MICROWAVE COMBUSTION
AND SINTERING WITHOUT
ISOSTATIC PRESSURE

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EXECUTIVE SUMMARY

This subtask focused on the development of an innovative technology to extract, deposit, and vitrify radioactive species in high-level waste. Uranium hexachloride and uranium (IV) borohydride are volatile compounds for which a chemical vapor infiltration procedure might be developed for the D&D of high-level waste. A hybrid heating technique will be adopted for firing materials, which possess low dielectric constants and loss factors, in order to convert hazardous waste into a dense vitrified form.

During FY98, conventional vitrification tests on surrogate materials of waste sludge were conducted. The non-radioactive sludge surrogate was composed of calcium carbonate, magnesium oxide, silica, alumina, and ferric oxide, which were mixed with amorphous silica and borax, the glass forming components. Tests were conducted to optimize the vitrification of this mixture in a high temperature furnace. Several runs were conducted to evaluate critical process parameters. A heating ramp rate of 8°C/minute and a peak temperature duration of one hour were adopted. The heated mixtures were then cooled in air. The vitrified product adhered to the crucible, and great weight loss and volume contraction were observed. The results of these tests indicated a vitrification temperature for the surrogate mixture in the range of 1150°-1200°C. This temperature was independent of the particle size of amorphous silica.

Based on the chemical similarity of titanium dioxide and uranium dioxide, and their applicability to chemical vapor infiltration, titanium dioxide has been selected as the best surrogate material for uranium dioxide in simulating laboratory experimental work. Titanium dioxide reacts with chlorine gas in the presence of a reducing agent, such as carbon black, at 900°C to produce titanium tetrachloride and carbon dioxide (Stage I). The volatile titanium tetrachloride reacts with oxygen gas to produce chlorine gas and titanium dioxide (Stage II). The residual chlorine gas in the effluent flow is scrubbed by the sodium hydroxide solution.

Experimental runs on the conventional chemical vapor infiltration process have been conducted using Hafnium tetrachloride as another surrogate for uranium. Borosilicate glass fiber was utilized as the preform matrix for the deposition of the surrogate species prior to vitrification. Leachability tests were conducted with the vitrified products in simulated ground water and sea water, respectively. The amount of metal that leaches into the solution will be determined in the future.

Vitrification of the deposited oxide was conducted in a high temperature box furnace at 750°, 800°, and 850°C for one hour. The vitrified product appeared to be completely glass-like for the 800° and 850°C runs when titanium dioxide was deposited on the borosilicate glass fiber matrix. At a temperature of 750°, hafnium dioxide did not precipitate on the borosilicate matrix.

One gram of surrogate mixture was utilized for each experimental run. The quantity of volatile product was assessed from the weight loss of the starting surrogate material. A maximum weight loss (mainly titanium tetrachloride and carbon dioxide) of 89 percent was achieved.
1.0 INTRODUCTION

In recent years, interest has grown rapidly in extending the application of microwave energy to the processing of ceramics, composites, polymers, and other materials. Advances in the understanding of microwave/materials interactions will facilitate the production of new ceramic materials with superior mechanical properties. One application of particular interest is the use of microwave energy for the mobilization of uranium for subsequent redeposition.

This investigation was conducted in three phases. Phase I (FY96) dealt with the determination of physical and chemical properties of fired pre-form matrices. Phase II (FY97) consisted of fundamental research on the conventional sintering and microwave sintering of various mixes of ceramics and dielectric constant determination. Phase III (FY98) focused on the microwave-assisted chemical vapor infiltration tests for mobilization and redeposition of radioactive species in the mixed sludge waste.

Uranium hexachloride and uranium (IV) borohydride are volatile compounds for which the chemical vapor infiltration procedure might be developed for the separation of uranium. Microwave heating, characterized by an inverse temperature profile within a preformed ceramic matrix, is utilized for CVI using a carrier gas. Matrix deposition is expected to commence from the inside of the sample where the highest temperature is present. The preform matrix materials which include aluminosilicate-based ceramics and silicon carbide-based ceramics are all amenable to extreme volume reduction, densification, and vitrification. Important parameters utilized in microwave sintering, such as frequency, power requirement, soaking temperature, and holding time, would be investigated in order to optimize process conditions for the volatilization of uranyl species using a reactive carrier gas in a microwave chamber.

