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**PHASE SEPARATION AND CRYSTALLIZATION OF COMPLEX
BOROSILICATE MELTS FOR GLASS-CERAMIC WASTE FORMS**

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Nicholas Roberts, Paul Porter, and Richard K. Brow

Missouri University of Science and Technology

Department of Materials Science & Engineering

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BACKGROUND INFORMATION

This work was principally done by two graduate students in the Materials Science & Engineering Department at Missouri University of Science & Technology. Nicholas Roberts received his Masters' Degree in MSE in December 2018, and this report is based on his thesis for that degree, "Microstructural Development and Its Effect on Aqueous Corrosion of a Borosilicate Glass Ceramic for Waste Vitrification." Paul Porter is a candidate for a PhD in MSE, expected in 2020. His contributions to this study are included in this report. Both students, as well as a number of undergraduate assistants, were supervised by Prof. Richard K. Brow.

The report is divided into four sections. The first provides background information about issues associated with waste vitrification, and the development of the borosilicate glass-ceramic materials. The second section describes the effects of thermal history on the phase transformations that occur in the borosilicate glass-ceramic and includes the results of both isothermal and continuous-cooling experiments. The third section describes the effects of thermal history on the chemical durability of the borosilicate glass-ceramics, and includes the results of product consistency tests performed at Missouri S&T and at Savannah River National Lab. Sections 2 and 3 are written in the form of journal articles. The fourth section is an appendix that includes a description of the hot thermocouple test assembly, as well as data sets used in the first two sections.

ABSTRACT

Waste loadings of reprocessed spent nuclear fuel vitrified into borosilicate glass can be increased by precipitating environmentally stable phases concentrated with waste components in a chemically stable glass matrix. The principal objective of this study was to characterize the development of crystalline powellite (CaMoO_4 and related phases) and oxyapatite ($\text{Ca}_2\text{LN}_8\text{Si}_6\text{O}_{26}$) in borosilicate glass-ceramics and to determine how the formation of those phases affected its chemical durability.

Borosilicate glasses provided by PNNL were re-melted and quenched at rates from over 300°C/s to $\sim 0.05^\circ\text{C/s}$. Isothermal heat treatment experiments were conducted by quenching melts in a molten tin bath at various temperatures, holding for periods of time, and then quenching in a water bath. Analytical electron microscopy and x-ray diffraction provided information about the kinetics of the phase separation and crystallization processes responsible for microstructural development. Powellite and oxyapatite crystals formed during slower quench rates and longer isothermal times, and time-temperature-transformation (TTT) diagrams were developed from the latter experiments. A hot thermocouple test (HTT) system was built to rapidly quench melts and hold them at set times to provide supplementary TTT information, but that system was not as sensitive to the phase transformations as other tests.

Corrosion tests were performed to understand how the individual phases in the glass-ceramic affect its overall chemical durability. Product consistency tests, performed at Missouri S&T and at Savannah River National Lab, provided release rates of major elements from samples as a function of cooling rate, and atomic force microscopy and profilometry measurements of surface topology determined the relative corrosion rates of

the residual glass and oxyapatite phases. Faster dissolution rates were measured from samples cooled more slowly and these were explained by the greater fractions of B₂O₃ in the residual glass phase after the formation of oxyapatite and powellite. Oxyapatite was found to be more durable than the residual glass.

ACKNOWLEDGMENTS

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1. BACKGROUND INFORMATION

1.1. RADIOACTIVE WASTE

With the notable discoveries of x-rays by Wilhelm Conrad Roentgen in 1895, radioactivity in 1896 by Henri Becquerel, radioactive materials by Pierre and Marie Curie in the early 1900's, and the nuclear reactor in 1942 by Enrico Fermi, artificial radioactive materials have been available for over a century [1-3]. Many industries have greatly benefited from the use of radioactive materials including, but not limited to, the power generation, agriculture, and medicine [4]. However, a major challenge in the nuclear field is determining what to do with the long-lived radioactive waste.

Radioactive waste is described by the U.S. Environmental Protection Agency as a hazardous waste under the criteria in Title 40 of the Code of Federal Regulation, part 261, and becomes exempt from this code if the waste is compliant with Title 40 of the Code of Federal Regulation part 266, Subpart N [5]. Under Title 10 of the Code of Federal Regulation Part 961, the United States Department of Energy, DOE, is contracted to manage the proper disposal of nuclear waste [6]. To determine whether radioactive waste forms are compliant and environmentally safe, the DOE developed the Waste Acceptance Product Specifications (WAPS) [7]. The WAPS considers the radionuclide inventory, chemical composition, product consistency, and phase stability of the waste form [7].

The International Atomic Energy Agency, IAEA, has classified radioactive waste into six distinct categories based on safety guidelines, summarized in Table 1.1 [8]. The main source of radioactive wastes comes from the Nuclear Fuel Cycle, NFC, used for power generation and military purposes, which produces low level waste (LLW), intermediate level waste (ILW), and high-level waste (HLW) [4]. Reprocessing techniques

recycle the radioactive waste, greatly reducing the amount of waste, and combine them into single HLW compositions for disposal.

Table 1.1. Radioactive waste types [8].

Waste Name	Characteristics	Disposal
Exempt Waste (EW)	Small concentrations of waste $\leq 10\mu\text{Sv/year}$	No disposal required
Very Short-Lived Waste (VSLW)	Waste with short half-life's ≤ 100 days.	Stored until acceptable
Very Low-Level Waste (VLLW)	Materials used in operating nuclear facilities that are slightly above EW waste concentrations	Landfill type storage
Low Level Waste (LLW)	Restricted waste with harmful levels	Near Surface Disposal
Intermediate Level Waste (ILW)	Higher concentrations than LLW that need more containment from the atmosphere.	Geological Disposal
High Level Waste (HLW)	Waste that generally produces 10^4 TBq/m^3 and generate significant quantities of heat.	Engineered Geological Disposal and Containment Canister

In the United States, programs such as the Fuel Cycle Research and Development, FCRD, program of the Office of Nuclear Energy in the DOE are tasked with developing spent nuclear fuel reprocessing technologies in efforts to greatly reduce the volume of radioactive waste. The fuel reprocessing technologies are intended to combine waste streams into groups based on composition to minimize volume [9]. To develop a single global waste form, long-lived, heat-generating, and volatile radioactive components would need to be further combined with lanthanides, minor actinides, and any remaining undissolved solids [9]. Existing HLWs, at sites like those in Hanford, WA, are generally from open nuclear cycles, unprocessed waste streams, which contain many different

components. Future nuclear fuel cycles are intended to be close-ended systems in which the waste is to be reprocessed many times and separated into groups with similar components allowing waste forms with higher waste loadings and more desirable properties to be developed [9].

The DOE developed the Used Fuel Disposition Campaign (UFDC) to characterize all the Spent Nuclear Fuel (SNF) and HLW, both defense and commercial, in the United States and this is compiled into an on-line waste library (OWL) database [10, 11]. In the US, as of 2012, the UFDC listed 69,500 metric tons of heavy metal (MTHM) commercial SNF, totaling ~23 billion curies of long-lived radioactivity, stored at 75 sites in 33 states [11]. In 2048 there is projected to be 142,000 MTHM of SNF. The study determined that there are 43 different waste types, and 50 waste forms that can be combined into 10 different waste groups, based on disposal needs, to be stored in a geological disposal, summarized in Table 1.2 [11].

Table 1.2. The ten waste groups [11].

Waste Group and Description
1- SNF packaged in purpose-built disposal containers
2- SNF packaged in dual-purpose canisters of existing design
3- Vitrified HLW - all types of canistered HLW glass, existing and projected
4- Other engineered HLW waste forms
5- Metallic and non-oxide spent fuels
6- Sodium-bonded fuels
7- DOE oxide fuels
8- Salt, granular solids, and powders
9- Coated-particle spent fuel
10- Spent Naval fuel

Many HLWs from open or “once through” fuel cycles in the United States have already been studied [11]. In some cases, like the West Valley Demonstration Project

(WVDP) in New York, the waste has been reprocessed into a borosilicate glass [11]. Due to advancements in the fuel cycle process, “once through” HLW waste can be recycled through the fuel cycle again to further produce energy and greatly reduce the amount of waste [12]. Once the products become nonfissionable, they are removed from the process stream using aqueous reprocessing techniques such as TRUEX⁺ or UREX⁺ and stored as secondary HLW. Nonfissionable waste streams from the reprocessing of a baseline “once through” HLW contain alkalis, alkaline earths, lanthanides, and transition metals. An approximation for a final secondary HLW composition containing large amounts of Mo, and Cs is shown in Table 1.3, and is denoted as Collins CLT [13]. The waste composition was determined from light-water reactor (LWR) fuel after a 40 GWe burn-up and a 10-year decay storage.

Table 1.3. Composition of Collins CLT waste [13].

Oxide	Content [mol%]	Oxide	Content [mol%]
MoO ₃	19.719	CdO	0.621
ZrO ₂	17.590	SeO ₂	0.534
Nd ₂ O ₃	11.279	Eu ₂ O ₃	0.354
BaO	10.443	Ag ₂ O	0.353
Cs ₂ O	7.416	SnO ₂	0.339
SrO	6.887	Gd ₂ O ₃	0.322
Ce ₂ O ₃	6.859	Rh ₂ O ₃	0.226
La ₂ O ₃	3.527	Br	0.205
Pr ₂ O ₃	3.187	PdO	0.100
TeO ₂	2.985	Pm ₂ O ₃	0.036
Sm ₂ O ₃	2.240	Sb ₂ O ₃	0.028
Y ₂ O ₃	2.019	In ₂ O ₃	0.007
Rb ₂ O	1.641	Tb ₂ O ₃	0.006
RuO ₂	1.076	Total	100.000

1.2. BOROSILICATE GLASS-CERAMICS FOR WASTE VITRIFICATION

HLW produced by reprocessing SNF is intended to be stored as acceptable waste forms. The WAPS was designed by the DOE to ensure that the waste is acceptable and compliant with US Code of Federal Regulations [7]. The standard waste form defined by the WAPS is vitrified waste in a borosilicate glass. In the United States, the only waste form currently used for HLW immobilization is borosilicate glass. The West Valley Demonstration Project and Savannah River site have both successfully developed borosilicate glass compositions for vitrifying HLW that is compliant with the WAPS [11].

Borosilicate glasses have desirable properties for the immobilization of nuclear waste, including excellent chemical durability, thermal and radiation stability, and mechanical integrity [4]. Also, many waste elements are soluble in borosilicate glasses allowing high levels of waste loadings. However, some elements such as molybdenum, are insoluble at levels >3 mol% [13, 14]. Typical borosilicate glass compositions according to the WAPS, are listed in the Table 1.4 [7, 15], and compared to the PNNL glass used in this study (Table 1.5).

Table 1.4. Oxide additions to develop borosilicate waste glasses [7, 15].

Oxide Additions		WAPS [wt.%]	PNNL [wt.%]
Formers	SiO ₂	33-65	30.99
	B ₂ O ₃	3-20	8.53
Intermediates	Al ₂ O ₃	3-20	4.80
Modifiers	Na ₂ O	4-22	2.73
	Metal Oxides	0-50, ~24	-
	Li ₂ O	-	1.05
	CaO	-	5.12

PNNL developed an alternative strategy to the use of homogeneous waste glass, and that was to use glass-ceramic materials with greater levels of CLT-loadings [15]. An example of one borosilicate glass-ceramic with 47.394 wt.%, Collins CLT is listed in Table 1.5 and this composition was used in this study [15]. Increasing the waste loading by allowing crystalline phases to form adds microstructural complexity which requires an understanding of phase separation and subsequent nucleation and growth of the crystals. Furthermore, the chemical durability of the individual phases in these new materials must be understood to ensure that the waste form meets corrosion requirements. The following sections outline relevant theories and studies from the literature related to this study on microstructural development and its effect on aqueous corrosion of a borosilicate glass-ceramic for waste vitrification.

Table 1.5. Centroid glass-ceramic target composition [15].

Oxide	Content [mol%]	Oxide	Content [mol%]
SiO ₂	47.674	TeO ₂	0.641
B ₂ O ₃	11.329	Sm ₂ O ₃	0.481
CaO	8.445	Y ₂ O ₃	0.433
MoO ₃	4.229	Rb ₂ O	0.353
Al ₂ O ₃	3.789	RuO ₂	0.232
Na ₂ O	4.069	CdO	0.135
ZrO ₂	3.774	SeO ₂	0.116
Li ₂ O	3.247	RhO ₂	0.089
Nd ₂ O ₃	2.427	Eu ₂ O ₃	0.076
BaO	2.241	Ag ₂ O	0.075
Cs ₂ O	1.591	SnO ₂	0.072
SrO	1.476	Gd ₂ O ₃	0.070
Ce ₂ O ₃	1.472	PdO	0.023
La ₂ O ₃	0.757	Total	100.000
Pr ₂ O ₃	0.684	Waste Loading	21.447

1.2.1. Glass Structure. Atomic arrangements affect the properties of a material, so it is important to understand the relationships between composition and structure. The materials under study here are glass-ceramics which possess both crystalline and amorphous phases [17]. Zachariasen described the structure of glass and related crystalline materials [18]. Crystalline materials possess periodic arrangements of unit cells with fixed atomic dimensions whereas glasses have an indefinitely large unit cell and lack atomic periodicity. These different arrangements are schematically shown in Figure 1.1. Zachariasen predicted that the local (polyhedral) arrangements of the atoms in a glass were similar to those of atoms in a crystal, and that the differences between the structure and morphologies of a glass and crystal result from how those polyhedra are arranged on large scales. He recognized that certain crystal structures were associated with ready glass formation when melts were quenched, and that these structures are relatively open [18]:

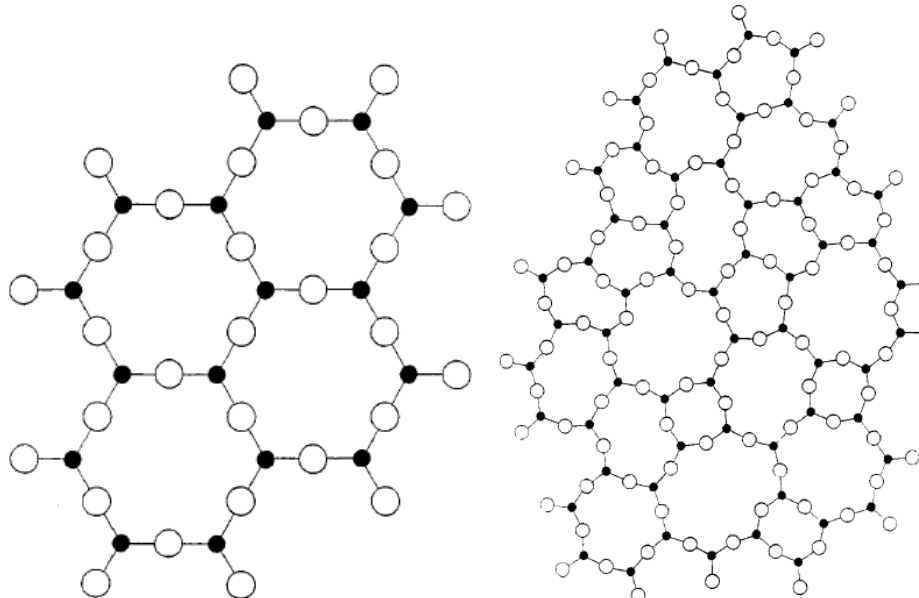


Figure 1.1. Atomic structure representations of (left) crystal and (right) glass [17].

Oxides with these open structures are classified as glass formers. In borosilicate glasses, silica and boron oxide are the glass formers. In silica glass, silicon is coordinated by four oxygen ions which then link to neighboring silica tetrahedra [17]. In vitreous B_2O_3 , boron has a coordination number of 3, forming BO_3 triangular structural units.

Not all oxides are glass formers, and Sun associated the cation-oxygen bond strength with glass forming tendency [19]. Glass forming oxides have the highest cation-oxygen bond strength and are the backbone of the glass structure. Glass modifying oxides have the lowest bond strengths and do not form glasses when quenched from a melt. Instead, these oxides are incorporated into the glass network, modifying the glass properties. Intermediate oxides also do not form a glass structure by themselves, but when combined with other formers and modifiers, they tend to develop strong bonds similar to the glass formers.

Additions of modifier and intermediate oxides to a glass affect properties, generally by increasing or decreasing connectivity between network forming polyhedra. Connectivity relates to properties associated with mass-transport such as viscosity, diffusion, electrical conductivity, and corrosion [17]. The connectivity is increased when greater fractions of glass formers and intermediates are added to a composition and the connectivity in silicate glasses is decreased when modifying oxides are added. Modifying cations are linked to glass network through weak bonds with non-bridging oxygen (NBO), replacing strong bridging oxygens that link network forming polyhedra. In borate glasses, modifying oxides can be incorporated into the glass structure by transforming borate triangles into borate tetrahedra, increasing the fraction of bridging oxygens, with concomitant changes in glass properties. Table 1.6 summarizes the effects of the additions

of intermediates and modifiers on the structures of simple glass systems [17]. It should be noted that the specific effects depend on particular composition ranges.

Table 1.6. Effects of intermediate and modifier oxide additions on glass structure [17].

Glass	Effect
Modifier Additions	
Alkali-Silicate	Creation of 1 NBO per alkali ion reducing connectivity.
Alkali-Alkaline Earth Silicate	Creation of 2 NBO's per alkaline earth ion, which stabilizes alkaline-silicate glasses maintaining connectivity.
Alkali-Borate	Creation of NBOs, reducing connectivity, or the conversion of Boron from 3 to 4 coordination, increasing connectivity.
Alkali-Borosilicate	Creation of NBO's if the alkali associates with Si. Conversion of Boron coordination from 3 to 4 if alkali associates with B.
Intermediate Additions	
Alkali-Aluminosilicate	$\text{Al}_2\text{O}_3/\text{Alkali} < 1$ removes NBO/Al -ion $\text{Al}_2\text{O}_3/\text{Alkali} = 1$ no NBO's $\text{Al}_2\text{O}_3/\text{Alkali} > 1$ addition of 1, 2, or 3 NBO's

The borosilicate centroid composition shown in, Table 1.5, has intermediate (Al_2O_3) and glass network modifier oxides (Na_2O , CaO , and Li_2O) that are added to control the properties of the glass. Some high-level wastes will already contain significant amounts of aluminum and sodium [20, 21]. Aluminum increases the viscosity at high temperatures, increasing homogenization time but reducing volatilization [20, 22]. Aluminum also significantly increases the chemical durability of the glass [22]. Additions of sodium and lithium reduce melting temperatures to enable glass production [20]. Calcium is added to the glass to increase the chemical durability [20].

1.2.2. Glass-Crystal Transformations. A characteristic of a glass forming melt is its resistance to crystallization upon cooling. In fact, all liquids can be vitrified into a glass if the cooling rate of the melt is fast enough [17]. For macroscopic crystallization to occur, crystal nuclei must first form and then, crystals must grow from the nuclei [23]. It is important to understand the nucleation and crystallization pathways to develop glass-ceramics with controlled properties.

Nucleation and crystallization occurs when melts are undercooled below the equilibrium melting temperature, shown in Figure 1.2 from Varshneya [17]. The relative area overlapping the nucleation and crystallization rate curves relates to the ability of an undercooled melt to be quenched without crystallization and so to form a glass. If there is little or no overlap between the nucleation and crystallization rate curves, then the material will easily form a glass. A large overlap, however, means fast crystallization of the nuclei that form, increasing the difficulty of forming a homogeneous glass upon melt quenching.

For glass-ceramics, crystal growth and the onset of crystallization are widely described using time-temperature transformation and constant-cooling transformation diagrams. A time-temperature-transformation (TTT) diagram maps out the temperatures and times where crystals form. Similarly, constant-cooling-transformation (CCT) diagrams also map out crystallization conditions. The difference between the two is that TTT diagrams describe isothermal heat treatments, and CCT diagrams describe constant cooling heat treatments. Crystallization conditions depicted in CCT diagrams are shifted to slower times compared to TTT diagrams and only reproduce the top half of the crystallization curve. Billings and Edwards performed an extensive study developing TTT diagrams for glasses containing projected HLW at the Savannah River site [24]. Figure 1.3 is a TTT

diagram for one of the borosilicate glasses they studied showing the times and temperatures when certain crystals have formed.

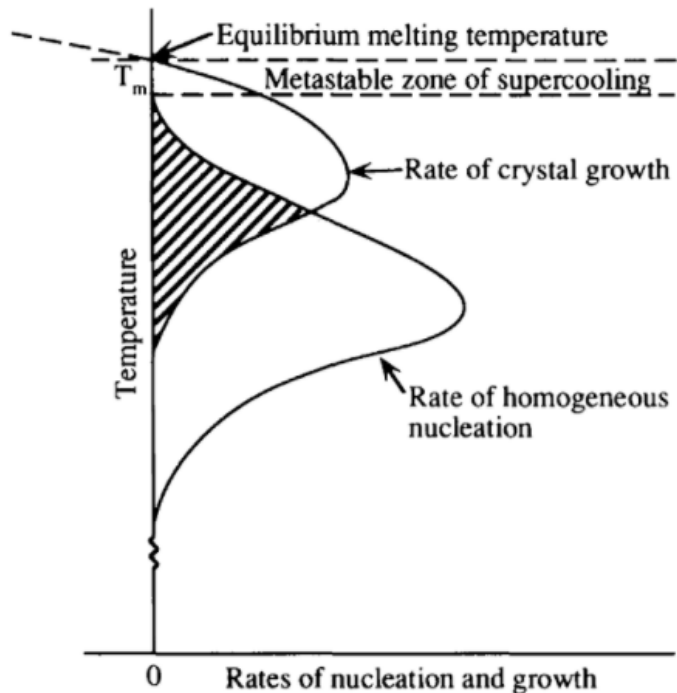


Figure 1.2. Crystal growth rate and nucleation rate as a function of temperature [17].

As stated by the Waste Acceptance Product Specifications (WAPS), TTT diagrams must be produced for waste forms [7]. The TTT diagram is needed to identify the times and temperatures where significant phase changes can occur. Along with the TTT diagram, the glass transition temperature for the waste glass must be known as well. Above the glass transition temperature, there is an increased likelihood for crystallization to occur, altering the properties of the waste form. A goal of the present study is to develop a TTT diagram,

similar to the diagrams produced by Billings and Edwards, for the centroid composition shown in Table 1.5 [25].

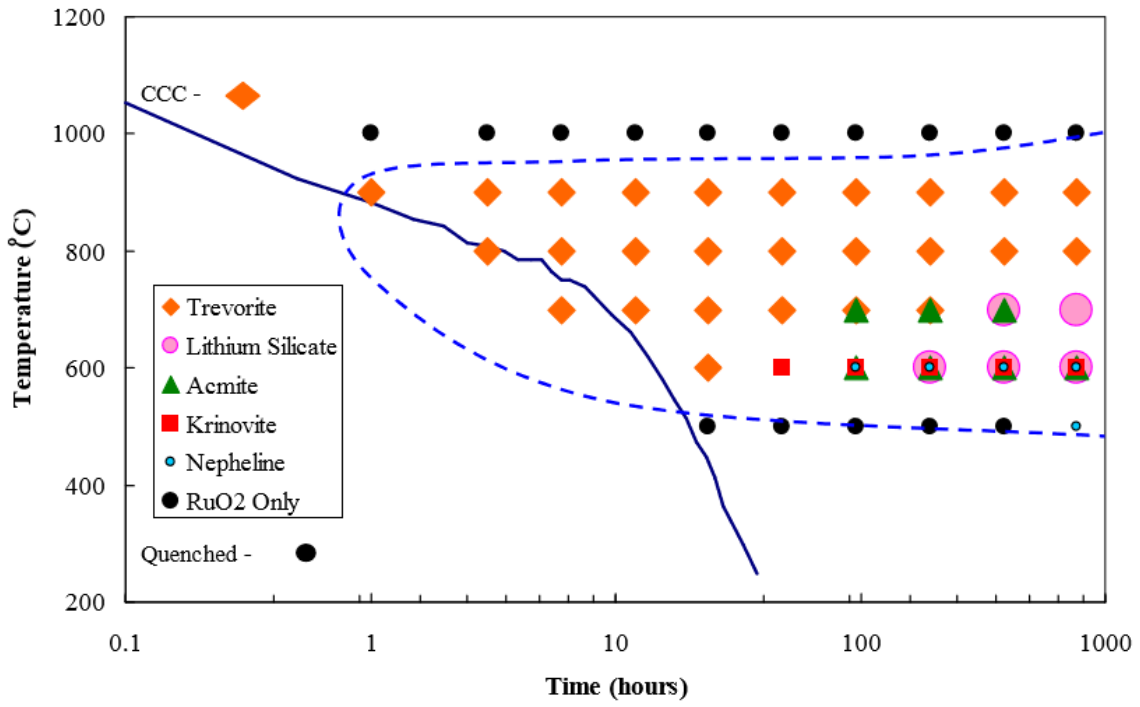


Figure 1.3. TTT diagram for the C2-510 glass studied by Billings and Edwards [24].

1.2.3. Glass-Crystal Transformations Experimental Methods. The temperature dependence of the phase stability of the waste form is important considering the amount of heat released from the radioactive components. The minimum limit for the glass transition temperature according to the WAPS is 400°C [7]. At the glass transition temperature, the structure relaxes, and heat is absorbed i.e., the glass transition is an endothermic process. Commonly used techniques for determining the glass transition temperature are differential scanning calorimetry (DSC) and differential thermal analysis (DTA) [17]. In both tests, the samples are heated or cooled through the transformation zones at fixed rates and compared

to standards with known transitions. In DSC, the power supplied to the sample is varied to keep the sample at the same temperature as the reference. For DSC, the reference is usually just an empty pan. In DTA, the difference in temperature between the reference and the sample is measured. The reference material used is usually a material with no transitions in the desired temperature range, like α -alumina. Along with the glass transition temperature, DTA and DSC can determine the onset of crystallization and the maximum crystallization rate temperatures due to the processes being exothermic.

For the development of CCT curves, samples are commonly heated or cooled at designated rates and later analyzed using microscopy and x-ray diffraction. For TTT diagrams, samples are melted then quenched to a pre-determined temperature and held for various times before being quenched to room temperature to be analyzed. This latter technique is common to describe the crystallization of systems where the onset time of crystallization is much greater than the cooling time transferring from the melt to the isothermal temperature. As the sample is cooled from the melt to the isothermal holding temperature and later to room temperature, the sample is still nucleating and crystallizing so maximizing these cooling rates is important. In the Billings and Edwards experiments, 30 grams of glass was placed in a crucible that was melted and simply transferred to another furnace at the desired heat-treatment temperature [24]. For their glass-ceramics, crystals took hours to form so the cooling rate from the melt furnace would have a minimal effect on the results, with temperatures equilibrating within minutes. For melts that quickly crystallize, techniques that quickly quench the melt to the isothermal holding temperature are necessary. Peterson developed TTT diagrams for mold fluxes by quenching samples from a melt into a molten tin bath held at the desired heat-treating temperature [26]. The

quenching step was necessary since some crystals formed within minutes. Peterson measured quench rates from the melt to the molten tin to be $\sim 65^{\circ}\text{C/s}$ and from the molten tin to the water bath at room temperature to be $\sim 30^{\circ}\text{C/s}$.

1.2.4. JMAK Crystallization Model [23, 27-32]. The Johnson-Mehl-Avrami-Kolmogorov (JMAK) model can be used to characterize the time and temperature dependence of transformation processes, commonly but not limited to characterizing the time and temperature dependence on crystallization. The overall transformation process (e.g. crystallization) at a specific time ($\alpha_n(t)$) can be simply defined as

$$\alpha_n(t) = \frac{V_n(t)}{V_0} \quad (1)$$

where $V_n(t)$ is the volume of crystals at time (t) and V_0 is the total volume. In the JMAK model, $\alpha_n(t)$ can be predicted knowing the nucleation rate (J), linear growth velocity (v), and the geometric shape factor (ω_n) in which J , v , and ω_n are constants independent of time. Accounting for the decrease in available volume for potential crystallization with increasing crystallization, the classic JMAK equation is described as

$$\alpha_n(t) = 1 - e^{(-\frac{\omega_n}{n+1} J v^n t^{n+1})} \quad (2)$$

where n , the Avrami exponent of the transformation, relates to the different growth and nucleation mechanisms. Needle-like features grow when $n=1$, disc or plate like features form when $n=2$, and spherical features form when $n=3$. If

$$k_n = \frac{\omega_n}{n+1} J v^n \quad (3)$$

then Equation (6) can be rewritten as

$$\ln[-\ln(\alpha_n(t))] = \ln(k_n) + (n + 1)\ln(t) \quad (4)$$

where $\ln(k_n)$ is the intercept and n is the slope of a double logarithmic $-\ln(1 - \alpha_n(t))$ versus t plot which can be extracted from the TTT diagrams. The Avrami kinetic coefficient (k_n) relates to the activation energy (E) using the Arrhenius equation,

$$k_n = k_o e^{\frac{-E}{RT}} \quad (5)$$

where k_o is a frequency factor and R is the gas constant. Equation 9 can be re-written as

$$\ln(k_n) = \ln(k_o) - \frac{E}{RT} \quad (6)$$

where the activation energy is obtained from the slope of the plot of $\ln(k_n)$ vs $1/T$.

The applicability of the JMAK model to describe crystallization kinetics for glasses was investigated by Malek for both non-isothermal and isothermal conditions [32]. To understand the validity of the JMAK model it is important to understand the assumptions about the model. There are four main assumptions for the JMAK model:

- 1.) Isothermal crystallization
- 2.) Homogeneous nucleation (heterogeneous nucleation if the foreign sites are randomly and uniformly dispersed)
- 3.) Growth rate of the new phase is independent of time
- 4.) Spherical or low anisotropic growth of the new crystalline phase

When applying the JMAK model, it is important to verify that these conditions are met, as outlined by Malek [32]. A simple way to confirm the applicability of the JMAK model is to verify that the double log of crystallization ($\ln[-\ln(\alpha_n(t))]$) vs the $\ln(t)$ is linear, and that the Avrami kinetic coefficient (k_n) exhibits Arrhenius temperature dependence. Malek found linear dependencies for the crystallization of chalcogenide glass, $(\text{GeS}_2)_{0.3}(\text{Sb}_2\text{S}_3)_{0.7}$, shown in Figure 1.4. Furthermore, the maximum of the $z(\alpha_n)$ vs α_n plot should occur when $\alpha_n \approx 0.632$, where

$$z(\alpha_n) = \phi t \text{ (isothermal test methods)} \quad (7)$$

$$z(\alpha_n) = \phi T^2 \text{ (non-isothermal test methods)} \quad (8)$$

and ϕ is the normalized specific heat flow per sample mass [32]. The maxima of the $z(\alpha_n)$ function vs (α_n) was found to be within the range predicted using the JMA model, shown in Figure 1.5 from Malek.

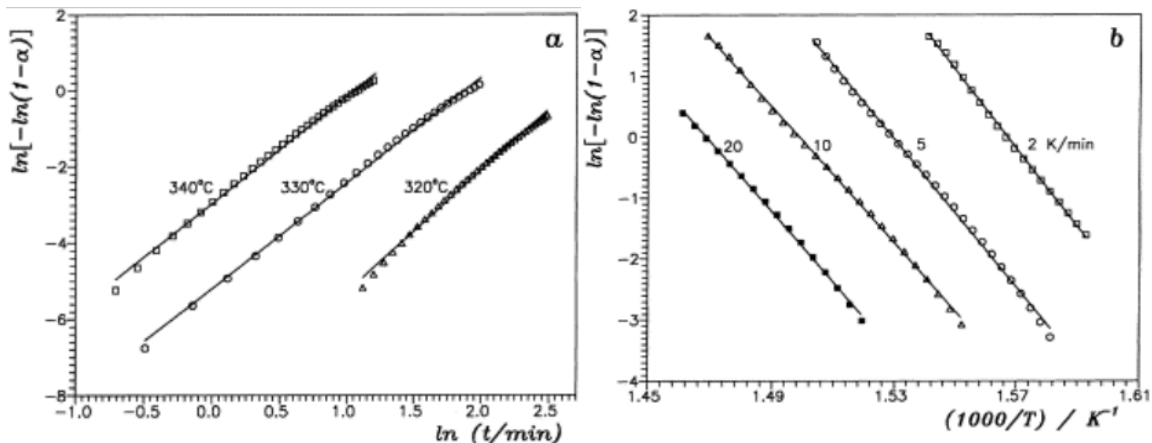


Figure 1.4. The double logarithmic plots obtained from DSC from Malek [32].

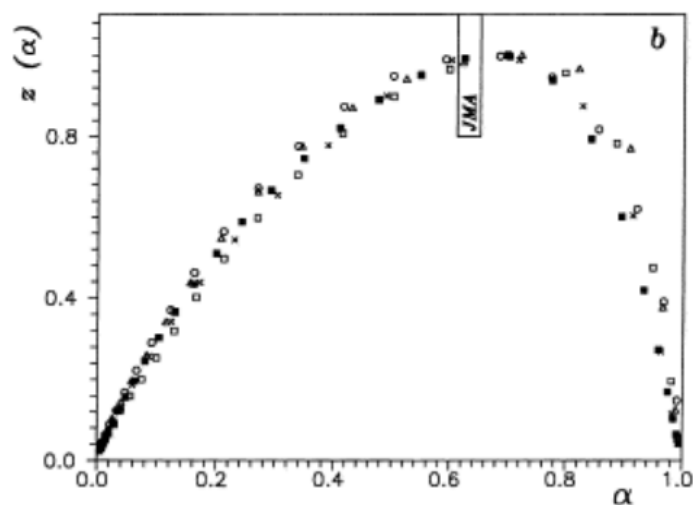


Figure 1.5. Normalized $z(\alpha_n)$ function for crystallization of bulk $(\text{GeS}_2)_{0.3}(\text{Sb}_2\text{S}_3)_{0.7}$ from Malek [32].

1.2.5. Chemical Durability of Borosilicate Glasses. The chemical durability of waste glasses must be understood to ensure the hazardous ions are not released into the biosphere. Chemical durability of glasses refers to the resistance of the glass to dissolution in aqueous solutions [33]. The rate of the aqueous dissolution of glass can be expressed as a function of thermodynamic and kinetic stability. Thermodynamics suggests that the reaction will continue until equilibrium is reduced. Kinetically, the reaction will occur if the system has enough energy to overcome the activation barrier.

Gin et al. outlined state-of-the-art dissolution (corrosion) processes for glasses [34]. There are many intertwined corrosion processes when glass surfaces come in contact with aqueous solutions; some of these are listed in Table 1.7.

For the reaction of waste glasses with groundwater, certain processes dominate at certain reaction times as shown in Figure 1.6 [34]. Stage I, the initial rate, is dominated by interdiffusion and hydrolysis reactions. Over time a gel layer may form, reducing the interdiffusion rate since the ions now have to diffuse through this gel layer before exchanging into solution. The dissolution rate is also known to drop to the “residual rate” or Stage II due to saturation of silica in the surrounding aqueous environment. Stage II of the corrosion of glass is controlled by many competing processes. Over long times, a resumption of dissolution (Stage III) can occur when the precipitation of mineral phases on the glass surface reduces the activity of soluble silicates in solution.

1.2.6. Chemical Durability Experimental Methods. Understanding the chemical durability is important for optimizing waste form compositions. Simple pH and weight loss tests are commonly used to understand the dissolution of glasses and these are often

combined with more advanced tests such as the analysis of reacted ions in leached solutions [33].

Table 1.7. Glass corrosion processes [34].

process	description
diffusion	diffusion of water into glass
interdiffusion	ion-exchange between ions in solution and weakly bound cations, usually alkali, in glass structure
hydrolysis	chemical breakdown of Si–O–M bonds (M = Si, Al, Zr, Fe, Zn, etc.)
condensation	condensation of the detached species
precipitation	crystallization of amorphous and soluble species

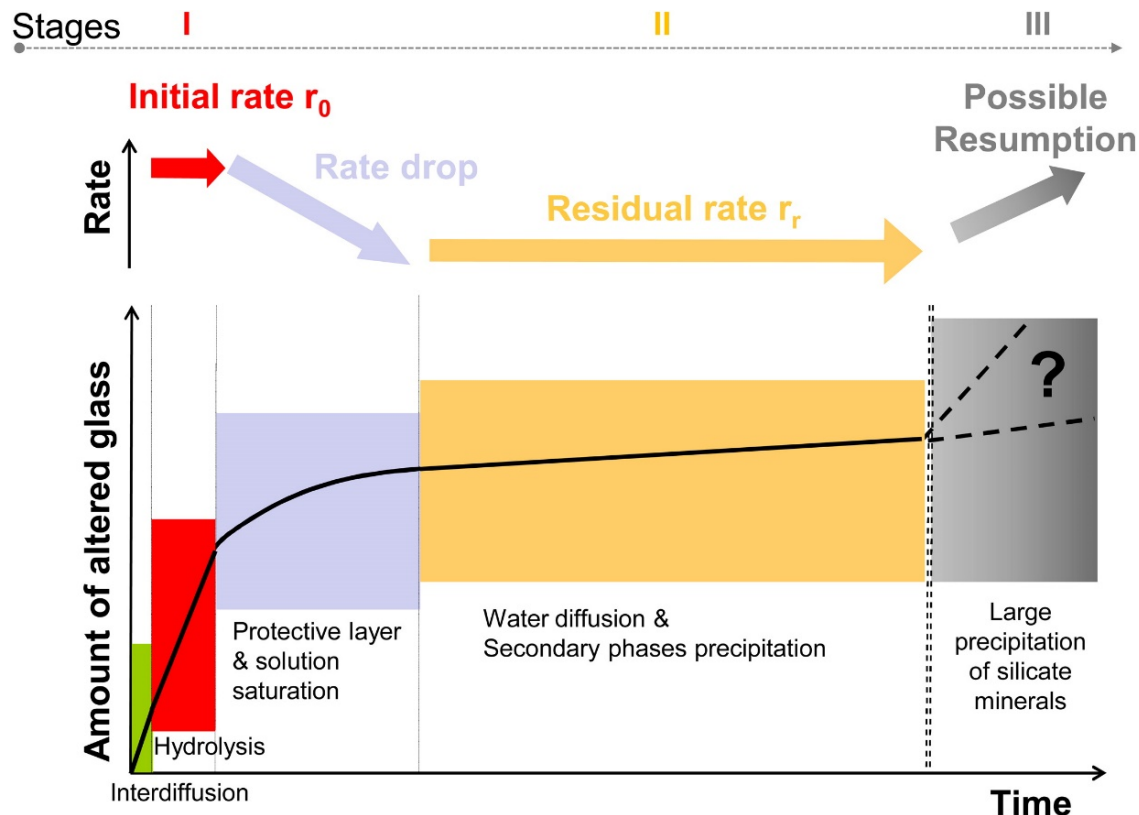


Figure 1.6. Long term dissolution processes of glass as a function of time [34].

The standard procedure for determining the chemical durability of waste glasses described by the waste acceptance product specifications, WAPS, is the Product Consistency Test, PCT [7, 35]. The PCT has rigorous test methods for determining the dissolution information in carefully controlled conditions. There are two versions of the PCT. Test method “A” is a specific seven-day test with no variability in experimental methods, allowing for cross-lab comparisons to be made. PCT test method “B” allows variability in test design to cater to the objectives of a study: for example, evaluating Stage I, II or III, dissolution behavior. As stated in the WAPS, the results of these tests are to be compared to the performance of the environmental assessment (EA) glass, the benchmark standard for acceptable waste glasses [36, 37].

1.3. BOROSILICATE GLASS-CERAMICS FOR IMMOBILIZING US COMMERCIAL REPROCESSED SNF

The current study is part of a much larger project that has been ongoing for over a decade involving many research groups to develop a borosilicate glass-ceramic for immobilizing the HLW stream generated from aqueous reprocessing of commercial SNF in the United States [25]. This section will review the previous work linked with this study, as well as other relevant literature.

The materials characterized in the present study and associated projects are intended to immobilize the Collins CLT waste stream from commercial reactors, Table 1.3. These studies have focused on immobilizing the waste using borosilicate glasses and glass-ceramics, although some other work has been done to evaluate other waste forms, such as iron-phosphate glasses [38].

PNNL has played a major role in the research on glass-ceramic development for immobilizing the Collins CLT waste [13, 15, 16, 25, 40-42]. Borosilicate glass-ceramics are of interest because they would allow greater loadings of insoluble components than would be possible for homogeneous glasses [39]. Compositions with up to 50 wt.% waste have been developed [16, 40]. These compositions have been melted using cold crucible induction melter technologies, demonstrating the possibility for full-scale processing of these glass-ceramics [41]. The initial experiments [13, 16, 40, 41] produced a compositional matrix for optimizing the glass-ceramics, considering both variations in the types and concentrations of glass components (Al_2O_3 , B_2O_3 , CaO , Li_2O , Na_2O , and SiO_2) and insoluble waste components, including lanthanide oxides (Ln_2O_3), ZrO_2 , and MoO_3 [25].

1.3.1. Phase Development. The crystalline phases that form in the glass-ceramics developed at PNNL for vitrifying the Collins CLT waste, include oxyapatite ($\text{Ca}_2\text{Nd}_8\text{Si}_6\text{O}_{26}$), Ln-borosilicate ($\text{Gd}_3\text{BSi}_2\text{O}_{10}$), powellite ($(\text{Ca},\text{Sr})\text{MoO}_4$) or ($\text{Ca},\text{Ba})\text{MoO}_4$), CsLiMoO_4 , $\text{La}_2\text{Mo}_3\text{O}_{12}$, pollucite (CsAlSiO_4), $\text{Zr}_{0.9}\text{Ce}_{0.1}\text{O}_2$, Y_2O_3 , and RuO_2 [25]. The centroid composition in the matrix, the composition used in the present study and given in Table 1.5, produced oxyapatite and powellite as the major phases.

The development of phases from the borosilicate melts is summarized in Figure 1.7 from Crum et al. [41]. Upon cooling, molybdenum-rich droplets separate from the surrounding glass, and that glass itself separated into and aluminum-silicon–cesium-rich droplets and a borosilicate matrix. When cooled slowly, the molybdenum-rich droplets crystallize into BaMoO_4 and CaMoO_4 , and oxyapatite crystallizes from the residual glass matrix. Quantitative x-ray diffraction determined that a sample with the centroid

composition cooled along the centerline cooling temperature profile of a waste canister ($\sim 0.005^\circ\text{C/s}$) included 18.6 wt.% oxyapatite ($\text{Ca}_2\text{Ln}_8\text{SiO}_2$), 9.1 wt.% $\text{Ca}_x\text{Sr}_{1-x}\text{MoO}_4$, and 0.4 wt.% $\text{Ca}_x\text{Ba}_{1-x}\text{MoO}_4$ [15]. A goal of this study is to determine at which temperature the phases in Figure 1.7 are forming to develop an updated phase transformation diagram as a function of time and temperature.

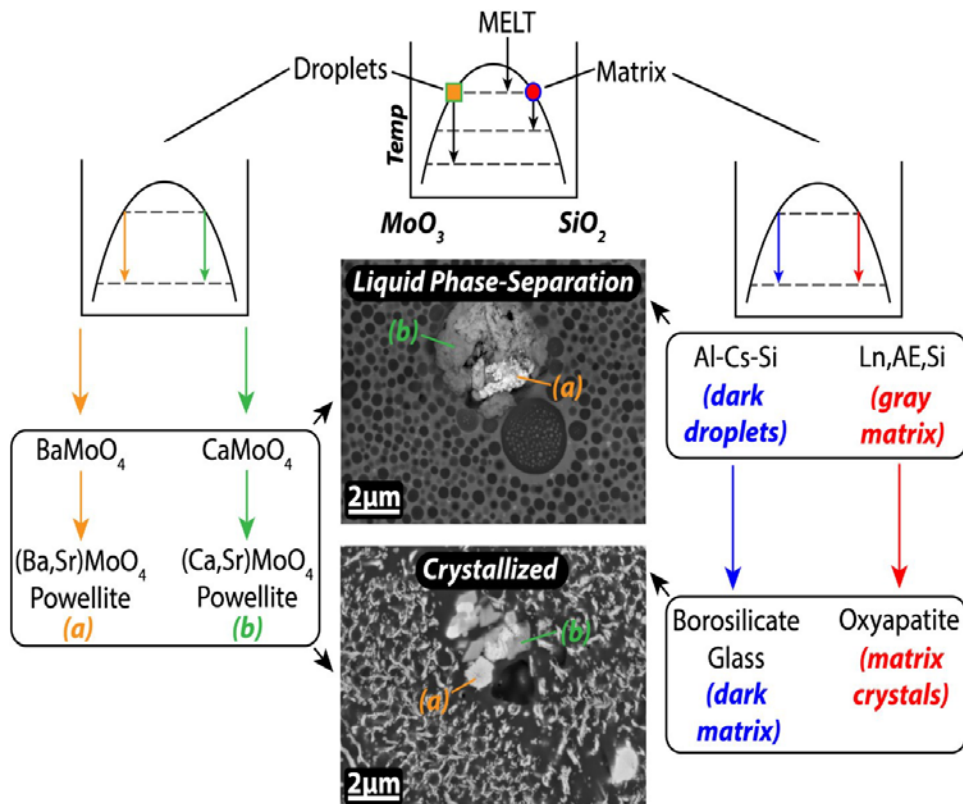


Figure 1.7. Proposed phase development [41].

