLCs cannot be avoided using the best mode of operation. Furthermore, the occurrences of defects may be attributed to the lack of porosity within the green body prior to depressurization supercritical conditions.
5.4 REFERENCES


CHAPTER 6

CONCLUSIONS AND FUTURE WORK
6.0 CONCLUSIONS

A combined supercritical extraction and thermal cycle has been developed to remove binder from green MLCs. This process has shown a positive effect on both cycle time and yield. A partial removal of the binder content in MLCs with supercritical carbon dioxide lead to an increase in the gas permeability and a decrease in the adhesion strength of the green body. Because the MLCs exposed to SCE could survive the subsequent rapid TC, the increase in the gas permeability had a more pronounced effect than did the reduction in adhesion strength. Furthermore, the upstream processing conditions showed a direct impact on MLC yield. For MLCs of comparable size and physical properties, a typical cycle used in industry would have lasted 50-150 hours, which is much longer than the rapid cycle of ~14 hours developed in this work.

The residual carbon content in MLCs was affected by removing the organic constituents with a combined SCE$_{CO_2}$/TC instead of a TC alone. For the PVB-based MLC samples with and without nickel metal electrodes, the combined cycle led to a 25-30% reduction in the residual carbon content and to much less sample-to-sample variation, as compared to a TC alone. Though a 93% weight loss was obtained after exposing the PVB-based MLC samples to supercritical hexane, the mean %RC was only reduced an additional 8.65% for the MLCs without nickel electrodes and 0.65% for the MLCs with nickel electrodes, as compared to the SCE$_{CO_2}$/TC results. For the acrylic-based MLC samples, the mean %RC was nearly identical for the TC and SCE$_{CO_2}$/TC processes, and in fact was very close to the sensitivity of the instrument. This was
attributed to nearly complete oxidation, as the thermal cycle for these samples was conducted in air.

The defects formed in the PPC-based and acrylic-based ceramic tapes occurred much more frequently while removing the organic content in the presence of oxygen. A best mode of operation was identified to enhance the quantity of organic material extracted and to minimize the occurrence of defect formation in green ceramic structures. This best mode employed the use of (a) a purge cycle to avoid oxidization, (b) thermal expansion of the carbon dioxide during the ramp, and (c) longer depressurization times to reduce mechanical vibrations. The acrylic-based MLCs always exhibited defects following SCE, regardless of the operating conditions. Also, the percent weight loss of the acrylic-based MLCs had a tendency to increase with decreasing sample size.
6.1 FUTURE WORK

The successful development of a combined supercritical extraction and thermal cycle has indicated that the increased porosity within a MLC sample following SCE was the major factor contributing to the survival of samples during a rapid thermal cycle. This may suggest that as the porosity within a ceramic green body is increased, the rate of thermal oxidization can also be increased. Thus, a valid follow-up study would be to assess the development of a short thermal cycle where the rate of oxidation is initially slow and then becomes more rapid as the porosity within the green body of the MLC increases.

The effects of a combined supercritical extraction and thermal cycle resulted in a 25-30% reduction in the residual carbon content. This was attributed to a 32% weight loss of the organic fraction after exposure to supercritical carbon dioxide at 90 °C, 30 MPa for three 1 hour cycles. After removing 93 wt% of the organic fraction with supercritical hexane, however, little to no change was exhibited in the %RC. Though these results were attributed to residual hexane possibly remaining in the sample, the exact mechanism controlling the %RC left in samples following a thermal cycle was never identified and needs to be further explored.

Finally, the formation of defects in the acrylic-based MLCs was always evident after exposure to supercritical carbon dioxide. The defects within the MLCs were attributed to the insignificant quantity of organic material removed prior to depressurization from supercritical conditions. Therefore, it may be beneficial to further
explore the solubility of the acrylic/adipate binder system in different types of supercritical fluids or at other supercritical conditions.