1.1 BACKGROUND OF THE PROBLEM

There is growing evidence to support the use of microwave energy in certain industrial processes and continued research and development for its use in many other materials-processing applications, such as radioactive waste disposal. In contrast to conventional furnaces, the material to be processed in a microwave furnace interacts with the “cold” microwaves instead of radiant heat.

In the application of mixed waste disposal, conventional heating is inferior to microwave heating. Conventional joule heating requires electrodes in direct contact with the mixed waste. Electrode corrosion can limit the service life of these smelters. In microwave treatment, microwave energy is absorbed by a large class of waste materials commonly found in many low-level waste streams, such as fabrics, rubber, concrete, metal powders, oxides, nitrides, sulfates, nonmetallic filtering media, water, carbon, and glass. Because microwave energy is absorbed directly, it has the considerable advantage of much higher efficiency and faster temperature control compared to conventional radiant heating. Microwave power can be transmitted through wave-guides from generators that can be located safely outside a radioactive processing area where routine system maintenance can be easily performed. Microwave heating is extremely flexible in that a wide range of processing temperatures are available in a single system. Microwave disposal systems can be designed to be small enough to be mobilized for on-site treatment of radioactive wastes. Heating profiles can be custom-tailored for each application by
exciting the appropriate microwave field configuration or “mode,” choosing the proper heating
frequency and material geometry, and understanding how the microwave penetration depth is
controlled by temperature and material composition of the waste form.

Microwave-assisted chemical vapor infiltration (MCVI), an innovative technique to fabricate
fiber-reinforced ceramic composites, has been proposed as a technique for treating radioactive
waste. Uranium hexachloride and uranium (IV) borohydride are volatile compounds for which
the chemical vapor infiltration procedure might be developed for the separation of uranium. The
radioactive species in mixed waste, after mixing with a reactant gas, produces volatile
compounds in a microwave cavity. These compounds are transported by a carrier/reactant gas to
a react tube, where the MCVI will take place. The volatilized uranyl species that is deposited
in the ceramic matrix will be vitrified in-situ.

1.2 PURPOSE OF THIS INVESTIGATION

This project is devoted to the development of an innovative technique for the disposal of mixed
waste utilizing microwave energy. Because most uranium and plutonium components as well as
most fission products have dielectric properties that allow excellent microwave and high-
frequency energy coupling, dielectric heating has the potential for application in many processes
for treating hazardous wastes. This technology, whether used on its own or as hybrid in
conjunction with a conventional process, has positive features, such as energy efficiency,
increased throughput, volume reduction, and reduction of disposal and transportation cost, and
provides a technique not feasible by conventional means. The hazardous waste will be converted
into a dense, stable, and vitrified form so that it may qualify for eventual off-site disposal. If
successful, this program will lead to major cost saving for the DOE system.

1.3 LITERATURE REVIEW

Chemical vapor infiltration (CVI) is an attractive technique for introducing matrix materials into
fibrous reinforcements as it minimizes stress during processing. Also, when compared to hot
pressing, relatively low temperatures are used that limit unwanted chemical attack on the
reinforcement. Difficulties associated with CVI processing using conventional heating are
preferential deposition in the substrate's outer regions leading to pore blockage, long processing
time, non-uniform composite density, high residual porosity, and limitations on substrate
geometry.

The use of microwave energy in the CVI processing has received recent attention due to the
volumetric heating and the resultant inverse temperature profile. With the internal region of the
substrate hot, cool reactant gases penetrate inwardly prior to the onset of the deposition reaction.
It should be noted that the inverse temperature gradients could also be established by “seeding”
the center (or one side) of the preform with a material that readily absorbs microwave energy.
The following advantages of MCVI over the conventional CVI have been well demonstrated by
several researchers: relatively short processing time, no limitations on the preform geometry,
products with spatially uniform and high density, deposition from the inside-out.
MCVI has been documented in the fabrication of Al₂O₃ fiber/Al₂O₃ matrix composites (Skamser et al. 1994) and the SiC fiber/SiC matrix composites (Deepak and Evans 1993; Day et al. 1994). Infiltration of Si₃N₄ in Nicalon™ cloth (Devlin et al. 1993), SiC and ZrO₂ into reticulated SiC foam, Nicalon™ cloth and ZrO₂ based porous fiber boards (Yin et al. 1997) were also experimentally demonstrated. Experiments conducted in these reports required the application of microwave radiation at a fixed frequency.
2.0 PROJECT DESCRIPTION

This project involves the following phases of investigations of the utilization of microwave energy: 1) sintering of aluminosilicate materials, 2) sintering of matrix materials for chemical vapor infiltration of radioactive materials, 3) investigation of microwave-assisted chemical vapor infiltration techniques.