Molybdates and oxyapatite crystals precipitate from the borosilicate waste glass due to the low solubilities of Mo and lanthanide oxides [13, 14, 43, 44]. Molybdate species like $[\text{MoO}_4]^{2-}$ are not strongly bonded to the silicate and borate network formers [14, 44].

Lanthanide ions are believed to be associated with the $[\text{MoO}_4]^{2-}$ units in the melts which are also found to slightly inhibit the crystallization of powellite [44, 45].

Depending on the glass composition, other molybdate phases can form, including Na_2MoO_4 [43]. Na_2MoO_4 has a lower chemical durability than CaMoO_4 and BaMoO_4 , so avoiding Na_2MoO_4 in the waste forms is desired [46]. It was found that increasing the $[\text{Ca}^{2+}]/[\text{Na}^+]$ ratio in the glass favored the formation of CaMoO_4 over Na_2MoO_4 [45, 46]. In addition, increasing the B_2O_3 content reduced the formation of Na_2MoO_4 by sequestering the Na^+ ions to charge balance the $[\text{BO}_4]^-$ units [46].

1.3.2. Chemical Durability. PCT studies at PNNL of glass-ceramics with low waste loadings (< 25 mass%) Si, B, Na, and Li release rates that were all an order of magnitude below those from the EA reference glass [13]. The release rate of Mo from a glass with 3.5 mass% Mo was much greater (1.06 g/L) when the melt was cooled slowly, following the canister center-line cooling (CCC) protocol than when the glass was quenched on a steel plate (0.3 g/L) [13]. No crystalline phases were present in the plate quenched sample and crystalline oxyapatite ($\text{Ca}_2\text{LN}_8\text{Si}_6\text{O}_{26}$) and a molybdate phases were found in the slow cooled sample. It was believed that the increase in the Mo release rate from the slow cooled sample was due to the preferential dissolution of the Mo-rich crystalline phase.

Glasses with high waste loadings, up to 50 wt.%, were cooled using the CCC profile and were reacted in static dissolution conditions for various times [40]. Mo release rates significantly increased (0.05 to 0.4 g/L) when the MoO_3 composition increased from 6.25 wt.% to 6.95 wt.%. It was believed that Mo was mostly contained in the crystalline phase (CaMoO_4) in the samples with lower Mo concentrations (6.25 wt.% and less) and the

increase in Mo leach rate from glasses with greater Mo-contents was due to the release of the residual Mo in the glass. Nd release rates were very low, indicating that oxyapatite is more durable than the residual glass. Also, in the higher waste loaded glass-ceramic, there was a significant increase in the release rates of boron and sodium, leading to the conclusion that the residual glass is less durable than the crystalline phases.

Dissolution tests were performed on the glasses from the test matrix study [25]. Three different dissolution tests were used to develop an understanding of the chemical durability of the glass-ceramic. One test, a modified PCT method “B”, static dissolution test, was used to analyze samples cooled following the CCC protocol after 7, 28, 119, 448 days on test. The second test was a single-pass flow-through, SPFT, test used on samples cooled at 4, 1, and 0.25 times the CCC-cooling rates [25, 42]. The third test was a modified single-pass flow-through, MSPFT, test used to further analyze the CCC cooled samples, combining weight loss measurements with corrosion depth measurements made using scanning electron microscopy [15, 25]. From the PCT tests, the effects of composition on the release of various components from the glass were determined; these are summarized in Table 1.8 [25]. Alkali-rich molybdates corrode faster than alkaline earth molybdates. B_2O_3 and Al_2O_3 tend to sequester the alkali ions in the glass phase as charge compensators, reducing the concentration of the alkali rich molybdate phases, increasing the overall durability.

Table 1.9 shows the release rates of different elements reported in a PCT study of the centroid sample, with B, Na, and Mo having the fastest leach rates [25]. The high Mo leach rate was attributed to the presence of low durability powellite phases. The SPFT dissolution tests showed that powellite dissolved faster than the residual glass, which

dissolved faster than the oxyapatite phases, regardless of thermal history, pH, or flow rate [25]. This conclusion was based on the relative release rates of Mo from powellite, B from the residual glass, and La from oxyapatite. On the other hand, the MSPFT dissolution test indicated that the residual glass dissolved fastest among these three microstructural features.

Table 1.8. Compositional effects on elemental release rates [25].

Glass Component	Trend increasing/decreasing release rates
Al ₂ O ₃ , ZrO ₂	decreases B, Li, Na, Cs, Si
SiO ₂	decreases B, Cs
B ₂ O ₃ , Na ₂ O, MoO ₃	increases B, Li, Na, Cs, Si
Ln ₂ O ₃	increases B, Li, Na, CS
Li	increases B, Si / decrease in Li
CaO	no effect

Table 1.9. Normalized release rates from PCT of the centroid glass composition cooled with CCC (g/m²) [25].

days	B	Ca	Na	Nd	Mo	Ba	Si
7	0.179	0.032	0.172	-	0.172	0.070	0.051
28	0.223	0.038	0.191	-	0.223	0.089	0.051
119	0.217	0.045	0.223	-	0.255	0.070	-
448	0.140	0.064	0.274	-	0.274	0.045	-

2. PHASE DEVELOPMENT IN A COMPLEX BOROSILICATE GLASS-CERAMIC WASTE FORM

ABSTRACT

A borosilicate glass provided by PNNL was melted and quenched with different thermal histories to produce samples with a range of microstructures that were then described by analytical electron microscopy and quantitative x-ray diffraction. Slower quench rates and longer isothermal treatment times produced samples with greater fractions of crystalline powellite (CaMoO_4 and related phases) and oxyapatite ($\text{Ca}_2\text{Ln}_8\text{Si}_6\text{O}_{26}$). A time-temperature-transformation, TTT, diagram was developed which shows that the fastest crystallization on cooling occurs at $\sim 1000^\circ\text{C}$. The growth of these crystalline phases is accompanied by changes in the composition and morphology of the residual glass. Oxyapatite was found to form hollow hexagonal crystals, more spaced out and larger at temperatures $\geq 1000^\circ\text{C}$, and these filled-in with time. Crystalline molybdate phases were present as lanthanide-containing needles in samples held above about 1000°C and as alkaline earth-containing droplets when grown from a phase separated residual glass at lower temperatures and times that precede oxyapatite crystal formation. Subsequently to the formation of oxyapatite and from a more homogeneous residual glass, molybdates phases were present as alkaline earth-containing crosses.

2.1. INTRODUCTION

A goal of the U.S. Department of Energy, D.O.E., Global Nuclear Energy Partnership, GNEP, is to develop spent nuclear fuel reprocessing technologies, such as the UREX process, to greatly reduce the volume of spent fuel that must be treated as a waste

material [1, 2]. The D.O.E. Fuel Cycle Research and Development (FCRD) program is tasked with developing viable storage options for the wastes resulting from fuel reprocessing [3]. There are manufacturing cost benefits in combining the reprocessed waste streams and incorporating them into a single waste glass formulation [4].

Borosilicate glasses are used worldwide and are understood by the FCRD to be a viable option to immobilize nuclear waste [3, 5]. Borosilicate glasses exhibit favorable properties regarding the immobilization of nuclear waste [6], including excellent chemically durability, thermal and radiation stability, and mechanical integrity. Also, many waste elements are soluble in borosilicate melts, allowing high waste loadings in the quenched glass. Some elements, such as molybdenum, however, have low solubility, <3 wt.%, in borosilicate melts [7, 8, 9], limiting loadings from Mo-rich combined waste streams into a homogeneous borosilicate glass. Waste loading levels can be increased in borosilicate glass-ceramics if precipitated phases and residual glass possess requisite chemical durability.

Recently, researchers at Pacific Northwest National Lab (PNNL) have developed a borosilicate glass-ceramic composition to immobilize a waste generated by aqueous spent nuclear fuel reprocessing techniques [8, 10-16]. One of the goals of this development program, described in the preliminary Technology Maturation Plan, TMP, is to develop processes for the full-scale production of glass-ceramics to immobilize high level waste [17]. One outcome of the TMP is to “predict the amounts, types, and compositions of phases formed in the final, canistered waste form [17].” More specifically, in order to improve formulation and processing to ensure that an optimized waste-loading glass-ceramic is developed, models accurately predicting phase growth and crystallization as a

function of thermal history are needed [12, 17]. The Waste Acceptance Product Specifications (WAPS) also require the development of time-temperature-transformation (TTT) diagrams to fully understand the crystallization pathways [18]. The WAPS is a set of acceptance specifications for high-level nuclear waste developed by the DOE.

Researchers at PNNL have developed production scale melting techniques for borosilicate glasses and have described crystallization and phase formation as a function of cooling rate for compositions similar to that in the present study [12, 15]. In that work, it was proposed that the melt would separate into Mo-rich and silicate-rich phases. Crystalline BaMoO_4 ($(\text{Ba}, \text{Sr})\text{MoO}_4$) and CaMoO_4 ($(\text{Ca}, \text{Sr})\text{MoO}_4$) would precipitate from the Mo-rich phase. The silicate-rich phase would further separate into a Cs-alumino-borosilicate liquid and an alkaline earth-lanthanide-silicate liquid, the latter of which would crystallize into oxyapatite. The present study is intended to determine the effects of time and temperature on these morphological and compositional transformations.

Prior isothermal heat treatment studies [10] of these borosilicate glass-ceramics involved samples that were quenched from the melt to room temperature, and then reheated and isothermally held at different temperatures for various times, as shown schematically in Figure 2.1a. In the present work, samples were either quenched from the melt directly to the isothermal temperature, held for desired times, then quenched again to room temperature, or were continuously cooled from the melt to room temperature at different rates, as shown in Figure 2.1b. Quenching directly from the melt more closely simulates the thermal history of a waste glass which will follow a nucleation and crystallization pathway significantly different from the more conventional quench and reheat experiments.

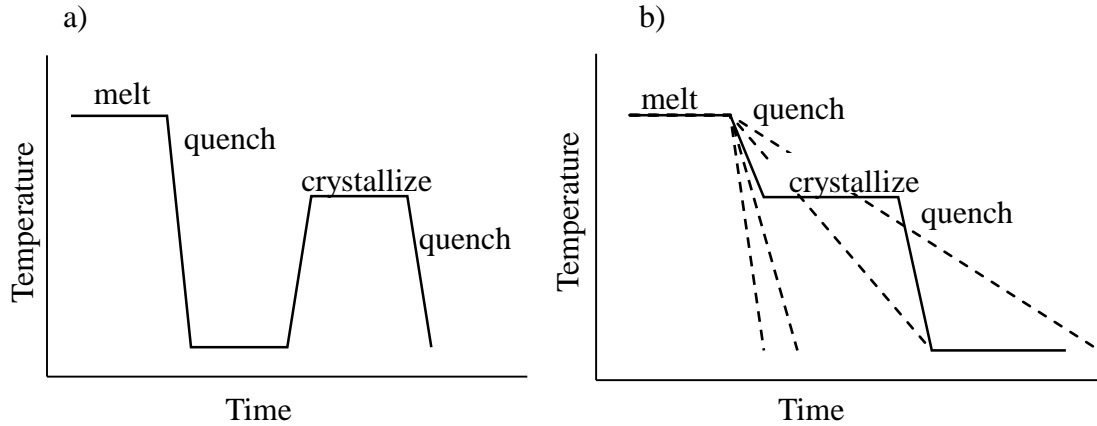


Figure 2.1. Schematic representations of isothermal experiments. a) Reheating quenched glasses to the treatment temperature. b) Quenching from the melt directly to the isothermal treatment temperature (solid lines) or quenching to room temperature at different fixed rates (dashed lines).

The focus of the present study is to characterize the effects of thermal history on the microstructural development of a borosilicate glass-ceramic waste form. Microstructures were characterized as a function of time and temperature through isothermal and constant cooling experiments. Time-Temperature-Transformation (TTT) and Continuous-Cooling-Transformation (CCT) diagrams, and activation energies for the growth of major phases that form in these samples, have been determined, with the intention of using this information to help engineer an optimized waste-loading glass-ceramic.

2.2. EXPERIMENTAL PROCEDURES

2.2.1. GLASS COMPOSITION

The glass-ceramic material used in these studies was produced at Pacific Northwestern National Lab (PNNL) as part of a larger material development program and was provided by Jarrod Crum (PNNL). Information about sample preparation can be found in reference [13]. Reagent grade raw materials were melted twice in covered Pt/10% Rh crucibles between 1250°C and 1450°C for an hour, then quenched on an Inconel plate in air. Figure 2.2 shows a picture of representative samples of these inhomogeneous materials, and Table 2.1 provides their nominal composition [13].



Figure 2.2. Optical image of the as-received glass. The darker side is the side of the melt that was in contact with the Inconel plate when the melt was quenched.

Table 2.1. Centroid glass-ceramic target composition [13].

Oxide	Content [mol%]	Oxide	Content [mol%]
SiO ₂	47.674	TeO ₂	0.641
B ₂ O ₃	11.329	Sm ₂ O ₃	0.481
CaO	8.445	Y ₂ O ₃	0.433
MoO ₃	4.229	Rb ₂ O	0.353
Al ₂ O ₃	3.789	RuO ₂	0.232
Na ₂ O	4.069	CdO	0.135
ZrO ₂	3.774	SeO ₂	0.116
Li ₂ O	3.247	RhO ₂	0.089
Nd ₂ O ₃	2.427	Eu ₂ O ₃	0.076
BaO	2.241	Ag ₂ O	0.075
Cs ₂ O	1.591	SnO ₂	0.072
SrO	1.476	Gd ₂ O ₃	0.070
Ce ₂ O ₃	1.472	PdO	0.023
La ₂ O ₃	0.757	Total	100.000
Pr ₂ O ₃	0.684	Waste Loading	47.394 wt.%

2.2.2. CONTINUOUS COOLING EXPERIMENTS - MELT QUENCHING TECHNIQUES

Fast quench rates, several hundred °C/s, were achieved using a custom-built dual roller system (Figure 2.3, left). The 8 inch (circumference), water-cooled stainless-steel rollers were set with a gap spacing of 0.25 mm and a rotation rate of 1140 rpm powered by a ½ HP motor. As-received glass was melted at 1300°C in a platinum crucible for 1 hour and the melt was poured directly between the rollers to produce glass ribbon ~0.25 mm thick (Figure 2.3, right).

Two wedge molds, made from different metals but with the same wedge dimensions, were used to quench melts at rates between about 4°C/s and 45°C/s, Figure 2.4. One mold was made from copper and was water-cooled to produce a faster range of quench rates than the second mold which was made of stainless steel and was air cooled.

Sheathed type-K thermocouples were inserted through the sides of the mold to the centers of each of the steps. Further information about the wedge mold experiments is provided by Maldonado and Peterson [19, 20]. About 100 grams of as-received glass was melted at 1300°C in a platinum crucible for 1 hour and then the melt was immediately poured down the center of the mold, onto each of the eight thermocouples to record individual cooling rate profiles using a portable data logger (Graphtec midi logger GL 220) that recorded temperatures ten times every second. Once the sample was cooled, the mold was opened, and samples were collected near each thermocouple junction for microstructural analyses.



Figure 2.3. Roller quenching system.

In another set of experiments, about ~30 grams of as-received glass was melted for one hour at 1300°C in a platinum crucible. The crucible was removed from the melting furnace and a K-type thermocouple was then inserted into the melt (Figure 2.5). One crucible was then allowed to cool to room temperature in air at an average rate of ~3.5°C/s. The second crucible was transferred to a preheated annealing furnace at 1100°C that was then shut-off to allow the glass to cool at a much slower rate (~0.1°C/s). A third sample

was transferred to an annealing furnace set at 1150°C that was programmed to cool following the canister centerline cooling (CCC) profile, summarized in Table 2.2 [8]. The CCC profile corresponded to an average cooling rate of ~0.005°C/s, from 1150°C to 600°C. At the completion of each of these three crucible experiments, material near the tip of the respective thermocouple was collected for microstructural characterization.

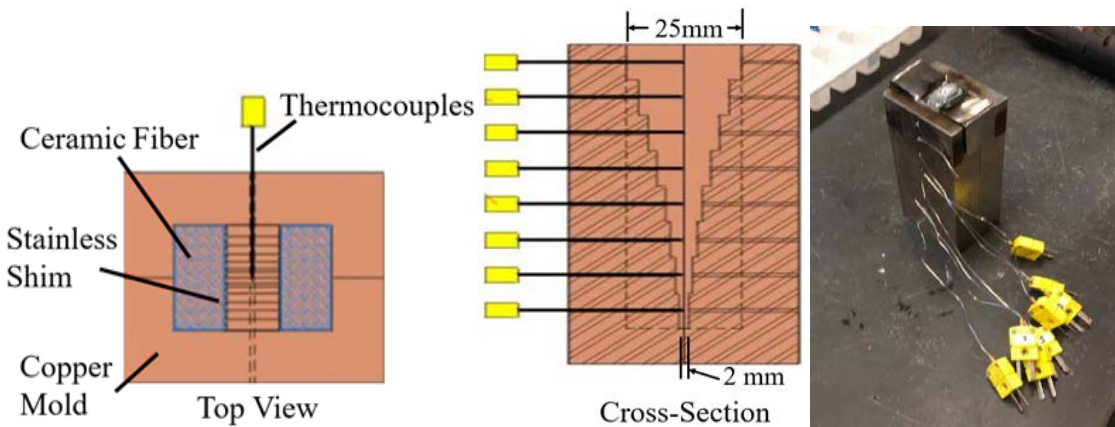


Figure 2.4. Schematic diagrams of assembled water-cooled copper wedge mold (left and center) and a photograph of the steel wedge mold (right).



Figure 2.5. Furnace cool experiment with embedded thermocouple.

Table 2.2. Treatment schedule comparable to the CCC of a waste storage canister [8].

Step	Start Temperature (°C)	Rate (°C/min)	Step Duration (hours)
1.	1200 - 1150	0	Preheat 0.5 hours then fast cool
2.	1150 - 1050	~ -7	0.2
3.	1050 - 950	-0.935	1.8
4.	950 - 886	-0.288	3.7
5.	886 - 845	-0.108	6.3
6.	845 - 626	-0.205	17.8
7.	626 - 400	-0.126	29.9
8.	400	0	Dwell 1 hour

2.3. ISOTHERMAL EXPERIMENTS - MELT QUENCHING TECHNIQUES

Molten tin was used to rapidly quench and thermally equilibrate melt samples in the isothermal experiments. The tin was contained in either an alumina or a graphite crucible that was then held in a furnace. Argon gas was flowed over the molten tin surface to minimize oxidation. A thermocouple in a closed-end alumina tube was immersed in the tin bath to record the temperature throughout the experiments. Tin bath temperatures changed with time when samples were first transferred from the melting furnace to the tin bath, and average temperatures over the course of an isothermal holding time are reported.

Three different tin bath configurations were used in these experiments, as shown in Figure 2.6. The four-crucible box furnace tin bath arrangement shown in Figures 2.6a and b, was used for isothermal experiments between 600 and 800°C. These alumina crucibles contained ~ 250 cm³ of molten tin, and temperatures fluctuated up to 15°C in some experiments. This configuration was also used by Peterson [20]. The second configuration used a larger graphite crucible to hold a larger volume of molten tin (500 cm³) to reduce temperature fluctuations and operated in a kiln that could go to temperatures as high as ~ 1050 °C, Figure 2.6c and d. The third configuration used the same larger volume of tin

but operated in an annealing furnace that reached 1100°C. Supplemental information about the different tin baths can be found in Appendix A.

As-received glass was crushed using a steel impact mortar and about 5 grams of powder was packed inside a crimped 304 stainless steel tube, 0.25" OD, 0.23" ID, ~6" length. These tubes were sealed at the open end with a stainless-steel rod which was then used to lower the tube into a melting furnace set at 1300°C and held here for 90 seconds to melt the glass. The tubes were removed from the furnace and immediately plunged into a molten tin bath set at the desired temperature and held for the specified treatment time, after which the sample tubes were then removed from the molten tin and immediately quenched in room temperature water. Once fully quenched, the tubes were sectioned using a diamond saw and material was prepared for characterization. Only material from the centers of the tube that were fully submerged in the tin bath were characterized.

In this report, samples that were held in tin baths between 600°C and 1100°C, from 0.5 to 60 minutes, are described. In addition, a sample that was quenched directly from 1300°C to room temperature water was prepared as a baseline against which the isothermally heat-treated samples are compared. Furthermore, a sample held at 900°C for 24 hours was prepared and characterized.

2.2.4. SAMPLE CHARACTERIZATION

2.2.4.1. X-ray Diffraction. Representative samples with different thermal histories from the continuous cooling and isothermal hold experiments were ground with a mortar and pestle and sieved to <45 µm. Some of these powders were mixed with 10% (by weight) SRM 674 standard rutile, (TiO₂), and used in the quantitative x-ray diffraction (XRD) study. Powders were pressed into standard 2.5 cm diameter powder mounts and analyzed

using a Philips X'pert multipurpose diffractometer (PIXcel detector, Cu K α radiation, at 45 kV and 40 mA, and a 0.38 fixed divergence slit). Hour long scans were taken from 9.995° to 70.000° 2 θ with a step size of 0.026°. PANalytical HighScore software was used to identify crystalline phases. If required, MDI RIQAS software was used to perform Rietveld analysis to compare peak intensities with the known intensities of the rutile standard. Comparable peak broadening and subtracted baseline factors were used on all patterns to determine the relative weight fractions of different crystalline phases and the residual glass from the Rietveld analyses. Samples were collected from several repeated isothermal heat treatment experiments, and other samples were prepared and repeated at least twice to determine the uncertainties of the quantitative XRD analyses. These repeat experiments indicate a relative uncertainty in the quantitative phase analyses of about 20% (Table A.2, Appendix A). The difference between the Rietveld fit and the measured pattern shows very little peaks proving the fit to strongly match the measured pattern, Figure A.9 in Appendix A.

2.2.4.2. Analytical Electron Microscopy. An FEI Helios NanoLab 600 FIB/FESEM scanning electron microscope (SEM) equipped with an Oxford energy dispersive spectrometer (EDS) was used to characterize the various phases that formed in heat-treated samples. Representative samples were mounted in epoxy, polished to a 0.25 μ m diamond suspension finish, masked with copper tape, and sputter coated with an Au/Pd film before loading into the SEM. Images were obtained at various magnifications, electron voltages, and beam currents. EDS spectra and maps were obtained using Oxford Instruments AZTEC software at various magnifications using 15 keV and 1.4 nA electron beam. ImageJ software was used to quantify microstructural features.

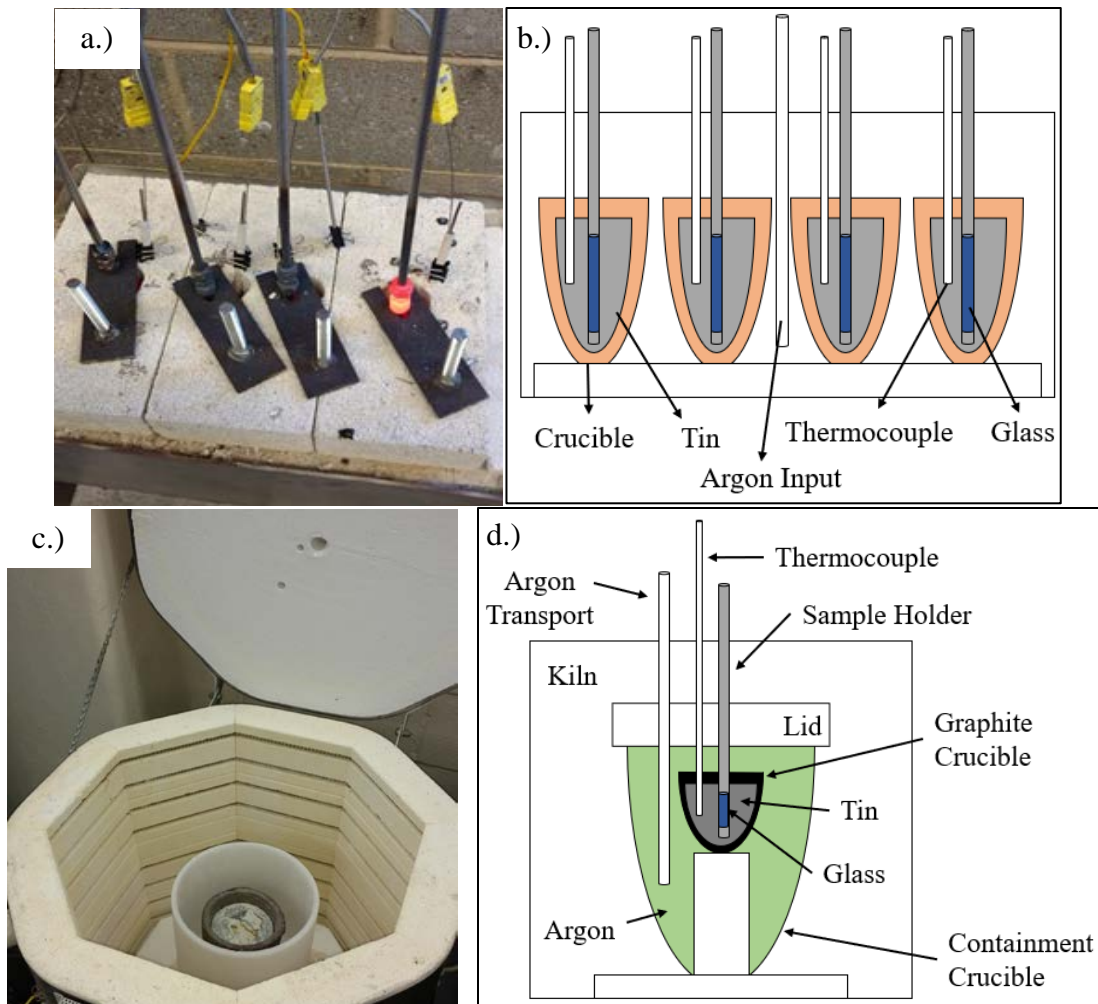


Figure 2.6. a) Photograph and b) schematic illustration of the set-up for the small volume isothermal tin bath. c) Photograph and d) schematic illustration of the set-up for the large volume isothermal tin bath.

Some samples were also characterized using a transmission electron microscope/scanning transmission electron microscope, TEM/STEM, (FEI Tecnai G2 F20 S-Twin). Samples were ground to a powder ($<45\text{ }\mu\text{m}$) using a mortar and pestle, then dispersed on a copper grid and loaded into the TEM. Another TEM sample was prepared by milling a section $\sim 100\text{ nm}$ thick from a polished bulk glass sample using the Focused Ion Beam (FIB) with the FEI Helios NanoLab 600 dual beam system. Before the final milling the sample was transferred to a copper grid using an Omniprobe and later the

specimen attached to the copper grid was loaded into the TEM. TEM examinations were performed at various magnifications, an acceleration voltage of 200 keV and an emission current of 80 μ A. Micrographs were collected using annular bright field/dark field detectors. Energy dispersive X-ray spectroscopy, EDX, spectra was collected using an Oxford ultra-thin window (EDX) detector. Selected area diffraction (SAD) patterns along zone axes were collected using a Gatan ORIUS wide-angle charge-coupled device (CCD).

2.2.4.3. Raman Spectroscopy. Raman spectra were collected from polished samples with a Horiba Jobin YVON LabRAM Aramis μ -Raman spectrometer, using a 632.8 nm HeNe laser under a 100x magnification objective with an 1800gr/mm grating. A silicon reference sample was used to calibrate the energy scale. Ten spectra were averaged over ten seconds and at least three areas of the desired phases were examined in every case to ensure that representative spectra were reported.

2.2.4.4. Differential Thermal Analysis. Differential thermal analysis (DTA) data was collected on some samples (Perkin Elmer Differential Thermal Analyzer DTA 7 with Pyris software). Powders samples (~25mg) were heated in Pt crucible at 10°C/min in a Nitrogen atmosphere to 1200°C. Thermal data was compared to an Al_2O_3 reference.

2.3. RESULTS AND DISCUSSION

2.3.1. SAMPLE OBSERVATIONS

The as-received glasses are visibly heterogeneous, and all glasses produced from the various quenching and isothermal experiments, including the roller quenched glass, are also visibly heterogeneous. For example, Figure 2.2 shows an optical micrograph of the

top side and the bottom side of the as-received glass; the latter was cooled faster with direct contact to the Inconel plate causing this surface to be darker in color.

Samples from the continuous cooling experiment were sent to Savannah River National Laboratories (SRNL) to determine their compositions after remelting in Pt at 1300°C for 60 min. Compositions were measured using ICP-AES and ICP-MS and are compared to the nominal composition, Figure A.10, A.11, and A.12 of Appendix A, showing a good fit with most components. Cs contents were measured to be slightly lower than the nominal compositions, likely due to Cs volatilization.

Figure 2.7 shows optical images of the cross-sections of representative samples sectioned from the stainless tubes from different isothermal experiments. The sample on the left was held at 1300°C for 90 seconds and then quenched into a tin bath at 800°C, held for 0.5 minutes and finally quenched into room temperature water. The center sample and the sample on the right were also isothermally heat-treated at 800°C, but held for 4 and 60 minutes, respectively. The cracks in these samples likely resulted from thermal shock induced in the glass during the water quench.

2.3.2. TIME-TEMPERATURE PROFILES

For the wedge-mold experiments, the melt was too viscous to flow into the narrowest (2mm) bottom (8th) step, so samples with seven different quench rates were obtained from each of these experiments. The thermal profiles measured from the wedge mold and crucible quench experiments are compared in Figure A.1 of Appendix A. The measured thermal profile of the CCC sample was very similar to the programmed furnace and appears to have converged after 20 minutes, as shown in Figure A.2 of Appendix A.

Quench rates were calculated from the respective times it took to cool each sample from their highest recorded temperature to 590°C, and these are given in Table 3 along with the maximum recorded temperature for each experiment. The cooling rates are calculated to 590°C because this is the glass transition temperature of the as-received glass that was measured using differential thermal analysis; the DTA data are shown in Figure A.3 of Appendix A.



Figure 2.7. Optical images of as-received glass melted in a stainless-steel tube at 1300°C for 90 seconds and then left) quenched into an isothermal tin bath at 800°C and held for 0.5 minutes before quenching in water; center) quenched into an isothermal tin bath at 800°C and held for 4 minutes before quenching in water; and right) quenched into an isothermal tin bath at 800°C and held for 60 minutes before quenching in water.

Figure 2.8 shows the quenching profile recorded from a thermocouple that was inserted into a stainless-steel tube, packed with as-received glass, placed into the 1300°C furnace, melted for 90 seconds, and then quenched into room temperature water. This profile indicates this baseline sample was quenched at rate nearly 300°C/s. Figure 2.8 also shows the measured temperature profile recorded from a thermocouple inserted into a stainless-steel tube packed with glass, placed in a furnace at 1300°C for 90 seconds quenched in a tin bath at 600°C, held for 4 minutes, and quenched to room temperature. Quench rates from these experiments are listed in Table 3. The quench rate of the roller

quenched samples was not measured, but based on the phases present in these samples, as discussed below, was at least as fast as the baseline quenched sample.

Table 2.3. Quench rates for continuously cooled and some isothermal samples, calculated from the time between the maximum recorded temperature and when the sample reached 590°C.

Continuous Cooling Experiments				
Quench Method	Maximum Temperature Recorded [°C]		Average Quench Rate [°C/s]	
CCC	1150		0.008	
Furnace Cool	1119		0.1	
Air Cool	1124		3.4	
Roller Quench	-		~300 - 1000	
Wedge Molds	Copper	Steel	Copper	Steel
Step 1 (25 mm)	1241	1190	5.1	4.1
Step 2 (20 mm)	1232	1168	5.8	4.3
Step 3 (16 mm)	1204	1181	8.7	6.3
Step 4 (12 mm)	1223	1197	12.1	9.4
Step 5 (8 mm)	1208	1179	18.1	15.3
Step 6 (6 mm)	1149	1102	24.6	20.5
Step 7 (4 mm)	1011	955	42.8	33.8
Isothermal Experiments				
Quench Description		Quench Rate [°C/s]		
1300°C to water quench (baseline)		294 ± 40		
1300°C to 600°C tin bath		43 ± 10		
600°C tin bath to water quench		50 ± 3		

2.3.3. PHASE IDENTIFICATION

Phases were identified in each sample using XRD, SEM EDS, and TEM EDS. In some samples, some phases were either x-ray amorphous or in amounts too small to be detected using XRD. TEM was used to determine phases too small to be analyzed with the SEM. TEM electron diffraction was also used on samples that were determined to be x-ray amorphous.

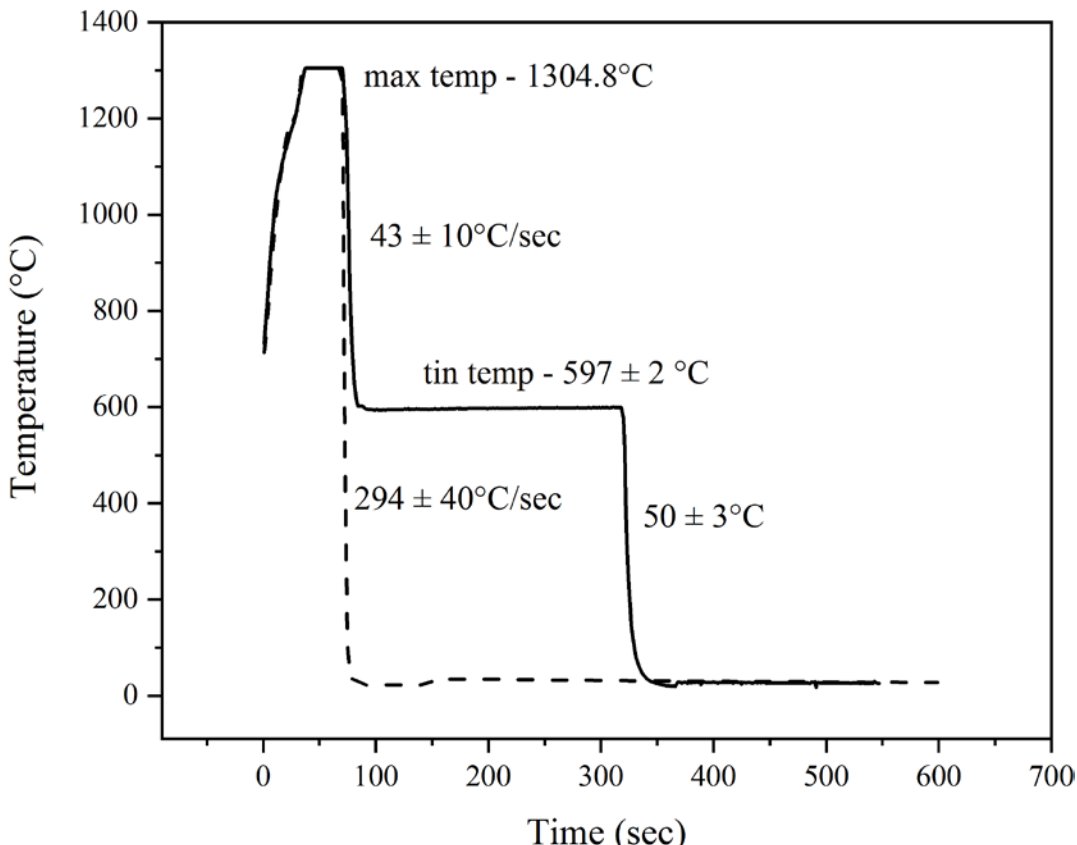


Figure 2.8. Cooling rate profiles for a thermocouple in a packed stainless-steel sample tube quenched in water bath (dotted line) and a thermocouple in a packed stainless-steel sample tube quenched in a tin bath set at 600°C, then quenched in water bath (solid line).

2.3.3.1. Phases Identified in the Continuously Cooled samples. Figure 2.9 shows the general trend in phase development from a mostly homogenous glass in a sample quenched at high rates, Figure 2.9a, to a developed glass-ceramic in the slower cooled samples. First, the glass is seen to separate into a droplet phase, dark circles, and a matrix phase with Mo-rich spherical particles, white circles, shown in Figure 2.9b. Next, oxyapatite, $\text{Ca}_2\text{Ln}_8\text{Si}_6\text{O}_{26}$, crystallizes in which different morphologies are found at different cooling rates. Oxyapatite is the hollow hexagonal features in Figure 2.9c and the dendritic type features in Figure 2.9d.

Scanning electron microscopy (Figure 2.10) provided evidence for a ruthenium-rich phase in every sample, including the roller quench samples. These phases were unevenly dispersed, and often agglomerated. Heavy metal phases are known to be insoluble in borosilicate glasses, at least to temperatures of 1300°C [8].

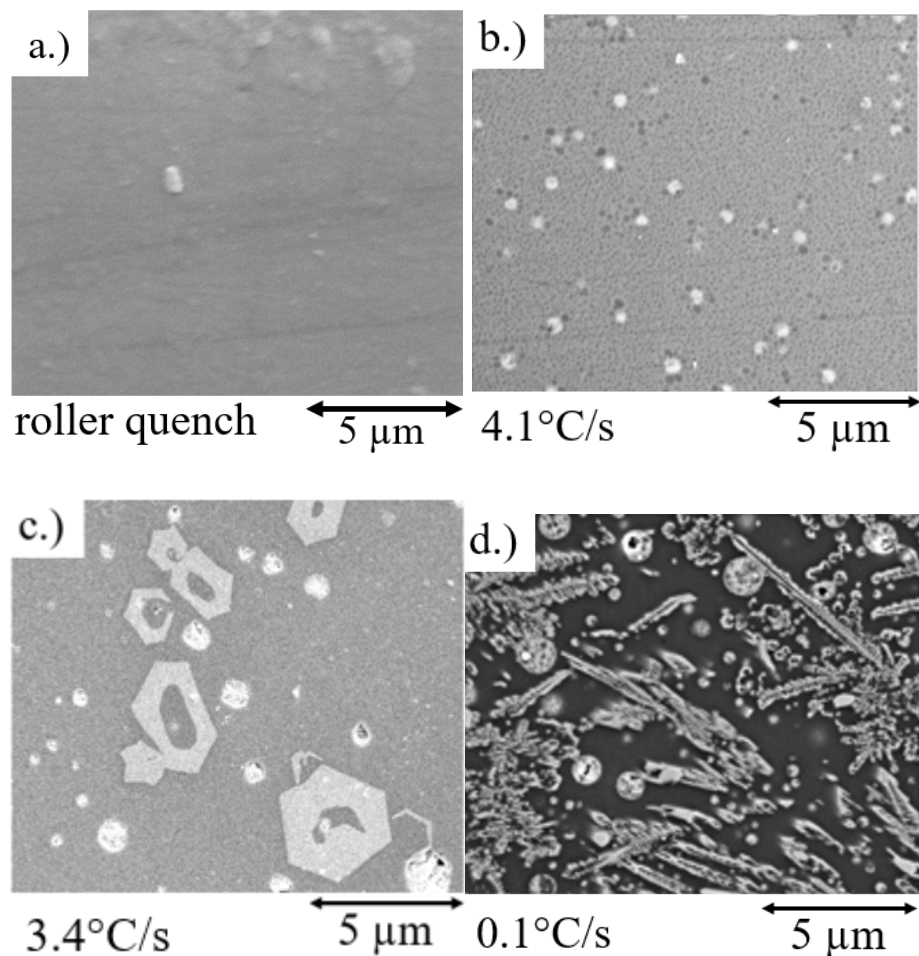


Figure 2.9. SEM micrographs of a) roller quench, b) $4.1\text{ }^{\circ}\text{C/s}$, c) $3.4\text{ }^{\circ}\text{C/s}$, and d) $0.1\text{ }^{\circ}\text{C/s}$.

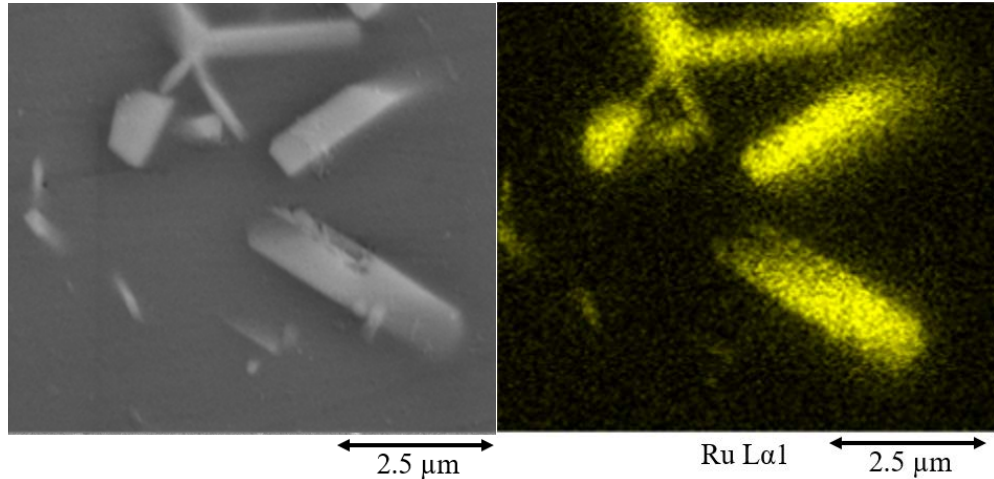


Figure 2.10. Scanning electron image (left) and Ru EDS map (right) of precious metal crystallites found in the roller quenched glass.

Figure 2.11 shows representative x-ray diffraction patterns for samples quenched at different rates. The CCC, 0.1°C/s and 3.4°C/s samples were doped with 10 wt.% SRM 674 rutile standard (PDF# 04-008-7848). Samples quenched at rates faster than about 4.1°C/s were mostly amorphous, showing only one significant peak near 28° (2θ) and three smaller peaks at 33° 45° 56° (2θ). These peaks closely match those associated with a strontium lanthanum molybdate phase ($\text{Sr}_{0.44}\text{La}_{0.39}(\text{MoO}_4)$, PDF# 04-002-4182) and this is further discussed in Appendix A. In the more slowly quenched samples, < 3.4°C/s, oxyapatite (e.g., $\text{Ca}_{1.1}\text{Nd}_{3.9}(\text{SiO}_4)_3\text{O}_{0.95}$, PDF# 04-007-5969) is clearly the major crystalline phase and crystalline powellite (e.g., CaMoO_4 , PDF# 04-013-6764) was also identified. A peak at ~27° (2θ) is also present in the patterns collected from the samples cooled at 3.4°C/s and 0.1°C/s, and this is assigned to a barium strontium molybdate phase (e.g., $\text{Ba}_{0.75}\text{Sr}_{0.25}(\text{MoO}_4)$, PDF# 04-019-9425).