2.1 OVERALL PROJECT GOALS

Based on the results of investigations on the microwave-assisted chemical vapor infiltration tests for the surrogates of waste, an innovative technique involving microwave energy may be developed for decontamination and decommissioning of radioactive waste.

In FY97, fundamental research was conducted on the microwave sintering of various aluminosilicate materials. The mineralogical development of these materials during microwave firing was evaluated, as were the changes of the dielectric constant with the temperature (up to 200°C). The process flowchart for the chemical vapor infiltration experiments was also outlined.

The overall project goals for FY98 were the following:

- To demonstrate the conventional chemical vapor infiltration technology with surrogates of radioactive species. A conventional furnace will be modified to meet this objective.
- To develop the microwave-assisted chemical vapor infiltration technology for the volatilization, mobilization, and deposition of the surrogate species in ceramic matrices.
- To apply the microwave-assisted chemical vapor infiltration technology for the decontamination and decommissioning of the high-level waste.

It is expected that the results of this investigation will be useful to the U.S. Department of Energy (DOE) for application in a microwave-assisted chemical vapor infiltration process. This technology can be used for the in-situ remediation of hazardous waste or as a hybrid microwave technology.

2.2 FY97 ACTIVITIES

During FY97, extrudates of various compositions of bauxite materials were fired in a conventional high temperature box furnace under air atmosphere and in a variable frequency microwave furnace. The weight loss, volume contraction, and true specific gravity, as well as the mineralogical composition, were determined. The dielectric constants of fired extrudates were also determined with a Hewlett Packard Network Analyzer.
FY97 activities included the following:

- Determination of the composition of the starting materials by chemical analysis.
- Firing the four types of bauxite extrudates—raw bauxite, beneficiated bauxite, tailing pond material, and capping material—under air atmosphere in a conventional high-temperature furnace at 1350°, 1450°, 1550°, and 1600°C for 4 hours and at 1450°, 1550°, 1650°, and 1700° for 1 hour.
- Firing the four types of bauxite extrudates in a 1.2KW variable frequency microwave furnace (VFMF) at a frequency of 5.5-5.75Ghz with a sweeping rate of 3 seconds at 1100°C for 1 minute and at 1350° and 1550°C for 30 minutes.
- Determination of the mineralogical composition of the fired products by powder X-ray diffraction (XRD).
- In-situ analysis of the changes of the dielectric constant of the four types of materials with the temperature up to 200°C.
- Determination of the dielectric constant of the extrudates that were fired at 500°, 800°, 1100°, 1350°, and 1600°C.
- Attending the First World Congress on Microwave Processing in Orlando, Florida, on January 5-9, 1997, and presentation of the project in the 32nd Microwave Power symposium on July 14-16, 1997, in Ottawa, Canada.
- Preparation of technical papers for presentation at the following conferences: the 32nd Microwave Power Symposium (July 14-16, 1997 in Ottawa, Canada) and the Second International Congress on Metallurgy and Materials Technology (October 12-17, 1997, in Sao Paulo, Brazil). In addition, a refereed journal paper derived from this project has been accepted by the Journal of Microwave & Electromagnetic Energy.
- A literature search to determine 1) a suitable surrogate for mixed waste, 2) a processing flowchart for the chemical vapor infiltration (CVI) and microwave-assisted chemical vapor infiltration (MCVI) processes, 3) instrumentation and supplies for conducting preliminary CVI experiments, 4) suitable carrier gas/reaction gas mixtures, and 4) types of ceramic matrix preform.

2.3 FY98 OBJECTIVES

During FY98, investigations focused on the following:

- **CONVENTIONAL VITRIFICATION OF THE MIXED WASTE SURROGATE:** Preliminary vitrification tests will be conducted on the conventional high-temperature furnace. The volume reduction, heating temperature, time required for vitrification, and the thermal and chemical stability of the products will be evaluated. This information is essential for accessing the efficiency of the CVI and MCVI processes.

- **CONVENTIONAL CHEMICAL VAPOR INFILTRATION FOR WASTE DISPOSAL:** CVI experiments will be conducted using a conventional furnace. The experimental setup to demonstrate the CVI process for treatment of mixed waste will be accomplished at Florida...
International University’s Hemispheric Center for Environmental Technology (FIU-HCET). A surrogate of the radioactive species in the mixed waste will be volatilized and mobilized by a carrier gas and then deposited in a ceramic matrix. The deposited products will be vitrified in the conventional high-temperature furnace.

- **Microwave-Assisted Chemical Vapor Infiltration for Waste Disposal:** Microwave energy will be utilized in the CVI processing to extract and deposit the radioactive species in the mixed waste. The MCVI system will consist of a microwave source and cavity, a gas delivery system, a quartz reactor tube with exhaust plumbing and vacuum connection, pressure control, and a scrubbing unit. The reacting/carrier gas 1) will react with the radioactive species present in the mixed waste in the volatilization chamber to produce volatile compounds. Reacting/carrier gas 2) will react with the volatile compounds, and infiltration deposition will proceed within the preform ceramic matrix located inside the microwave cavity. The products will be vitrified in-situ.
3.0 PROCESS DESCRIPTION

Although it is still in an early stage of development, the use of microwave energy to process a wide variety of ceramic materials offers many new and exciting opportunities. There are several reasons for the growing interest in microwave processing versus conventional processing methods. These include the potential for significant reductions in manufacturing costs due to energy savings and shorter processing times, improved product uniformity and yields, improved or unique microstructures and properties, and the synthesis of new materials. In the case of MCVI, this would overcome the main problems encountered in conventional CVI where preferential deposition in the substrate’s outer regions leads to pore blockage, long processing time, non-uniform composite density, high residual porosity, and limitations on substrate geometry.

3.1 FUNDAMENTAL CONCEPTS AND THEORIES

Based upon the literature search conducted, a flow chart has been developed for a microwave-assisted chemical vapor infiltration (MCVI) process for the treatment and decontamination of mixed radioactive waste. As shown in the following charts, the MCVI system will consist of a microwave source and cavity, a gas delivery system, a quartz reactor tube with exhaust plumbing and vacuum connection, pressure control, and a scrubbing unit. The reacting/carrier gas 1 will react with the radioactive species present in the mixed waste in the volatilization chamber to produce volatile compounds. Reacting/carrier gas 2 will react with the volatile compounds and infiltration deposition will proceed within the preform located inside of the microwave cavity.
A schematic diagram of the MCVI reactor is shown in Figure 2. It will consist of a cylindrical microwave cavity through which a quartz reactor tube is inserted. A ceramic matrix preform will be loaded co-axially within the quartz tube. Since the heating configuration produces primarily radial temperature gradients, deposition is expected to commence at the preform axis and grow outward.
3.2 SAFETY AND REGULATORY CONCERNS

The U.S. Department of Energy (DOE) has numerous regulatory requirements that must be followed by any DOE operation or contractor working at a DOE site in the United States. Since the Microwave Project involves radioactive wastes, the following regulations apply:

- **DOE Order 5400.5, Radiation Protection of the Public and the Environment.** This order constitutes a list of standards and requirements that DOE and its contractors must abide by to operate the facility while protecting the public, work crews, and the environment against unnecessary risks from radiation.

- **DOE Order 1332.1, Uniform Reporting System.** This order applies to any operation related to the DOE. It establishes a uniform reporting system for contracts, loans, and loan guarantees and provides instructions, forms, and procedures for effective management.

- The U.S. Environmental Protection Agency (EPA) was established to reduce and control pollution by setting standards and enforcing regulatory policy. The regulatory requirements that must be followed include the Clean Air Act of 1970 (CAA), the Clean Water Act of 1977 (CWA), the Safe Drinking Water Act of 1974 (SDWA), and the Resource Conservation and Recovery Act (RCRA). Other laws that apply to hazardous waste include the Toxic Substances Control Act of 1976 (TSCA) and the Comprehensive Environmental Response, Compensation, and Liability Act (CERDRA), which addresses the cleanup of hazardous waste contamination from previous operations.
Lambda Technologies provided a Microwave Leak Detector (MLD) to monitor any possible microwave leakage. The MLD is interfaced with an interlock circuitry, which provides immediate shutdown of the unit if hazardous levels of microwave energy are detected.