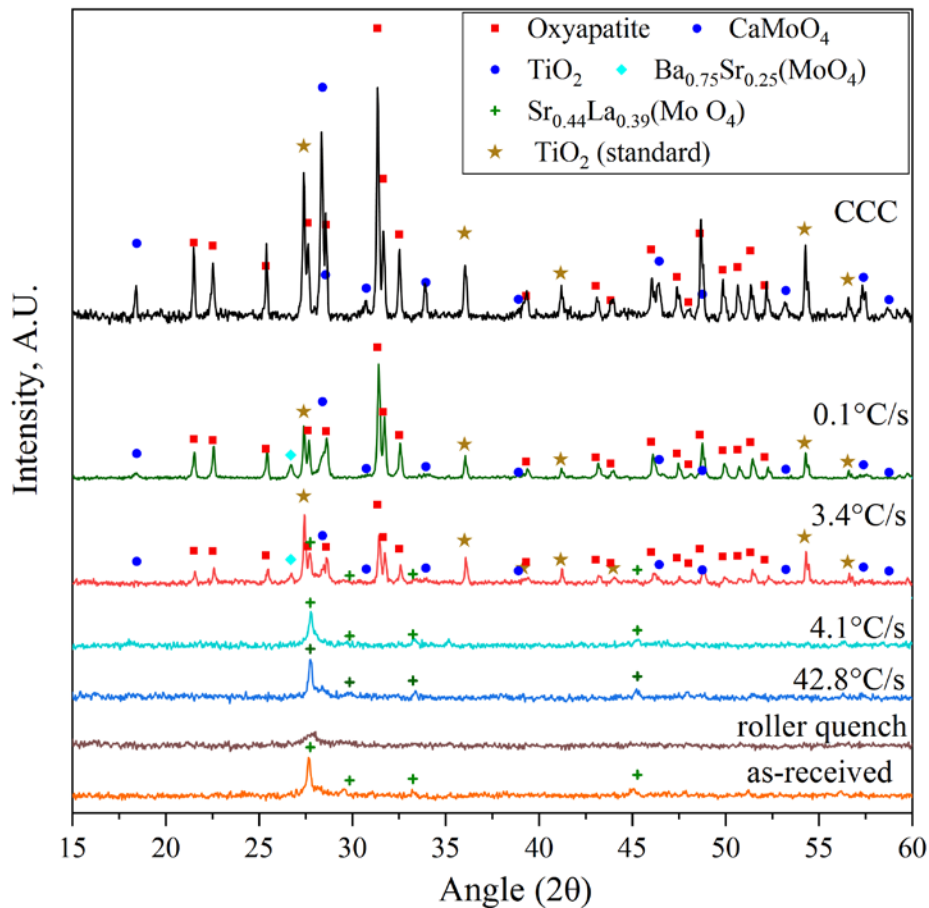


Figure 2.11. XRD patterns from the as-received glass cooled at various rates with the peaks assigned to the identified phases.

Figure 2.12 shows examples of X-ray maps from the EDS analyses used to identify the elements in the different phases that form in a quenched sample, including Ca and Nd in the oxyapatite crystals and Mo, Ba, and Ca in the powellite crystals. Sodium does not appear to be a major component in the latter phases, a preferred condition due to the poor chemical durability of NaMoO_4 [15]. Figure 2.13 shows an example where the powellite have separated into Ba-rich molybdate and Ca-rich molybdate phases. Crystalline powellite only could be detected by x-ray diffraction in samples cooled at 3.4°C/s and slower (Figure 2.11). Similar Mo-rich phases were found by SEM EDS in samples quenched faster than 4.1°C/s , and up to at least 42.8°C/s , and, TEM electron diffraction

confirmed that the Mo-rich droplet phase in a sample quenched at 4.1°C/s was crystalline, as shown in Figure 2.14. Additional information on the selected area diffraction (SAD) pattern, Figure 2.14 is given in Appendix A.

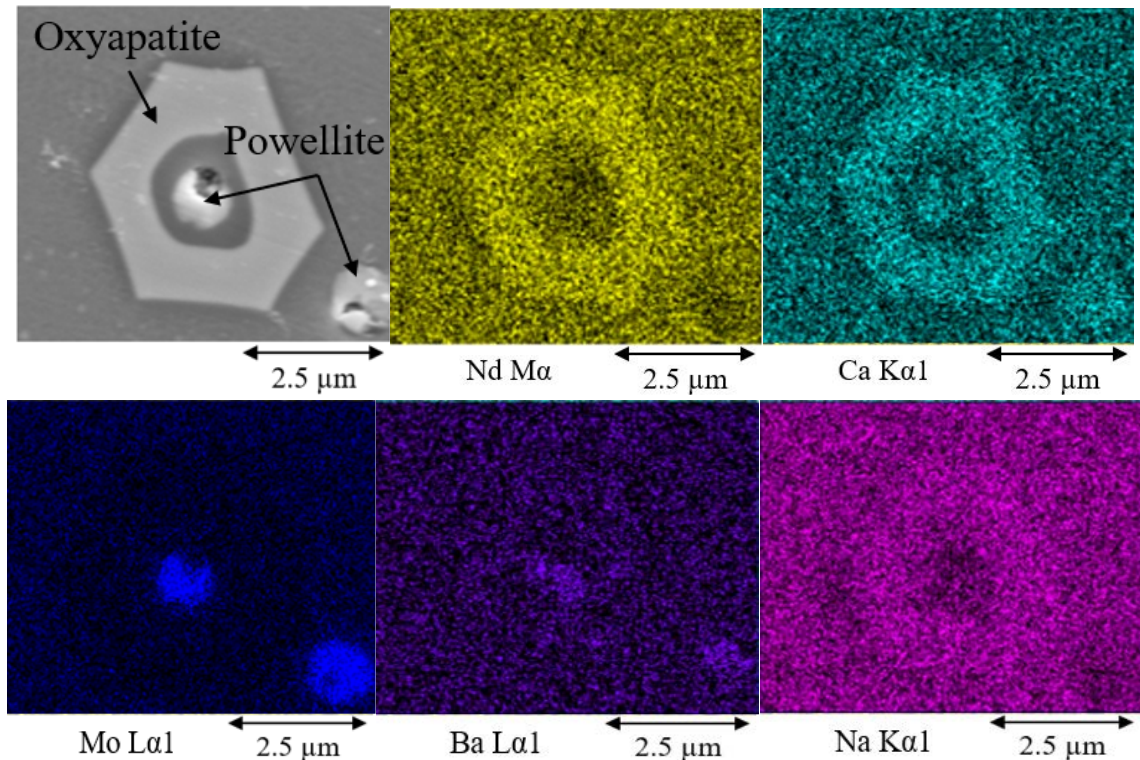


Figure 2.12. An SEM image and Nd, Ca, Mo, Ba, and Na X-ray maps collected from oxyapatite and powellite particles formed in a sample quenched at 3.4°C/s.

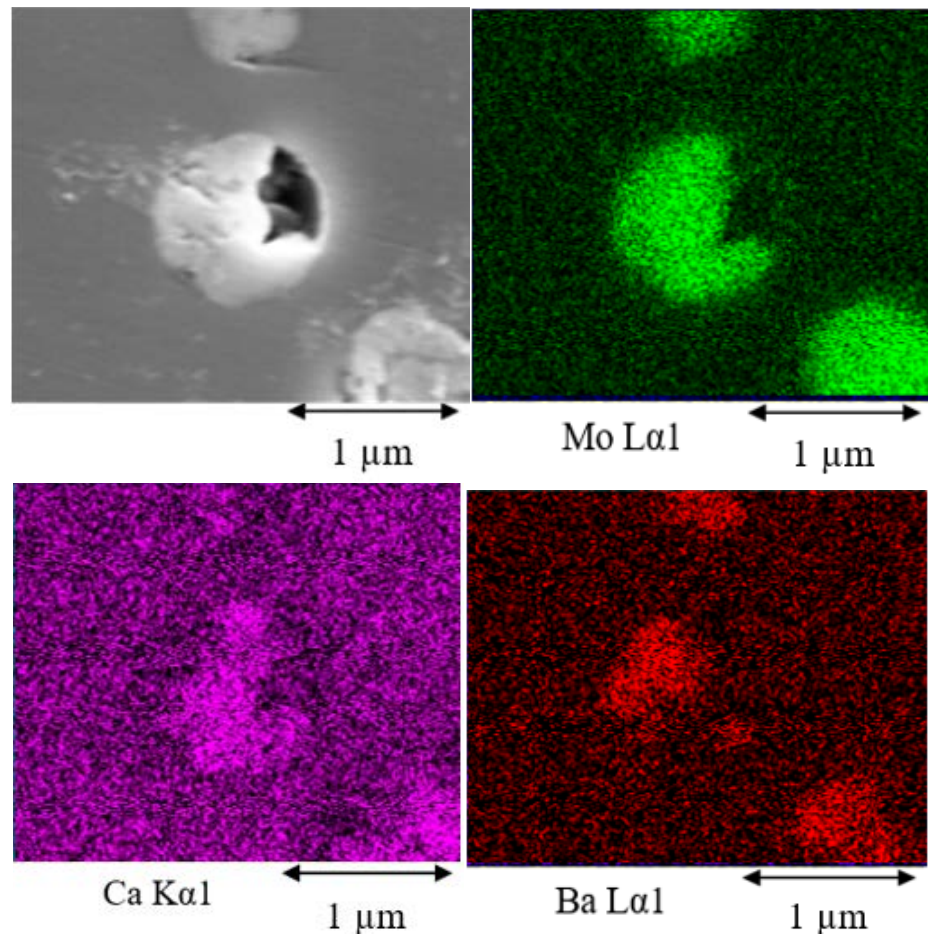


Figure 2.13. Electron image and Mo, Ba, and Ca X-ray maps collected of a molybdate particle formed in a sample quenched at 3.4°C/s . The particle is separated into a Ba-rich crystal and a Ca-rich crystal.

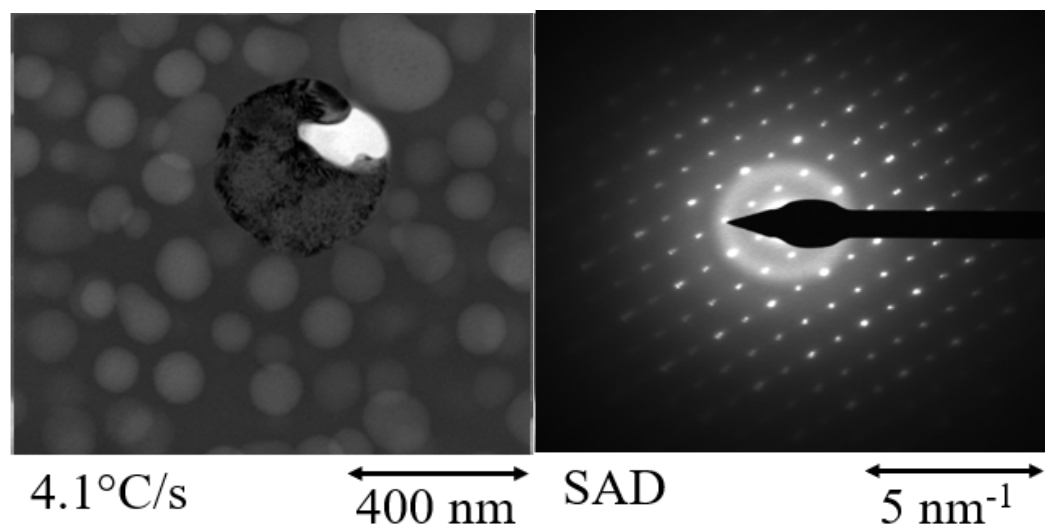


Figure 2.14. TEM electron image and selected area diffraction (SAD) pattern of a Mo-rich particle in the 4.1°C/s sample showing diffraction pattern.

2.3.3.2. Phases Identified in the Isothermal Samples. Similar phases were identified in the samples from the isothermal quench experiments. For example, Figure 2.15 compares the XRD patterns collected from the as-received glass, the glass that was quenched at $\sim 300^\circ\text{C/s}$ directly from 1300°C to room temperature water, and the samples quenched from 1300°C to 900°C , held for various times, and then quenched in room temperature water. An XRD pattern from a sample melted at 1300°C and held isothermally at 1100°C for 60 min before water quenching is also shown for comparison. The isothermal samples were also doped with 10% SRM 674 standard rutile for the quantitative x-ray analysis. The pattern from the as-received sample has peaks assigned to the $\text{Sr}_{0.44}\text{La}_{0.39}(\text{MoO}_4)$ (PDF# 04-002-4182) phase that do not appear to be present in the pattern from the “baseline” quenched sample; the pattern from this latter sample possesses only a broad, amorphous hump. Electron microscopy revealed that sub-micron Mo-rich droplets could still be detected in the rapidly quenched baseline sample, as shown in Figure A.5 of Appendix A. The difference in XRD patterns between the “as received” and “baseline” samples, and the smaller Mo-rich droplets in the baseline sample, discussed in Section 2.3.4.4, suggests that melting the samples at 1300°C for 90 seconds successfully “reset” the glass microstructure before the start of each isothermal experiment. There are peaks in the XRD patterns from the isothermal samples near 41° (2θ) that do not appear in the patterns from the continuously cooled samples (Figure 2.11), and this may be due to some contamination from the stainless-steel tubes used in the former experiments. Some Cr, Fe, and Ni from the steel was concentrated in the glasses to a depth of about $20\ \mu\text{m}$ from the tube wall, and it is possible that some transition metal oxide, like FeCr_2O_4 , has

contaminated isothermal samples after longer times at greater temperatures. Figures A.6 and A.7 in Appendix A provides additional information about this contamination.

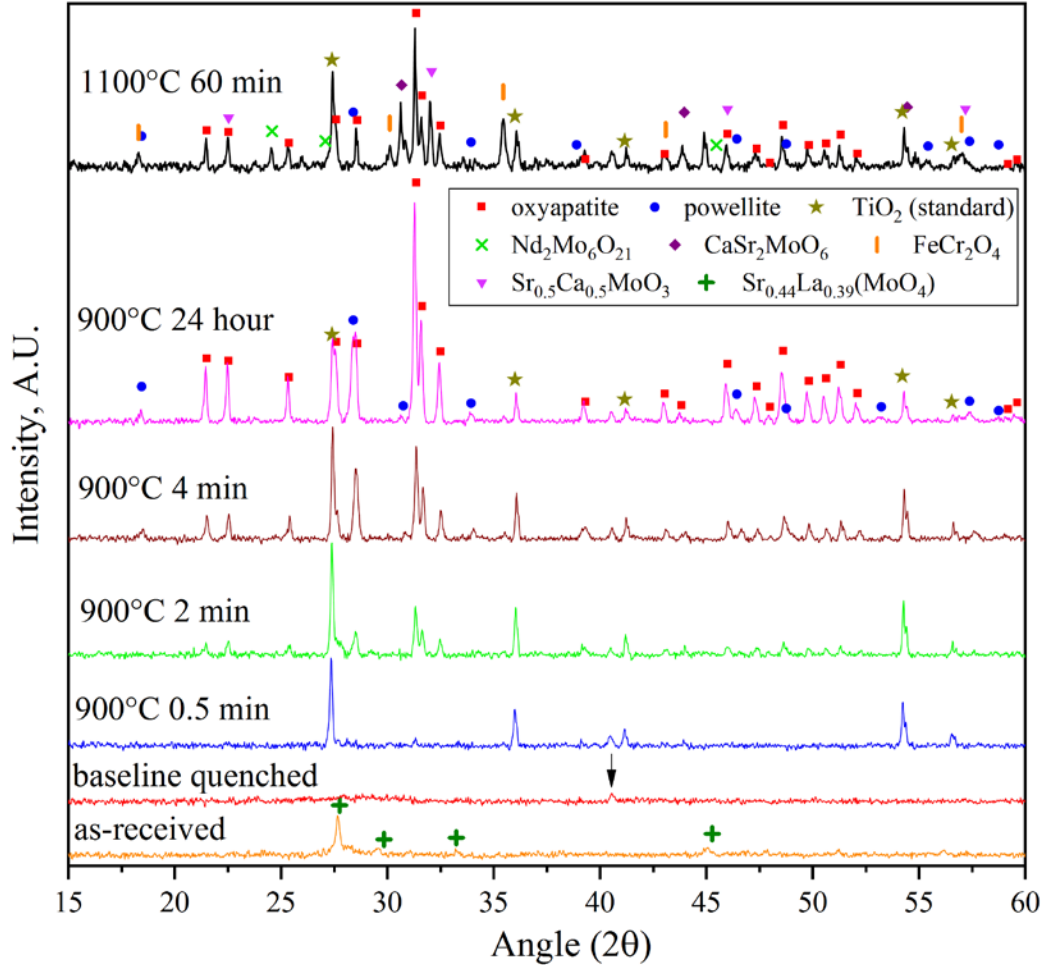


Figure 2.15. XRD patterns of as-received glass, baseline quenched sample (~300°C/s), samples heat-treated for various temperatures at 900°C, and heat treated at 1100°C for 60 min. The peak at 41° (2θ) marked with a black arrow is due to an unidentified phase that does not appear in the continuously cooled samples.

The first unambiguous new crystalline phase that formed in the isothermal samples is oxyapatite (e.g., $\text{Ca}_{1.1}\text{Nd}_{3.9}(\text{SiO}_4)_3\text{O}_{0.95}$, PDF# 04-007-5969). Oxyapatite was found after an isothermal hold for at least 1 minute at 900°C and is clearly present in the XRD pattern of the sample heat treated for 2 minutes at 900°C, as shown in Figure 2.15. Crystalline

powellite (e.g., CaMoO₄, PDF# 04-013-6764) was detected in the isothermal samples held for 4 minutes at 900°C, as indicated by the peak at 18.5° (2θ).

The XRD patterns from samples heat treated at 1100°C reveal several different peaks than what were found in the patterns of samples held at 900°C. Oxyapatite (e.g., Ca_{1.1}Nd_{3.9}(SiO₄)₃O_{0.95}, PDF# 04-007-5969) was detected but crystalline powellite (e.g., CaMoO₄, PDF# 04-013-6764) was not. Instead, there is evidence for the presence of other Mo-rich phases, including CaSr₂MoO₆ (PDF# 00-048-0800, 30.625° 2θ) and Sr_{0.5}Ca_{0.5}MoO₃ phase (PDF# 04-021-8196, 32° 2θ). Needle shaped Mo-rich crystals were found in samples held at 1100°C, and these needles were also rich in lanthanides, as shown in Figure 2.16. The peak at 24.5° 2θ in the XRD pattern from the sample heated at 1100°C for 60 minutes could indicate the presence of Nd₂Mo₆O₂₁ (PDF# 00-032-0675).

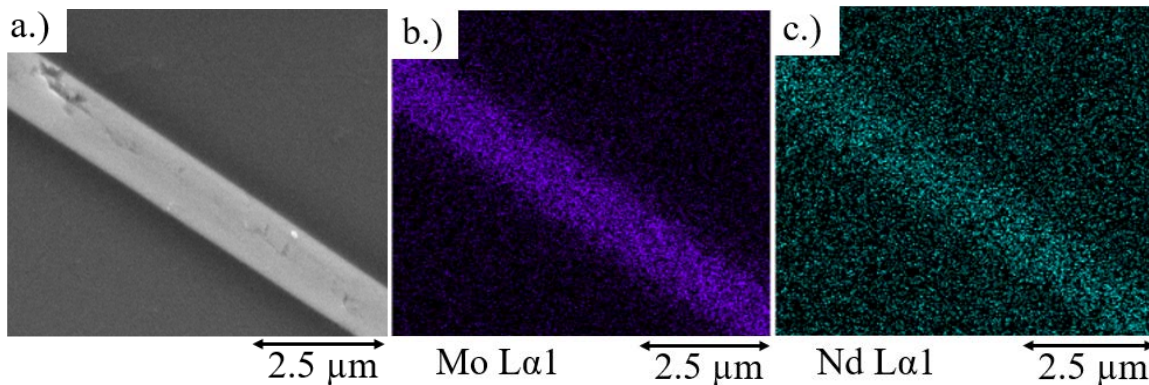


Figure 2.16. a) SEM electron image of the sample heat treated at 1100°C for 60 min of a Mo-rich phase also containing lanthanides, b) Mo EDS map, and c) Nd EDS map.

2.3.4. PHASE MORPHOLOGY AND COMPOSITION

Analytical SEM, TEM, and Raman spectroscopy were used to characterize the morphology and composition of the various phases that form in samples with different

thermal histories. In the following sections, the effects of heat treatment on the microstructures and compositions of the residual glass, oxyapatite, and molybdate phases are described.

2.

3.4.1. Residual Glass Morphology and Composition. The complex borosilicate glass studied here formed a droplet-in-matrix phase separated morphology. For example, Figure 2.17a shows a TEM image of a roller quenched sample with droplets $<0.1\text{ }\mu\text{m}$ in diameter. A similar morphology exists in the wedge mold sample cooled at 4.1°C/s shown in Figure 2.17b, although the droplets in this sample were larger ($0.13 \pm 0.03\text{ }\mu\text{m}$) than in the roller-quenched sample. Figure 2.17c shows an SEM image of the microstructure of a sample quenched at 3.4°C/s , where larger Mo-rich droplets (light phase) can be seen in the phase separated residual glass. The droplets in this residual glass were smaller, $0.05 \pm 0.02\text{ }\mu\text{m}$.

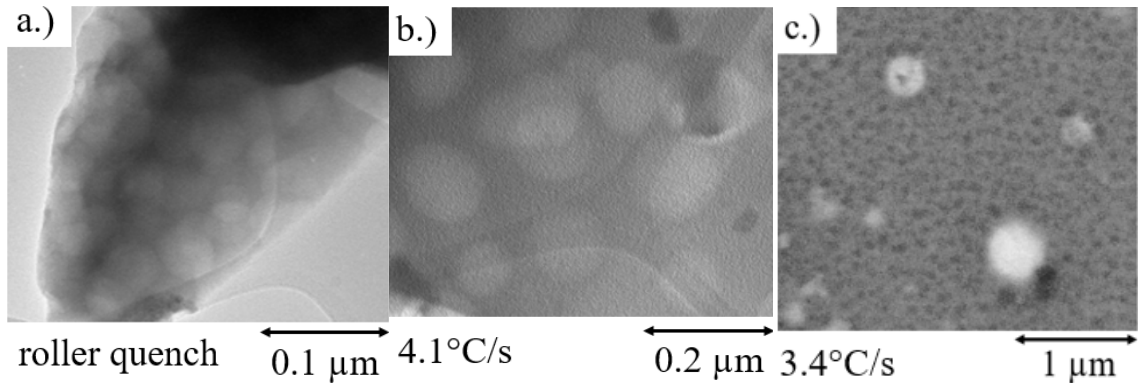


Figure 2.17. Electron micrographs of the PNNL glass after different continuous cooling experiments. a) TEM micrograph of the roller quench. b) TEM micrograph of sample cooled at 4.1°C/s . c) SEM micrograph of sample cooled at 3.4°C/s .

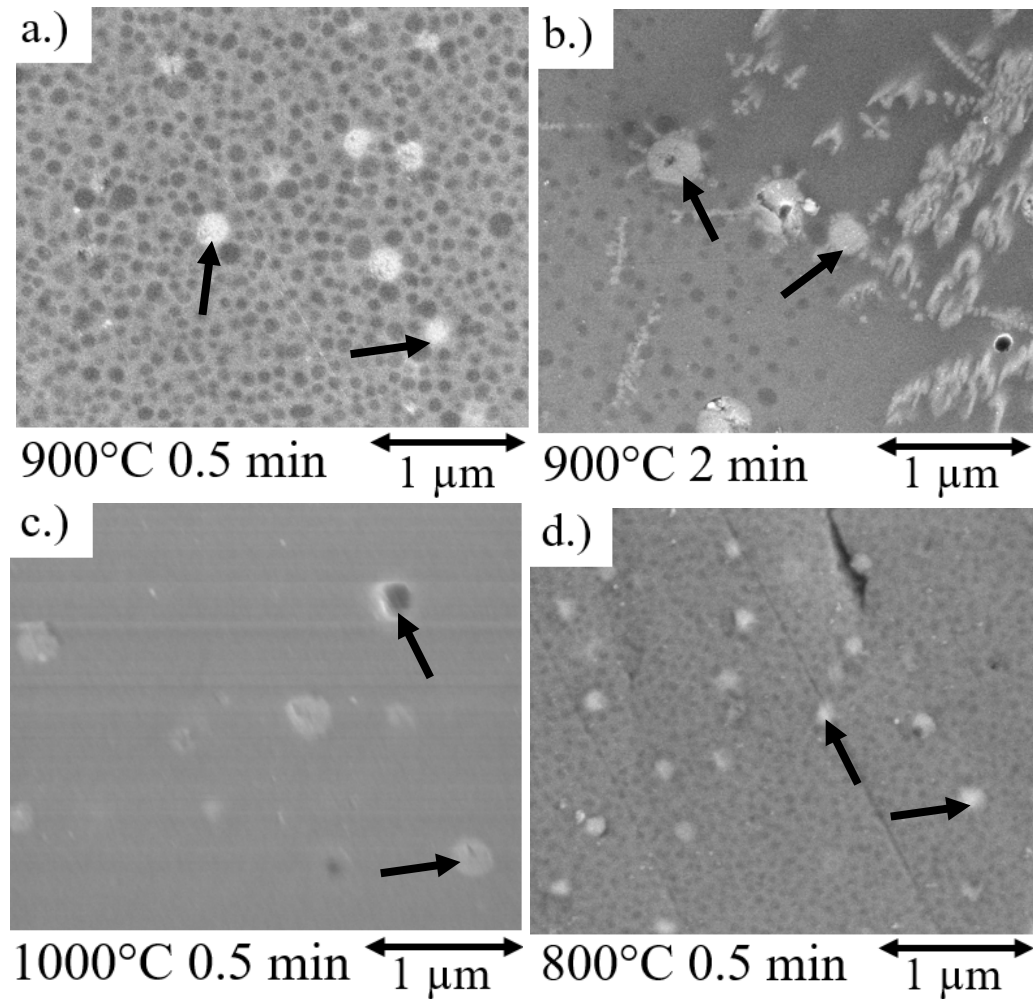


Figure 18. SEM micrographs of the a) 900°C isothermal sample after 0.5 min, b) 900°C isothermal sample after 2 min, c) 1000°C isothermal sample after 0.5 min, and d) 900°C isothermal sample after 0.5 min. Arrows show some of the Mo-rich droplets that have formed in these samples.

Figure 2.18 shows scanning electron micrographs from samples held isothermally at 900°C for 0.5 minutes (a) and 2 minutes (b). The former image shows Mo-rich droplets (arrows) embedded in the phase-separated residual glass matrix, similar to what was found for samples from the continuous-cooling experiments. After two minutes at 900°C, crystalline oxyapatite is evident and confirmed by XRD (Figure 2.14). It is noteworthy that in regions where the oxyapatite crystals have formed, the residual glassy phase appears to be much more homogeneous than the phase-separated morphology that is evident in

regions where the oxyapatite crystals have not yet formed. Figure 2.18c and d are SEM images from samples held for 0.5 min at 1000°C and 800°C, respectively, clearly showing that the size of the phase separated droplets is much smaller, if existant at all, in these samples.

EDS analysis of the TEM sample in Figure 2.17b indicates that the droplets are relatively rich in Al and Si, whereas the matrix phase is enriched with Ca, Zr, Mo, Cs, Ba, and the rare earth elements (Table 2.4). B and Li could not be detected by this EDS system and Sr and Rb had overlapping peaks but, Crum et al. [12] have shown that these elements are also concentrated in the matrix region of the phase separated glass. Crum et al. also found Cs to be more concentrated in the droplet phase. The growth of the oxyapatite crystals alters the composition of the surrounding glass by sequestering ions like Ca and the rare earths, and this appears to allow the remaining borosilicate liquid to form a more homogeneous glass upon quenching.

Figure 2.19 shows that the average droplet size in the phase-separated residual glass increased from ~30 nm to about ~150 nm with decreasing quench rate (~300°C/s to 4.1°C/s), but then decreased for quench rates slower than ~4°C/s. No droplets could be detected in samples cooled 0.1°C/s and slower using SEM. Also shown is the droplet size ($0.14 \pm 0.05 \mu\text{m}$) found in the as-received glass near the surface (~1 mm) that was in contact with air upon being plate quenched. The residual glass in samples cooled at the slowest rates appears to be homogeneous. This change in glass morphology is believed to be associated with the growth of the oxyapatite crystals. Crum et al. [12] reports similar morphological changes in the residual glass and suggested that when oxyapatite crystallize from the lanthanide, alkaline-earth-silicate matrix, the Cs-alumino-borosilicate droplet

phase dissolves into the matrix to form a homogeneous liquid that quenches into a homogeneous glass.

Table 2.4. Metal atom compositions of the different phases in the phase-separated residual glass of the 4.1°C/s sample, determined by TEM EDS.

Element	Droplet [at. %]	Matrix [at. %]
Na	1	2 ± 1
Al	15 ± 3	3 ± 2
Si	70 ± 9	42 ± 18
Ca	1	8 ± 4
Y	1 ± 1	2 ± 1
Zr	4 ± 2	5 ± 2
Mo	1 ± 1	5 ± 2
Ru	-	-
Rh	-	-
Te	1 ± 1	2 ± 2
Cs	1 ± 1	1 ± 1
Ba	-	3 ± 2
La	1 ± 1	3 ± 2
Ce	2 ± 1	6 ± 3
Pr	-	4 ± 2
Nd	2 ± 2	10 ± 5
Sm	1 ± 1	3 ± 2

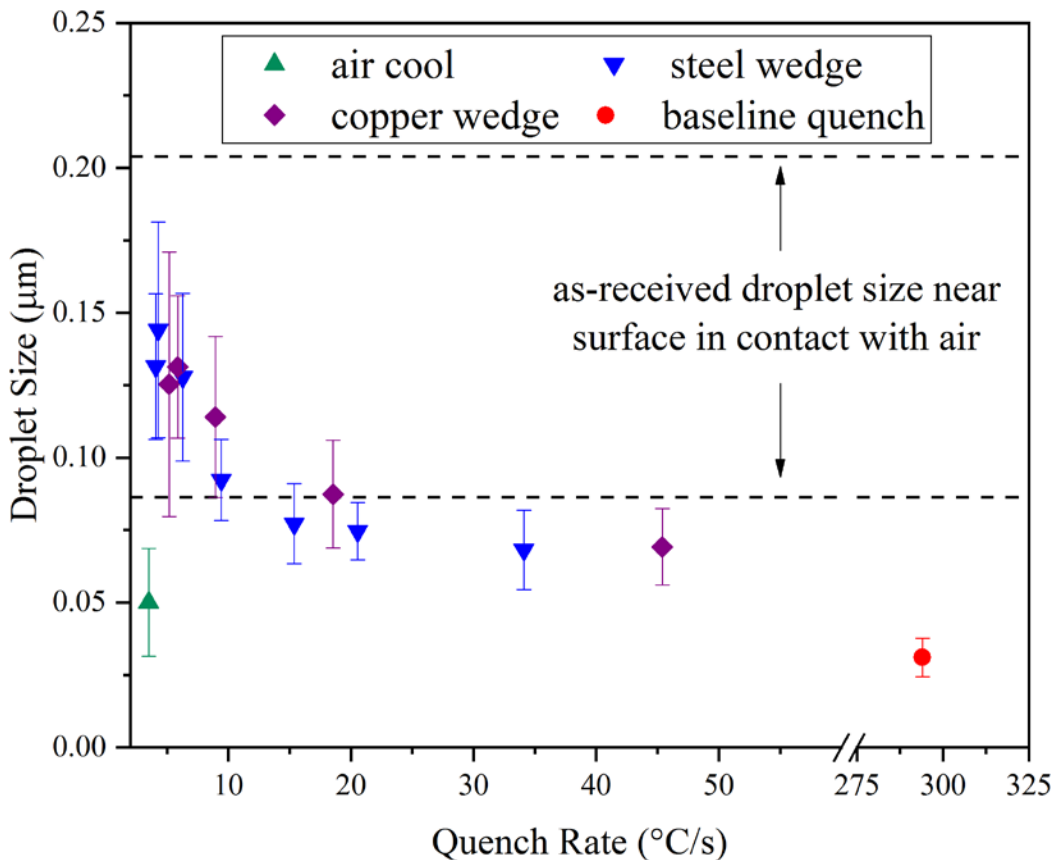


Figure 19. Phase-separated droplet size in the PNNL glass quenched at different rates.

2.3.4.2. Oxyapatite Morphology and Composition. The morphology of the oxyapatite crystals depends on thermal history. No oxyapatite crystals formed in wedge-mold samples cooled faster than 4.1°C/s (step one, steel wedge), nor in isothermal samples held at 600°C, or for short times (<1-2 minutes) at higher temperatures.

There does not appear to be differences in the peak positions from the XRD patterns, Figure 2.11, and 2.15, which are assigned to oxyapatite. These peaks are assigned to oxyapatite with an expected stoichiometry of $\text{Ca}_{1.1}\text{Nd}_{3.9}\text{Si}_3\text{O}_{12.95}$ that belongs to the hexagonal crystal system and P63/m space group (#176).

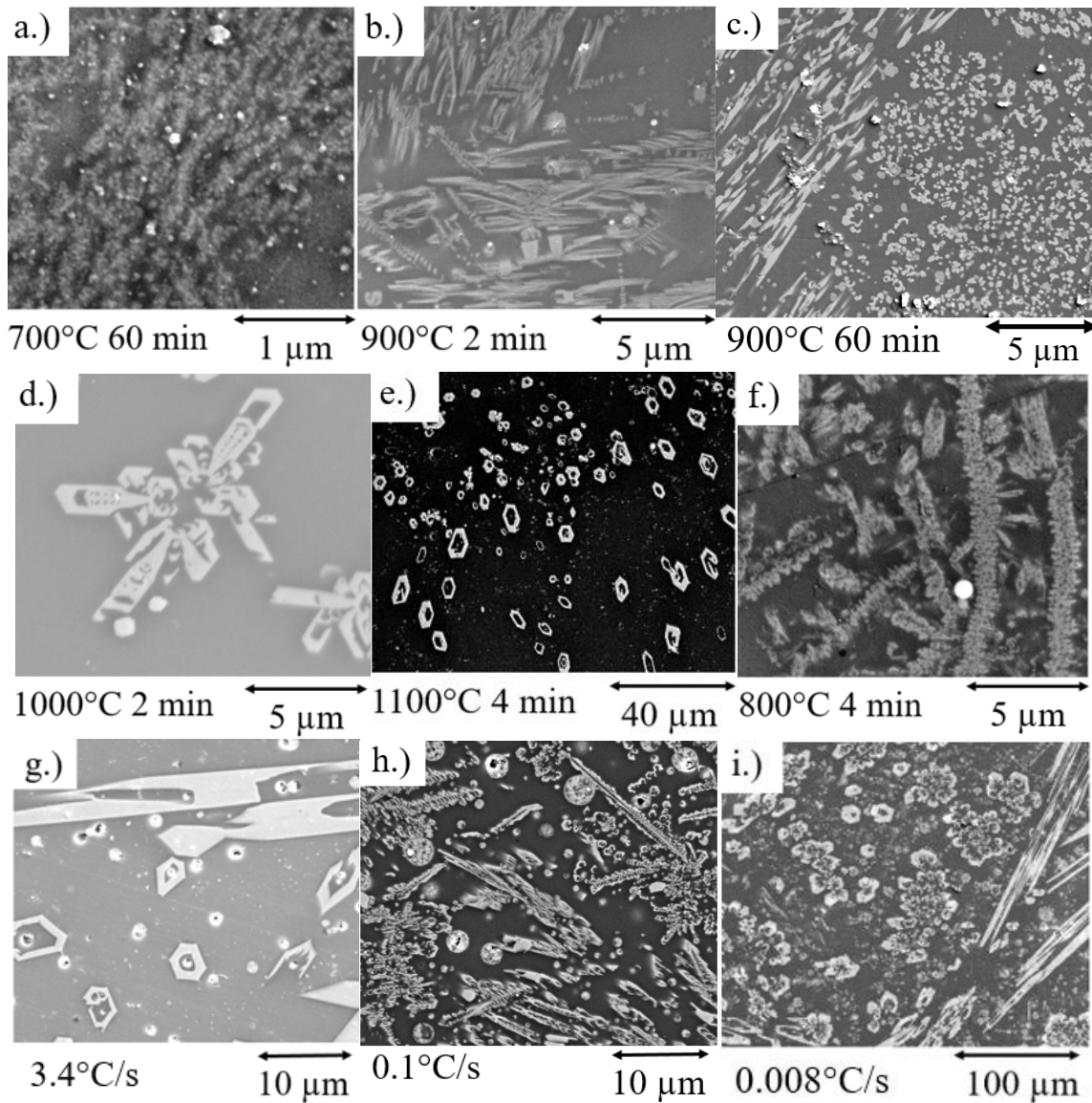


Figure 20. SEM micrograph of oxyapatite in samples heat treated at a) 1100°C for 4 min, b) 1000°C for 2 min, c) 900°C for 2 min, d) 800°C for 4 min, e) 700°C for 60 min, f) 900°C for 60 min, g) 3.4°C/s , h) 0.1°C/s and i) 0.008°C/s .

The SEM images and optical micrographs in Figure 2.20 show representative oxyapatite morphologies for a select number of the heat treatment conditions. For example, at 700°C , oxyapatite forms very small, $<1\text{ }\mu\text{m}$, crystals that cluster in dendritic formations (Figure 20a). At 900°C , oxyapatite forms the hollow hexagonal particles shown in Figure 2.20b, that over time form larger, seemingly solid hexagonal crystals, Figure 2.20c. The

crystals are also more dispersed, more defined, and less dendritic at higher temperatures, as shown Figure 2.20d, e, and f. Crum et al. described this hollow hexagonal form as a Hopper morphology and found that over time, the crystals grow inward [14]. At 1100°C, the hollow hexagonal oxyapatite crystals appear to be, in some cases, an order of magnitude larger than the crystals that form at 1000°C, Figure 2.20d.

The morphology of the oxyapatite crystals that formed in the continuous cooling samples were consistent with the morphologies of the crystals in the isothermal samples. The samples cooled at 3.4°C/s had dispersed oxyapatite crystals (Figure 2.20g) that were similar in size and morphology to samples held isothermally at 900°C or 1000°C for short (≤ 2 min) times (Figure 2.20b, d). The samples cooled at 0.1°C/s developed uniformly dispersed collections of crystals (Figure 2.20h), which grew to larger sizes in the sample cooled under the CCC-conditions (Figure 2.20i).

The fractions of oxyapatite crystals in the isothermal samples were determined from the quantitative XRD measurements and this information is summarized in the time-temperature-transformation diagram in Figure 2.21. Heat treatment conditions where oxyapatite was not detected are represented by the unfilled squares, and the black lines represent the conditions where the indicated fractions of oxyapatite were detected. The maximum fraction of oxyapatite (28.5 wt.%) was found in the sample held at 900°C for 24 hours. The solid blue lines represent the continuous cooling curves for several samples. It is interesting that the amount of oxyapatite was lower in the 0.008°C/s (15.2 wt.%) than the 0.1°C/s (21.4 wt.%). Asmussen et al. reported samples with similar compositions cooled along the CCC profile to contain 18.6 wt.% oxyapatite [16]. The 3.4°C/s sample does not appear to cross the region where oxyapatite formed in the isothermal experiments,

however the CCT curves are shifted to longer times and lower temperatures when compared to TTT curves.

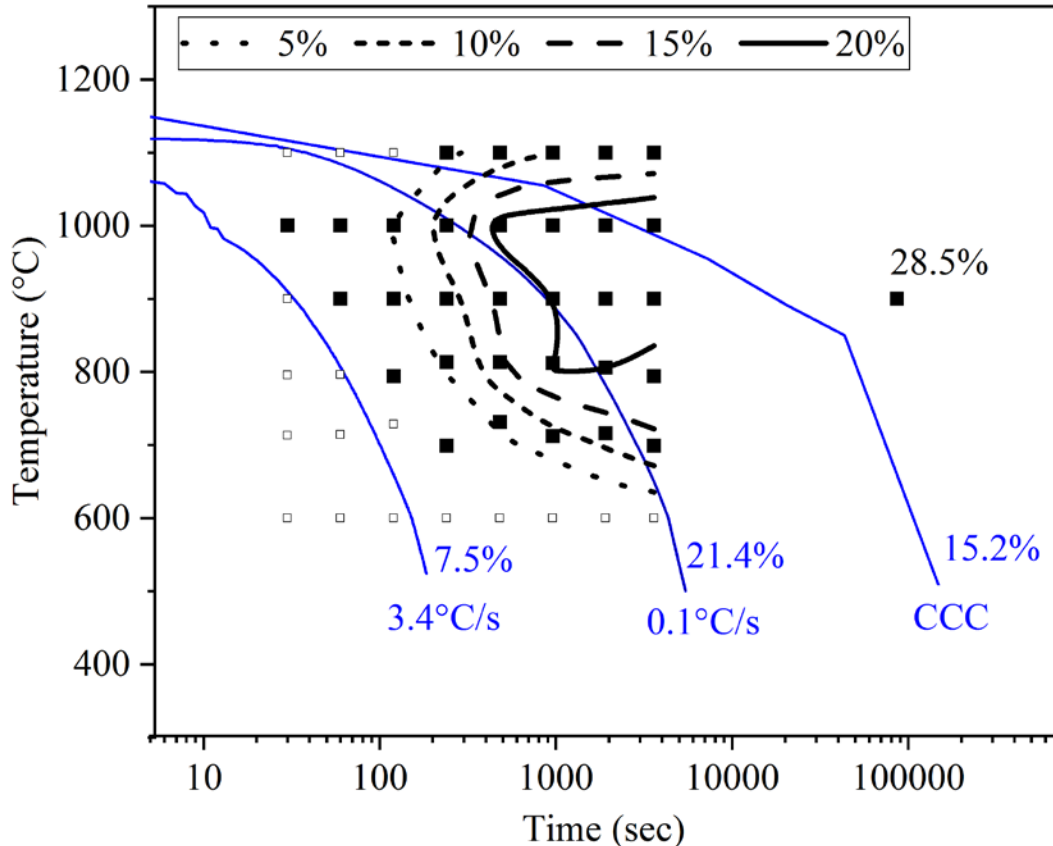


Figure 2.21. TTT and CCT diagrams for oxyapatite formation.

In some instances, the oxyapatite crystals were found to grow outward from a single point. For example, Figure 2.22 shows crystals that have grown from an area with a group of RuO_2 crystals. This might explain the differences in morphologies of oxyapatite across a sample. Shown in Figure 2.20c, the oxyapatite in the left side of the image are found elongated in similar directions verse the oxyapatite on the right which are mostly equal in all directions. Chouard et. al found that RuO_2 crystals act as nucleating agents for the oxyapatite crystals [21].

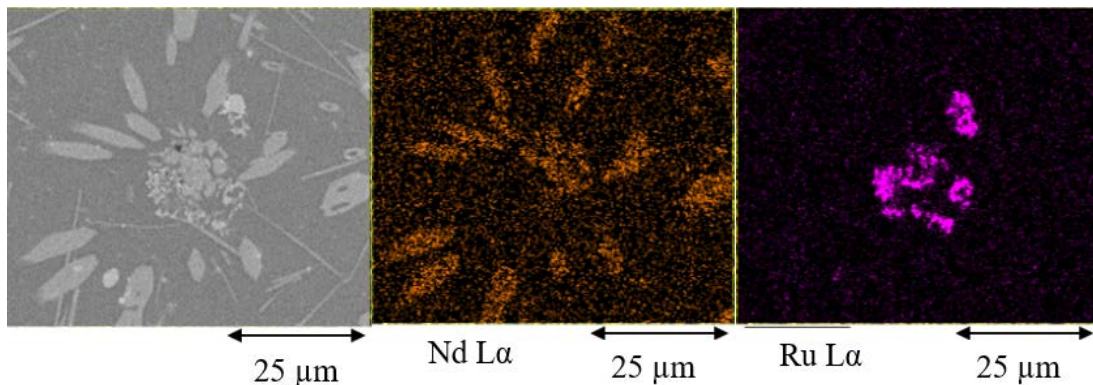


Figure 2.22. SEM micrograph and EDS maps showing the outward growth of oxyapatite from a RuO_2 crystal cluster in the sample heat treated at 1100°C for 32 minutes.

The compositions of oxyapatite crystals formed in a variety of samples were determined using SEM EDS and are summarized in Table 2.5. The major components of these crystals include Si, Ca, and the lanthanide elements, mainly Nd and Ce. The compositions do not appear to differ significantly in samples with different thermal histories and morphologies. The composition of oxyapatite crystals described by Crum et al. is also listed in Table 5 [13]. The lanthanide (LN) to silicon and Ca/Si ratio for each heat treatment are similar. The measured LN/Si ratio of oxyapatite is slightly greater and the Ca/Si ratio is slightly lower than the expected ratio ($\text{Ca}_{1.1}\text{Nd}_{3.9}\text{Si}_3\text{O}_{19.95}$) determined from XRD pattern assignments.

Table 2.5. Cation fractions in oxyapatite crystals in samples with different heat treatments with oxygen normalized to 0, compared with those reported by Crum et al. [13].

	3.5°C/s [At.%]	CCC [At.%]	1100°C 60 min [At.%]	Crum et al. [13] [At.%]
Na	-	-	-	2 ± 3
Al	3 ± 1	-	1 ± 1	
Si	36 ± 5	37 ± 1	37 ± 1	39 ± 4
Ca	18 ± 2	15 ± 2	14 ± 2	18 ± 2
Rb	3 ± 3	1 ± 1	-	-
Sr	1 ± 1	1 ± 1	2 ± 2	1 ± 1
Y	1 ± 1	2	2 ± 3	2
Zr	1 ± 1	-	-	NR*
Mo	-	-	-	0
Ru	-	-	-	NR*
Rh	-	-	-	NR*
Te	-	-	-	NR*
Cs	1	1	1 ± 1	0
Ba	2 ± 1	-	2 ± 1	NR*
La	4 ± 2	4	5 ± 1	4 ± 2
Ce	9 ± 2	11 ± 1	9	11 ± 4
Pr	3 ± 2	6 ± 1	3	4
Nd	15 ± 2	19 ± 1	19	17 ± 7
Sm	3 ± 2	4 ± 1	3 ± 1	2
LN/Si	1.0 ± 0.2	1.2 ± 0.1	1.1	1.0 ± 0.4
Ca/Si	0.5 ± 0.1	0.4	0.4	0.5 ± 0.1

* NR = values not reported

3.4.3. Oxyapatite Crystallization Kinetics. Kinetic information for formation of oxyapatite was obtained from the quantitative XRD data on samples collected from the tin bath isothermal experiments. Crystal fractions were normalized by dividing the weight fraction of oxyapatite for an isothermal sample by the weight fraction of oxyapatite in a sample crystallized by holding it at 900°C for 24 hours, conditions believed to produce the maximum fraction of oxyapatite (ω_{\max}). Normalized weight fractions ($\alpha = \omega(t) / \omega_{\max}$) are plotted vs time for each temperature in Figure 2.23. Some samples were run multiple times to determine experimental variability, as represented by the relevant error bars. The fastest

oxyapatite crystallization rate was determined to be $\sim 1000^{\circ}\text{C}$, and rates decreased at 1100°C .

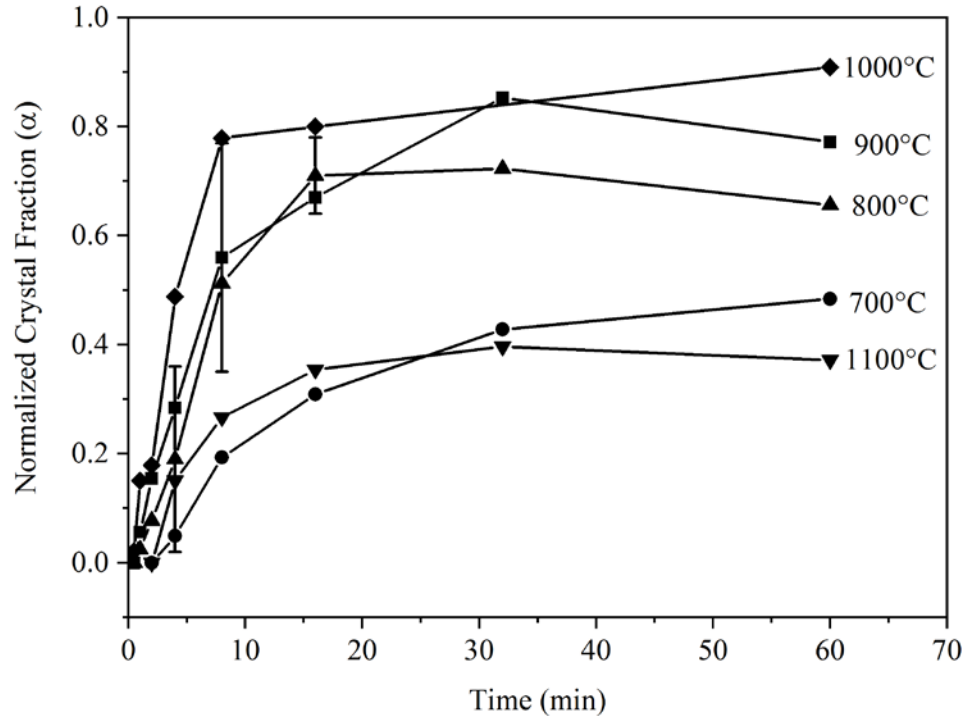


Figure 23. Crystal fraction of oxyapatite vs time for samples from the isothermal experiments.

The Johnson–Mehl–Avrami–Kolmogorov (JMAK) model was used to analyze the crystal growth kinetics [22, 23]. The classic JMAK equation is

$$\alpha_n(t) = 1 - e^{[-(k_n t)^n]} \quad (1)$$

where t is time, n relates to the different nucleation and growth mechanisms, and k_n is the dissolution constant [22]. k_n is related to temperature through the Arrhenius equation,

$$k_n = k_o e^{\frac{-E}{RT}} \quad (2)$$

where k_o is a frequency factor, E is the activation energy, R is the gas constant, and T is the absolute temperature. Equation (1) can be rewritten as

$$\ln[-\ln(\alpha_n(t))] = n \ln(k_n) + n \ln(t) \quad (3)$$

where the term $n \ln(k_n)$ is the intercept and n (Avrami exponent) is the slope of a double logarithmic $-\ln(1 - \alpha_n(t))$ versus t plot. Figure 2.24 shows the Avrami plots for oxyapatite formation between 700°C and 1000°C

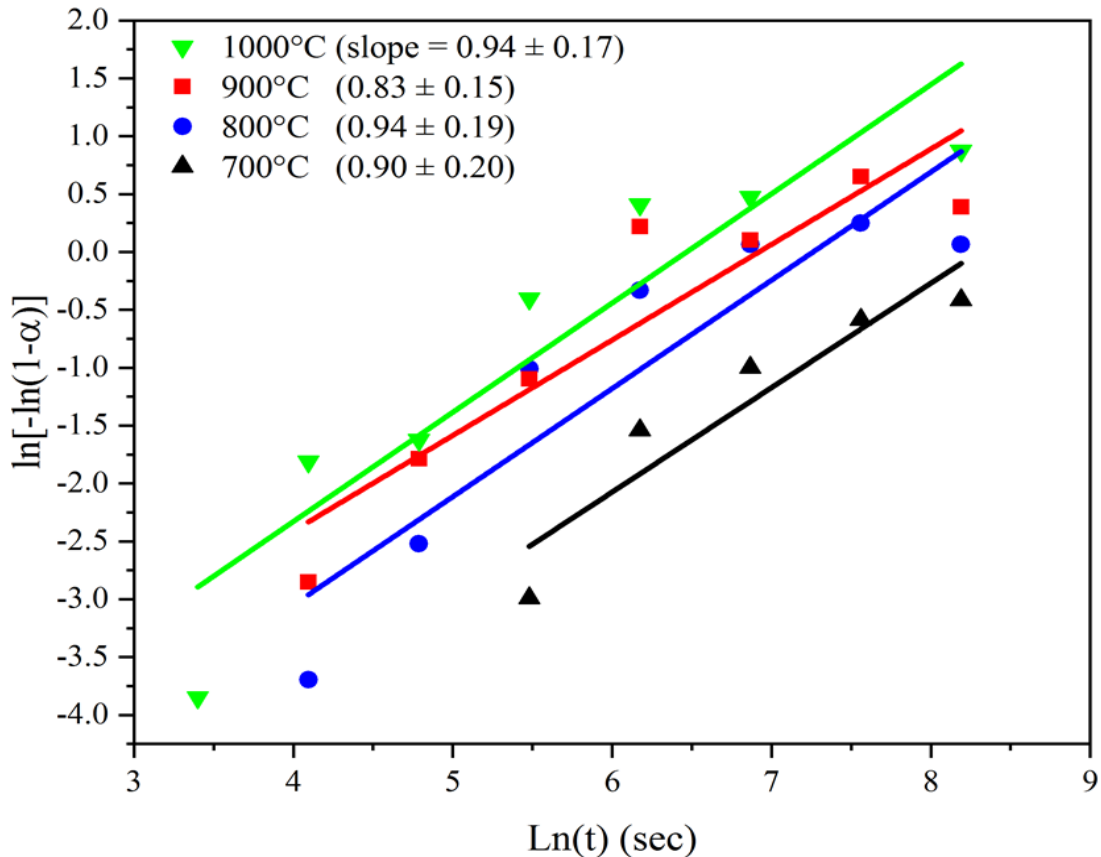


Figure 2.24. Avrami plot for oxyapatite isothermally held at various temperatures.

Equation (2) can be rewritten as

$$\ln(k_n) = \ln(k_0) - \frac{E}{RT} \quad (4)$$

where $-E/R$ is the slope of the $\ln(k_n)$ vs $1/T$ plot. The $\ln(k_n)$ values extracted from the slopes and intercepts from Figure 2.24 are plotted vs. $1000/T$ in Figure 2.25. A slope of -7.1 ± 1.0 was determined and this corresponds to an activation energy of 58.8 ± 8.16 kJ/mol. The Avrami exponent (n), the slopes of the Avrami plot (Figure 2.24), was calculated to be 0.9 ± 0.2 and this relates to the growth mechanism.

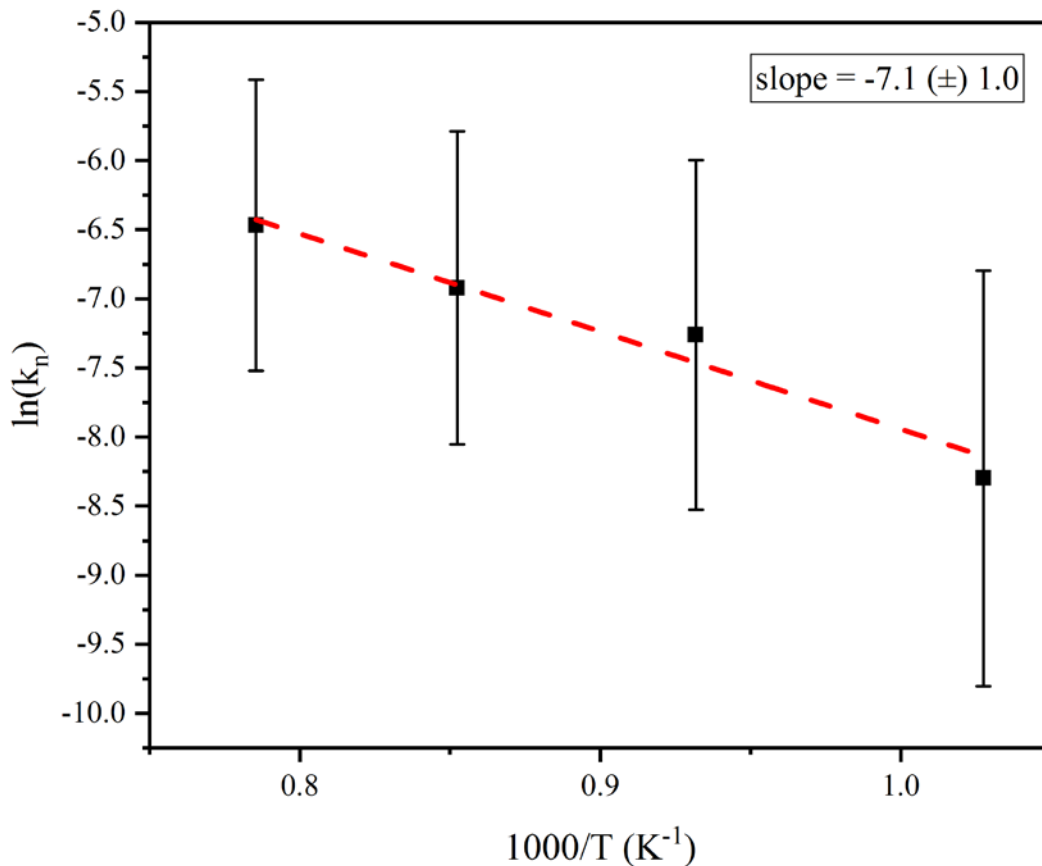


Figure 2.25. $\ln(k_n)$ vs $1000/T$ for oxyapatite growth from isothermal hold experiments between 700 and 1000°C.

Fournier et al., described JMAK studies of the growth of rare earth oxyapatite crystals ($\text{Ca}_2\text{RE}_8\text{Si}_6\text{O}_{26}$) in borosilicate glasses and reported similar n values, 0.65 [24] and 0.84 [25] which they related to diffusion-controlled crystallization mechanisms. However, they also reported much greater activation energies 475 ± 90 kJ/mol [24] and 496 ± 46 kJ/mol [25] for crystal growth. A greater activation energy means there is a much greater increase in crystal growth rates with increasing temperature. Fournier et al. used less complex glass compositions in their studies while our studies have found a change in the residual glass with temperature prior to the crystallization of oxyapatite, Figure 2.17a, c, and d. This change in the residual glass could influence diffusion properties effecting the

crystallization of oxyapatite causing a reduction in the activation energy. Crum found that residual glass phase separation affected the melt rheology increasing viscosity [14].

2.3.4.4. Mo-rich Phase Morphology and Composition. The formation of Mo-rich crystalline phases appears to be much more complex than the formation of crystalline oxyapatite. From x-ray diffraction, Section 2.3.3.1 and 2.3.3.2, there are many molybdate phases assigned to the various patterns. From the continuous cooling patterns peaks are assigned with $\text{Sr}_{0.44}\text{La}_{0.39}(\text{MoO}_4)$, CaMoO_4 , and $\text{Ba}_{0.75}\text{Sr}_{0.25}(\text{MoO}_4)$, in which all belong to the tetragonal crystal system I 4₁/a space group (#88). They also all have a 1:1 alkaline earth (AE) and lanthanide (LN) to molybdenum ratio. In the isothermal experiments preformed at 1100°C, there are assignments of a $\text{CaSr}_2\text{MoO}_6$, $\text{Sr}_{0.5}\text{Ca}_{0.5}\text{MoO}_3$, and $\text{Nd}_2\text{Mo}_6\text{O}_{21}$ belonging to different crystal systems and space groups.

In the continuous cooling experiments, spherical morphologies were found in all samples cooled at least 0.1°C/s; viz., Figures 2.26a and b. In the CCC sample, the Mo-rich droplets appear to be less spherical and many have formed cross-type morphologies, as shown in Figure 2.26c. Mo-rich droplets were not detected by SEM in the roller quenched glass but were found in the baseline quenched glass (~300°C/s, from the isothermal study) and the droplets grow with decreasing quench rate (Figure 2.27). The droplet size distributions were similar for samples from both the copper wedge mold and the steel wedge mold experiments. A large increase in the size of the Mo-rich droplets occurred for the air-cooled sample (3.4°C/s), which also had oxyapatite crystals. The isothermal experiments showed that oxyapatite would crystallize before crystalline powellite could be detected using XRD; e.g., Figure 2.15. The rapid growth of the Mo-rich phase after the

precipitation of the lanthanum silicate oxyapatite phase indicates that the lanthanide ions influence the solubility of Mo in the glass melt. This is discussed in more detail in Section 2.3.6.

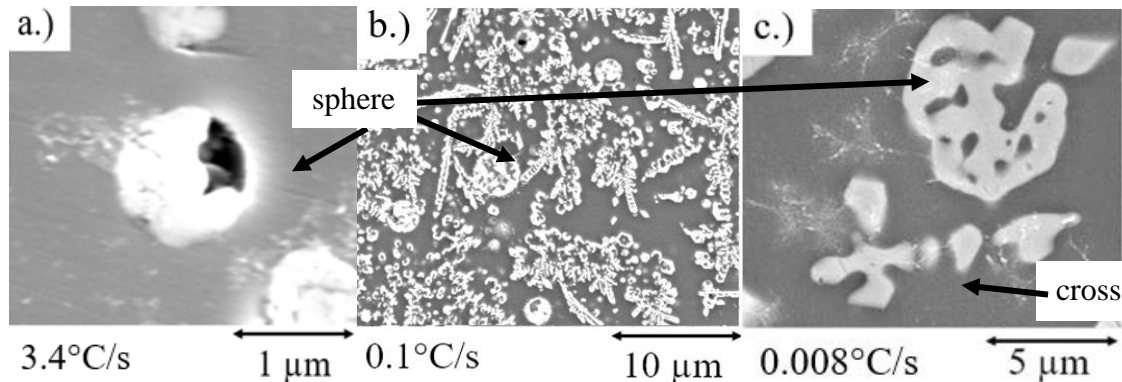


Figure 2.26. SEM images of powellite in samples continuously cooled a) 3.4°C/s , b) 0.1°C/s , and c) 0.008°C/s .

The Mo-rich droplets in the as-received glass sample were found to be $0.24 \pm 0.05 \mu\text{m}$ represented with the dashed horizontal lines shown in Figure 2.27. If these droplet sizes are an indication for the quench rate of a sample, then the estimated quench rate of the as received glasses likely falls in the range $10\text{-}25^{\circ}\text{C/s}$.

The isothermal experiments reveal that the morphology of the Mo-rich phases changes significantly with time and temperature. Figure 2.28 shows examples of Mo-rich phases with spherical, dendritic, cross-like, and needle-like morphologies, with the latter being more abundant at higher temperatures and longer times. The dendritic structures appear to grow between the aluminosilicate-rich droplets in the phase separated residual glass and, in many cases, the dendrites grow from the spherical droplets, as shown in Figure 28a. The Mo-rich crosses appear to form in the more homogenous residual glass and are not associated with other microstructural features; viz., Figure 2.28a. These cross-like structures are described by Delattre et al. [26] and Li et al. [27] as possessing perpendicular

arms of equal length. At 1100°C, where slow nucleation rates and fast crystal growth rates are expected, needles with lengths many orders of magnitude larger than the crosses and dendrites are the dominant Mo-rich phase (Figure 2.28b and c). Many powellite needles are oriented in similar directions or from similar points.

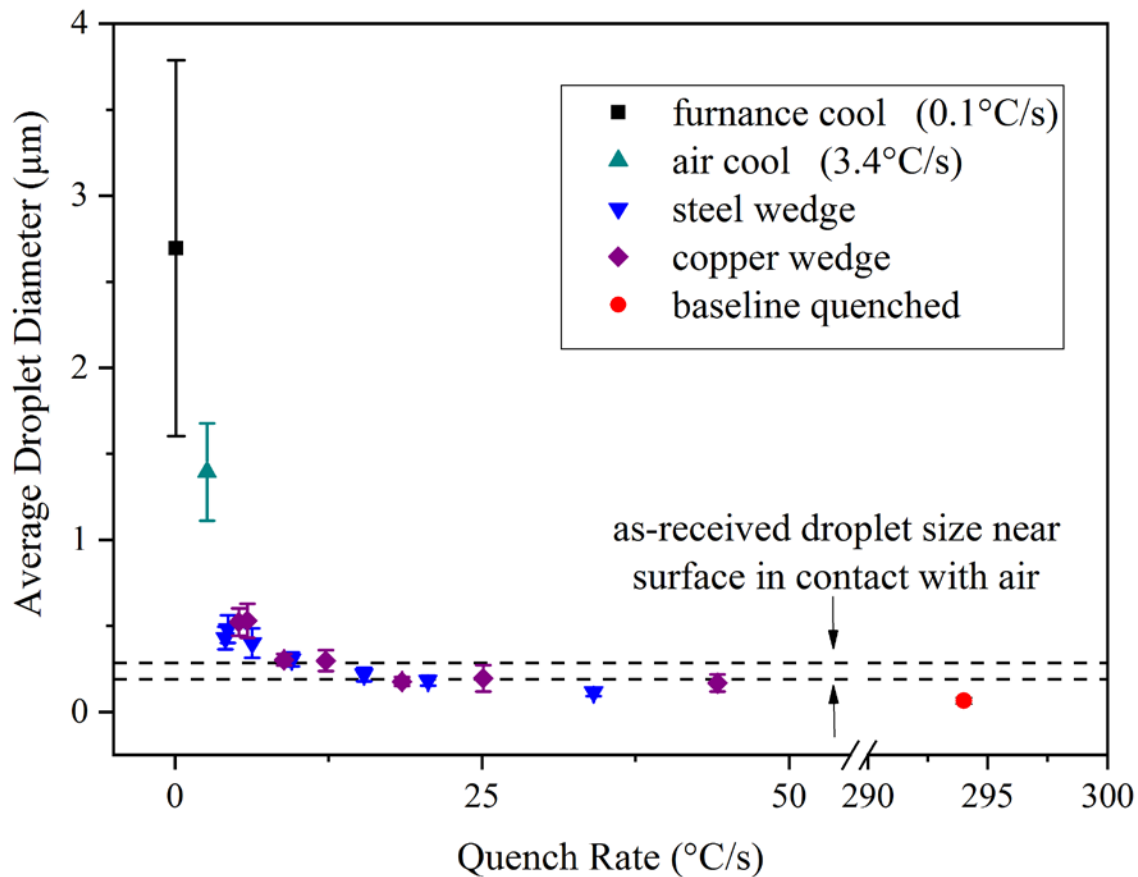


Figure 2.27. Molybdenum droplet diameter vs quench rate for samples from the continuous cooling experiments. The horizontal dashed lines represent the average diameter with one standard deviation of droplets from the as-received glass 1 mm from the surface that was in contact with air.

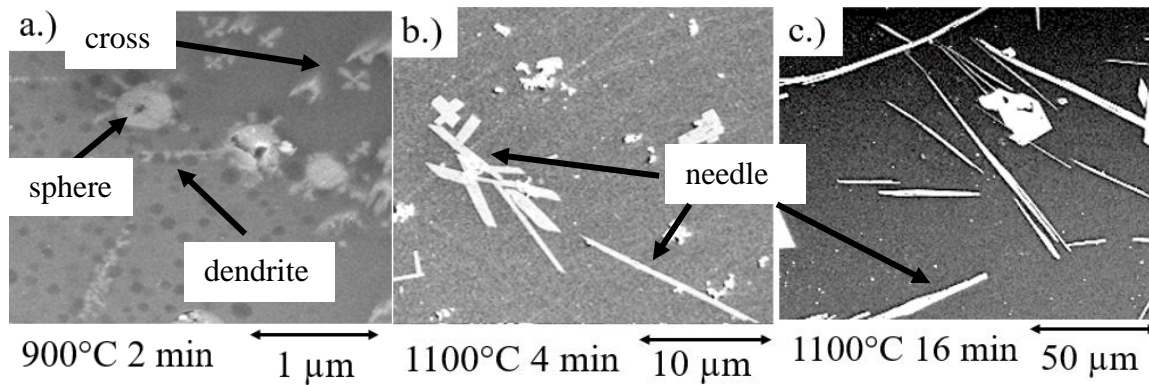


Figure 2.28. SEM images of Mo-rich phases in samples heat treated isothermally a) 900°C for 2 min, b) 1100°C for 4 min, c) 1100°C for 16 min.

The average compositions of different Mo-rich phases were determined using SEM EDS and TEM EDS and are summarized in Table 2.6. The spherical morphologies in samples prepared using quench rates faster than the CCC treatment separated into Ba-rich and Ca-rich molybdate crystals, as shown in Figure 2.13. Crum et al. found that the spherical molybdate particles separated into spherical clusters of small BaMoO_4 and CaMoO_4 crystals [12]. Later studies found these crystals to be based on two types of solid solutions, $\text{Ca}_x\text{Sr}_{1-x}\text{MoO}_4$ and $\text{Ca}_x\text{Ba}_{1-x}\text{MoO}_4$ [13, 16]. Crum et al. found similar compositions for these phases in samples with the same composition [13]. In the CCC sample, Ca/Sr-molybdate particles were found but no Ba/Sr-molybdate particles. The cross-like molybdates have similar compositions to the spheres, but the needles have significantly greater lanthanide concentrations, as discussed earlier with Figure 2.16. The alkaline earth (AE) and lanthanide (LN) to molybdenum concentrations are also listed in Table 2.6 and have a 1:1 ratio comparable to the expected ratios from the XRD pattern assignments.

Table 6. Cation fractions of Mo-rich crystals in samples with different heat treatments with oxygen normalized to 0, compared with those reported by Crum et al. [13].

Element	4.1°C/s (Ba,Sr,Mo)	4.1°C/s (Ca,Sr,Mo)	CCC (cross)	CCC (droplet)	1100°C 60 min (needle)	Crum et al. [13]
	TEM [At.%]	TEM [At.%]	SEM [At.%]	SEM [At.%]	SEM [At.%]	SEM [At.%]
Na	10 ± 2	6 ± 1	4 ± 1	3 ± 1	1 ± 1	3 ± 4
Al	4 ± 1	-	1 ± 1	2 ± 1	1 ± 1	-
Si	-	13 ± 2	5 ± 4	8 ± 5	31 ± 7	-
Ca	20 ± 2	27 ± 3	26 ± 5	27 ± 2	8 ± 2	33 ± 11
Rb	-	-	-	-	-	3 ± 4
Sr	7 ± 3	5 ± 3	7	7	1	8 ± 4
Y	-	-	-	-	-	-
Zr	-	-	1 ± 1	-	3 ± 1	NR*
Mo	41 ± 8	41 ± 10	46 ± 3	46 ± 5	36 ± 11	55 ± 7
Ru	-	-	1 ± 1	-	-	NR*
Rh	-	-	-	-	-	NR*
Te	-	-	1 ± 1	1 ± 1	-	NR*
Cs	-	-	-	1 ± 1	-	-
Ba	14 ± 5	2 ± 3	1 ± 1	1	1	NR*
La	-	-	1	1 ± 1	3 ± 1	-
Ce	-	2 ± 4	1	1 ± 1	5 ± 1	-
Pr	-	-	1 ± 1	0	2 ± 1	-
Nd	3 ± 3	3 ± 4	2	2 ± 1	6 ± 1	-
Sm	-	1 ± 4	1 ± 1	0	-	-
(AE+LN)/Mo	1.0	1.1	0.9 ± 0.1	0.9 ± 0.2	0.8 ± 0.4	0.7 ± 0.2

* NR = values not reported

A TTT/CCT diagram for the formation of the molybdate phases is shown in Figure 2.29. At low temperatures, <900°C, molybdates are only found as spheres. These spheres form before the crystallization of oxyapatite and this is discussed later in Section 3.6. At temperatures in the range from 900°C to 1000°C, Mo-rich phases with spherical, dendritic, cross-like, and needle-like morphologies have been found, whereas at 1100°C, the molybdate phases were found to have spherical and needle morphologies.

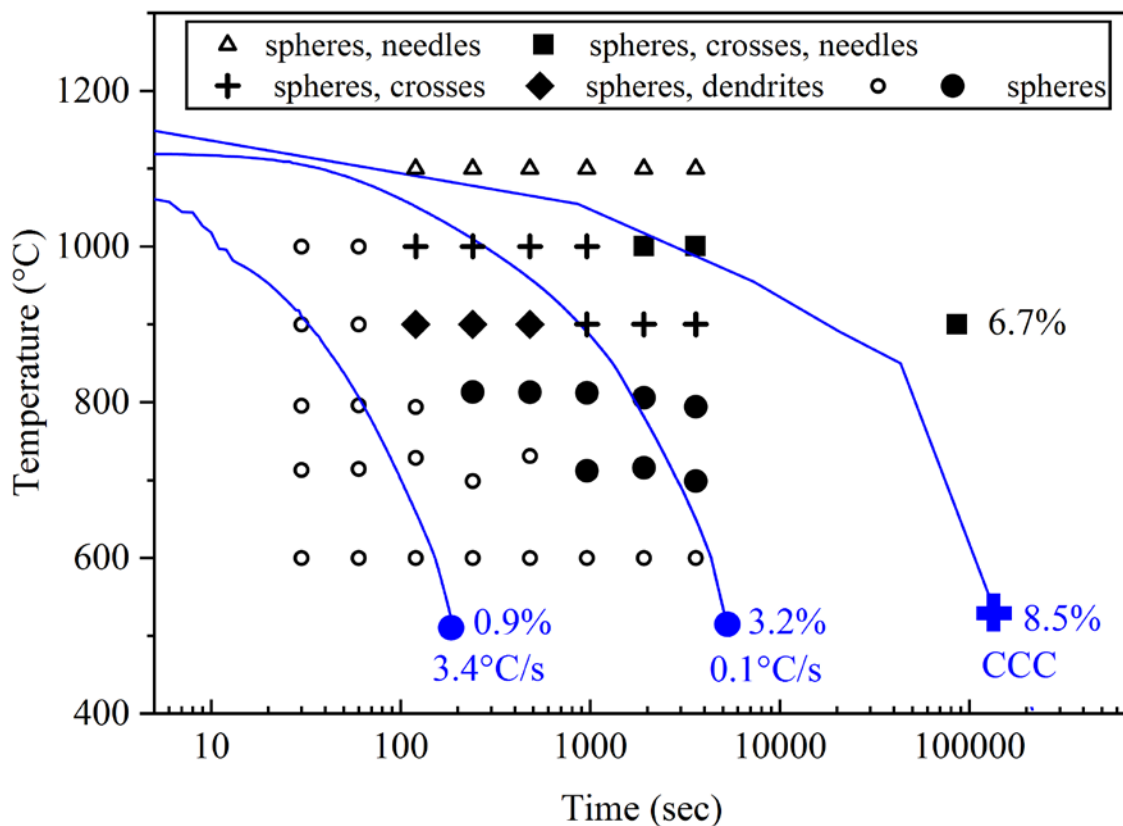


Figure 2.29. TTT (black symbols) and CCT (blue lines) for the formation of different Mo-rich crystals. The open symbols indicate x-ray amorphous Mo-rich phases. The morphologies were determined using SEM and the crystal fractions were determined by quantitative XRD.

The results of the continuous cooling experiments align with those from the isothermal experiments. In the samples cooled at 3.4°C/s and 0.1°C/s, spherical powellite particles were found with sizes and compositions similar to those found in isothermal samples held at $\leq 900^{\circ}\text{C}$. No needles, dendrites, or crosses were found in the 3.4°C/s and 0.1°C/s samples because they were cooled quickly through the temperature ranges where those morphologies developed in the isothermal samples (Figure 2.29). In the sample cooled according to the CCC profile (0.008°C/s), cross-type powellite was the dominant phase, with some droplets. This is consistent with the longer times spent at temperatures where cross-type powellite was found to form in the isothermal experiments. The variations

in the compositions and morphologies of the Mo-rich phases with temperature invalidated the use of the JMAK kinetic analysis, as discussed in Appendix A.

Raman spectroscopy was useful to differentiate the various Mo-rich phases and confirmed that the needles have different structures and compositions than the spheres and crosses. Figure 2.30 shows representative Raman spectra of the various morphological features found in these samples. Figure 2.31 shows optical micrographs the features from where the Raman spectra were collected. The peaks in the residual glass spectrum correlate to a symmetric internal $[\text{MoO}_4]^{2-}$ tetrahedral Mo-O stretching mode (910 cm^{-1}) [28, 29] and another internal $[\text{MoO}_4]^{2-}$ tetrahedral O-Mo-O bending mode (320 cm^{-1}) [29, 30]. The peaks from spectra of the oxyapatite crystals correlate to $[\text{SiO}_4]^{4-}$ tetrahedral stretching (860 cm^{-1}) and bending modes (529 cm^{-1} and 400 cm^{-1}), as well as a La-O stretching mode (298 cm^{-1}) [31]. The spectra from the cross and droplet features contain peaks assigned to internal $[\text{MoO}_4]^{2-}$ tetrahedral vibrational modes ($324, 385, 795, 843$, and 880 cm^{-1}) and external (vibrations between the lattices) $[\text{MoO}_4]^{2-}$ tetrahedral rotational modes (197 cm^{-1}) [32]. The spectra collected from the Mo-rich needles are likely due to a reduced molybdate, e.g., MoO_2 .

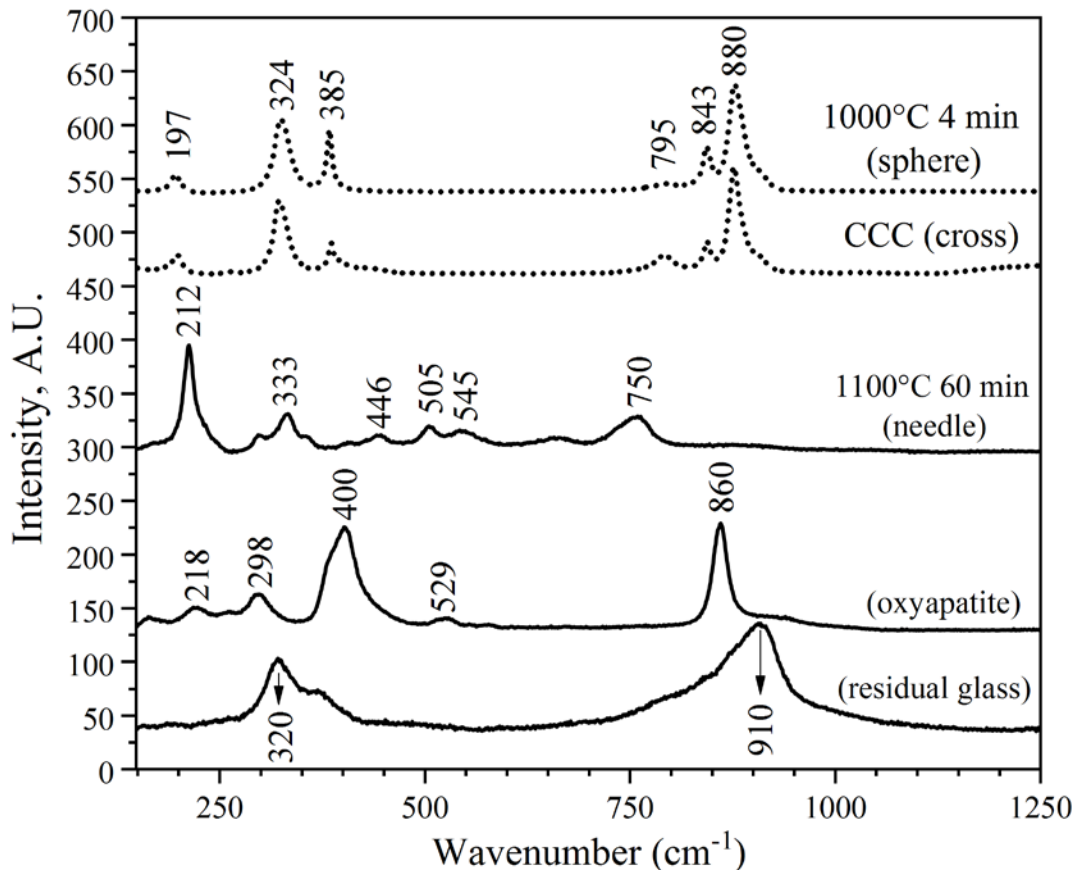


Figure 2.30. μ -Raman spectra of (dashed lines) a powellite sphere and cross in the sample heat treated at 1000°C for 4 min and (solid lines) spectra of a powellite needle, oxyapatite, and the residual glass in the 1100°C 60 min sample.

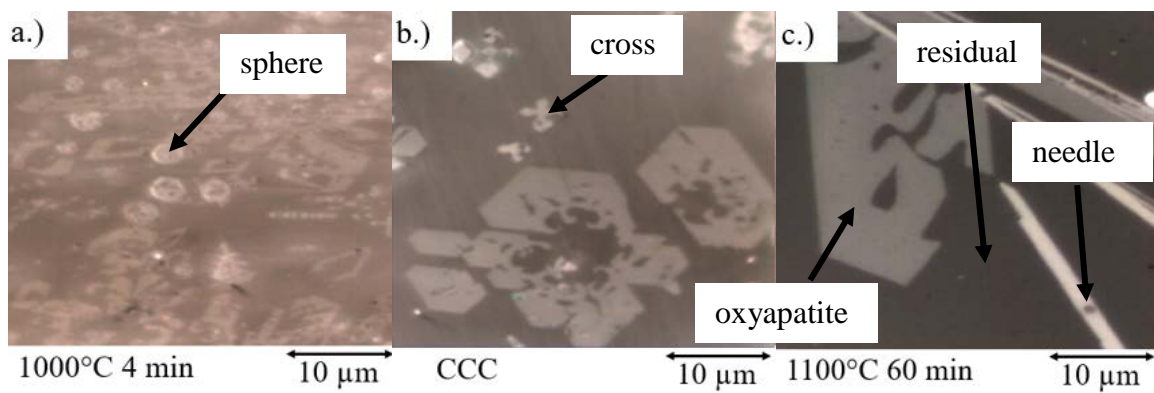


Figure 2.31. Micrographs showing locations of where the Raman spectra in Figure 30 was collected.

2.3.5. TIME-TEMPERATURE-TRANSFORMATION DIAGRAM

A Time-Temperature-Transformation, TTT, diagram that summarizes the overall crystallization of the borosilicate waste glass was developed using the information from the isothermal tin bath experiments and is shown in Figure 2.32. Scanning electron microscopy identified the morphologies of the phases present in each sample, which are indicated by the different symbols. Quantitative XRD only detected and identified two phases, oxyapatite, represented by the large symbols, and powellite, represented by large closed symbols. The quantitative XRD information was used to develop the lines that represent the overall crystallization fractions (5, 10, 20, and 25 wt. %), determined by linear interpolation of two neighboring measured data points and these are also plotted in Figure 2.32.

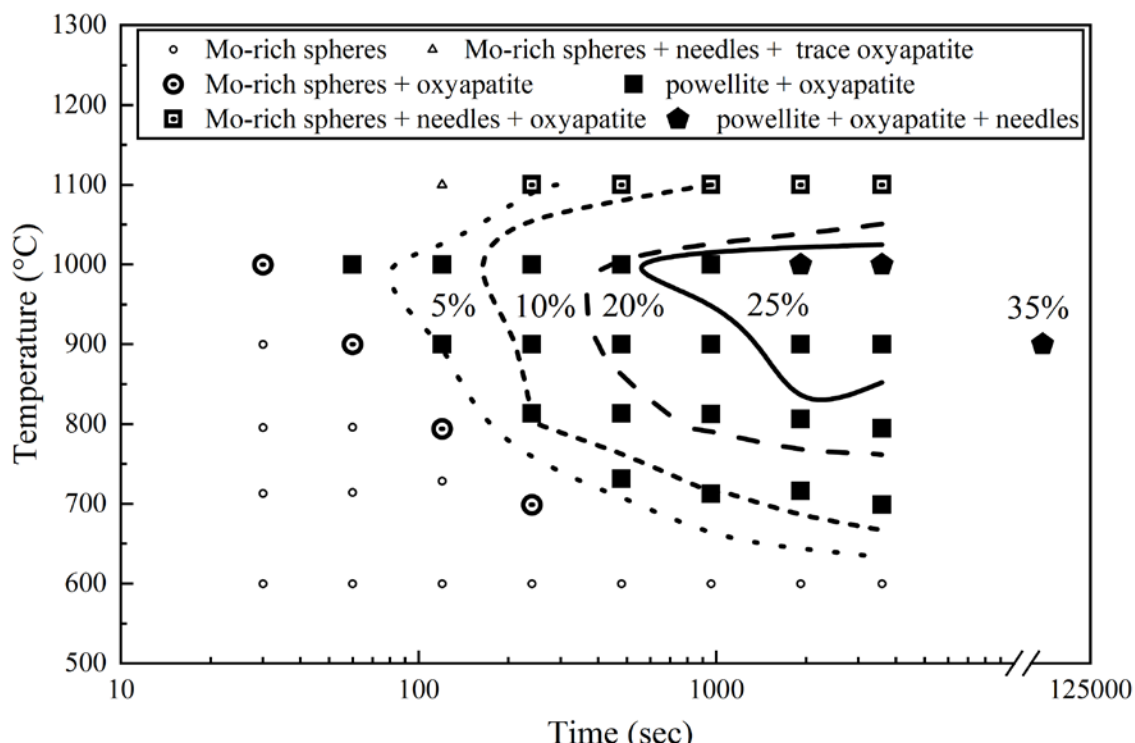


Figure 2.32. TTT diagram describing the fraction of crystallized material that forms in the PNNL borosilicate waste glass.

Mo-rich spheres were detected by SEM in all samples. The phases that form after short times are x-ray amorphous, but TEM determined that these spheres were indeed crystalline and possessed similar compositions to the crystalline powellite detected by XRD in samples heated for longer times. Raman spectroscopy also found very similar vibrational modes for all Mo-rich spheres. The dendritic and cross molybdate phases were also found to be very similar to the spheres and all are believed to be the powellite phase detected by XRD.

Powellite was not detected in the XRD patterns of samples heated at 1100°C; these samples instead possess Mo-rich needles with large concentrations of lanthanide ions. This phase is labeled as “needles” in Figure 2.32 and these needles were also found using SEM and Raman spectroscopy in samples after long times at 1000°C and 900°C. SEM images of the 1100°C/2 minute sample indicated the presence of some oxyapatite crystals, even though they could not be detected by XRD; these samples are labeled “trace oxyapatite” in Figure 2.32. Samples held at 1100°C for short times, 0.5 and 1 min, were not recorded in this diagram because it is uncertain to whether the Mo-rich spheres that were found in the samples formed at this temperature or if they formed while the samples were quenched in water.

The first crystalline phase to be detected by XRD at every temperature was oxyapatite, with crystalline powellite detected at longer times. The weight fraction of oxyapatite was always greater than that of powellite, with the former approaching 30 wt.% of the sample, and powellite not exceeding 10 wt.%. The fastest crystallization occurs around 1000°C. The sample held at 900°C for 24 hours reached an overall crystal

percentage of 35.2%, the largest fraction of any sample studied. The data from the Rietveld analyses for the crystalline percentages can be found in Table A.3 of Appendix A. Not shown in the diagram is the RuO₂ crystals that were found in every sample using SEM and are known to be insoluble in this borosilicate glass.

2.3.6. GLASS-CERAMIC FORMATION PATHWAYS

The isothermal and constant cooling rate experiments provided important information on the crystallization pathways, and the types and sizes of the phases that form are summarized in Figure 2.33. A transformation diagram was developed to show how these phases evolve, as shown in Figure 2.34. It should be noted that the heavy metal (e.g., RuO₂) phase that is present in all samples and is known to be insoluble in these melts even at 1300°C [8] is not included in this analysis. The following section will describe the phase transformations as a function of the relative nucleation and crystallization rates expected at different temperatures.

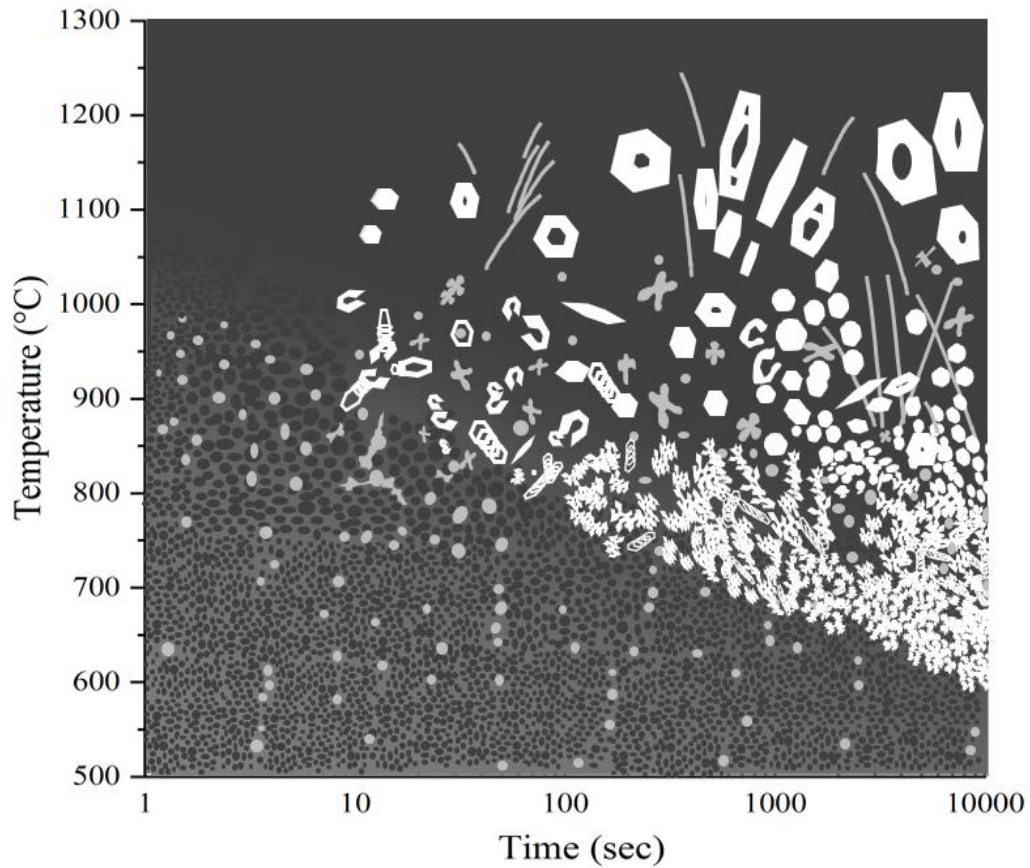


Figure 2.33. Schematic figure of the effects of time and temperature on the development of microstructural features in the borosilicate waste glass-ceramic. Sizes are not to scale but show general trends.