The microwave furnace was used in a heavily ventilated laboratory equipped with HEPA-filtered exhaust fume hoods. This reduces the level of metal vapor in the laboratory as well as that of the exhaust fumes released into the atmosphere.
4.0 EXPERIMENTAL INVESTIGATION

Experimental work completed in FY97 included the firing of four types of bauxite materials under air atmosphere in a conventional high-temperature furnace and in a variable frequency microwave furnace. The weight loss, volume contraction, true specific gravity, as well as the mineralogical composition, of the fired products were determined. The dielectric constants of the materials at various temperatures were also analyzed using a Hewlett-Packard Network Analyzer.

4.1 FIRING TESTS

Bauxite extrudates have been fired at 1450°, 1550°, 1650°, and 1700°C for 1 hour under an air atmosphere. Extrudates were also fired for a duration of 4 hours at 1350°, 1450°, 1550°, and 1600°C in an air atmosphere. Two conventional furnaces were utilized to conduct the firing. Pellets were preheated to about 800°C in the low-temperature furnace and rapidly transferred to the high-temperature furnace. The drop in the temperature was estimated to be less than 20°C during the transfer. The pellets were then heated at a rate of 5°C/min to the expected temperatures in the conventional high-temperature box furnace and held at the required temperature. These pellets were then allowed to cool to room temperature. In order to measure the physical and mineralogical properties of the sintered specimens, 16 batches of firing tests were conducted.

Microwave firing of bauxite extrudates was conducted in a 1.2 Kw variable frequency microwave furnace (VFMF) at a frequency of 5.5 - 5.75 Ghz with a sweeping rate of 3 seconds at Lambda Technologies, Inc. Three firing regimes were adopted with the microwave furnace. The first regime involved heating the extrudates to 1100°C and maintaining that temperature for 1 minute. The other two regimes involved heating the extrudates up to 1350°C and 1550°C and maintaining that temperature for 30 minutes, respectively. These tests revealed that with hybrid heating using silicon carbide as a susceptor only 20 minutes were required to achieve a firing temperature. The fired extrudates were found to be void of visible cracks.

4.2 MEASUREMENT OF PHYSICAL PROPERTIES

The weight loss, volume contraction, and true specific gravity of the fired extrudates were measured. The weight loss was determined by the difference in the sample before and after firing. The dimension of each sample (radius and length) was measured prior to and proceeding the firing. The volume contraction was then estimated. To determine the true specific gravity, the fired samples were ground to a fine powder so that the individual pieces were too small to contain pores. The true specific gravity of the powder was determined by means of a specific gravity bottle (pycnometer).

4.3 ANALYSIS OF MINERALOGICAL COMPOSITION

The mineralogical compositions of the fired extrudates were analyzed using powder X-ray diffractometer with an internal standard method. Commercial micron powders of mullite,
corundum, and cristobalite were used as the reference materials. Calcium fluoride was applied as the internal standard material. The internal standard material was mixed with the powders of the fired extrudates in an agate motor. The powders were covered with 10 milliliters of ethyl ether as the mixing solvent. The slug was stirred with a pestle until the liquid evaporated. Stirring considerably increases the degree of mixing. The mixture was exposed under X-ray, and the relative contents of mullite, corundum, and cristobalite were calculated according to the diffraction peaks.

4.4 MEASUREMENT OF DIELECTRIC CONSTANTS

In-situ measurement of the dielectric constants for various bauxite materials was taken using a Hewlett Packard Network Analyzer along with a dielectric probe. In order to obtain the readings, the four samples (raw bauxite, beneficiated bauxite, tailing pond material, and capping material) were heated up to 200°C, and then the measurements were taken. A thermocouple was used to monitor the temperature of the sample while the readings were taken. Three readings at three distinct temperatures were taken for each of the four samples. This gives a total of twelve curves to compare with each other. In order to avoid rapid cooling of the sample, the samples were wrapped in an insulating foam.

Ex-situ measurement of the dielectric constants for the extrudates fired at various temperatures was also taken. Extrudates were heated up to 500°C, 800°C, 1100°C, 1350°C, and 1600°C, respectively, and held at those temperatures for 1 hour and then cooled down to the ambient temperature. The measurements were taken at ambient temperature, and three readings were taken for each of the four samples. A total of 20 curves were generated for comparison of dielectric properties as a function of firing temperature and frequency.
5.0 EXPERIMENTAL RESULTS

The results of the experiments described above are reported in the following sections.