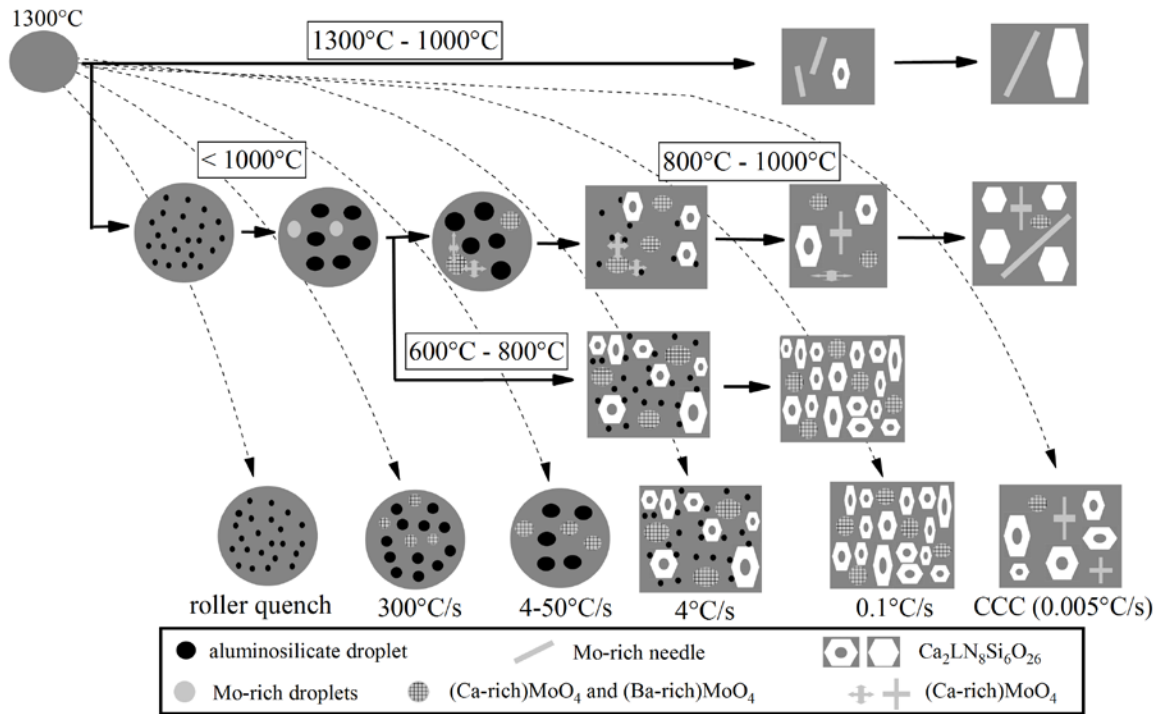


Figure 2.34. Phase transformation pathways identified in the isothermal and continuous cooling experiments. Crystalline features are below the detection limit of the XRD in the circular maps.

At high temperatures (~ 1000 to 1300°C , the melting temperature (T_{melt}) used here), where there are expected to be fewer nuclei and fast crystal growth rates, a small number of oxyapatite and needle-like molybdate crystals rapidly grow in size. As indicated in Figure 2.34, the oxyapatite and Mo-rich needles crystallize directly from the melt around the same time. The phase assemblage of the glass heat-treated for short times is uncertain due to the fast growth of the phase separated glass and the Mo-rich spheres at $\sim 900^{\circ}\text{C}$, Figure 2.17a. It is believed that any phase separation in the samples held for short times occurred upon quenching the glass in water instead of forming at the high temperatures. At high temperatures, $>1000^{\circ}\text{C}$, the oxyapatite crystals are large, hexagonal-shaped crystals (colored white in Figure 2.33 and Figure 2.34) sometime with a hollow center and the molybdate phase is found as mostly long needles (colored light grey in Figure 2.33 and

Figure 2.34). Often these crystals are found to grow from a central point, also displayed in Figure 2.33.

In the temperature range from $\sim 800^{\circ}\text{C}$ to $\sim 1000^{\circ}\text{C}$, the nucleation and crystallization rates are likely similar and so crystallization is fastest at these temperatures. On cooling, the melt first separates into an aluminosilicate-rich droplet phase and an alkali and alkaline earth (AE)-lanthanide (LN)-Mo-rich borosilicate matrix phase, as shown in Figure 2.34. Mo-rich droplets that also concentrate Ca and Ba ions then grow from the AE-LN-Mo-borosilicate phase, and these crystallize to form the two phases with the powellite crystal structure, $(\text{Ca-rich})\text{MoO}_4$ and $(\text{Ba-rich})\text{MoO}_4$. At longer times, oxyapatite $(\text{Ca}_2\text{LN}_8\text{Si}_6\text{O}_{26})$ crystallizes from the matrix portion of the phase-separated residual glass, and this causes the droplet phase to dissolve into the modified matrix to form what appears in SEM to be a homogeneous glass. In this temperature range, oxyapatite crystallizes fast enough so that there is sufficient Mo still left in the newly homogenized residual glass that will then crystallize out as $(\text{Ca-rich})\text{MoO}_4$ crosses. It appears that when Mo crystallizes out directly from a homogeneous residual glass phase, it forms crosses and needles, but if the glass has a phase separated morphology, Mo-rich spheres separate and subsequently crystallize. The needles were found to be a much different material, one containing greater concentrations of lanthanides and lower concentrations of alkaline earth, most likely due to the composition of the glass from which they crystallize. The crosses were found to be only $(\text{Ca-rich})\text{MoO}_4$. The spheres were found to contain amounts of Ba unlike the crosses which crystallizes into both $(\text{Ca-rich})\text{MoO}_4$ and $(\text{Ba-rich})\text{MoO}_4$ with the former to be more abundant.

In the temperature range from $\sim 600^{\circ}\text{C}$ to $\sim 800^{\circ}\text{C}$, one expects that the relative nucleation rates are much greater than the crystal growth rates, and this leads to the formation of dendritic clusters of smaller oxyapatite crystals, as represented in Figure 2.33. Since the glass transition temperature of this material was $\sim 590^{\circ}\text{C}$, no significant phase transformations are expected below this final temperature range.

Mo is incorporated in borosilicate glasses as MoO_4^{2-} units, and Chouard et al. found that samples with increased Nd had a higher Mo solubility. He believed that lanthanide ions, like Nd^{3+} , increased molybdenum solubility through blocking the transport of the MoO_4^{2-} units through also bonding with the silica network [21]. At high temperatures this effect will be minimal due to the lower viscosity allowing the MoO_4^{2-} and Nd^{3+} units to combine with alkaline earths and precipitate molybdate and oxyapatite crystals, respectively.

At lower temperatures, instead of the Nd^{3+} blocking the transport of the MoO_4^{2-} through the bondings with the silica network [21], the MoO_4^{2-} and Nd^{3+} ions are located in the AE-LN-Mo-rich borosilicate matrix phase. Brehault et al. [9] found this combination also increased the solubility of MoO_3 . The increased local concentration of MoO_4^{2-} units is still great enough to cause some of the MoO_4^{2-} ions to precipitate out into powellite. With longer times Nd^{3+} ions precipitate into oxyapatite. The close association of powellite and oxyapatite crystal formation, both temporal and spatial, is consistent with the concentration of lanthanide ions and molybdate tetrahedra in the matrix portion of the phase separated liquid that first forms on cooling. Powellite crystals are generally found near oxyapatite crystals, and there are many examples of powellite spheres that have formed within the hollow oxyapatite hexagons; i.e., Figure 2.12. Also, because there's increased solubility of

Mo in the AE-LN-Mo-rich borosilicate matrix phase, the precipitation of powellite crosses may be influenced by the decrease in Mo solubility with the sequestration of Nd in the oxyapatite crystals and reduction of the phase separation.

Table 2.7. Estimated compositions of the residual glass in samples prepared at various cooling rates, with the analyzed composition for the CCC sample.

Oxide	as-received [At. %]	3.4°C/s [At. %]	0.1°C/s [At. %]	CCC [At. %]	CCC EMPA*
Al ₂ O ₃	3.8	3.9 ± 0.1	3.8 ± 0.5	4.5 ± 0.2	5.62
B ₂ O ₃	11.3	12.1 ± 0.2	14.5 ± 1.2	13.4 ± 0.5	16.41
BaO	2.2	2.2 ± 0.1	2.2 ± 0.5	2.5 ± 0.1	3.09
CaO	8.4	7.6 ± 0.3	5.7 ± 1.5	5.6 ± 0.9	3.66
Ce ₂ O ₃	1.5	1.2 ± 0.1	1.2 ± 1.1	0.9 ± 0.2	0.23
Cs ₂ O	1.6	1.6	1.8 ± 0.2	1.9 ± 0.01	1.29
La ₂ O ₃	0.8	0.64 ± 0.1	0.6 ± 0.5	0.6 ± 0.1	0.26
Li ₂ O	3.2	3.5	4.2 ± 0.4	3.9 ± 0.1	--
MoO ₃	4.2	4.2 ± 0.1	3.6 ± 0.6	1.2 ± 0.9	0.22
Na ₂ O	4.1	4.3 ± 0.1	4.1 ± 1.6	4.7 ± 0.2	5.32
Nd ₂ O ₃	2.4	2.0 ± 0.1	1.7 ± 0.5	1.3 ± 0.3	0.32
Pr ₂ O ₃	0.7	0.6 ± 0.1	0.5 ± 0.5	0.3 ± 0.1	--
SiO ₂	47.7	48.0 ± 1.0	47.7 ± 5.9	51.1 ± 2.1	57.49
SrO	1.5	1.5 ± 0.1	1.4 ± 0.5	1.0 ± 0.2	1.00
ZrO ₂	3.8	3.9 ± 0.1	4.2 ± 0.4	4.4 ± 0.2	5.11
OTHER	4	2.7 ± 0.1	3.0 ± 0.3	2.5 ± 0.2	
TOTAL	100	100	100	100	100

*from Washington State University

The composition of the residual glass for some continuously cooled samples was estimated by subtracting out the crystal phases from the as-received composition and are summarized in Table 2.7. The composition of the phases determined by EDS were normalized to their respective crystal fractions determined by quantitative XRD and subtracted from the nominal glass composition. Also shown are the results of the EMPA (Washington State University) analyses of the residual glass, including the direct detection

of boron (but not lithium). The lanthanides and Mo decrease with decreasing cooling rate due to oxyapatite and powellite crystallizing out of the glass. B was found to only be contained in the residual glass and increased in relative concentration with slower cooling rates.

The phase development pathways identified in this study are useful for explaining the microstructural development of oxyapatite reported in other studies. For example, Delattre et al. [25] found that the oxyapatite crystals that formed in a borosilicate glass after treatment times >1 hour at temperatures from 700°C -775°C were hollow hexagonal dendrites and needles, whereas solid hexagonal dendrites/needles formed at temperatures from 700°C -840°C. In the present study, hollow oxyapatite crystals formed at isothermal temperatures as high as 1100°C, and these hollow oxyapatite forms are favored at shorter times for all temperature, whereas the filled oxyapatite crystals were more common at longer times. Delattre et al. nucleated their samples for 6 hours at 630°C prior to their isothermal heat treatments, whereas in the present study, samples were isothermally heat treated immediately after melting, avoiding any prior nucleation or crystallization events. Delattre et al. did not find any hollow crystals at the higher temperatures, since at those temperatures the crystallization rate is faster, and the residual melt is less viscous, so the pre-nucleated samples would quickly transform to the filled hexagonal crystals.

The phase development pathways identified in this study also expand on results reported by Asmussen et al. [16] by providing a range of cooling rates ~4 magnitudes wider and Crum et al. [12] providing more precise times and temperatures for the phase transitions. Asmussen et al. found clusters of much smaller oxyapatite crystals formed in samples cooled four times faster than the cooling rate of the CCC sample, ~0.03°C/s, a

morphology similar to what was created in the sample cooled at 0.1°C/s in this study. Crystals produced in their samples cooled $\frac{1}{4}X$ the CCC sample were also much larger aligning with the large crystals found in the samples heat treated at high temperatures ($>1000^{\circ}\text{C}$) in which this sample would spend sufficient time at those temperatures. Crum et al. [12] described similar phase transformations in which the glass would phase separate into an aluminosilicate droplet and matrix phase and from this matrix phase powellite and oxyapatite would crystallize out. The TTT and CCT diagrams developed in this study provides useful data describing times, temperatures, and cooling rates for these transformations.

2.4. CONCLUSIONS

Experimental protocols to quantitatively characterize the microstructural transformations that occur as functions of time and temperature for a complex borosilicate glass-ceramic designed for nuclear waste remediation have been developed. Oxyapatite, $\text{Ca}_2\text{LN}_8\text{Si}_6\text{O}_{26}$, and powellite, $(\text{Ca-rich})\text{MoO}_4$, are the two major crystalline phases that form when these melts are cooled. Molybdenum and rare earth ions are concentrated in the matrix liquid which surrounds aluminosilicate rich droplets, when the melts initially phase separate on cooling. Molybdenum-rich droplets separate from the matrix to form powellite-like crystalline phases, followed by the formation of the oxyapatite crystals. The changes in the liquid composition with the precipitation of these crystals causes the residual glass microstructure to become more homogeneous upon quenching.

Molybdenum-rich droplets were found by electron microscopy in samples cooled slower than about 45°C/s. From phase separated residual glass, Mo-rich spheres would separate and subsequently crystallize while from a more homogenous residual glass Mo-rich phases would form cross-shaped and needle-like morphologies, the latter two morphologies occurring at temperatures 1000°C and above or after the formation of oxyapatite at temperatures 900°C and above. The composition of the needles was determined to be much different than the cross and droplet shaped Mo-rich phases. Due to the changing mechanisms and low crystal percentages of the Mo-rich phases (up to ~10 wt.%), the JMAK kinetic model was unable to accurately describe the crystallization of powellite.

Oxyapatite formed hollow hexagonal-shaped crystals that over time fill in to form more solid features. The fastest growth of oxyapatite in the isothermal experiments occurred at 1000°C, and crystalline oxyapatite formed in samples continuously cooled from the melt at rates slower than about 4°C/s. A sample with about 30 wt.% oxyapatite was created after an isothermal hold at 900°C for 24 hours. A JMAK analysis of the crystallization kinetics for oxyapatite indicate a diffusion-controlled growth mechanism, but the resulting activation energy, 58.8 ± 8.16 kJ/mol, was an order of magnitude lower than what was reported elsewhere for growth over a much narrower temperature range, indicating some significant changes in the growth conditions over the 700-1000°C experimental range in the present study.

This report provides data on the microstructural development of a borosilicate glass-ceramics as a function of time and temperature through isothermal and constant cooling experiments. Understanding the crystallization pathways is essential in developing

acceptable waste forms to vitrify hazardous nuclear waste. Information from this study will be considered when further developing the glass-ceramic composition and processing techniques.

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3. THE EFFECTS OF MICROSTRUCTURE ON THE DISSOLUTION BEHAVIOR OF A COMPLEX GLASS-CERAMIC WASTE FORM

ABSTRACT

Borosilicate glass-ceramics developed by Pacific Northwest National Laboratories (PNNL) are intended to vitrify reprocessed spent nuclear fuel into environmentally safe waste forms. The multi-phase glass-ceramics exhibit complicated corrosion behavior. Borosilicate glass ceramics provided by PNNL were re-melted and quenched with different thermal histories to produce samples with a range of microstructures, described by analytical electron microscopy and x-ray diffraction. In particular, slower quench rates produced samples with greater fractions of crystalline powellite (CaMoO_4 , up to ~10 wt. %) and oxyapatite ($\text{Ca}_2\text{Ln}_8\text{Si}_6\text{O}_{26}$, up to ~30 wt. %). Product consistency tests (PCT) were performed to characterize the effects of microstructural development on elemental release rates. Atomic force microscopy (AFM), profilometry, and electron microscopy studies of polished, monolithic samples provide information about the relative recession rates of the residual glass and major phases, and reveal powellite to be the least durable phase, followed by the residual glass, and the oxyapatite. However, the thermal history dependent composition of the residual glass phase appears to have the greatest effect on the overall dissolution kinetics of this glass-ceramic.

3.1. INTRODUCTION

Efforts around the world are being made to reduce the volume of spent nuclear fuel and to develop viable waste storage options [1-3]. In the United States, the waste form must meet the requirements described in the Waste Acceptance Product Specifications (WAPS)

[4]. Chemically durable borosilicate glasses are used worldwide and are understood to be a viable option to immobilize nuclear waste [3,5].

In order to understand the chemical durability of waste glasses, it is important to first understand the potential driving forces associated with dissolution. There are several stages for the dissolution of waste glasses in contact with groundwater in a geological depository, and these are summarized in Figure 3.1 [6]. In Stage I, the initial dissolution rate is dominated first by the exchange of ions in solution with weakly bonded alkalis in the glass and then by the hydrolysis of the glass network until a protective layer is formed. In Stage II, the dissolution rate decreases to the “residual rate” due to the saturation of silica in the surrounding aqueous environment, in equilibrium with the protective layer that formed in Stage I. If these equilibrium conditions change and crystals precipitate from the saturated solutions, then glass dissolution can accelerate in Stage III.

Section 1.3 of the WAPS states that the waste form needs to be more chemically durable than the environmental assessment (EA) reference glass, as determined using the Product Consistency Test (PCT) [4,7,8]. The product consistency test is a standard test procedure for determining the dissolution rates of waste glasses. For test method “A”, a seven-day test, the stage I and possibly stage II dissolution reactions would dominate the overall release rates. For the present study, only stage I and possibly stage II dissolution behavior will be evaluated.

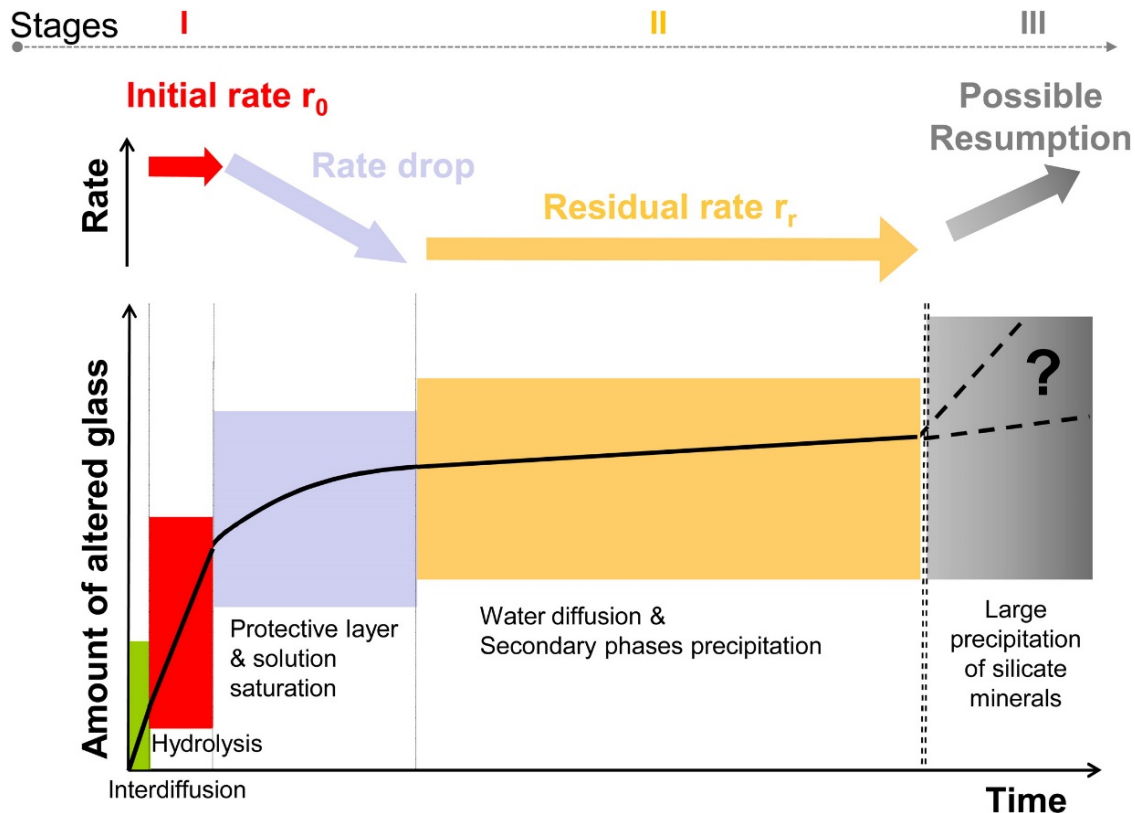


Figure 3.1. Long-term dissolution behavior of a waste glass [6].

Over the past decade, researchers at Pacific Northwest National Laboratory (PNNL) have done extensive work for the D.O.E Fuel Cycle Research and Development (FCRD) program developing borosilicate glass-ceramic compositions to immobilize wastes generated from aqueous spent nuclear fuel (SNF) reprocessing techniques [9-16]. The Collins CLT waste (Table 3.1) describes the non-radioactive components expected from reprocessed SNF and has large concentrations of transition metals, lanthanides and Cs-oxides that have relatively low solubilities in borosilicate glasses. To increase the loadings of waste like Collins CLT, PNNL researchers have developed glass-ceramic formulations that retain acceptable durabilities with greater waste loadings [9,16].

Table 3.1. Composition of Collins CLT waste [9].

Oxide	Content [mol%]	Oxide	Content [mol %]
MoO ₃	19.719	CdO	0.621
ZrO ₂	17.590	SeO ₂	0.534
Nd ₂ O ₃	11.279	Eu ₂ O ₃	0.354
BaO	10.443	Ag ₂ O	0.353
Cs ₂ O	7.416	SnO ₂	0.339
SrO	6.887	Gd ₂ O ₃	0.322
Ce ₂ O ₃	6.859	Rh ₂ O ₃	0.226
La ₂ O ₃	3.527	Br	0.205
Pr ₂ O ₃	3.187	PdO	0.100
TeO ₂	2.985	Pm ₂ O ₃	0.036
Sm ₂ O ₃	2.240	Sb ₂ O ₃	0.028
Y ₂ O ₃	2.019	In ₂ O ₃	0.007
Rb ₂ O	1.641	Tb ₂ O ₃	0.006
RuO ₂	1.076	Total	100.000

A variety of different crystal phases have been detected in the glass-ceramics developed at PNNL for vitrifying the Collins CLT waste, including oxyapatite ($\text{Ca}_2\text{Nd}_8\text{Si}_6\text{O}_{26}$), Ln-borosilicate ($\text{Gd}_3\text{BSi}_2\text{O}_{10}$), powellite ((Ca,Sr)MoO₄ or (Ca,Ba)MoO₄), CsLiMoO₄, La₂Mo₃O₁₂, pollucite (CsAlSiO_4), $\text{Zr}_{0.9}\text{Ce}_{0.1}\text{O}_2$, Y₂O₃, and RuO₂ [16]. The effects of thermal history on the growth and morphology of the oxyapatite and powellite phases in a glass developed by PNNL are described in an earlier study [17]. In this paper, the relationship between microstructure and aqueous corrosion processes are described

In preliminary PCT studies at PNNL, glass-ceramics based on low (< 25 mass%) waste-loadings, the release rates of Si, B, Na, and Li were all found to be an order of magnitude below the respective benchmarks established with the EA reference glass [9]. The release rate of Mo from this composition (3.5 mass% Mo, nominal) was substantially lower (0.3 g/L) for samples rapidly quenched from the melt, compared to slowly cooled

(1.06 g/L), following the center-line canister cooling (CCC) protocol shown in Table 3.2. No crystalline features were found in the plate-quenched sample, whereas oxyapatite and powellite were identified in the CCC-sample. It was believed that the greater release of Mo from the slowly cooled sample was due to the preferential dissolution of the molybdenum-rich phase.

Table 3.2. Treatment schedule comparable to the CCC of a waste storage canister [9].

Step	Start Temperature (°C)	Rate (°C/min)	Step Duration (hours)
1.	1200 - 1150	0	Preheat 0.5 hours then fast cool
2.	1150 - 1050	~ -7	0.2
3.	1050 - 950	-0.935	1.8
4.	950 - 886	-0.288	3.7
5.	886 - 845	-0.108	6.3
6.	845 - 626	-0.205	17.8
7.	626 - 400	-0.126	29.9
8.	400	0	Dwell 1 hour

In studies of glasses with waste loadings up to 50 wt.%, samples cooled using the CCC profile were tested under static dissolution conditions for various times [10]. Mo release rates significantly increased (0.05 to 0.4 g/L) when the MoO_3 composition increased from 6.25 wt.% to 6.95 wt.%. It was believed that Mo was mostly contained in the powellite phase in the samples with lower Mo concentrations (6.25 wt.% and less) and the increase in Mo leach rate was due to the release of Mo in the residual glass phase. Nd release rates were very low, indicating that oxyapatite crystals were more durable than the residual glass. The release rates of boron and sodium from the glass-ceramic with 50

mass% CLT waste were relatively higher (+0.1g/L), leading to the conclusion that the residual glass is less durable than the crystalline phases.

To aid the development of borosilicate glass-ceramics at PNNL, a statistical compositional matrix was designed to determine the effects of composition on the important properties of these materials [16]. The centroid composition from the test matrix is the glass-ceramic composition used in this report. Three different dissolution tests were used in that study to characterize the chemical durability of these glass-ceramics. One test, a modified PCT method “B”, static dissolution test, was used to analyze samples cooled according to the CCC after 7, 28, 119, 448 days. The second test was a single-pass flow-through, SPFT, test used on samples cooled at 4, 1, and 0.25 times the CCC cooling rate [15,16]. The third test was a modified single-pass flow-through, MSPFT, test used to further analyze the CCC-cooled samples by weight loss measurements and dimensional changes using scanning electron microscopy [13,16]. From the PCT tests, the effects of composition on the release of various components from the glass were determined and are summarized in Table 3.3 [16]. B, Na, and Mo were found to have the fastest release rates; Mo was released primarily from powellite crystals and Na and B were released from the residual glass. The SPFT dissolution tests showed that regardless of cooling rate, pH or flow rate, the powellite crystals corroded fastest, followed by the residual glass phase, with the oxyapatite phase the most stable. The MSPFT dissolution determined that the residual glass dissolved fastest.

These earlier studies indicate that changes in microstructure and composition of the different phases have a significant effect on the overall dissolution rates of these complex borosilicate glass-ceramics. This work describes the effects of cooling rate of the centroid

composition from the PNNL test matrix [16] on chemical durability, to better understand the relationships between microstructural and dissolution rates. Slower cooling rates changed the types and percentages of the crystalline phases and altered the composition of the residual glass to affect the overall dissolution rates of the waste material.

Table 3.3. Compositional effects on elemental release rates [16].

Glass Component	Trend increasing/decreasing release rates
Al ₂ O ₃ , ZrO ₂	decrease in B, Li, Na, Cs, Si
SiO ₂	decrease in B, Cs
B ₂ O ₃ , Na ₂ O, MoO ₃	increase in B, Li, Na, Cs, Si
Ln ₂ O ₃	increase in B, Li, Na, Cs
Li ₂ O	increase in B, Si / decrease in Li
CaO	no effect

3.2. EXPERIMENTAL METHODS

3.2.1. GLASS-CERAMIC COMPOSITION

The nominal molar composition of the as-received centroid composition glass, with a waste loading of 47 wt.%, is shown in Table 3.4 [16]. Sample preparation information can be found in reference [16]. 500 g batches from reagent-grade oxides were melted twice in a lidded Pt/10% Rh crucible between 1250°C and 1450°C for 1 hour and quenched on an Inconel plate in air.

2.2. THERMAL TREATMENTS

Samples were prepared with different microstructures by melting the centroid composition at 1300°C for one hour, then quenching at different rates, using several

techniques described in [17]. Table 3.5 summarizes the conditions used to prepare samples in this study.

Table 3.4. Glass composition [16].

Oxide	Content [mol%]	Oxide	Content [mol%]
SiO ₂	47.674	TeO ₂	0.641
B ₂ O ₃	11.329	Sm ₂ O ₃	0.481
CaO	8.445	Y ₂ O ₃	0.433
MoO ₃	4.229	Rb ₂ O	0.353
Al ₂ O ₃	3.789	RuO ₂	0.232
Na ₂ O	4.069	CdO	0.135
ZrO ₂	3.774	SeO ₂	0.116
Li ₂ O	3.247	RhO ₂	0.089
Nd ₂ O ₃	2.427	Eu ₂ O ₃	0.076
BaO	2.241	Ag ₂ O	0.075
Cs ₂ O	1.591	SnO ₂	0.072
SrO	1.476	Gd ₂ O ₃	0.070
Ce ₂ O ₃	1.472	PdO	0.023
La ₂ O ₃	0.757	Total	100.000
Pr ₂ O ₃	0.684		

Table 3.5. Quench rates and phases for each sample [17].

Quenching Technique	Cooling Rate [°C/s]	Phases (Quantitative XRD) [wt. %]	
roller quench	>100	x-ray amorphous	
copper wedge step 2	5.8	x-ray amorphous	
copper wedge step 1	5.1	x-ray amorphous	
steel wedge step 1	4.1	x-ray amorphous	
air-cool	3.4	Ca ₂ Nd ₈ Si ₆ O ₂₆ CaMoO ₄	8.4 ± 1.3 1.2 ± 0.4
furnace-cool	0.1	Ca ₂ Nd ₈ Si ₆ O ₂₆ CaMoO ₄	21.4 3.2
CCC	0.008	Ca ₂ Nd ₈ Si ₆ O ₂₆ CaMoO ₄	15.2 8.5

3.2.3. PRODUCT CONSISTENCY TEST

The product consistency test (PCT) was used to measure elemental release rates from glasses with various cooling profiles. Sample preparation was done according to the procedures described for method “A” [7]. Powders were produced with an agate mortar and pestle and sieved between 75 and 150 μm , then reacted with DI water in Teflon vessels at $90 \pm 2^\circ\text{C}$ for 7 days. All experiments were performed in triplicate. Samples were then filtered (4 μm syringe filter) and leachate concentrations were evaluated using an inductively coupled plasma optical emission spectrometer, ICP-OES, (PerkinElmer Optima 2000 DV, Norwalk, USA) to measure the release rates of Si, B, Ca, Na, Ba, Nd, and Mo. The release rates were normalized to the concentrations of each element in the target composition and reported as g/m^2 , as outlined in the ASTM procedure [7].

In addition, PCT evaluations were conducted on similar samples at Savannah River National Lab and those results are reported in Appendix B.

2.4. DIFFERENTIAL DISSOLUTION TEST

The relative dissolution rates of different microstructural features were determined using techniques to measure changes in local topology. Samples with different thermal histories were cut and ground to sizes $\sim 1 \times 1 \times 0.1 \text{ cm}$, then polished using a 0.25 μm diamond suspension. A titanium layer ($\sim 150 \text{ nm}$ thick, Figure 3.2) was deposited on part of the surface of the polished samples using electron beam evaporation to provide a reference for sample height, Figure 3.2. Samples were then placed, Ti-layer facing up, in Teflon containers filled with $\sim 500 \text{ mL}$ of deionized water (90°C) for various times, and weight changes were recorded after drying the samples in room temperature air.

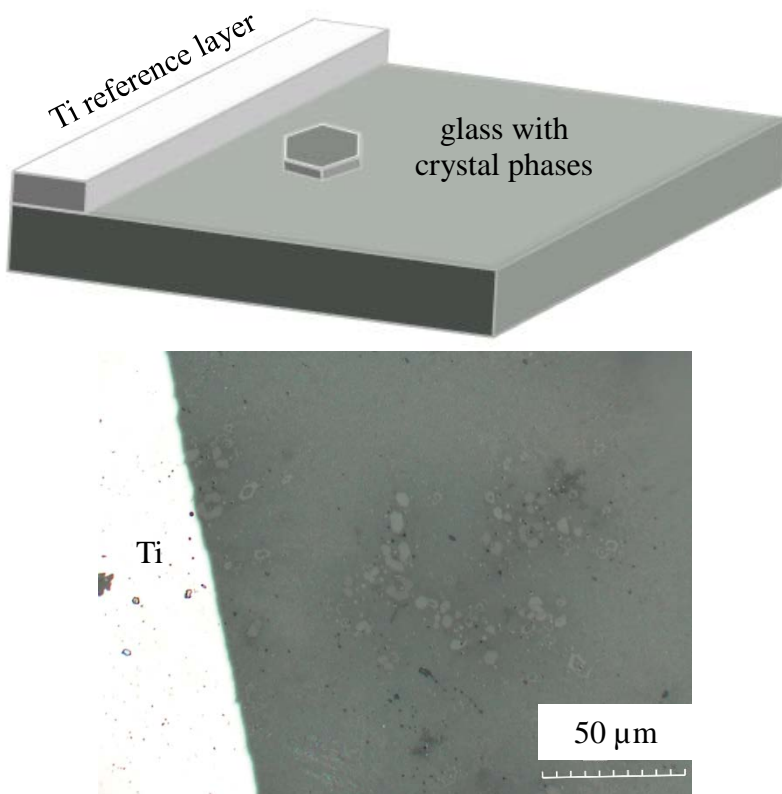


Figure 3.2. Top) schematic diagram of a sample from a differential dissolution test, with the titanium reference layer and a crystal exposed after dissolving away the surrounding glass. Bottom) an optical image of the air-cooled sample coated with ~150 nm of titanium before a dissolution test.

Atomic force microscopy, AFM, and profilometry were used to measure changes in the heights of different phases in the glass microstructure relative to the titanium layer, after dissolution of monolith samples at 90°C for various times. A Digital Instrument Nanoscope IIIA scanning probe microscope was used for the AFM experiments. The samples were scanned in tapping mode at 2 Hz with an AppNano ACTA-10 probe. The scans produced 3D maps of the surface and at least 10 measurements were used to determine the relative recession rates of the different phases. A KLA Tencor P-17 stylus profiler with Profile 8.1 software was also used to measure changes in surface topology of

corroded glasses. Scan lengths 200 – 1000 μm were analyzed at a rate of at 10 $\mu\text{m/sec}$ sampling at 50 Hz.

3.2.5. MICROSTRUCTURAL CHARACTERIZATION

A dual beam scanning electron microscope (SEM, FEI Helios NanoLab 600) with energy-dispersive x-ray spectroscopy (EDS) was used to characterize samples before and after the corrosion tests. The samples were sputter-coated with AuPd before they were loaded into the SEM. Micrographs were collected at different magnifications, electron voltages, and currents. EDS maps were obtained using Oxford Instruments AZTEC software at various magnifications using 15 keV and 1.4 nA.

3.3. RESULTS

3.3.1. EFFECTS OF THERMAL HISTORY ON MICROSTRUCTURE

In the first part of this study, the effects of thermal history on the development of the microstructure of the centroid composition are described [17]. Figure 3.3 shows representative microstructures from that study of the samples examined here. When melts were cooled by roller quenching, an aluminosilicate droplet phase separates from a borosilicate matrix that has large concentrations of alkalis (Na, Li, Cs), alkaline earths (Ca, Ba), lanthanides, and transition metals, including Mo, Figure 3.3a. The aluminosilicate droplets increase in size in samples cooled at slower rates to 4.1°C/s (~150 nm), shrink and seem to disappear from samples cooled at 0.1°C/s and slower. Table 3.6 shows the compositions of the droplet and matrix phases from the sample cooled at 4.1°C/s. Samples

cooled in the wedge mold experiments (5.8 , 5.1 , 4.1°C/s) have Mo-rich droplets (Figure. 3.3b) that are x-ray amorphous but were found by electron diffraction to be related to crystalline powellite (CaMoO_4). The composition of the powellite droplets from the sample cooled at 4.1°C/s is given in Table 3.6. The sample cooled at 3.4°C/s has larger powellite droplets that do diffract x-rays, as well as hexagonal oxyapatite ($\text{Ca}_2\text{LN}_8\text{Si}_6\text{O}_{26}$) crystals (Figure 3.3c), and the fraction of the latter has increased in the sample quenched at 0.1°C/s (Figure 3.3d). The sample cooled using the CCC profile (on average, 0.008°C/s), had the largest oxyapatite crystals, and the principal morphology of the powellite crystals changed from spheres to crosses. However, the composition of these cross-like crystals is similar to what was determined for the spherical particles in the slower quenched samples (Table 3.6). Table 6 also gives the composition of the oxyapatite crystals in the CCC-cooled sample.

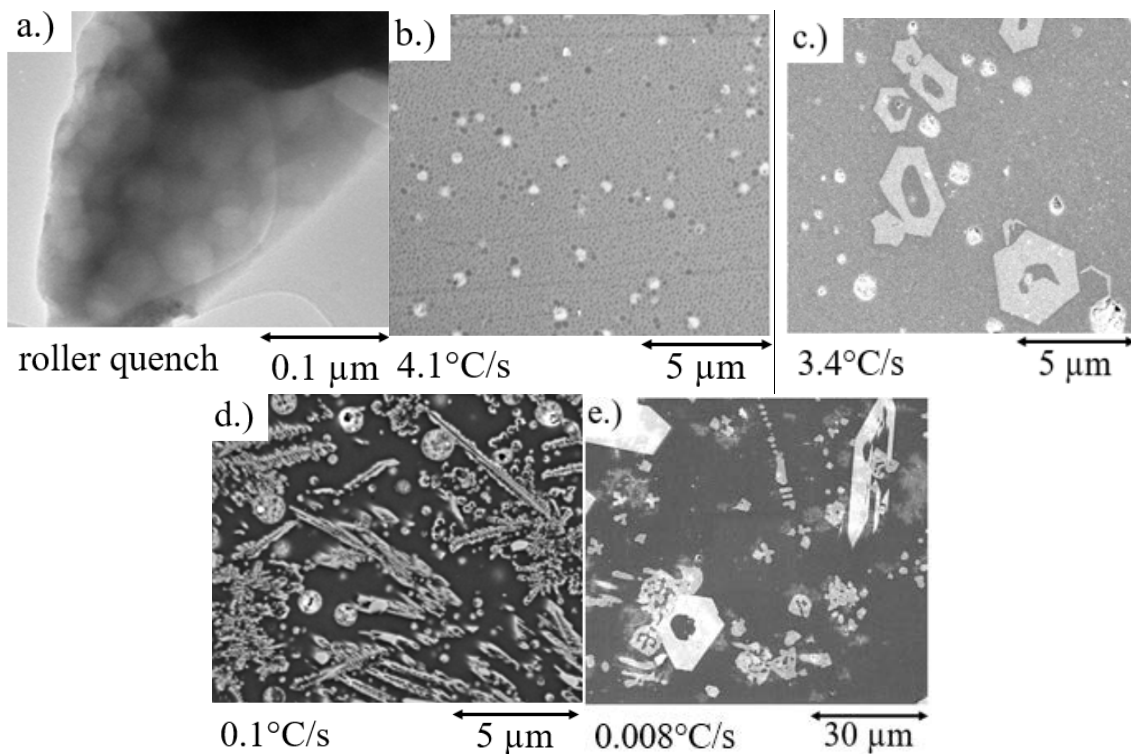


Figure 3.3. Electron images from samples quenched at different rates; a) roller quench (TEM), b) 4.1°C/s , c) 3.4°C/s , d) 0.1°C/s , and e) 0.008°C/s .

Table 3.6. Compositions of the major phases in the glass.

	residual glass (TEM) 4.1°C/s droplet	4.1°C/s matrix	4.1°C/s - sphere	powellite (TEM) CCC - cross	oxyapatite (SEM) CCC
	[At%]	[At%]	[At%]	[At%]	[At%]
Al ₂ O ₃	8	2	0	1	0
BaO	0	4	2	2	0
CaO	1	10	29	27	19
Ce ₂ O ₃	1	3	1	1	7
Cs ₂ O	0	1	0	0	0
La ₂ O ₃	1	2	0	1	2
MoO ₃	1	6	44	48	0
Na ₂ O	0	1	3	2	0
Nd ₂ O ₃	1	6	2	1	13
Pr ₂ O ₃	0	2	0	0	4
Rb ₂ O	0	0	0	0	0
RhO ₂	0	0	0	0	0
RuO ₂	0	0	0	1	0
SiO ₂	79	50	14	5	48
Sm ₂ O ₃	1	2	1	0	3
SrO	0	0	5	8	1
TeO ₂	1	3	0	1	0
Y ₂ O ₃	1	1	0	0	1
ZrO ₂	5	7	0	1	0
TOTAL	100	100	100	100	100

3.3.2. ION RELEASE VALUES FROM PCT

Ion release rates from the PCT tests on samples cooled at different rates are shown in Figure 3.4. The data from these analyses are given in Table B.1 of Appendix B. In general, the release rate for Ba, Ca, Mo, Na, and B all increase with decreasing quench rates, with maxima for each element at 0.1°C/s. The rates decrease again from the CCC-treated sample. Si and Nd release rates do not differ significantly from samples with different thermal histories, although the former has its lowest value from the CCC-treated

sample. The roller quench, 0.1°C/s , and CCC samples were sent to Savannah River National Laboratories and the PCT test preformed on them. Values were generally lower, however the same trend with cooling rate was seen. Additional data from the SRNL tests can be found in Appendix B.

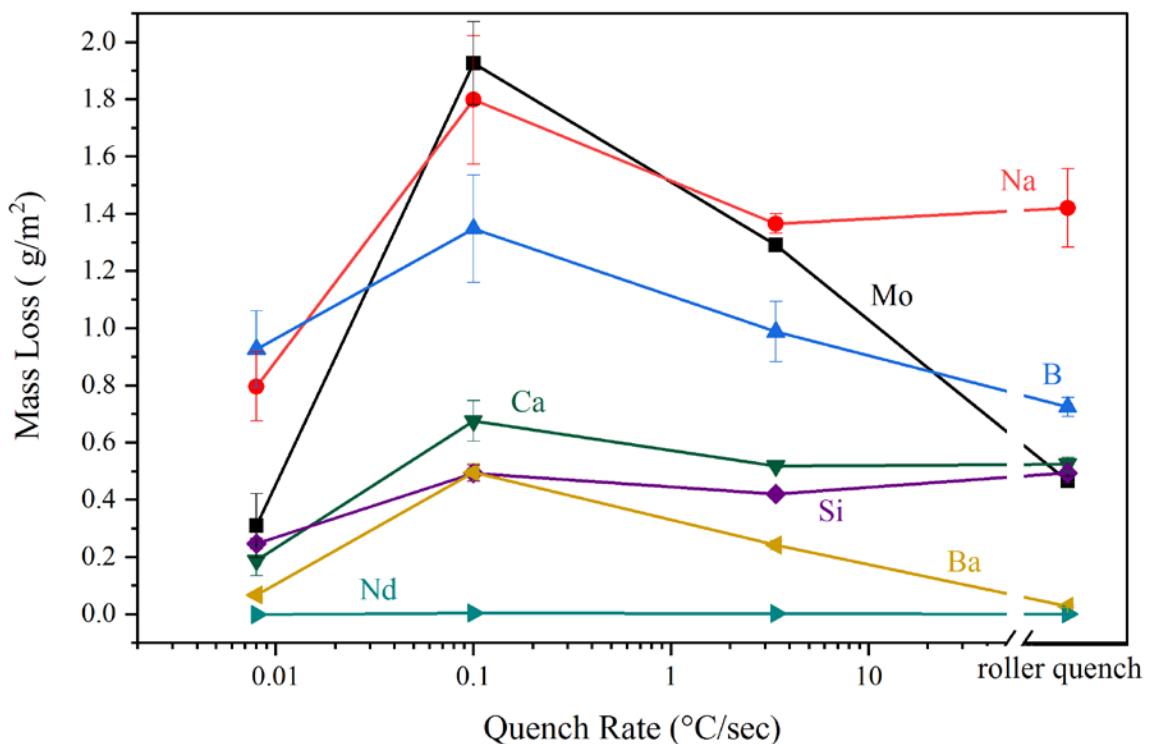


Figure 3.4. Seven-day PCT method “A” results for the centroid composition as a function of cooling rate. The lines are guides for the eye.