5.1 PHYSICAL PROPERTIES

The weight loss of the fired extrudates is shown in Figures 3 and 4 and Table 1.

Figure 3. Weight loss upon sintering of bauxite materials under an air atmosphere with the duration of 1 hour.

Figure 4. Weight loss upon sintering of bauxite material for 4 hours in an air atmosphere.
Table 1.
Weight loss of extrudates fired in a variable frequency microwave furnace at 1350°C for 30 minutes

<table>
<thead>
<tr>
<th></th>
<th>Weight loss (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw bauxite</td>
<td>32.2</td>
</tr>
<tr>
<td>Beneficiated bauxite</td>
<td>32.4</td>
</tr>
<tr>
<td>Tailing pond material</td>
<td>30.4</td>
</tr>
<tr>
<td>Capping material</td>
<td>25.8</td>
</tr>
</tbody>
</table>

The volume contractions of the fired extrudates are shown in Figures 5 and 6 and Table 2.

Figure 5. Volume contraction upon sintering of bauxite materials under an air atmosphere with the duration of 1 hour.
Microwave Combustion and Sintering Without Isostatic Pressure

Figure 6. Volume contraction upon sintering of bauxite materials for 4 hours in an air atmosphere.

Table 2.
Volume contraction of extrudates fired in a variable frequency microwave furnace at 1350°C for 30 minutes

<table>
<thead>
<tr>
<th>Material</th>
<th>Volume contraction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw bauxite</td>
<td>39.0</td>
</tr>
<tr>
<td>Beneficiated bauxite</td>
<td>41.1</td>
</tr>
<tr>
<td>Tailing pond material</td>
<td>41.3</td>
</tr>
<tr>
<td>Capping material</td>
<td>32.8</td>
</tr>
</tbody>
</table>

The specific gravity of the fired extrudates is shown in Figures 7 and 8.
Considerable weight loss and volume contraction occurred during the firing of extrudates. Figures 3 and 4 show that the weight loss of beneficiated bauxite (≈32.1%) was greater than that of the capping material (≈25.5%). The weight loss for each type of material was independent of the type of furnace used (microwave or conventional) as well as the processing parameters (temperature and time). Figures 5 and 6 show that the volume contraction generally increased with increasing firing temperatures. At temperatures of 1550°C and above, beneficiated bauxite experienced greater volume contraction (up to 67.3%) than that of capping material (≈58.7%). However, the volume contraction of extrudates fired in the microwave furnace was significantly lower than that of extrudates fired conventionally at the same temperature. It should be noted that
this observation may have been due to a shorter holding time of 30 minutes in the microwave furnace as compared to 1 hour and 4 hours, respectively, in the conventional furnace.

Weight loss in the firing of bauxite materials was mainly attributed to the decomposition of chemically bonded water, whereas volume contraction was primarily due to the mineralogical transformation and densification (Takeshita et al. 1993). A significant increase of volume contraction occurred when the soaking time was prolonged. This indicated that mineralogical transformation resulted in densification of the extrudates.

The specific gravity of fired extrudates ranged between 2.97 and 3.51 g/cm$^3$, as indicated by Figures 7 and 8. The change in the specific gravity is indicative of the relative proportions of the phases developed in the fired products. The specific gravity for the corundum, mullite, cristobalite, and amorphous silica are 4.03, 3.03, 2.32, and 2.2 g/cm$^3$ respectively (Grimshaw 1971). The higher specific gravity of the fired beneficiated bauxite extrudates indicated that a higher proportion of corundum developed during the firing. This was verified by X-ray diffraction analysis.

### 5.2 MINERALOGICAL DEVELOPMENT

It was found that all the fired extrudates contained a small amount of cristobalite. The amounts of the corundum and mullite developed in the fired extrudates varied with the firing temperatures and the soaking time. The results are shown in Figures 9 through 12 and Table 3.

![Figure 9. Corundum developed in the extrudates fired conventionally for 1 hour.](image-url)
Figure 10. Corundum developed in the extrudates fired conventionally for 4 hours.

Figure 11. Mullite developed in the extrudates fired conventionally for 1 hour.
Figure 12. Mullite developed in the extrudates fired conventionally for 4 hours.