3.3.3. RECESSION RATE VALUES FROM THE DIFFERENTIAL DISSOLUTION TESTS

Figure 3.5 shows SEM/EDS images from the surface of a sample cooled at 3.4°C/s exposed after exposing the surface to 90°C deionized water for 12.5 days. Cross-sections of oxyapatite crystals, rich in Nd and Ca, can clearly be seen, along with smaller Mo-rich

spheres. The SEM image of the sample cracks believed to be created when the corroded sample was dried.

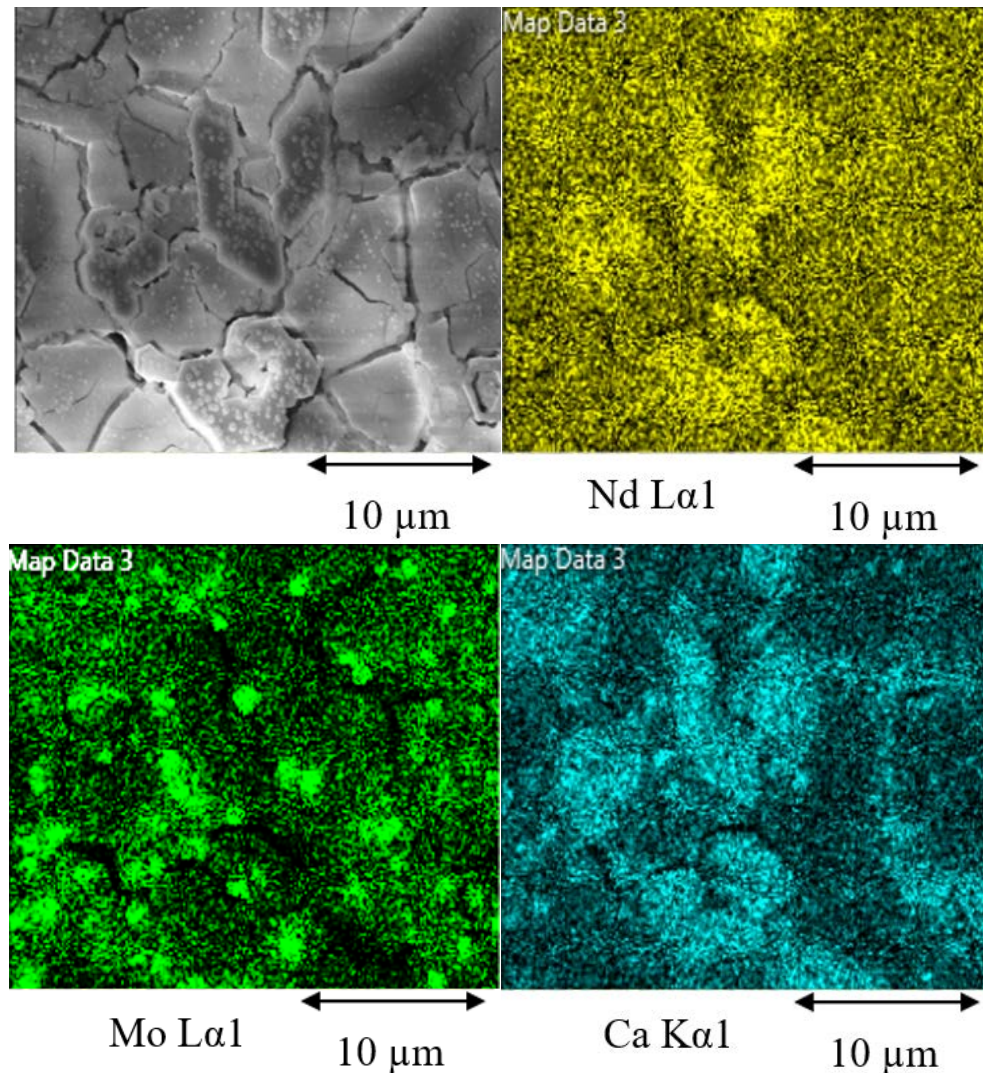


Figure 3.5. SEM micrograph and EDS maps from a sample cooled at 3.4°C/s , then reacted for how long in deionized water at 90°C .

Figures 3.6 and 3.7 show AFM scans around the same oxyapatite crystals shown in Figure 3.5. Clearly, the glass around the crystals has been etched away at a faster rate than the crystals themselves, leaving the latter at greater relative heights. Also shown in Figure 3.7 is an example of the height scan across the corroded surface which shows that the

average change in height (Δh) from the top of the crystal to the glass is 240 ± 90 nm. For the measurements from the Ti-layer to the glass surface (Figure 3.7, right), the Δh determined by subtracting the deposited Ti layer step-height (150 nm) from the step-height of the reacted sample had an average value of 260 ± 70 nm. The same sample was further reacted in 90°C DI water for a total of 18.75 days and the relative heights of the same areas and crystals were analyzed again.

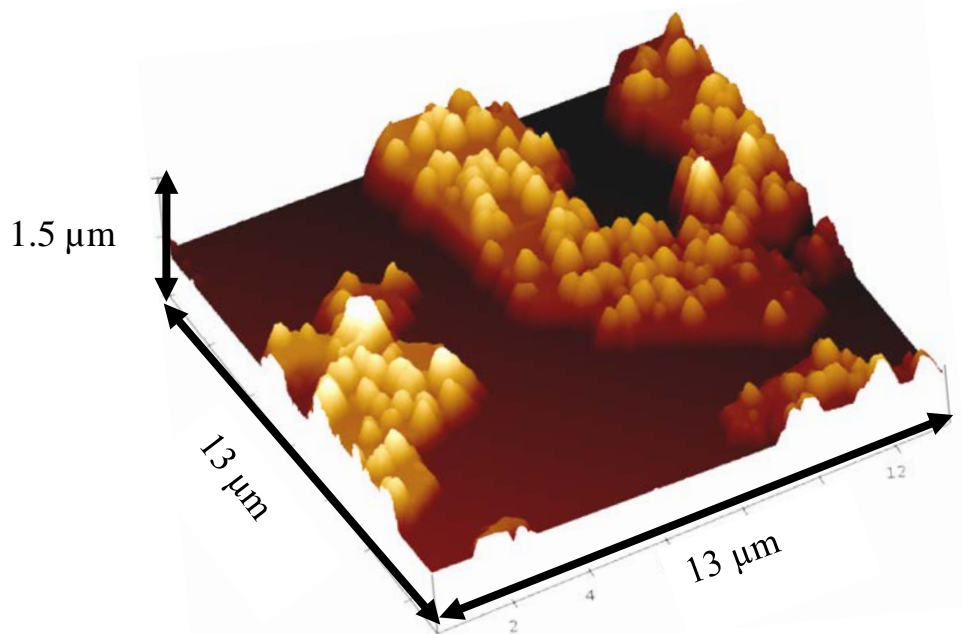


Figure 3.6. AFM scan of an oxyapatite crystal in the sample cooled at 3.4°C/s after 12.5 days in deionized water at 90°C.

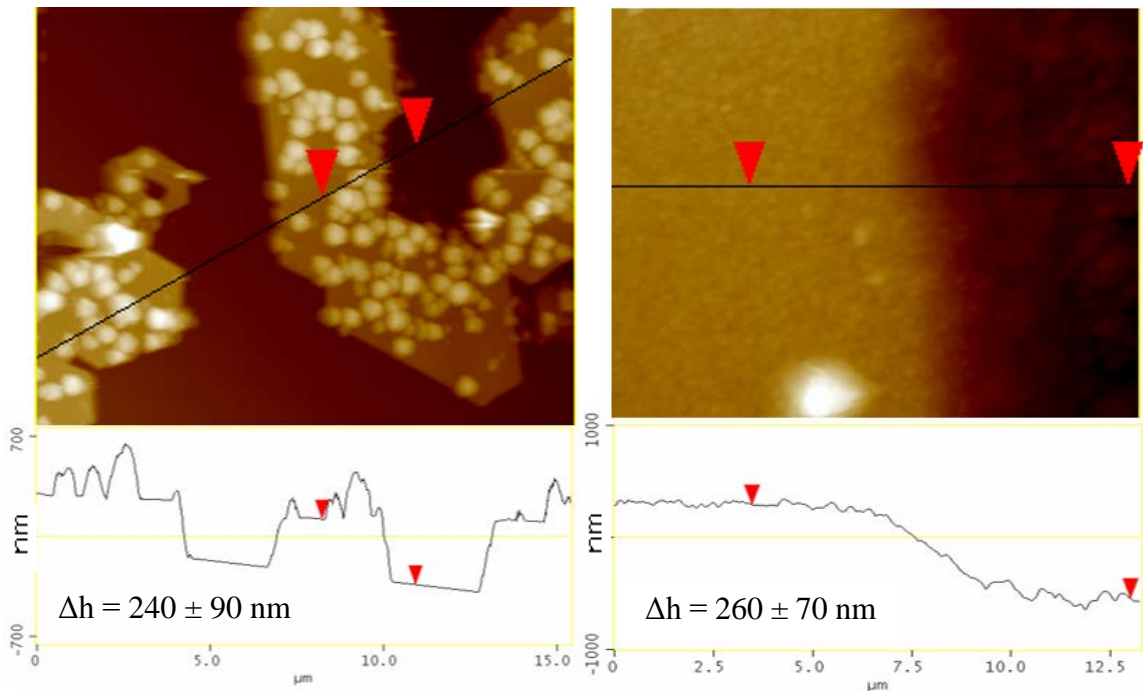


Figure 3.7. Left, example measurement of sample cooled at 3.4°C/s showing crystal to the glass measurements and, right, Ti coating to the glass measurements for the sample after 12.5 days in deionized water at 90°C.

Profilometry was used to measure the relative recession rates of different phases that formed in samples quenched at different rates, using the titanium layer deposited on the original sample as a height reference. This layer thickness was determined to be $160 \pm 10 \text{ nm}$ and was assumed to be constant for all corrosion samples. Figure 3.8 shows example profilometer scans of a sample cooled at 3.4°C/s, before and after reaction in 90°C water for eight days. This sample showed a change of height (Δh) of $250 \pm 30 \text{ nm}$.

The results from the profilometry and AFM experiments are shown in Figure 3.9 and reported in Appendix B, Table B.2. The AFM measurements of the 3.4°C/s follow the same trend as the profilometry test. The durability of the glasses decreases with decreasing quench rate. There appears to be a much larger relative recession rate of the sample cooled using the CCC profile (0.008°C/s).

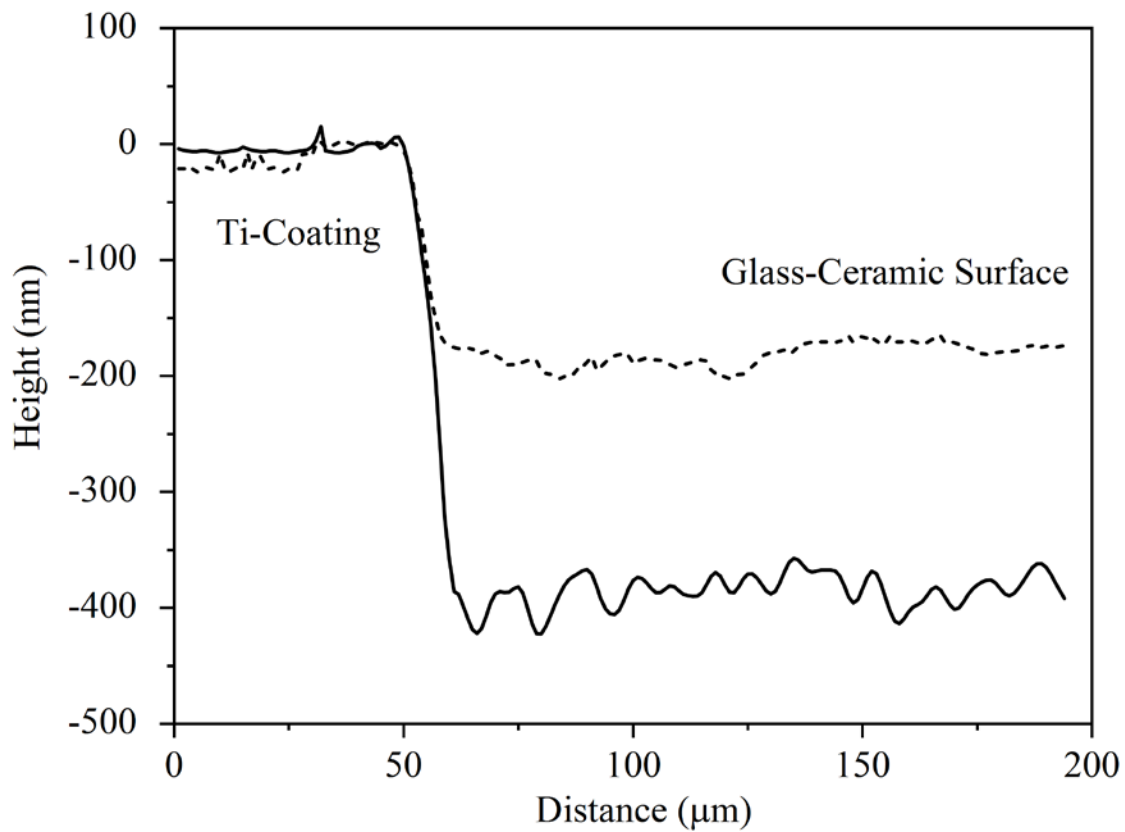


Figure 8. Example profilometry line scans of the sample quenched at 3.4°C/s before (dashed) and after (solid) reacting in 90°C deionized water for eight days.

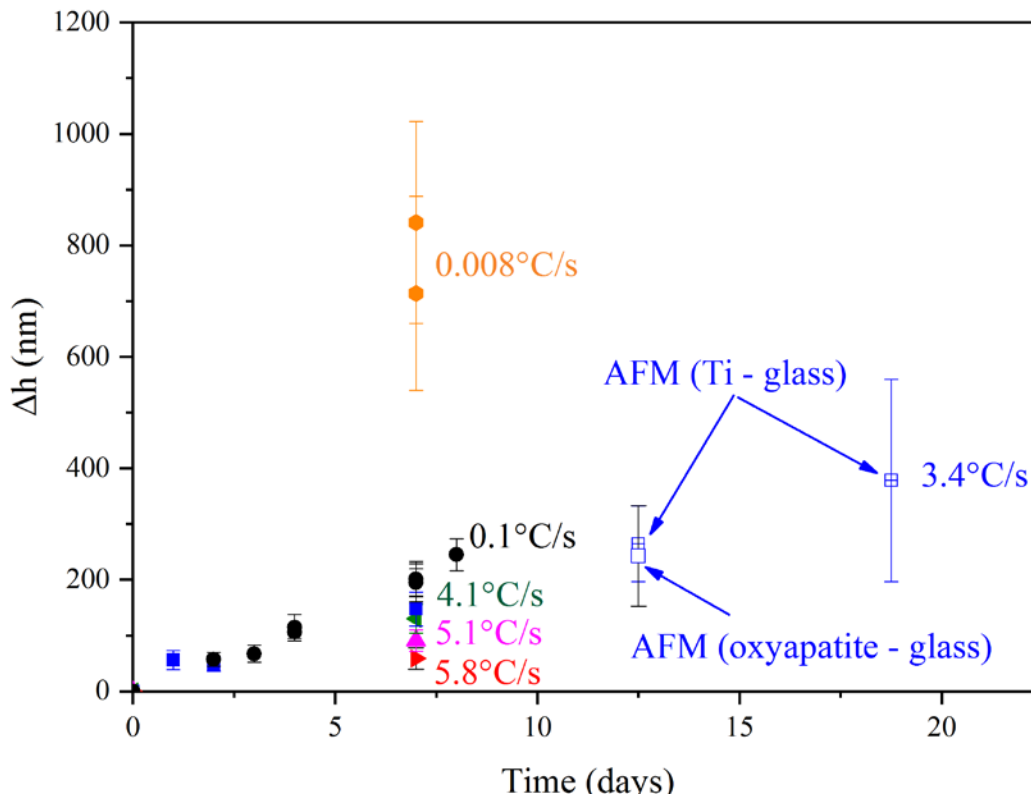


Figure 9. Profilometry (solid symbols) and AFM (open symbols) results in Δh as a function of time.

Simple weight measurements, normalized to surface area, made before and after the reaction shows a similar trend with increasing dissolution rates from samples quenched at slower rates, as shown in Figure 3.10. However, this is only up to the sample at 0.1°C/s and at 0.008°C/s the weight loss rate decreases, similar to the PCT test. The data for the 0.1°C/s and 3.4°C/s were fit to a parabolic function with the equation also listed in Figure 10. The weight loss measurements can be found in Table B.3 Appendix B.

Figure 3.11 shows micrographs of the surface of a CCC-quenched sample after one week in 90°C deionized water that clearly show pits left after the removal of a cross-like powellite crystals. The oxyapatite crystals remain. EDS analyses (not shown) reveal that the sub-micron crystals on the surface of this sample are rich in Zr.

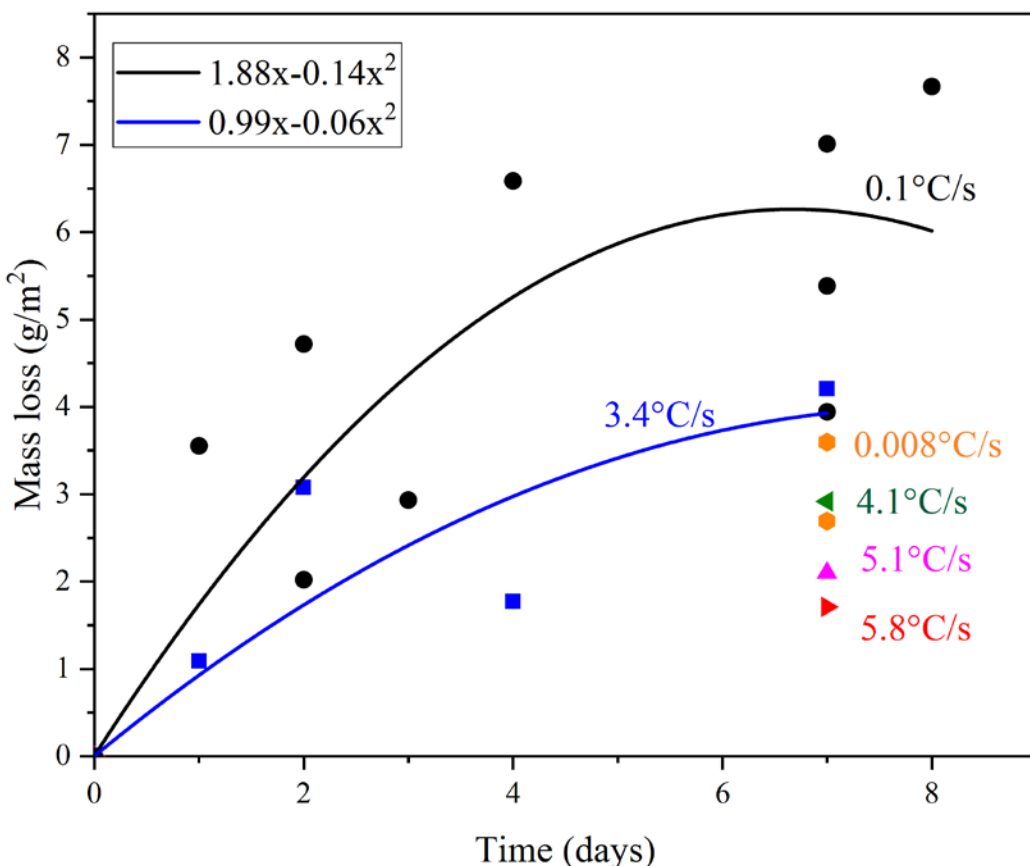


Figure 3.10. Mass loss from sample weight measurements in grams. The data for the 0.1°C/s and 3.4°C/s samples were fitted to a parabolic.

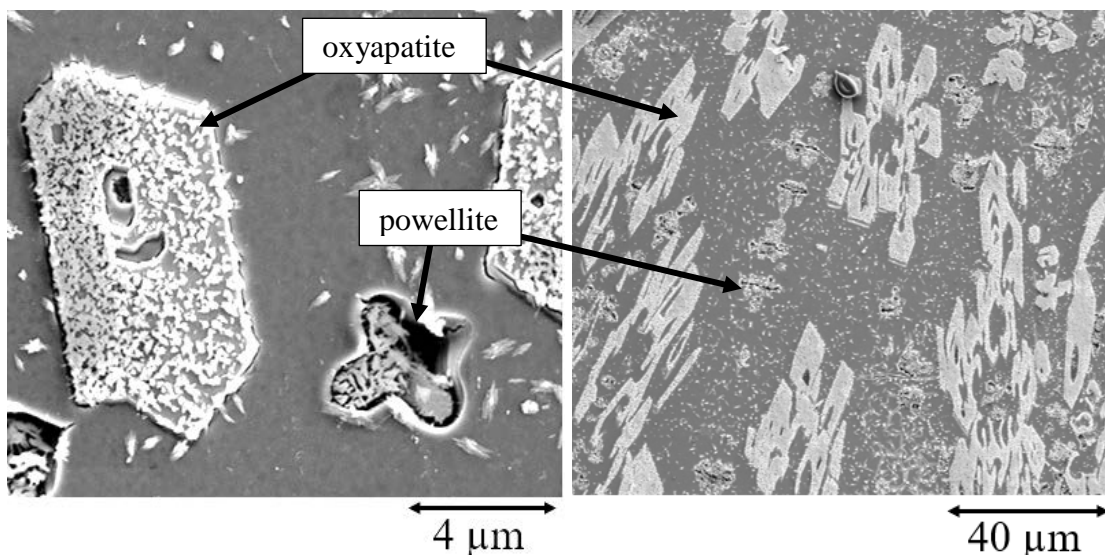


Figure 3.11. SEM images of a CCC sample after seven days in 90°C deionized water, showing left) a powellite and a oxyapatite crystal and right) a lower magnification image showing more representative sizes of the powellite and oxyapatite crystals.

The difference in height between the Ti layer, oxyapatite, residual glass, and molybdate phase was believed to be found using profilometry in the CCC sample due to the relatively large crystals, Figure 3.12. The step heights from the Ti layer (initially 150 nm) to the respective crystals post dissolution in DI water at 90°C for 7 days for the CCC sample were also calculated and are shown in Figure 3.12. The Ti layer is represented by the grey area shown being 150 nm thick. The step height from the Ti-layer to the oxyapatite crystals is shown with the red area, to the residual glass with the green area, and to powellite with the blue area. These measurements again show that oxyapatite dissolved out slowest followed by the residual glass followed by powellite. The change in height from the residual glass to the powellite crystals was found to be $\sim 1 \mu\text{m}$ which is in range with the size of a powellite crystal. The widths of the peaks match well with the size of oxyapatite and the width of the low valleys match well with the size of the powellite crystals as well shown in Figure 3.13 and Figure 3.11 right.

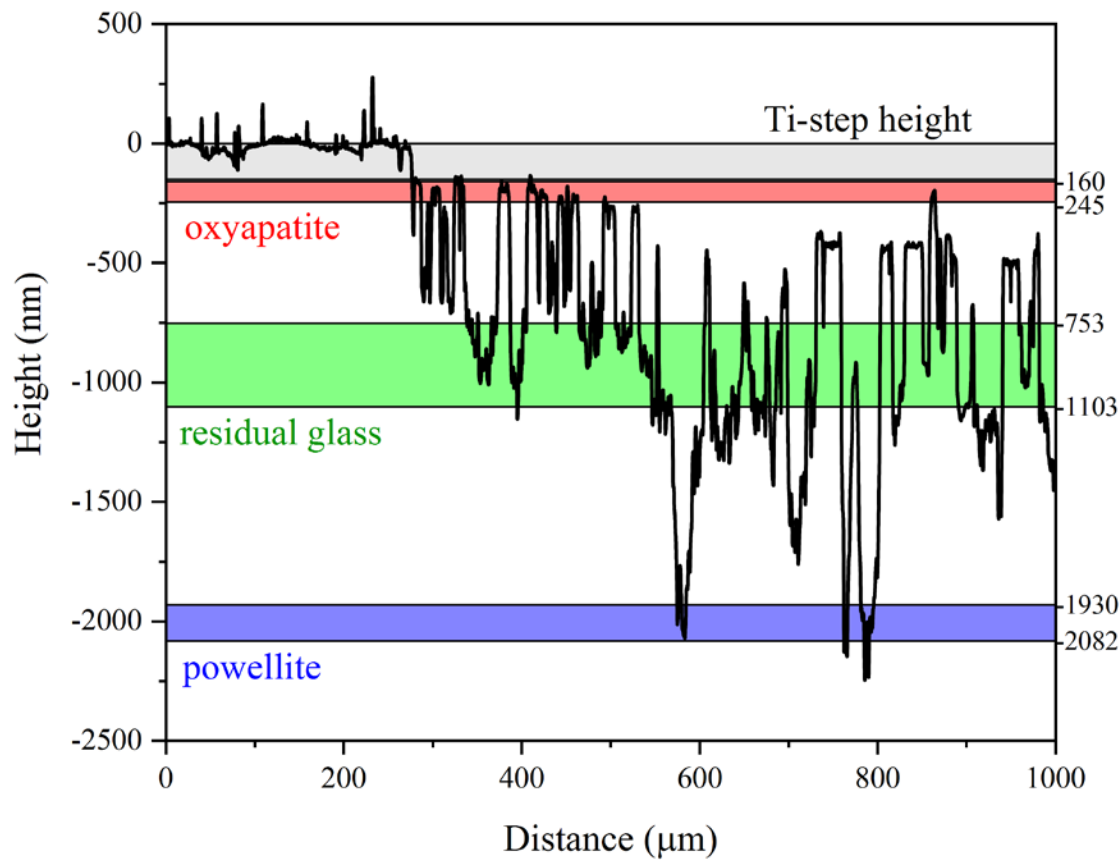


Figure 3.12. Profilometry scan of the CCC sample after 7 days in 90°C DI water.

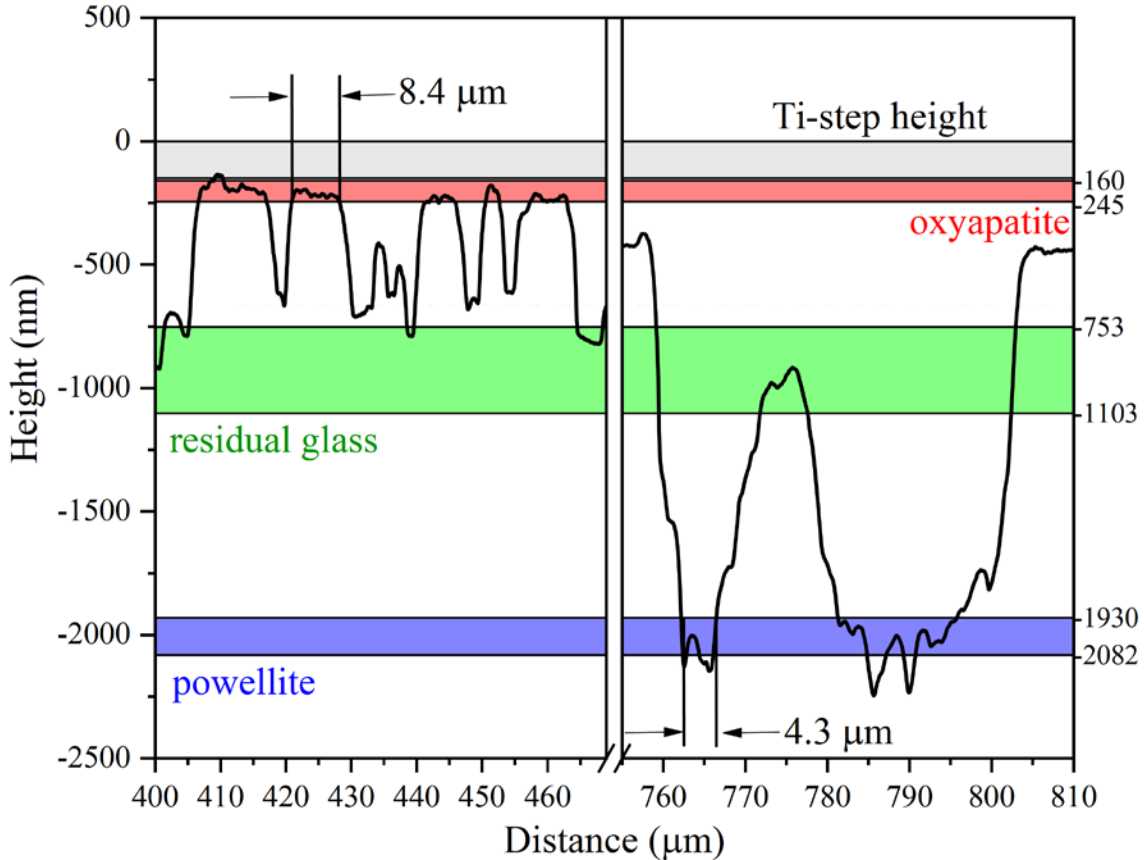


Figure 3.13. Expanded regions from Figure 3.12.

4. DISCUSSION

The absolute ion-release rates from the PCT experiments (Figure 3.4) were about five times faster than those reported by Crum et al. [16], listed in Table B.1 of Appendix B. However, the relative ion-release rates were found to follow a similar trend with ion release rates of $\text{Ba} \approx \text{Si} \approx \text{Ca} < \text{Mo} \approx \text{Na} < \text{B}$.

The change in the relative recession rate with quench rate can be associated with changes in the microstructure and associated changes in the composition of the residual glass. The residual glass compositions were calculated by subtracting from the nominal composition (Table 4) the oxide components associated with the powellite and oxyapatite

crystalline phases determined by quantitative XRD [17]; these estimated residual glass compositions are given in Table 3.7. The composition of the roller quench glass is estimated to have the same composition as the as-received glass. Asmussen et al. [15] reported a similar residual glass composition of the CCC sample, as shown in Table 3.7. B (glass former) and Na (glass modifier) are predominantly retained in the residual glass and increase in relative concentrations in samples that are cooled more slowly and so have greater crystal fractions. The growth of oxyapatite crystals removes lanthanides (Nd) and some Ca and Si from the residual glass; little to no Nd is expected to remain in the residual glass of the 0.1°C/s and CCC samples, the latter in agreement with Asmussen et al. [15]. Increasing fractions of powellite, (Ca-rich)MoO₄, crystallize from the melts with decreasing cooling rate, pulling mostly Mo and some of the alkaline earth ions from the residual glass. Based on the quantitative XRD results, some Mo remains in the residual glass of the sample cooled at 0.1°C/s but is mostly sequestered from the glass phase in the CCC sample (Table 3.7).

Table 3.7. Estimated compositions of the residual glass phase in glass-ceramic samples created with different quenching rates, the analyzed composition from the CCC sample, and the estimated composition reported by Asmussen et al. [15].

Oxide	roller quench [At. %]	3.4°C/s [At. %]	0.1°C/s [At. %]	CCC [At. %]	CCC EMPA* (At. %)	Asmussen [15] CCC [At. %]
Al ₂ O ₃	3.8	3.9 ± 0.1	3.8 ± 0.5	4.5 ± 0.2	5.62	4.7
B ₂ O ₃	11.3	12.1 ± 0.2	14.5 ± 1.2	13.4 ± 0.5	16.41	14.2
BaO	2.2	2.2 ± 0.1	2.2 ± 0.5	2.5 ± 0.1	3.09	2.8
CaO	8.4	7.6 ± 0.3	5.7 ± 1.5	5.6 ± 0.9	3.66	4.3
Ce ₂ O ₃	1.5	1.2 ± 0.1	1.2 ± 1.1	0.9 ± 0.2	0.23	0.5
Cs ₂ O	1.6	1.6	1.8 ± 0.2	1.9 ± 0.01	1.29	2
La ₂ O ₃	0.8	0.64 ± 0.1	0.6 ± 0.5	0.6 ± 0.1	0.26	0.4
Li ₂ O	3.2	3.5	4.2 ± 0.4	3.9 ± 0.1	--	4.1
MoO ₃	4.2	4.2 ± 0.1	3.6 ± 0.6	1.2 ± 0.9	0.22	0.0
Na ₂ O	4.1	4.3 ± 0.1	4.1 ± 1.6	4.7 ± 0.2	5.32	4.8
Nd ₂ O ₃	2.4	2.0 ± 0.1	1.7 ± 0.5	1.3 ± 0.3	0.32	0.9
Pr ₂ O ₃	0.7	0.6 ± 0.1	0.5 ± 0.5	0.3 ± 0.1	--	0.5
SiO ₂	47.7	48.0 ± 1.0	47.7 ± 5.9	51.1 ± 2.1	57.49	52.5
SrO	1.5	1.5 ± 0.1	1.4 ± 0.5	1.0 ± 0.2	1.00	0.5
ZrO ₂	3.8	3.9 ± 0.1	4.2 ± 0.4	4.4 ± 0.2	5.11	4.7
other	2.8	2.7 ± 0.1	3.0 ± 0.3	2.5 ± 0.2	--	3.1
TOTAL	100	100	100	100	100	100

*Analyses provided by Washington State University

Nd has the lowest relative release rates, Figure 3.4, in these samples, indicating that the oxyapatite phase is more chemically durable than the other phases. As a function of quench rate, the release of Nd did not appear to change much, if any at all due to the release rates being near the detection limit of the ICP-OES, even in the sample where Nd was situated in the residual glass, Figure 3.14. Because Nd was also not released into solution when incorporated in the less durable residual glass, the release rate of Nd would be most likely low compared to the actual release rate of oxyapatite, in agreement with Asmussen et al [15]. The release rate of ions from the oxyapatite crystal may not be congruent with each other, allowing certain elements to preferentially dissolve out resulting in an increased dissolution rate of oxyapatite compared to Nd release rate. AFM scans, Figure 3.6, and SEM micrographs, Figure 3.5 and Figure 3.11 of oxyapatite crystals post dissolution

clearly show oxyapatite to be more durable than the residual glass. Also, it is believed that the profilometry scans show peaks where oxyapatite crystals are located, Figure 3.12 and Figure 3.13, suggesting this is the most chemically durable phase.

Aside from Nd, from the fast-cooled roller quenched glass sample to the sample cooled at 0.1°C/s, ion release rates were found to increase, described by the PCT (Figure 3.4). Also, Δh profilometry values were found to increase, (Figure 3.9), and the mass loss was found to increase (Figure 3.10). These three tests confirm that the chemical durability of the glass-ceramic decreases with decreasing quench rate. Results from the profilometry experiments suggests this trend continues with even slower release rates. However, the ion release rates and mass loss were found to decrease from the 0.1°C/s to the 0.008°C/s sample. This could partly due to the change in the components of the residual glass. For example, Crum et al. [16] found increases in MoO_3 and Ln_2O_3 in the glass to always increase the dissolution of B and Na, Table 3.3. Therefore, in comparing the decrease in ion release rates and mass loss with quench rate from the sample cooled at 0.1°C/s to the sample cooled at 0.008°C/s, as more Mo and lanthanides crystallize out into powellite and oxyapatite respectively, B and Na release rates would decrease.

The general trends discussed by Crum et al and summarized in Table 3.3 can be simplified by comparing the change in B_2O_3 , Al_2O_3 , and SiO_2 in the residual glass composition with the change in B ion-release rates [16]. Shown in Figure 3.14, the B ion release rate was found to increase with the increase in B_2O_3 . As stated before, B is situated in the residual glass regardless of cooling rate as an indicator for the recession rate of the residual phase. The increase in the release rate from the PCT test can be described by the B_2O_3 ratio as a function of quench rate, listed in Table 3.7. From the roller quench to the

0.1°C/s sample the B_2O_3 concentration in the residual glass significantly increases which then attributes to the higher B release rate and less durable glass.

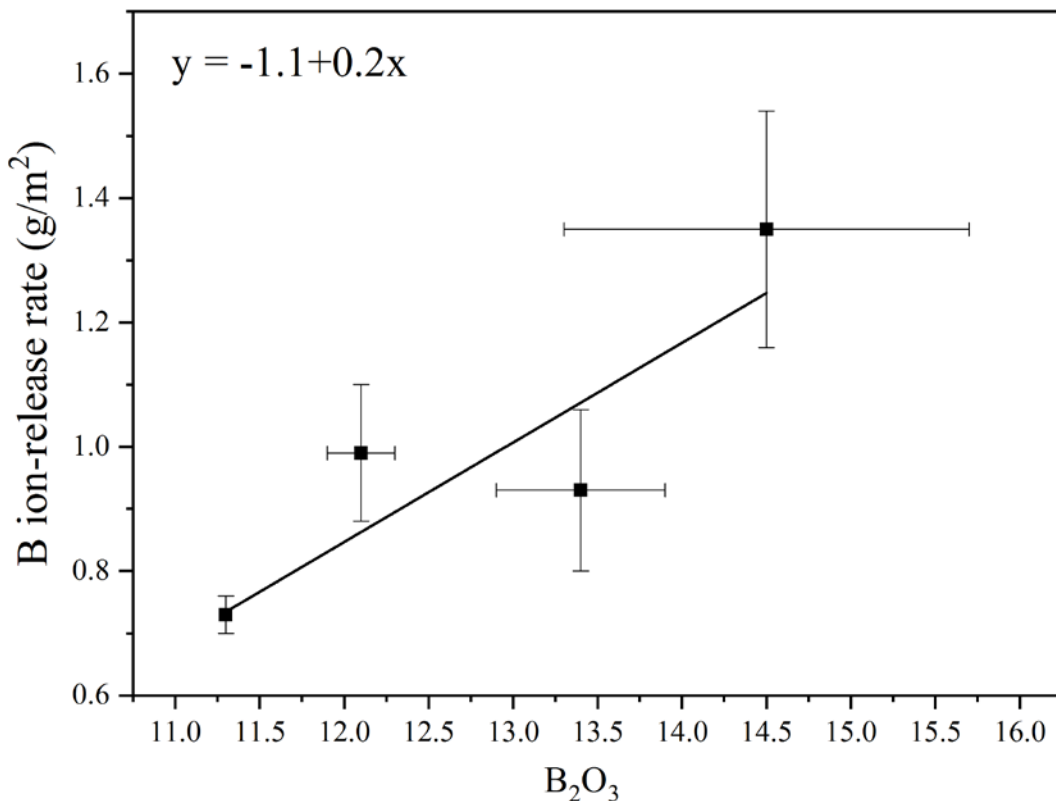


Figure 3.14. B ion release rate vs B_2O_3 concentration in the residual glass.

A less durable glass would recede faster, increasing water exposure to the molybdate phases. This added exposure allows more molybdate phases to dissolve which were described by Crum et al. [16] to not be a durable phase. Due to the relatively high release rates of Mo ions into solution, Mo release rates can act as indicators of the release rate for the molybdate phase. Other components such as Ba and Sr are also located in the molybdate phases but not always. Ba molybdates were found in most continuous cooling experiments, except the CCC sample, the reason for the increase in Ba concentration in the residual glass from the 0.1°C/s to the CCC sample, Table 3.7.

The reason for the decrease in the Mo ion-release rate from the 0.1°C/s to the CCC sample can be explained using the B₂O₃ concentration in the residual glass. With decreasing cooling rate, from 0.1°C/s to 0.008°C/s, the concentration of B₂O₃ greatly decreases which would cause an increase in the chemical durability. Slight differences in the morphologies and compositions of the molybdate phases between these samples could also attribute to changes in the dissolution behavior. However, it is believed that the composition of the residual glass, has the greatest influence on the chemical durability. In the CCC sample, the relatively durable residual glass does not recede as fast, limiting exposure of water to only the powellite crystals near the surface, which were found to completely dissolve out (Figure 3.11). Due to the powellite crystals being relatively large in the CCC sample, the dissolved particles left behind deep pits that greatly increased the step height measured with profilometry, Figure 3.9.

At quench rates, from the roller quench to the 0.1°C/s sample, ion-release rates increase due to the precipitation of oxyapatite crystals which increased the B₂O₃ concentrations by removing components like silica in the residual glass [16]. A decrease in the ion-release rates and sample mass is seen from the 0.1°C/s to 0.008°C/s samples described by composition of the residual glass having a reduced amount of B₂O₃.

This study provided additional information about the complex chemical durability of this borosilicate glass ceramic as a function of thermal history. The dissolution behavior of a wider range of quench rates and microstructures was examined. This study is intended to be used to help optimize the glass ceramic.

3.5. CONCLUSION

Ion release rates and recession rates were determined as a function of microstructure of a borosilicate high-level waste glass-ceramic. Eight samples were continuously cooled with quench rates ranging by about five orders in magnitude. The residual glass was found to change as crystals precipitated resulting in differing dissolution behaviors described by the B_2O_3 concentration in the residual glass. Mo release rates increased from the roller quenched glass, to the sample cooled at $0.1^{\circ}C/s$ but then decreased and was lowest in the CCC sample ($\sim 0.005^{\circ}C/s$). Results from this study clearly showed Nd was found to have very low release rates correlating the high relative chemical durability of oxyapatite, confirmed by SEM and AFM. SEM also found whole powellite crystals completely dissolved out in the CCC leading to believe the durability of powellite is less than the residual glass. Information from this experiment about the dissolution behavior of a borosilicate high-level waste glass-ceramic is intended to help assist in developing an optimized composition.

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4. SUMMARY AND FUTURE WORK

The objective of this study was to understand the effects of thermal history on the microstructural development and dissolution behavior of a borosilicate glass-ceramic loaded with waste from reprocessed spent nuclear fuel, and then to use this information to help optimize the design of these materials.

Continuous cooling and isothermal experiments provided data about the effects of thermal history on microstructure development, including the compositions, morphologies, and crystal percentages of the major crystalline phases. The major crystalline phase (up to 30 wt. %) was oxyapatite ($\text{Ca}_2\text{Ln}_8\text{Si}_6\text{O}_{26}$) and its formation affected the development of the second most abundant (up to 10 wt. %) crystalline phase, powellite ($\text{Ca}_{1-x-y}\text{Sr}_x\text{Ba}_y\text{MoO}_4$), and the composition of the residual glass. Insoluble heavy metal crystals, predominately Ru-rich, were also detected in trace amounts in the as-received and heat-treated composition.

On quenching from 1300°C, the waste-loaded melt first separates into aluminosilicate droplets and a matrix borosilicate liquid that contains high concentrations of lanthanides and transition metals, including Mo. This phase separated microstructure is clearly evident in glasses quenched from melts above about 1000°C at rates faster than about 4°C/s. Mo-rich droplets separate from the matrix liquid when melts are cooled slower than about 300°C/s, and these droplets grow in size with slower quenches and longer isothermal heat-treatment times. The droplets crystallize to form two types of powellite, a Ca-rich form ($\text{Ca}_{1-x}\text{Sr}_x\text{MoO}_4$) and a Ba-rich form ($\text{Ba}_{1-x}\text{Sr}_x\text{MoO}_4$). Needle-shaped lanthanide-rich molybdate crystals form in samples isothermally held at 1100°C, along with the drop-shaped alkaline earth molybdate crystals.