Table 3. Mineralogical development in extrudates fired in a variable frequency microwave furnace (%)

<table>
<thead>
<tr>
<th></th>
<th>Corundum</th>
<th>Mullite</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1100°C</td>
<td>1350°C</td>
</tr>
<tr>
<td>Raw bauxite</td>
<td>57.9</td>
<td>49.3</td>
</tr>
<tr>
<td>Beneficiated bauxite</td>
<td>61.7</td>
<td>53.8</td>
</tr>
<tr>
<td>Tailing pond material</td>
<td>59.1</td>
<td>50.8</td>
</tr>
<tr>
<td>Capping material</td>
<td>25.3</td>
<td>12.8</td>
</tr>
</tbody>
</table>

* Three firing regimes were: 1100°C for 1 minute, 1350°C for 30 minutes, and 1550°C for 30 minutes.

X-ray diffraction analysis of the fired products revealed that gibbsite was completely converted to corundum in all extrudates. The predominant crystalline mineral phases in the fired extrudates were corundum and mullite. Only minor amounts of cristobalite were detected in all fired extrudates. This may be due to the fact that most of the silica had been consumed in the formation of mullite or existed as glassy phases that are not detectable by XRD.

As expected, the firing of beneficiated bauxite produced the highest content of corundum. At certain processing conditions, corundum development in capping material could be completely suppressed. For example, the higher content of corundum that was found in extrudates fired at 1100°C resulted from complete dehydration of gibbsite and a slower reaction rate for mullite formation. At 1350°C, the higher rate of mullite formation from corundum and silica resulted in a lower corundum content. However, the highest mullite production occurred in the firing of capping material. Approximately 100% conversion to mullite occurred in the firing of capping
a lower corundum content. However, the highest mullite production occurred in the firing of capping material. Approximately 100% conversion to mullite occurred in the firing of capping material, whereas approximately 20wt% mullite development occurred in the firing of beneficiated bauxite. The amount of mullite developed in microwave firing was comparable to that produced by conventional firing at the same temperature. However, a greater amount of mullite developed with increased soaking time when extrudates were fired conventionally in a high-temperature furnace. The same result is anticipated with a microwave furnace.

5.3 MICROWAVE EFFECT

The results obtained from this investigation have indicated that at 1350°C, the weight loss of extrudates fired conventionally was similar to that obtained in the microwave furnace. However, less volume contraction occurred in extrudates fired in the microwave furnace as compared to conventionally fired extrudates. There was no significant difference in the amount of corundum and mullite that developed in extrudates fired in each type of furnace, providing sufficient soaking time was allowed.

No significant difference in physical properties was detected in extrudates fired at the same temperature of 1350°C for 30 minutes in the microwave furnace, as compared with 1 hour in the conventional furnace. During microwave firing, the forward and reflected microwave power levels were measured using two directional couplers during the microwave firing. The electrical energy requirement was estimated by taking into consideration the efficiency of the microwave tube (travelling wave tube). The processing time and the electrical power requirement for these two batches are compared in Table 4. The electrical power requirement for firing the extrudates in the microwave furnace during the ramp and soak periods was 1.2 and 0.8KW, respectively, while that for conventional furnace was 5.0 and 2.0KW, respectively. It is clear from these results that the use of a microwave furnace would result in a significant saving in energy, as well as reduced processing time.

<table>
<thead>
<tr>
<th>Time (Minutes)</th>
<th>Electrical energy (KWh)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ramp</td>
</tr>
<tr>
<td>Microwave</td>
<td>20</td>
</tr>
<tr>
<td>Conventional</td>
<td>270</td>
</tr>
</tbody>
</table>
5.4 DIELECTRIC CONSTANTS

The dielectric constants of the materials vary with the temperature and were measured in-situ at three temperatures, as shown in Figures 13 through 16.

The dielectric properties of extrudates that were fired at temperatures of 500°C, 800°C, 1100°C, 1350°C, and 1600°C and allowed to cool to ambient conditions before measurement have been determined using an HP Network Analyzer with a sample probe. The results are shown in Figures 17 through 20.

The results of the measurement show that below 200°C, the dielectric constant of each type of material is a function of the temperature and the microwave frequency. The values of the dielectric constants of the materials are around 3. This indicates that these materials do not couple with the microwave field very well; thus, hybrid heating method, using microwave susceptor to assist heating, has to be applied.