Oxyapatite crystals form in melts continuously cooled slower than about 4°C/s, or isothermally held for several minutes at temperatures between 700 and 1100°C. The growth of the oxyapatite crystals is accompanied by the evolution of the microstructure of the residual glass from a droplet-in-matrix morphology to a more homogeneous appearance. In addition, alkaline earth powellite ($\text{Ca}_{1-x}\text{Sr}_x\text{Ba}_y\text{MoO}_4$) crystals with cross-like morphologies form with the oxyapatite crystals, although it is unclear if the cross-like powellite precipitates from the residual glass or results from the transformation of pre-existing powellite droplets. The cross-like powellite crystals were present only in samples held isothermally from 900°C to 1100°C, and in the sample cooled most slowly, following the canister centerline cooling (CCC) protocol. From the isothermal experiments, crystallization was fastest at 1000°C, where crystalline powellite and oxyapatite could be detected after a one-minute isothermal hold.

The effects of microstructure on dissolution behavior were characterized using several short-term static dissolution tests. By using the Product Consistency Test (PCT), method A, the leach rates for Na, B, Ca, Mo, and Ba were found to all increase from samples with cooling rates that decreased from about 100°C/s to 0.1°C/s. These rates then decreased for the sample cooled most slowly (~0.008°C/s), following the CCC protocols. The Si leach rates did not change significantly from samples cooled at different rates, although it was lowest from the CCC sample. The Nd leach rates were lowest from all samples, remaining near the detection limit of the analytical system used.

The relative ion release rates indicate that the powellite phases dissolve most quickly from the glass-ceramic microstructures, followed by the residual glass phase, and then the oxyapatite phase. Topological analyses of corroded surfaces using electron

microscopy, atomic force microscopy, and profilometry were consistent with the trends in ion release rates. Changes in the composition of the residual glass with the precipitation of the powellite and oxyapatite phases affect the dissolution rate of the residual glass. As the B_2O_3 concentration in the residual glass was found to increase, the chemical durability of the glass decreased.

The following paragraphs describe technical issues identified in this work that require further study to resolve.

1. The distributions of boron in the residual glass and any crystalline phases need to be better determined. B_2O_3 is a significant (11.3 mole%) component of the starting glass but does not appear to be present in any of the major crystalline phases and could not be detected in the residual glass using the EDS systems at Missouri S&T. EMPA analyses done at Washington State University did show trends in the B-contents of the residual glass phase that were consistent with predictions from the EDS analyses.
2. The lanthanide-rich molybdate needles detected in the high temperature ($>1000^\circ C$) isothermal experiments need to be fully characterized. No specific diffraction data has been associated with these needles, and Raman spectra are inconsistent with the spectra collected from the other molybdate phases, despite the presence of significant concentrations of Mo from the EDS data. Preliminary association of these needles to a reduced molybdate phase needs to be confirmed.
3. Additional isothermal heat treatments should be performed, including tests for long times, >24 hours, to try to produce samples with the maximum amount crystallization at different temperatures. These tests will also confirm if there are any additional slow growing crystals.

4. A more in-depth analysis of the Raman spectra coupled with a more in-depth analysis on the residual glass composition may provide some reasons for the shifts in the vibrational frequencies for certain samples, which may offer some additional insight to the phase transformation pathways.
5. More ion-release tests should be performed, especially on samples that were isothermally treated at higher temperatures, where larger crystals would mean more quantitative individual recession rates.
6. The relative release rates of Mo from the residual glass and from the molybdate phases still should be determined.
7. It would also be helpful to understand the dissolution rates as a function of solution pH.

APPENDIX A: SUPPLEMENTAL INFORMATION FROM THE PHASE TRANSFORMATION STUDY

Measured Cooling Profiles

The measured cooling profile for the constant cooling experiments, except the roller quench are shown in Figure A.1. The measured cooling profiles never reach 1300°C, the temperature at which the samples were melted. This is because the samples cooled fast when the crucibles were moved from the furnace to the mold with the thermocouples. The cooling profile of the air-cooled sample was not as linear as the other cooling profiles. The uneven cooling profile in the first 25 seconds of the air-cooled sample is due to movement of the thermocouple in the sample. Figure A.2 shows the comparison between the furnace program temperature and the measured temperature of a thermocouple held closely to the glass surface of the CCC sample. The temperature appears to have fully converged with the program temperature after 20 minutes and at ~1050°C. The sharp jumps in temperature near ~1 min are from transferring the sample from the melting furnace to the cooling furnace. The sharp jumps are not associated with the temperature of the sample, just the temperature of the thermocouple since the thermocouple was brought away from the sample in order to transfer through the furnace door.

Differential Thermal Analysis

Differential Thermal Analysis, DTA, (Perkin Elmer DTA 7) was used to determine the glass transition temperature and exothermic transitions that occur upon heating. Figure A.3 shows the glass transition temperature (T_g) occurring at 590°C and the onset

temperature three exothermic reactions (T_{x1} , T_{x2} , T_{x3}) occurring at 710°C, 815°C, and 915°C.

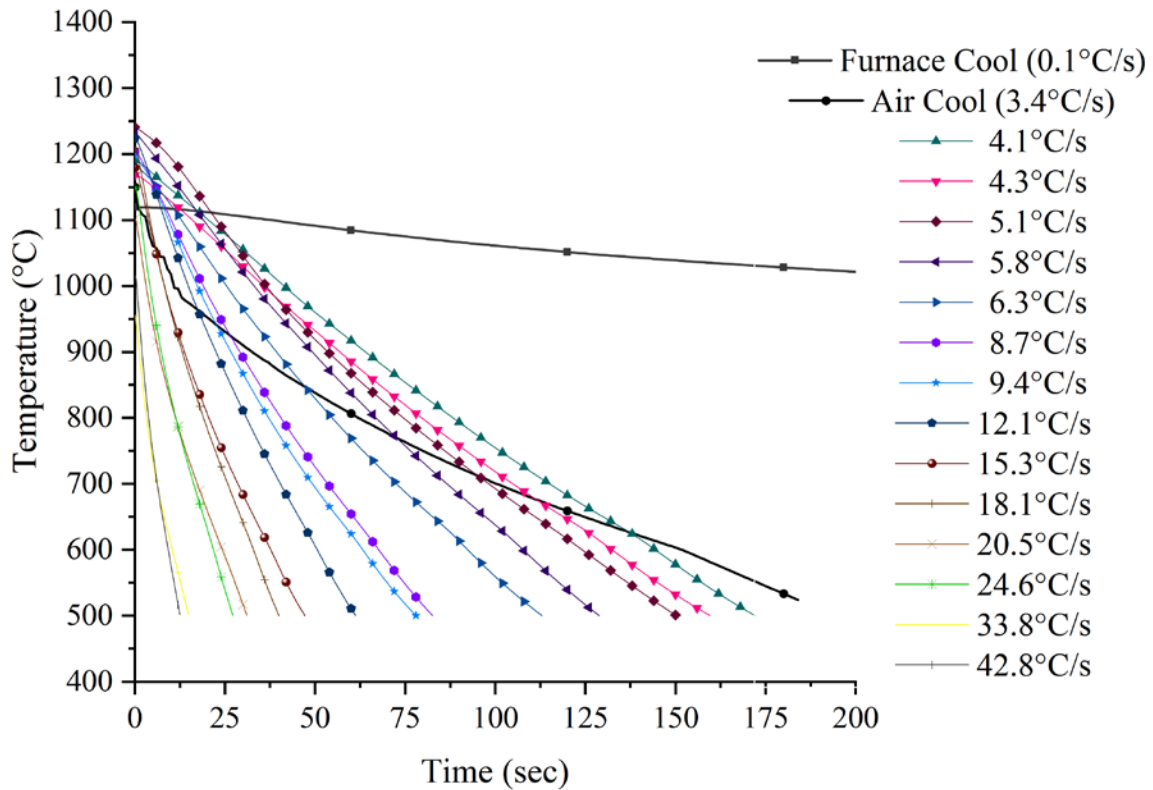


Figure A.1. Measured cooling profiles for the samples from the wedge mold, air cool, and furnace cool cooling methods.

X-ray Diffraction

Samples with fast quench rates ($\geq 4.1^{\circ}\text{C/s}$) and short times in the isothermal experiments have a lone peak at 27.65° (2θ) making peak assignment difficult, Figure A.4. This peak is believed to be however associated with a molybdate phase instead of RuO_2 (the heavy metal crystal that do not dissolve in the glass). Additional shorter peaks at 29.58, 33.2, and 45.05° (2θ) align much closer with the molybdate phase (e.g., $\text{Sr}_{0.44}\text{La}_{0.39}(\text{MoO}_4)$, PDF# 04-002-4182) vs the RuO_2 phase (PDF# 00-040-1290) making this phase the assignment for the 27.65° (2θ).

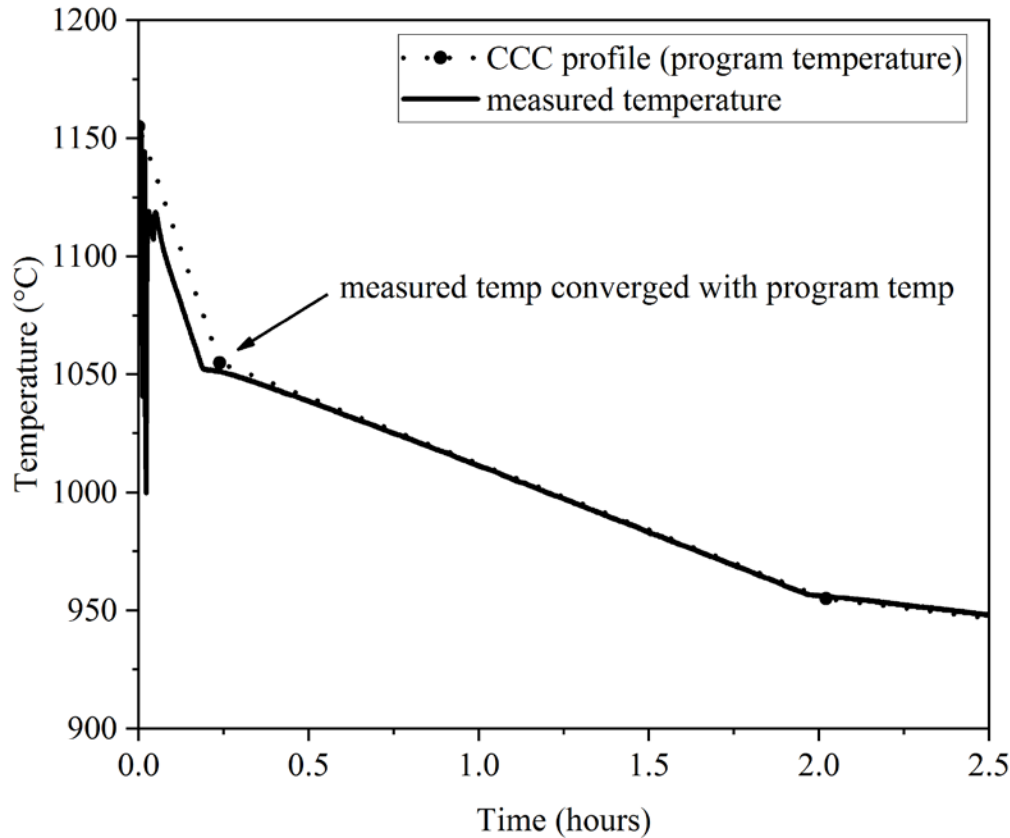


Figure A.2. Cooling profile comparison for the CCC sample between the (solid line) measured cooling profile from a thermocouple held above the sample and (dotted line) furnace program temperature set to follow the CCC profile.

The peak at 27.65° (2θ) also appears to grow in the isothermal samples heat treated at 600°C , also shown in Figure A.4. This peak could be associated with the Mo-rich droplets since the size of the Mo-rich droplets in the baseline sample was measured to be $0.07 \pm 0.02 \mu\text{m}$ and measured to be $0.14 \pm 0.03 \mu\text{m}$ after being held for 16 minutes at 600°C . The peak at 40.5° 2θ is believed to be due to contamination from melting the samples in stainless steel tubes.

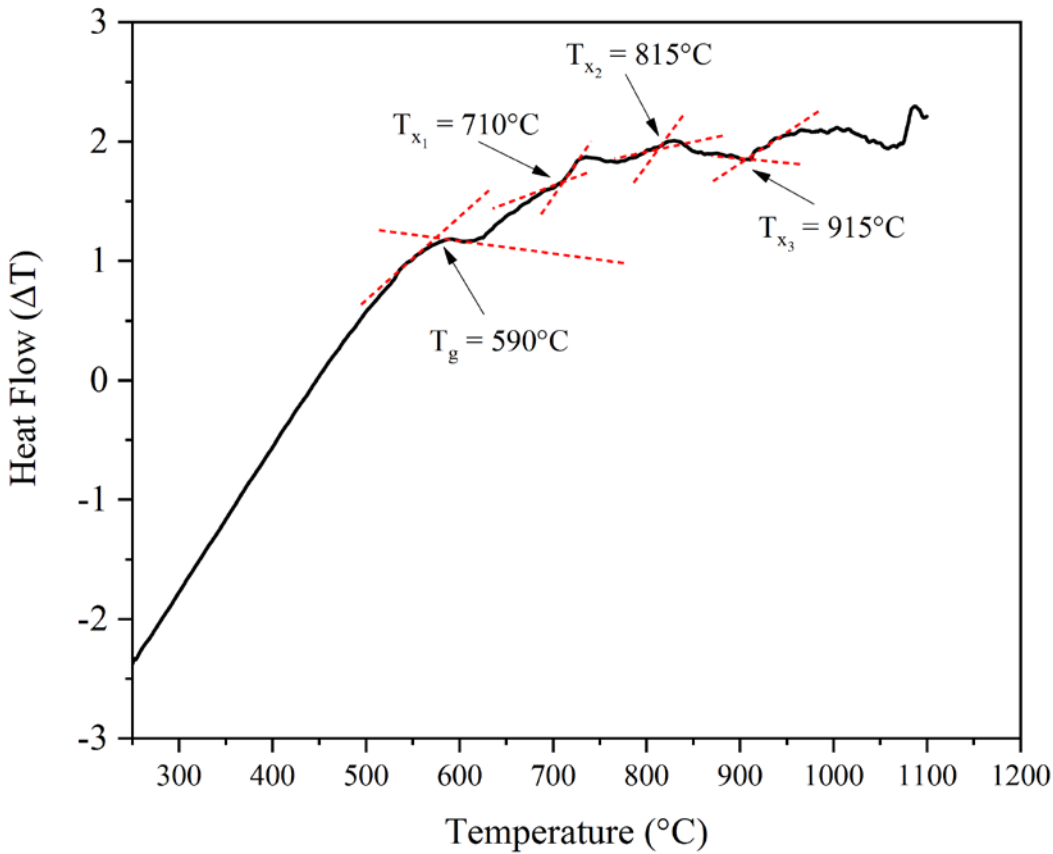


Figure A.3. DTA data collected of the as-received glass with labeled glass transitions (T_g) and onset of crystallization temperatures (T_x).

Tin Bath

Further tests were done to determine whether or not the microstructures were "reset" by only melting at 1300°C for 90 seconds in the isothermal experiments compared to 60 minutes in the continuous cooling experiments. XRD results of a quenched sample held at 1300°C for 90 seconds showed no additional peaks, actually showing less peaks than the as-received sample, Figure A.4. However, SEM did reveal that there Mo-rich droplets (Figure A.5) but image analysis determined the droplets to be smaller than the droplets in samples quenched at slower rates. There is a slight shift between the electron image and the molybdenum map in Figure A.5 due to surface charging from the electron

beam. The droplets in the sample melted for 1300°C for 90 seconds and quenched at ~300°C/s were measured to be 0.07 ± 0.02 μm in diameter and a sample melted at 1300°C for 1 hour and quenched at 42.8°C/s was measured to be 0.17 ± 0.05 μm . The reduction in peaks from XRD and the smaller size Mo-rich droplets confirm that the microstructures in the samples quenched for 90 seconds at 1300°C were "reset".

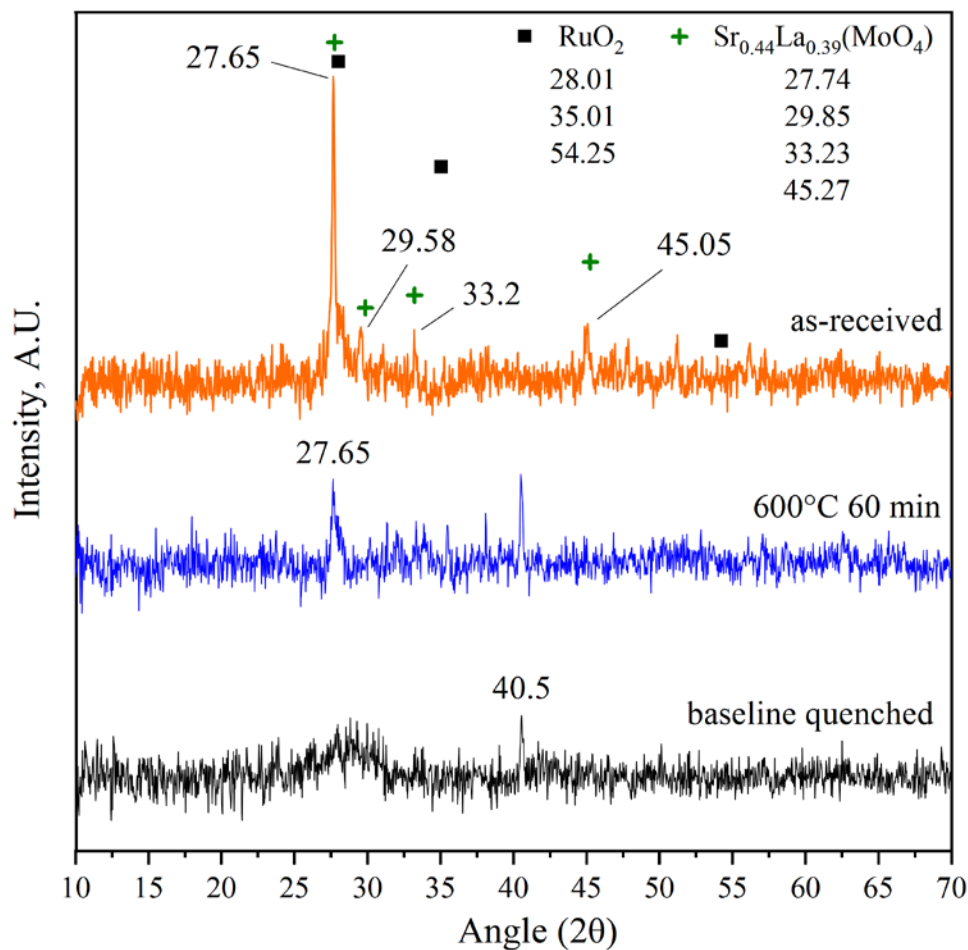


Figure A.4. XRD of the as-received glass, baseline sample, and sample heat treated at 600°C for 60 min with assessments.

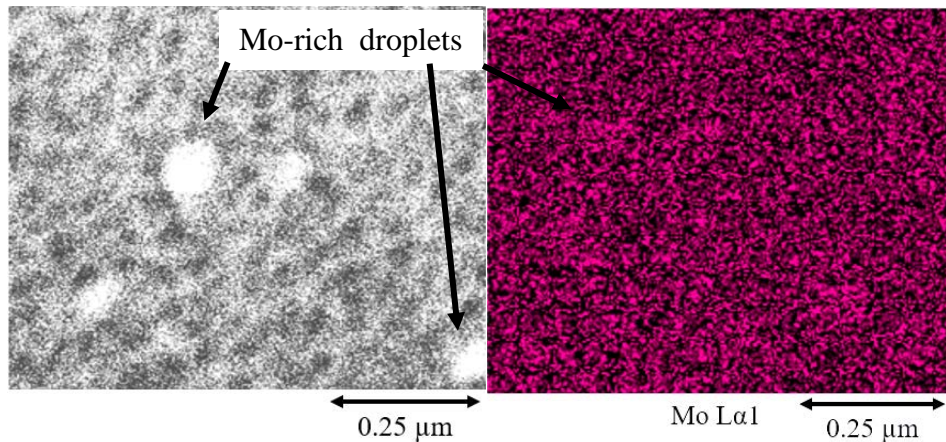


Figure A.5. SEM micrograph and molybdenum EDS maps for the baseline quenched ($\sim 300^{\circ}\text{C/s}$) sample.

It is believed that there was some contamination from the stainless-steel tubes since there is an extra peak in the XRD patterns from all the samples melted in stainless-steel versus samples melted in platinum. The small peak in the x-ray diffraction pattern at $\sim 40.5^{\circ}\Theta$ (Figure A.6) was found in all samples melted in stainless-steel and does not appear in the as-received or any samples from the continuously cooling experiments that melted in Pt crucibles. Although this phase does not correlate with XRD spectra from a used 304 stainless-steel tubes it could result from the stainless-steel species being in oxide form. Also, this contamination could be due to diffusion of Cr, Fe, and Ni into the glass. SEM EDS in the sample treated at 900°C for 24 hours, determined that these stainless-steel components diffused $\sim 20 \mu\text{m}$ into the sample, Figure A.7.

With the use of three different tin baths, additional experiments were done to determine variability in the results. One noticeable difference between the tin baths was the fluctuations in temperature of the small tin bath upon quenching the stainless steel samples. Figure A.8 shows an example of the temperature fluctuation that occurred in the

small volume tin bath ($\sim 250 \text{ cm}^3$) when a quench tube was immersed from the 1300°C melting furnace. The temperature initially increased by 15°C upon immersion, then decreased over the course of this 32-minute experiment. Over the entire experiment, the average temperature was 712°C , and this is the temperature reported for this experiment. The thermal fluctuations in the large volume tin baths were typically less than 1°C . To compare tin baths, samples were repeated using the same times and temperatures in each tin bath. Despite the different tin baths set-ups, the crystalline percentages of each sample were similar with only ~ 2 wt.% stdev, Table A.1. Variability in the Rietveld analysis and quantitative XRD procedures was also tested by running the same sample twice. Table A.2 shows the wt.% of oxyapatite and powellite detected in the air cooled sample run twice using the same XRD and Rietveld analysis. Figure A.9 shows an example of the matched patterns in the reitveld analysis in the sample isothermally heat treated at 900°C fro 24 hours. The subtracted background shows very little peaks proving the Reiveld analysis provides a good fit with the measured pattern. Table A.3 summarises the quantitative data from all the samples that contained crystals produced with the tin bath.

Composition Check

Samples from the continuous cooling experiment were sent to Jake Amoroso at Savannah River National Laboratories to measure the compositions after remelting in Pt at 1300°C for 60 min. Compositions were measured using ICP-AES and ICP-MS and are compared to the nominal composition. Cs was measured to be lower due to it volatilizing upon melting.

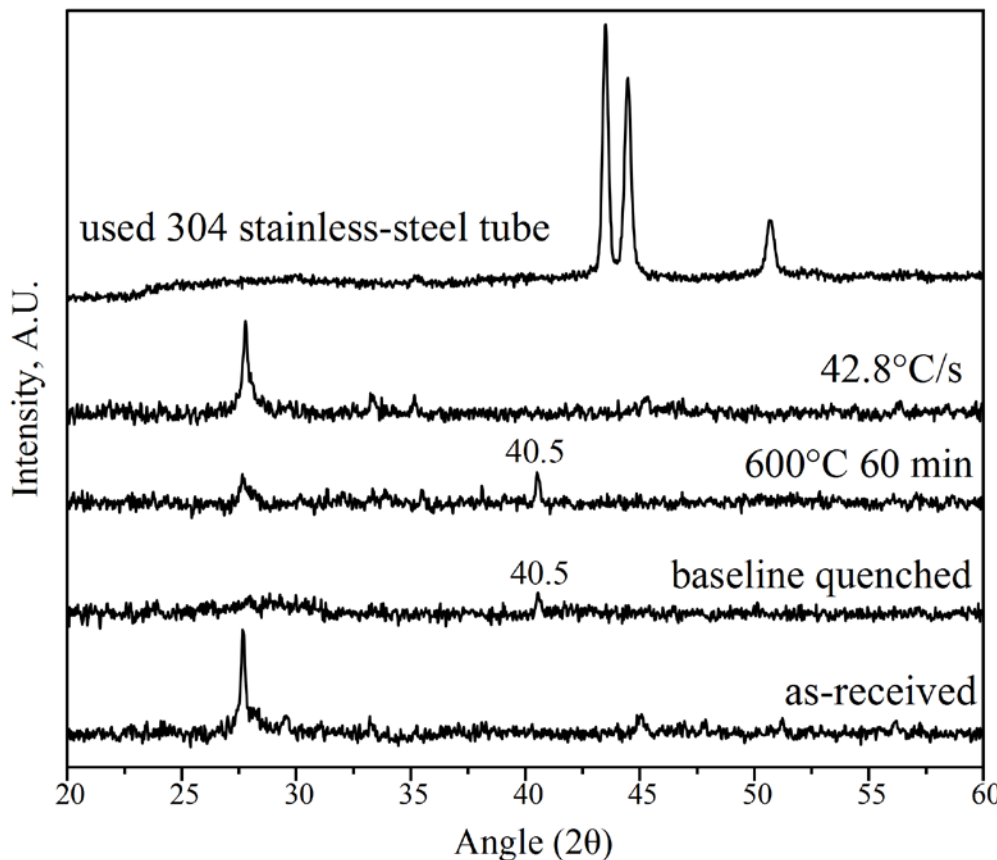


Figure A.6. XRD comparing the samples melted in stainless steel tubes, the baseline quenched sample and the sample heat treated at 600°C for 60 min, with the samples melted in Pt crucibles, the as-received glass and sample cooled at 42.8°C/s. An XRD pattern from the used stainless steel tube is also shown for comparison.

Powellite Crystallization Kinetics

The crystal fraction, normalized to the crystal fraction in the sample heat treated at 900°C for 24 hours, was determined by quantitative XRD and is plotted vs time in Figure A13. The Johnson–Mehl–Avrami–Kolmogorov (JMAK) model described in Section 3.4.3. was again used to determine the crystallization kinetics of the powellite crystals within the glass. The average slope of the double log Avrami plot was determined to be 0.52 ± 0.08 Figure A.14. A slope of -17.2 ± 9.2 was determined for $\ln(k_n)$ vs $1000/T$ plot from which

the activation energy was calculated to be 143.0 ± 76.8 kJ/mol, Figure A.15. Large error in the powellite activation energy correlates to the lower crystalline amounts, around the detection limits of the XRD and Rietveld analysis in most samples. The highest crystal percentage for powellite was measured to be only 6.7 wt.% by quantitative XRD for the isothermal samples. Also, there is a large increase in the size of powellite after the formation of oxyapatite further suggesting a change in mechanism for the growth of powellite with time. There also appears to be a change in growth mechanism with temperature due to the change in morphologies discussed in Paper I. Due to the lower crystal percentages and variations in growth mechanism the crystallization of powellite most likely cannot be described using the JMAK model.

TEM Diffraction Analysis

Fully indexed diffraction patterns of the 2 different zone axes of the Mo-rich droplets in the 4.1°C/s are shown in Figure A.16 and A.17. Figure A.18 compares the diffraction pattern with $\text{Sr}_{0.44}\text{Ln}_{0.39}(\text{MoO}_4)$ (blue), $\text{Ba}_{0.75}\text{Sr}_{0.25}(\text{MoO}_4)$ (green), and CaMoO_4 (red) which were assigned to the XRD patterns (Section 3.3.1.). The c/a lattice parameter ratios between the diffraction patterns are listed in Table A.4. These crystals all belong to the tetragonal crystal system ($I\ 4_1/a$ space group). The deviations in the lattice parameters most likely correlate to differences in alkaline earth and LN amounts within the crystal.

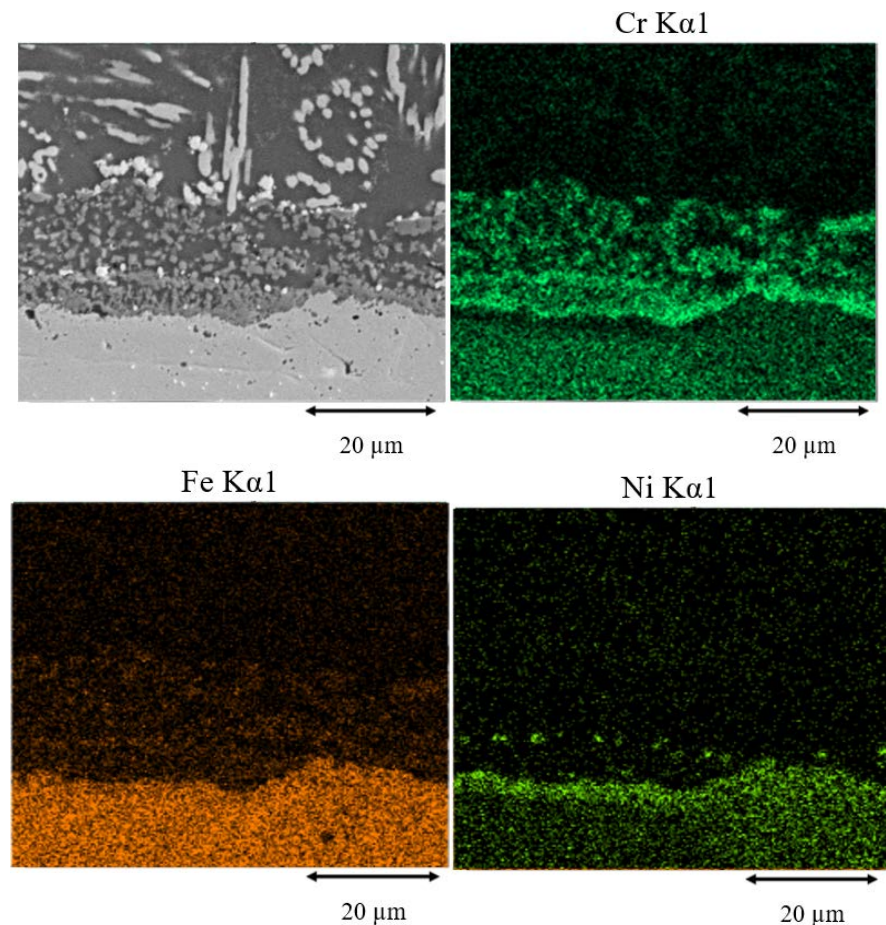


Figure A.7. SEM micrograph and EDS maps of the stainless steel diffusion into the glass in the sample heat treated at 900°C for 24 hours.

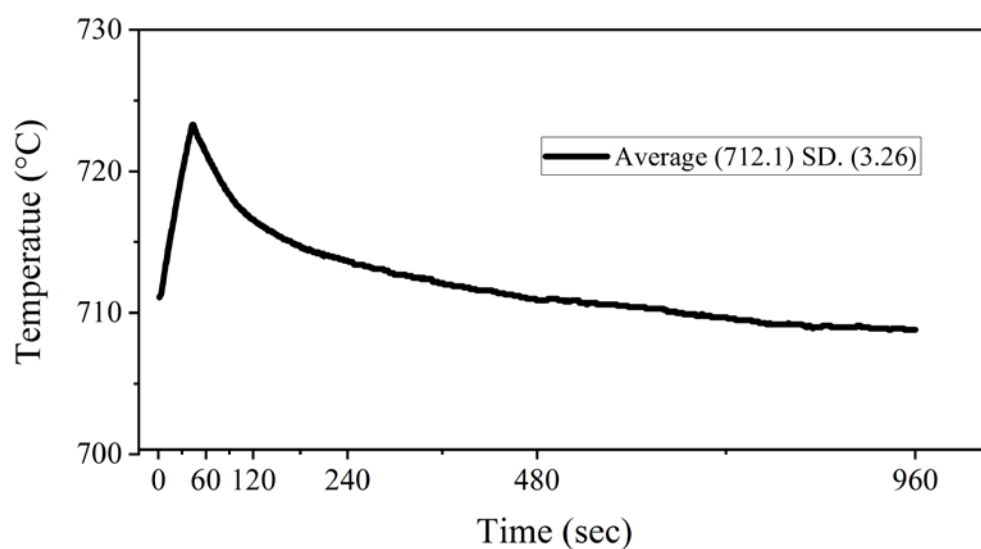


Figure A.8. Temperature-time profile during a quenching experiment in the low temperature tin bath.

Table A.1. Wt.% for oxyapatite and powellite for a sample held at ~800°C for 16 min in each tin bath.

Tin Bath	Oxyapatite [wt.%]	Powellite [wt.%]	Total [wt.%]
800°C for 2 min			
Small	8.7	1.4	10.1
Kiln	2.0	0.0	2.0
Average	5.4	0.7	6.0
Std. Dev.	4.7	1.0	5.7
800°C for 16 min			
Small	18.7	4.2	22.8
Kiln	19.2	0.9	20.1
Annealer	22.5	1.3	23.8
Average	20.1	2.1	22.2
Std. Dev.	2.1	1.8	1.9

Table A.2. Wt.% for oxyapatite and powellite in the air-cooled sample re-run twice with the XRD and Rietveld analysis.

Sample	Oxyapatite [wt.%]	Powellite [wt.%]	Total [wt.%]
Air cooled -1	9.3	1.5	10.8
Air cooled -2	7.5	0.9	8.4
Average	8.4	1.2	9.6
Std. Dev.	1.3	0.4	1.7
900°C 4 min -1	20.3	3.9	24.2
900°C 4 min -1	11.8	2.5	14.3
Average	16.1	3.2	19.3
Std. Dev.	6.0	1.0	7.00

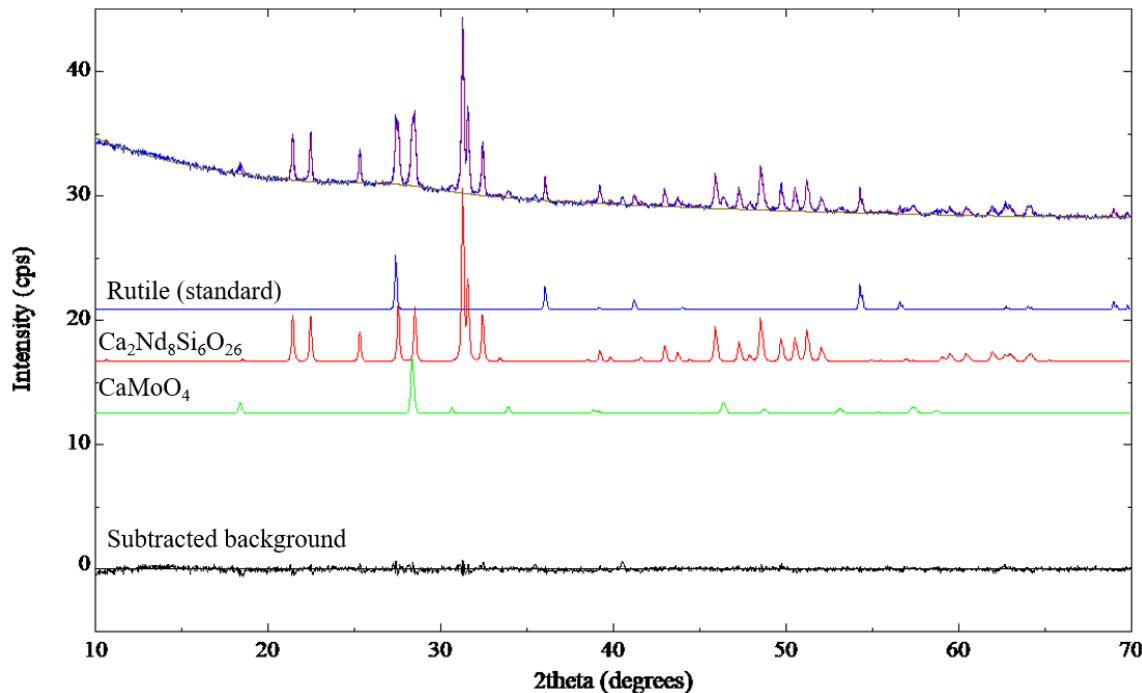


Figure A.9. Pattern matching and subtracted background from the Reitveld analysis.

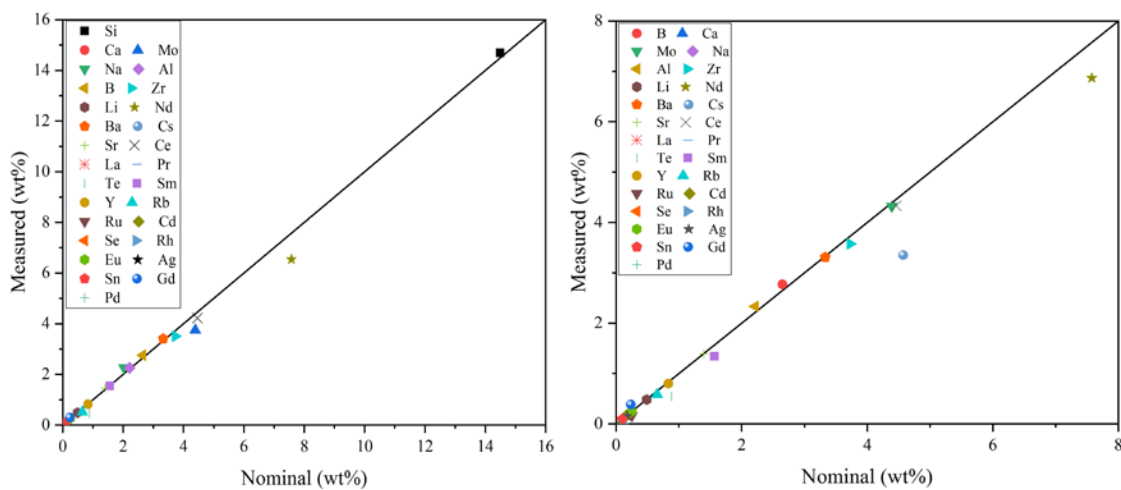


Figure A.10. Measured vs nominal composition of the roller quenched sample. Left) measured with ICP-AES and right) measured with ICP-MS

Table A.3. Quantitative XRD results.

Temperature (°C)	Time (sec)	Ca ₂ LN ₈ Si ₆ O ₂₆	Ca (MoO ₄)	Amorphous Content	Crystal Content
698.83	240	1.4	-	98.6	1.4
731.03	480	5.5	0.5	94.0	6.0
712.1	960	8.8	0.8	90.4	9.6
715.96	1920	12.2	1.2	86.6	13.4
698.83	3600	13.8	1.0	85.2	14.8
794.06	120	2.2	-	97.8	2.2
813.30	240	8.7	1.4	89.9	10.1
800	240	2.0	0.0	98.0	2.0
813.10	480	14.6	1.9	83.5	16.5
812.240	960	18.7	4.2	77.1	22.9
800	960	22.50	1.30	76.2	23.8
800	960	19.20	0.90	79.9	20.1
805.90	1920	20.6	4.1	75.3	24.7
794.28	3600	18.7	4.0	77.3	22.7
900	60	1.6	-	98.4	1.6
900	120	4.4	0.7	94.9	5.1
900	240	8.1	3.5	88.4	11.6
900	480	20.3	3.9	75.8	24.2
900	480	11.8	2.5	85.7	14.3
900	960	19.1	1.6	79.3	20.7
900	1920	24.3	5.5	70.2	29.8
900	3600	22.0	4.9	73.1	26.9
900	86400	28.5	6.7	64.8	35.2
1000	30	0.6	0	99.4	0.6
1000	60	4.3	0.6	95.1	4.9
1000	120	5.1	0.9	94.0	6.0
1000	240	13.9	2.9	83.2	16.8
1000	480	22.2	5.1	72.7	27.3
1000	960	22.8	5.5	71.7	28.3
1000	3600	25.9	3.9	70.2	29.8
1100	240	4.3	-	95.7	4.3
1100	480	7.6	-	92.4	7.6
1100	960	10.1	-	89.9	10.1
1100	1920	11.3	-	88.7	11.3
1100	3600	10.6	-	89.4	10.6
Sample	Cooling rate [°C/s]	Ca ₂ LN ₈ Si ₆ O ₂₆	Ca (MoO ₄)	Amorphous Content	Crystal Content
Air cooled	3.5	7.5	0.9	91.6	8.4
Furnace Cooled	0.1	21.4	3.2	75.4	24.6
CCC	0.005	15.2	8.5	76.3	23.7

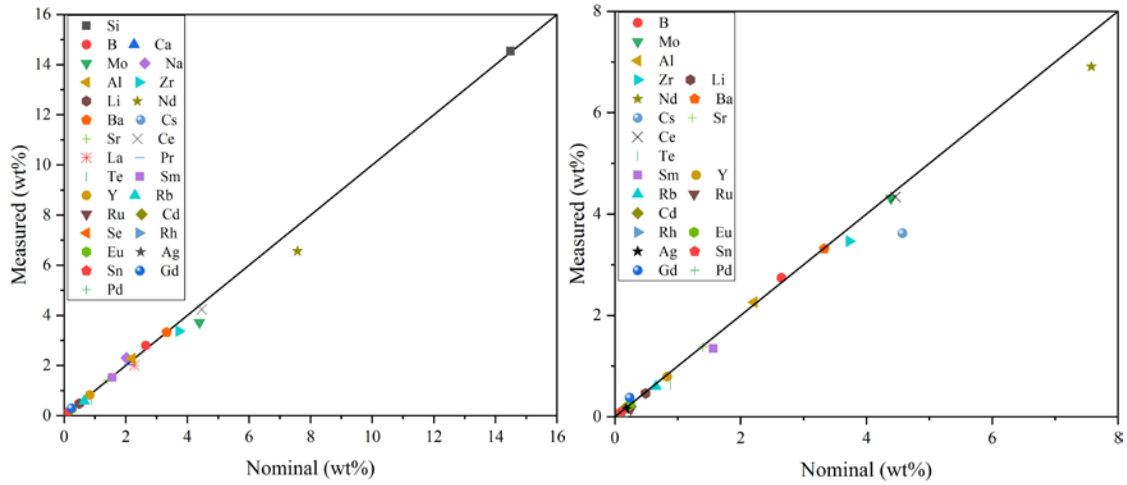


Figure A.11. Measured vs nominal composition of the 0.1°C/s sample. Left) measured with ICP-AES and right) measured with ICP-MS

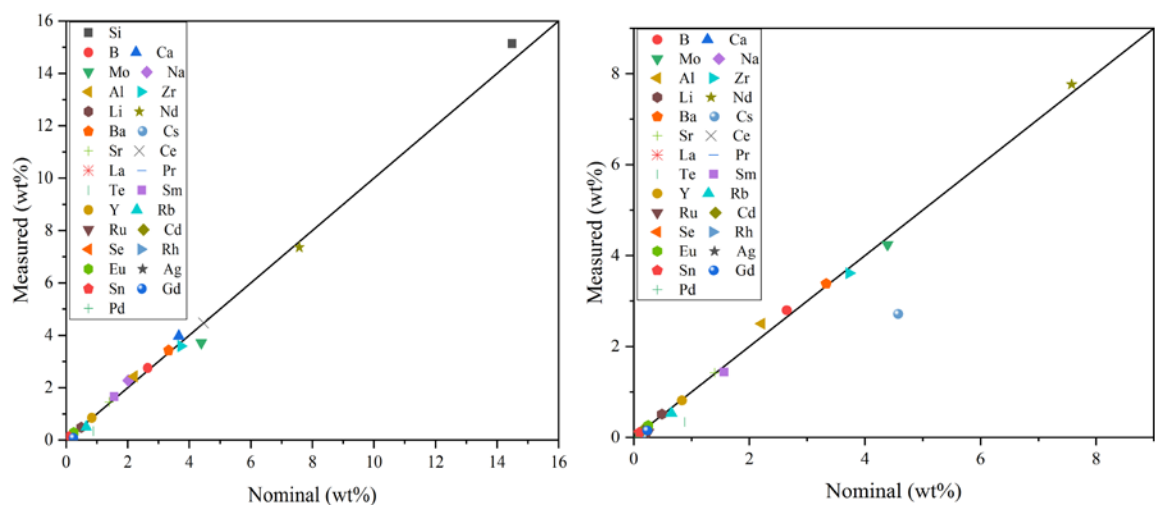


Figure A.12. Measured vs nominal composition of the CCC sample. Left) measured with ICP-AES and right) measured with ICP-MS

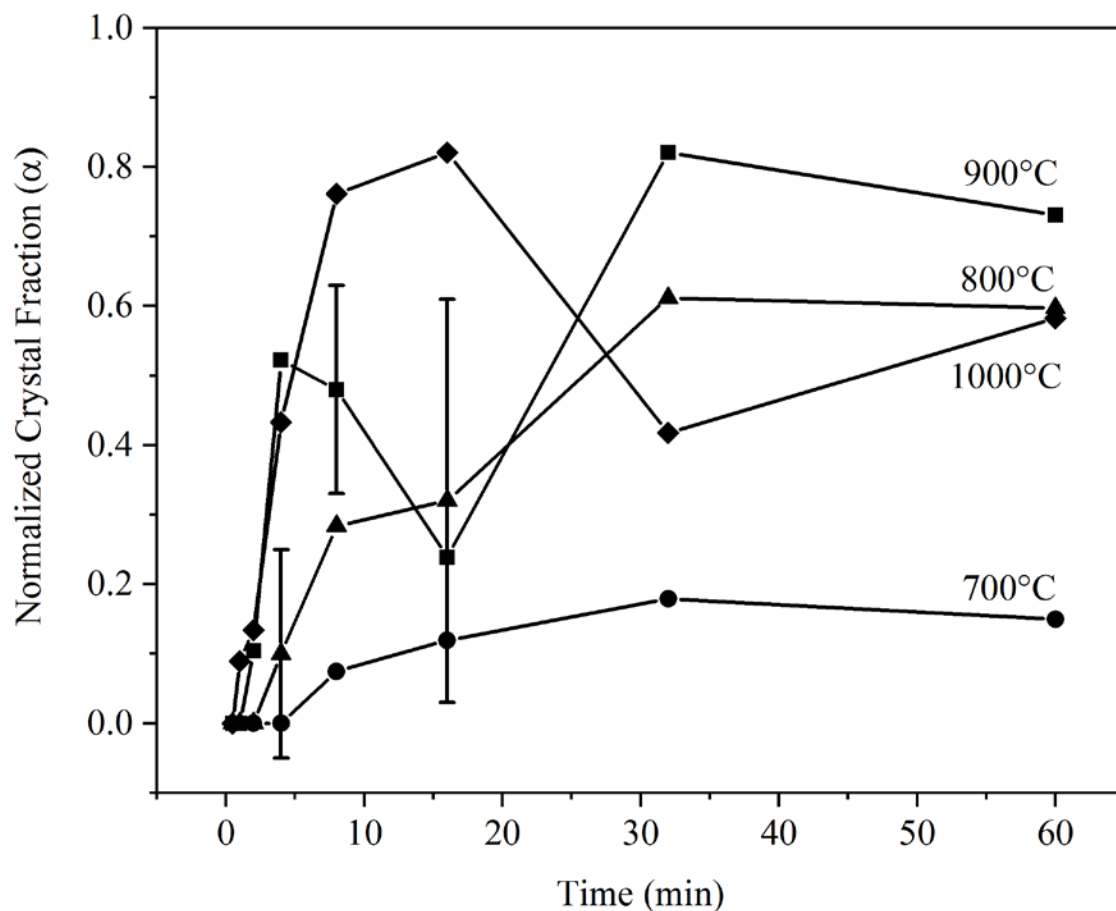


Figure A.13. Normalized crystal fraction vs time for powellite determined by XRD.