Figure 13. Dielectric constant of beneficiated bauxite at different temperatures.
Figure 14. Dielectric constant of capping material at different temperatures.

Figure 15. Dielectric constant of tailing pond material at different temperatures.
Figure 16. Dielectric constant of traditional bauxite at different temperatures.

Figure 17. Dielectric constants of the products fired at 500°C.
Figure 18. Dielectric constants of the products fired at 1100 °C.

Figure 19. Dielectric constants of the products fired at 1350 °C.
Figure 20. Dielectric constants of the products fired at 1600 °C.
6.0 MAJOR ACCOMPLISHMENTS

- A 1.2 KW variable frequency microwave furnace (VFMF), utilizing a frequency of 4.55 - 4.75GHz, has been used to fire alumino-silicate extrudates at 1100°C, 1350°C, and 1550°C at Lambda Technologies, Inc. For the purpose of comparison, extrudates were also fired in a graphite-lined furnace under helium and in a conventional high-temperature furnace under an air atmosphere at a series of temperatures (1350°C, 1450°C, 1550°C, 1600°C, 1650°C, and 1700°C) for soaking periods of 1 hour and 4 hours, respectively (November 1996 – March 1997).

- Quantitative analyses of the mineralogical compositions, as well as the determination of physical properties such as weight loss, porosity, volume contraction, and specific gravity, of the fired products have been accomplished. These results have lead to the conclusion that the firing of ceramic materials in a microwave furnace can be successfully accomplished with a substantial saving in energy and processing time (April 1997 – May 1997).

- The dielectric constant of four types of alumino-silicate materials were measured in-situ at different temperatures ranging from approximately 40°C to 200°C and at various frequencies. Determinations of the dielectric properties of extrudates ex-situ after firing the extrudates to temperatures of 500°C, 800°C, 1100°C, 1350°C and 1600°C and cooling to ambient conditions before measurement have also been accomplished (June 1997 – August 1997).

- Two technical papers have been published and presented at the following conferences: the 32nd Microwave Power Symposium (July 14-16, 1997, in Ottawa, Canada) and the Second International Congress on Metallurgy and Materials Technology (October 12-17, 1997, in Sao Paulo, Brazil). In addition, a refereed journal paper derived from this project has been accepted by the Journal of Microwave & Electromagnetic Energy (June 1997 – October 1997).

- A chemical vapor infiltration experimental system setup shown in Figure 21 has been established. The reaction in stage 1 occurs at 550°C within the quartz tube, which is heated indirectly by the tube furnace. The assembly for the influent gas, which consists of gas regulators, mass flow meters, and an inlet connection to the quartz tube, is shown in Figure 22. The effluent gas connection of the quartz tube to the exhaust gas pump/scrubber unit is shown in Figure 23.
Figure 21. The chemical vapor infiltration system experimental setup.

Figure 22. The assembly of the influent gas stream of the experimental setup.

Figure 23. The assembly of the effluent gas stream for the experimental setup.
The efficiency with which volatilization and deposition of surrogate product (assessed from the weight loss of the starting surrogate material) occurs was established at approximately 89% efficiency. The results of weight loss of surrogate mixtures, which corresponds to the formation of volatile species and carbon dioxide under various conditions, are listed in Table 5.

### Table 5. Results for Stage I of the conventional chemical vapor infiltration tests

<table>
<thead>
<tr>
<th>Runs</th>
<th>Mole ratio (TiO₂:C)</th>
<th>Temperature (°C)</th>
<th>Time (hours)</th>
<th>Weight loss (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1:1</td>
<td>550</td>
<td>4</td>
<td>23</td>
</tr>
<tr>
<td>2</td>
<td>1:1</td>
<td>900</td>
<td>3</td>
<td>57</td>
</tr>
<tr>
<td>3</td>
<td>1:2</td>
<td>900</td>
<td>3</td>
<td>87</td>
</tr>
<tr>
<td>4</td>
<td>1:2</td>
<td>900</td>
<td>2</td>
<td>81</td>
</tr>
<tr>
<td>5</td>
<td>1:3</td>
<td>900</td>
<td>3</td>
<td>89</td>
</tr>
</tbody>
</table>

Determination of the appropriate temperature profile for the CVI process is shown in Figure 24.

**Figure 24. Temperature/time process regime.**
7.0 REFERENCES