Table A.4. Measured composition of the furnace cool, CCC, and roller quench samples from Savannah River National Laboratories.

	Furnace Cool (0.1°C/s)		CCC (0.05°C/s)		Roller Quench (>100°C/s)	
	ICP-AES	ICP-MS	ICP-AES	ICP-MS	ICP-AES	ICP-MS
Si	14.54		15.14		14.69	
B	2.80	2.74	2.75	2.80	2.76	2.77
Ca	6.14		3.97		6.28	
Mo	3.72	4.31	3.71	4.24	3.75	4.32
Na	2.31		2.27		2.27	
Al	2.28	2.26	2.43	2.50	2.25	2.33
Zr	3.37	3.46	3.59	3.61	3.51	3.58
Li	0.48	0.46	0.48	0.51	0.49	0.48
Nd	6.56	6.91	7.36	7.76	6.54	6.87
Ba	3.32	3.32	3.43	3.38	3.41	3.31
Cs		3.62		2.71		3.35
Sr	1.42	1.38	1.45	1.42	1.46	1.39
Ce	4.23	4.34	4.47	4.65	4.22	4.33
La	2.00	1.99	2.10	2.08	2.04	1.98
	2.04	2.08	2.12	2.13	2.08	2.09
Te	0.61	0.62	0.34	0.34	0.47	0.55
Sm	1.52	1.35	1.66	1.44	1.56	1.34
Y	0.82	0.79	0.84	0.82	0.82	0.80
Rb	0.58	0.60	0.50	0.53	0.52	0.58
Ru		0.15		0.11		0.17
Cd	0.16	0.17	0.16	0.16	0.16	0.17
Se						
Rh		0.00		0.00		0.00
Eu	0.27	0.24	0.29	0.26	0.27	0.25
Ag		0.15		0.15		0.15
Sn	0.11	0.09	0.12	0.11	0.10	0.09
Gd	0.30	0.37	0.08	0.15	0.29	0.38
Pd		0.01		0.01		0.01

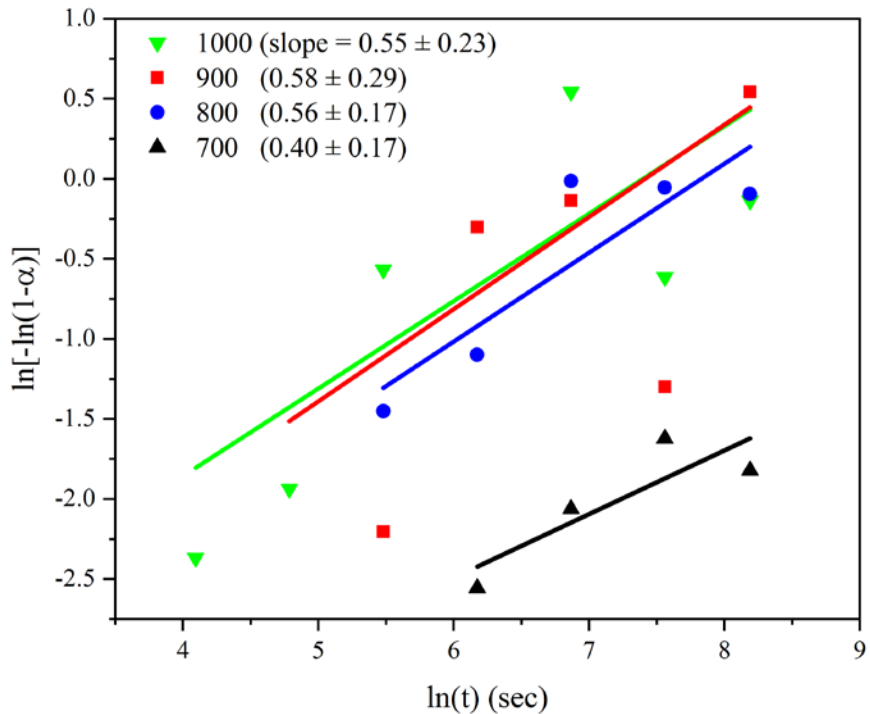


Figure A.14. Avrami plot for powellite formation in the isothermal experiments $\ln(k_n)$ vs $1000/T$ for powellite formation in the isothermal experiments.

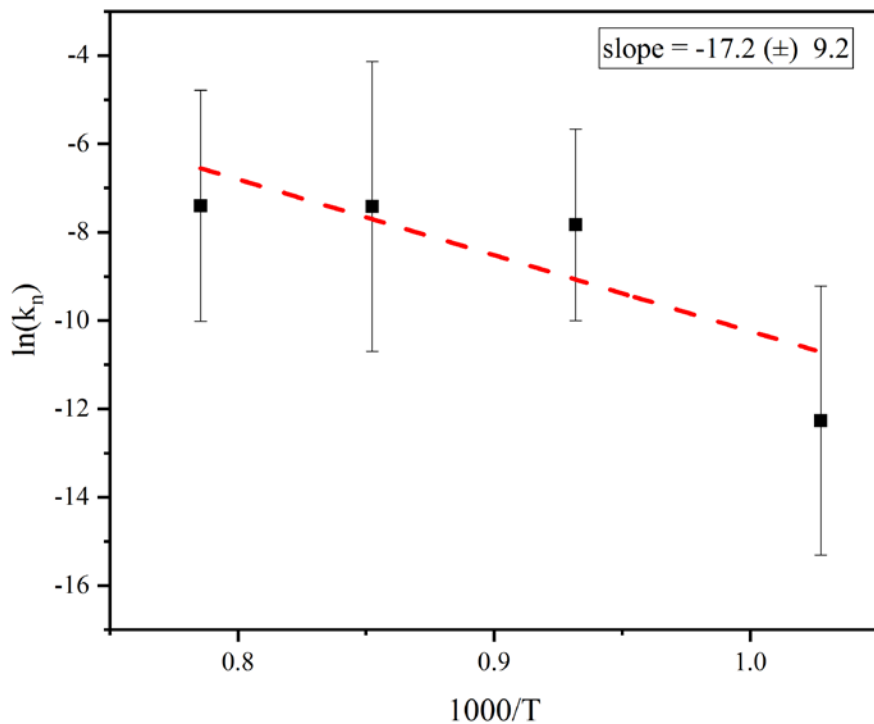


Figure A.15. $\ln(k_n)$ vs $1000/T$ for powellite formation in the isothermal experiments.

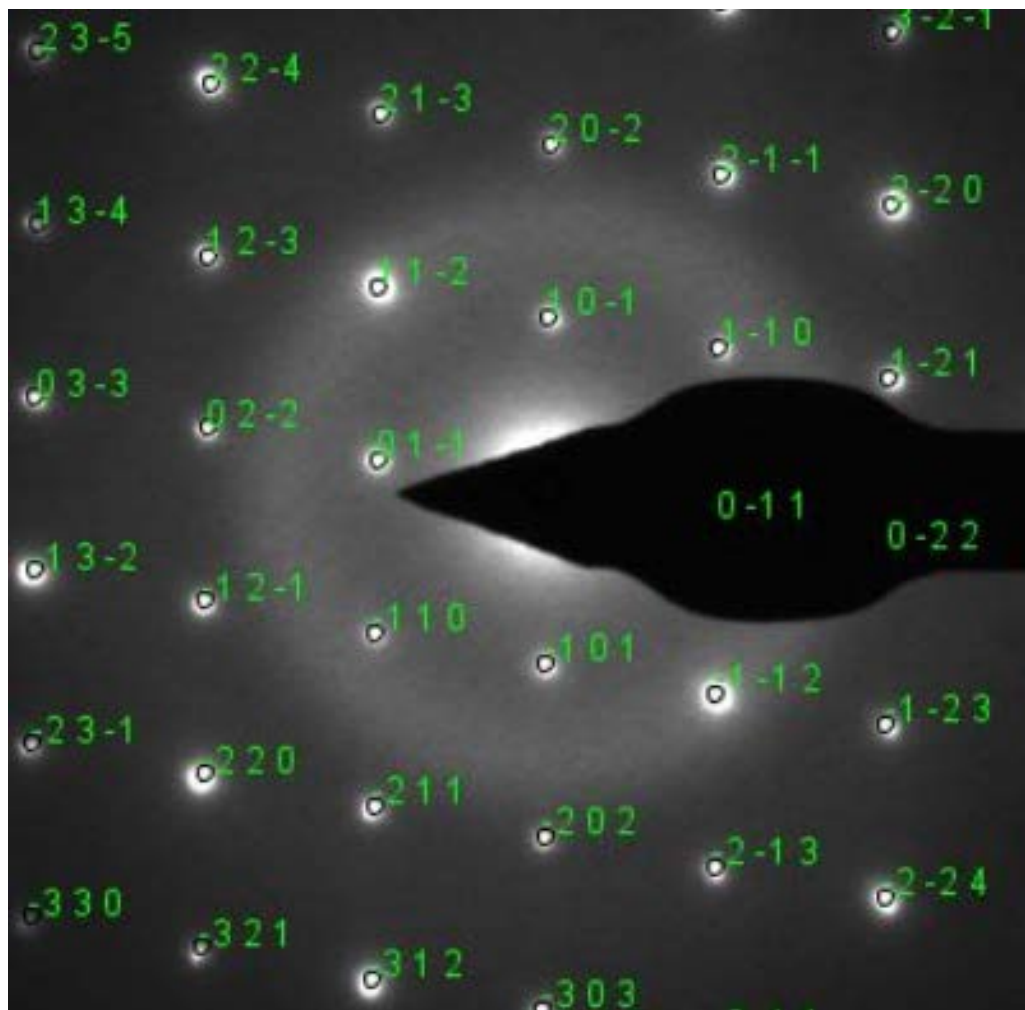


Figure A.16. Fully indexed SAD pattern from the Mo-rich droplet in the 4.1°C/s sample along zone 1

Table A.5. Comparison of the lattice parameters (c/a) for the various crystal.

Sr _{0.44} Ln _{0.39} (MoO ₄) (c/a)	Ba _{0.75} Sr _{0.25} (MoO ₄) (c/a)	CaMoO ₄ (c/a)	Measured (c/a)
2.22	2.28	2.20	2.26 ± 0.01

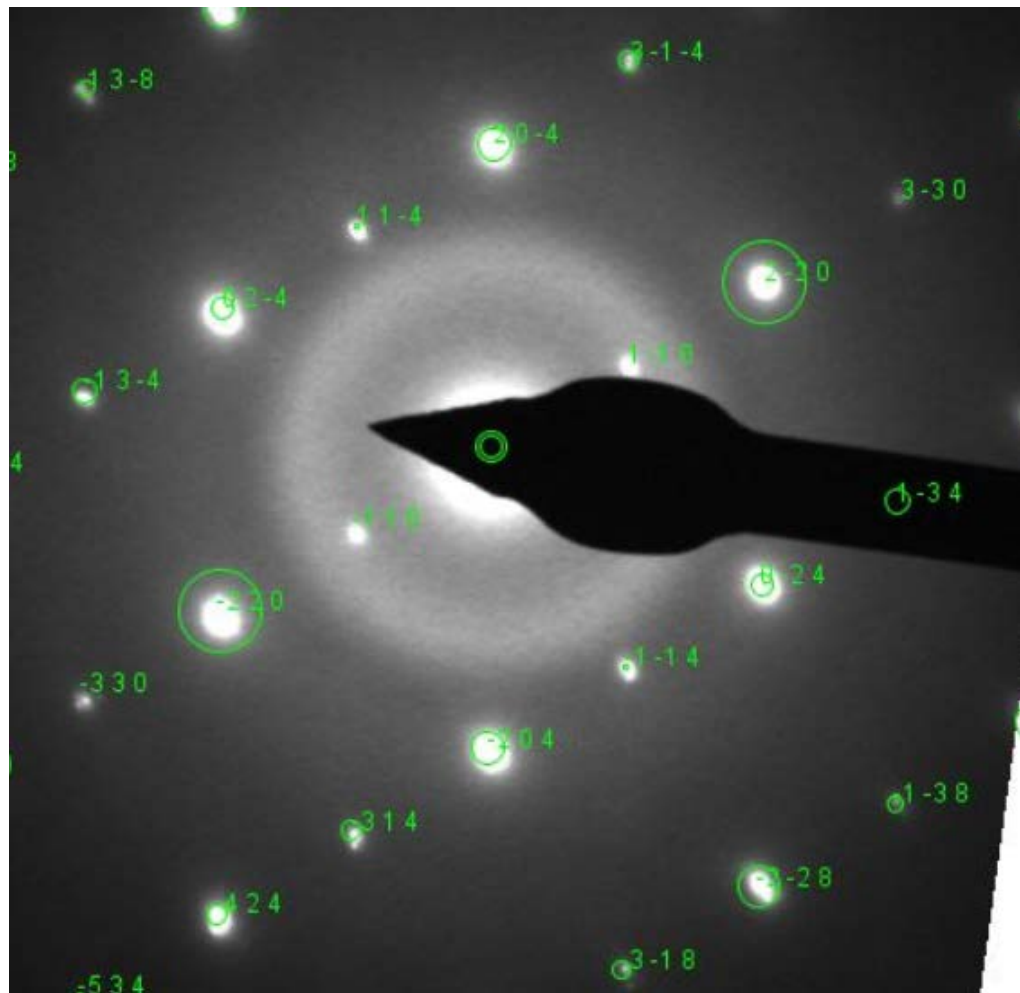


Figure A.17. Fully indexed SAD pattern from the Mo-rich droplet in the 4.1°C/s sample along zone 2

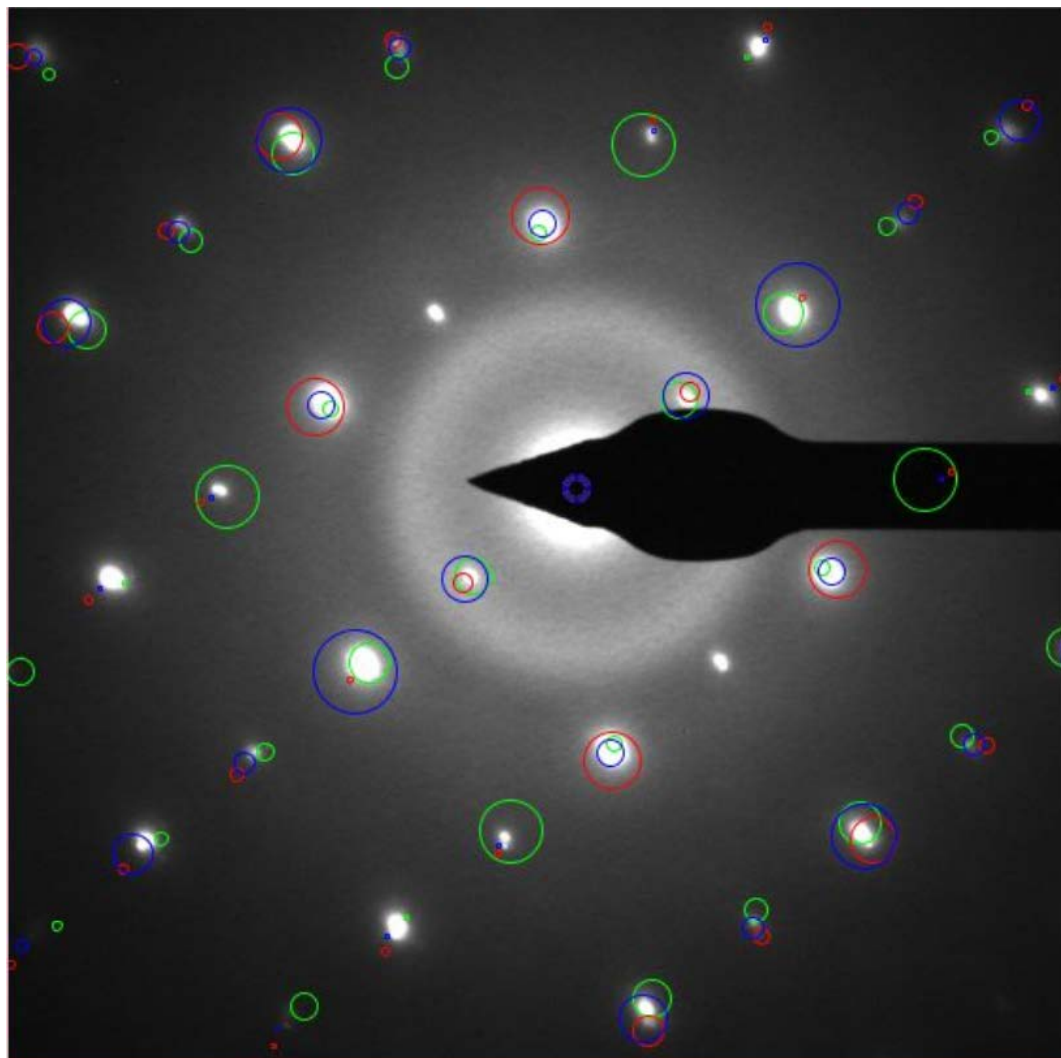


Figure A.18. Comparison to with Sr_{0.44}Ln_{0.39}(MoO₄) (blue), Ba_{0.75}Sr_{0.25}(MoO₄) (green), and CaMoO₄ (red) diffraction patterns.

APPENDIX B: SUPPLEMENTAL INFORMATION FROM THE CORROSION STUDIES

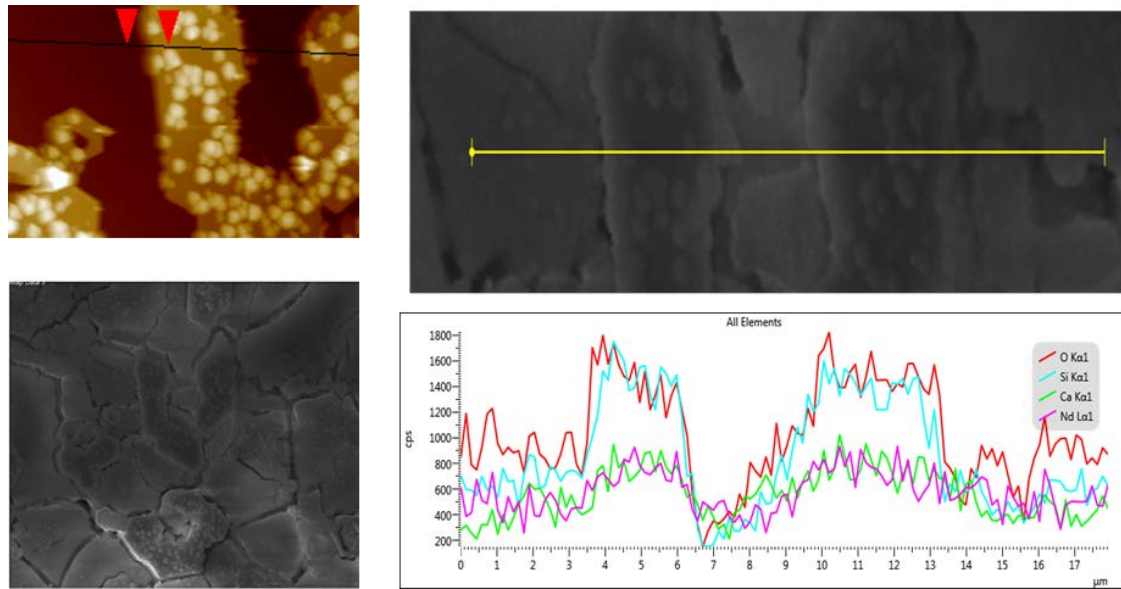


Figure B.1. Air cooled sample SEM micrographs and EDS line scan of the same area scanned with the AFM.

Table B.1. Normalized release rates for PCT dissolution experiments also including results reported by Crum et al. [16].

	Roller Quench [g/m ²]	Air Cool [g/m ²]	Furnace Cool [g/m ²]	CCC [g/m ²]	Crum et al. [16] CCC [g/m ²]
B	0.73 ± 0.03	0.99 ± 0.11	1.35 ± 0.19	0.93 ± 0.13	0.179
Ca	0.53 ± 0.02	0.52 ± 0.01	0.68 ± 0.07	0.19 ± 0.05	0.032
Na	1.42 ± 0.14	1.37 ± 0.03	1.80 ± 0.22	0.80 ± 0.12	0.172
Nd	-	-	0.01	-	NR*
Mo	0.47 ± 0.02	1.29 ± 0.01	1.93 ± 0.15	0.31 ± 0.11	0.172
Ba	0.03 ± 0.01	0.24 ± 0.02	0.50 ± 0.03	0.07 ± 0.01	0.070
Si	0.49 ± 0.03	0.42 ± 0.01	0.49 ± 0.03	0.25 ± 0.02	0.051

* NR – value is not reported

Table B.2. Normalized release rates for PCT dissolution experiments from Savannah River National Laboratories [16].

Sample	Furnace Cool (0.1°C/s) [g/m ²]	CCC (0.05°C/s) [g/m ²]	Roller Quench (>100°C/s) [g/m ²]
Si	1.01E-01	8.03E-02	1.05E-01
B	3.04E-01	1.97E-01	1.30E-01
Ca	6.36E-02	4.14E-02	3.86E-02
Mo	3.42E-01	6.51E-02	4.97E-02
Na	1.68E+00	1.47E+00	1.51E+00
Al	1.88E-02	3.19E-02	3.77E-02
Zr	7.80E-05	8.44E-04	1.10E-04
Li	4.68E-01	2.25E-01	2.32E-01
Nd	7.80E-05	4.25E-04	6.90E-05
Ba	7.50E-02	1.36E-02	8.30E-03
Cs	2.68E-01	1.14E-01	1.60E-01
Sr	1.15E-01	3.66E-02	3.43E-02
Ce	7.60E-05	4.41E-04	7.50E-05
La	7.60E-05	5.27E-04	5.90E-05
Pr	7.30E-05	4.29E-04	6.00E-05
Te	6.15E-02	7.32E-03	3.81E-03
Sm	8.30E-05	3.93E-04	6.60E-05
Y	8.20E-05	4.67E-04	6.40E-05
Rb	2.61E-01	1.12E-01	1.37E-01
Ru	4.46E-04	4.60E-04	2.46E-04
Cd	4.00E-03	1.75E-03	6.72E-04
Se	<1.049325	<1.056108	<1.055529
Rh	2.39E-02	1.17E-02	1.69E-02
Eu	4.27E-04	4.93E-04	4.25E-04
Ag	1.52E-03	2.88E-03	2.74E-04

Table B.3. Δh measurements for profilometry and AFM test.

device	Sample	Cooling Rate [°C/s]	Time [days]	Δh [nm]	St. dev.
profiler	copper step 2	5.8	7	59	19
	copper step 1	5.1	7	91	19
	steel step 1	4.1	7	130	26
	air cool	3.4	1	56	17
	air cool	3.4	2	49	13
	air cool	3.4	7	148	30
	furnace cool	0.1	2	58	12
	furnace cool	0.1	3	67	15
	furnace cool	0.1	4	115	24
	furnace cool	0.1	4	106	11
	furnace cool	0.1	7	201	31
	furnace cool	0.1	7	195	24
	furnace cool	0.1	7	195	34
	furnace cool	0.1	8	245.2	29
AFM	CCC	0.008	7	841	181
	CCC	0.008	7	714	174
	air cool (oxyapatite – glass)	3.4	12.5	243	90
	air cool	3.4	12.5	264	68
	air cool	3.4	18.75	378	181

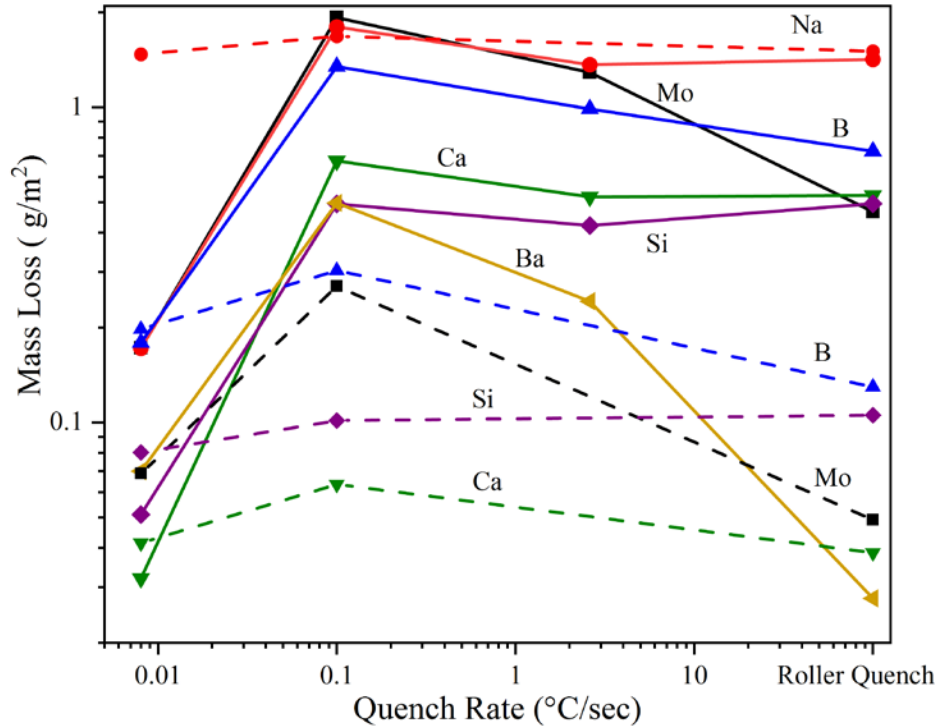


Figure B.2. PCT results vs quench rates coming this study (solid lines) and results from Savannah River National Laboratories (dotted line).

Table B.4. Mass loss measurements normalized to surface area.

Sample	Cooling Rate [°C/s]	Time [days]	Weight loss [g/m²]
copper step 2	5.8	7	1.7
copper step 1	5.1	7	2.1
steel step 1	4.1	7	2.9
air cool	3.4	1	1.1
air cool	3.4	2	3.1
air cool	3.4	4	1.8
air cool	3.4	7	4.1
furnace cool	0.1	1	3.6
furnace cool	0.1	2	2.0
furnace cool	0.1	2	4.7
furnace cool	0.1	3	2.9
furnace cool	0.1	4	6.6
furnace cool	0.1	7	7.0
furnace cool	0.1	7	5.4
furnace cool	0.1	7	3.9
furnace cool	0.1	8	7.7
CCC	0.008	7	2.7
CCC	0.008	7	3.6

APPENDIX C: HOT THERMOCOUPLE TEST SYSTEM

Introduction

The Hot Thermocouple Technique was originally developed by Ordway in 1952 for the synthesis of single-crystal samples of refractory ceramics for X-Ray Diffraction at NIST, and has since been improved and used by metallurgists as a technique for the in-situ characterization of molten slags and mold fluxes [1], [2]. The technique operates on the principle of using a single thermocouple as an electromotive sensor, resistive heater, and sample mount. By repeatedly applying a small heating pulse to the thermocouple and measuring the resulting temperature, a feedback loop can be constructed to control the temperature of the thermocouple over time. This was accomplished by supplying power as a half-rectified AC waveform to heat the thermocouple, with measurement events taking place during times at which the applied power was zero, schematic diagram shown in Figure 1. After each measurement event, the temperature of the thermocouple is compared to the desired temperature at that point in time as defined by the parameters of that specific test, and the power of the next heating pulse is adjusted accordingly.

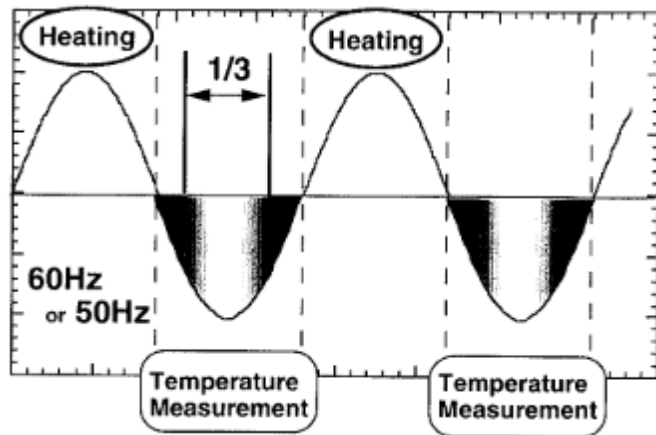


Figure 1 - Operating principle of the Hot Thermocouple Technique controller used by Kashiwaya et al. Reproduced from [2].

The device designed for this work was built using a DC-DC operating principle using MOSFET switching proposed by Esfahani et al., rather than the half-rectified AC waveform used by Ordway and Kashiwaya et al [3]. This design enables much faster switching between measurement and heating, operating at a switching frequency of 240Hz, and results in a smoother temperature profile with no measurable oscillations due to cooling between heating pulses.

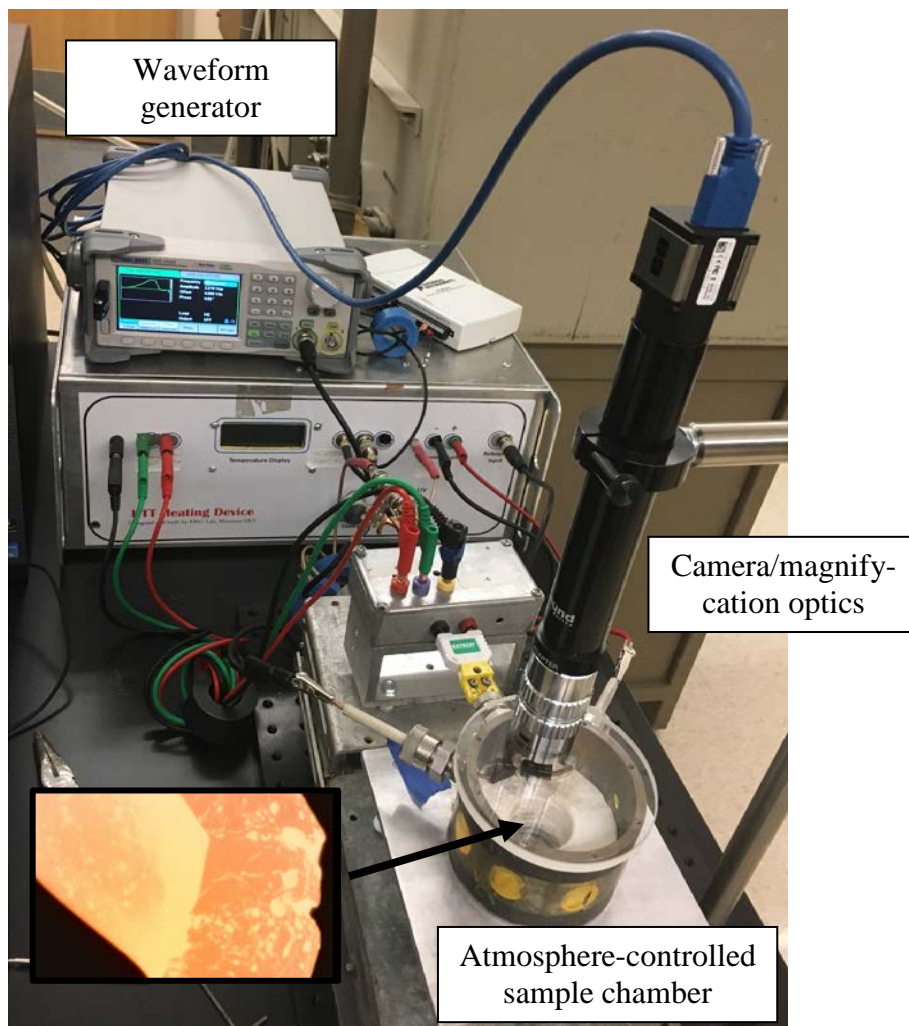


Figure 2: The hot thermocouple test system built at Missouri S&T; inset shows an image of a molten sample.

Figure 2 is a picture of the HTT system. The signal generator and associated control electronics provide power to heat the thermocouple and melt the sample (inset) on a half-cycle, then record the temperature at the thermocouple junction on the other half-cycle. Temperature is recorded as a function of time and is synchronized with the images captured through the magnifying optics. (The microscope system can record a 4192 x 3684 pixel resolution video at 21 frames per second and can resolve features 2-5 microns in size, depending on the objective used.

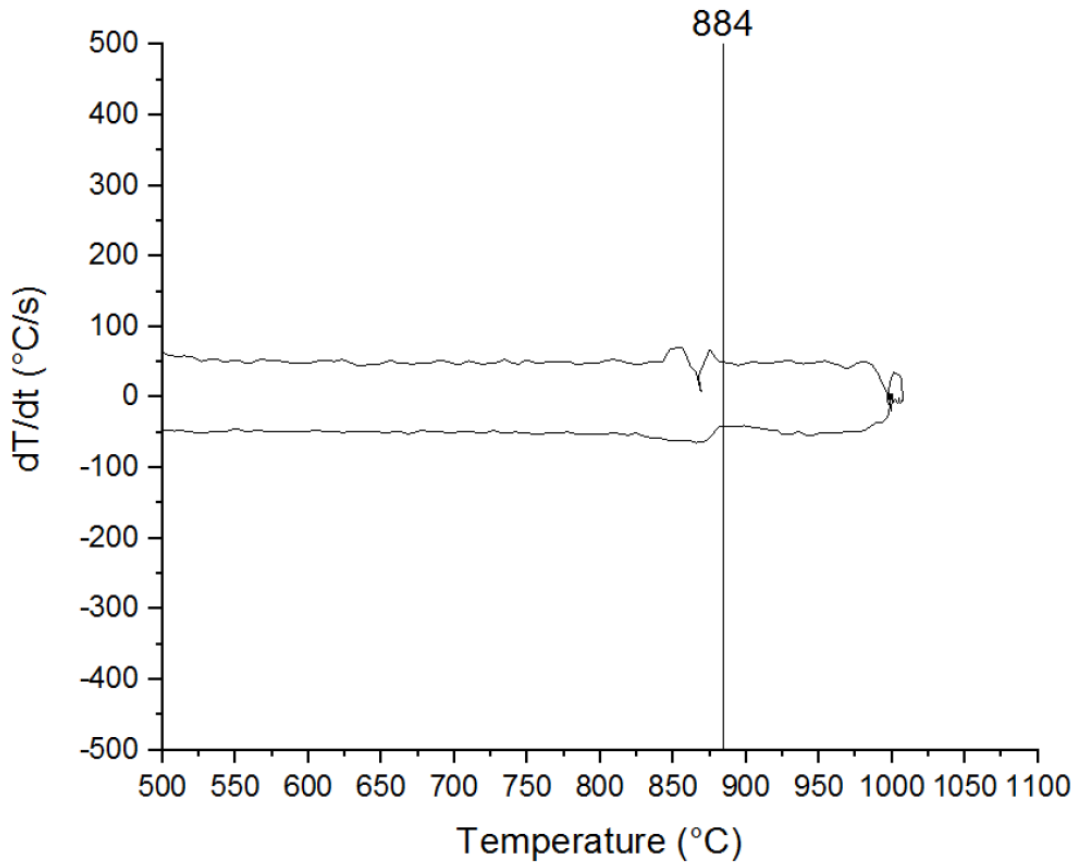


Figure 3: Detection of the endothermic melting and exothermic crystallization of Na_2SO_4 .

Figure 3 shows the results of a temperature-calibration test. Here, Na_2SO_4 (M.P. 884°C) was heated to 1000°C at a rate of 50°C/s , held for 10 seconds, then cooled to room temperature at a rate of 50°C/s . This is represented here as a 'loop' of heating from 500 to

1000°C with a dT/dt of 50°C/s. Endothermic melting could be detected on heating and exothermic crystallization could be detected on cooling. These small signals, however, preclude the use of the HTT to thermally detect phase transformations in more slowly crystallizing systems, like the borosilicate glass-ceramics designed for waste encapsulation.

The HTT system has been evaluated by testing a series of simple silicate glasses with well-characterized crystallization rates and appropriate viscosities [4-7], including Li-disilicate, Li-metasilicate, Ca-disilicate, Ba-disilicate, and diopside (CaO-MgO-2SiO_2). Quenched glasses were crushed and appropriately sized shards (~20 mg) were mounted to the Type B thermocouple by placing the fragment on the thermocouple and heating it using the HTT device, or by first heating the thermocouple and touching the shard to it using tweezers. Once mounted, samples were heated, isothermally held, and quenched as desired. During the various tests, thermal data was recorded by the HTT device and images/video of the sample were captured as desired.

To measure linear crystal growth rates, samples were heated at 40°C/s to temperatures at which no residual crystal could be seen and isothermally held for 20 seconds. Samples were then quenched at 100°C/s to the specified treatment temperature and held until the test is cancelled. Video was captured using the highest resolution capable at frame rates ranging from 9.5 to 20 frames per second based on the size of area of interest, objective lens used, and exposure time necessary for image clarity. Due to CPU and data throughput limitations, frames were saved with JPG compression at a quality level of 75% in uEye camera control software. After capture, individual frames were extracted from

videos and ImageJ software was used to measure the distance traveled by the crystal-glass interface over ranges of 50 to 100 frames.

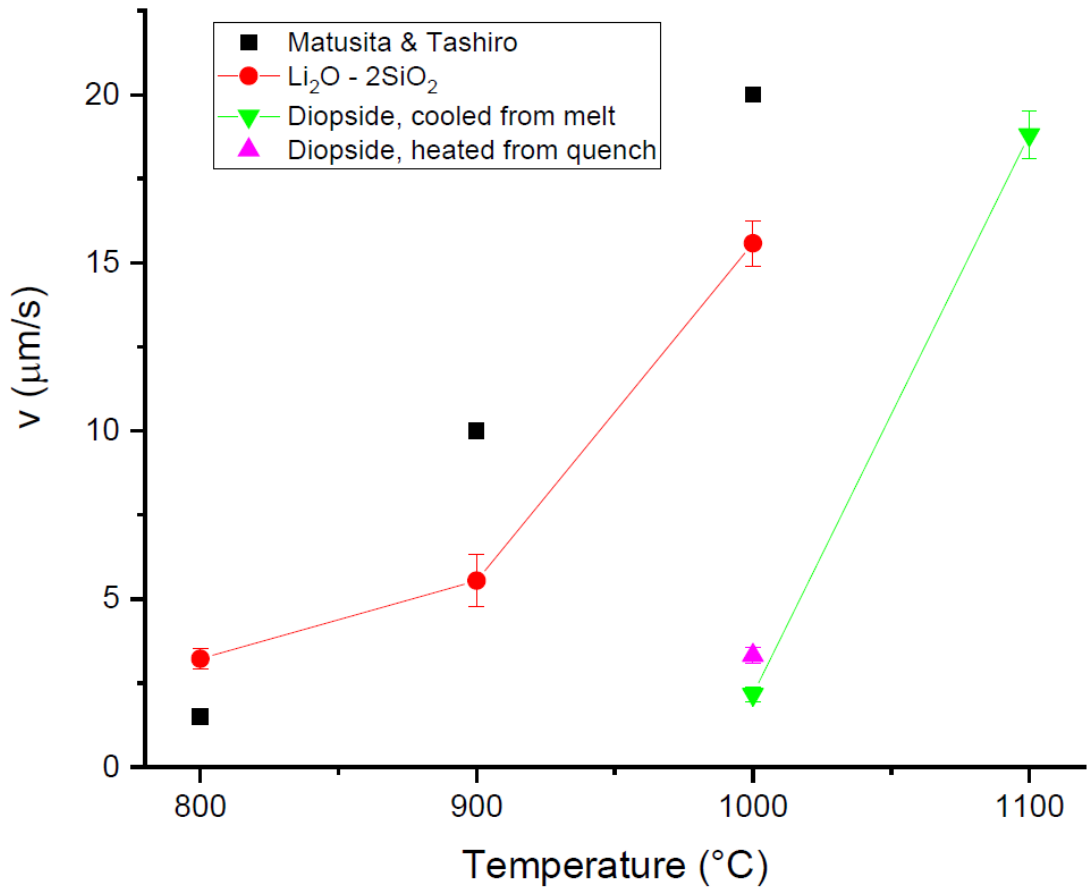


Figure 4: Linear crystal growth rates for Li-disilicate and diopside melts measured with the HTT under isothermal conditions.

Growth rate measurements have so far been performed on two of the systems investigated, Li₂O – 2SiO₂ and CaO – MgO – SiO₂ Diopside glass. The results of these experiments along with some values measured by Matusita and Tashiro (reproduced by analysis of plots [8]) are given in Figure 4. Diopside glass held isothermally at 900°C yielded no detectable crystallization after 10 minutes. As an additional point of comparison, the growth rate of Diopside was also measured by heating to 1000°C at a rate of 40°C/s after quenching at 100°C/s from a fully melted 1500°C sample. This was done to investigate potential impacts from differences in experimental setup

between this experiment and the prevalent method of reheating glass samples to a treatment temperature after quenching from a melt, which would pre-nucleate crystals.

During the measurement for crystal growth rate in Diopside glass at 1100°C, no visible crystals formed until over 5 minutes into the test, after which point the crystal-glass interface rapidly propagated throughout the sample, shown in Figure 5. This indicates that even though certain experiments at resulted in no detectable crystal growth within 10 minutes, the growth rates may still be significant once nuclei form. This behavior is to be expected from basic kinetic theory of nucleation and growth, but the timescale associated with the formation of a viable nucleus under these conditions appears large enough to necessitate an alternative experimental setup. Possible solutions include pre-nucleation, which appears to affect the linear growth rate as well as the “incubation time” of formation of a stable nucleus, or a shift in image capture technique from continuous video recording to high-quality image capture at specified time intervals.

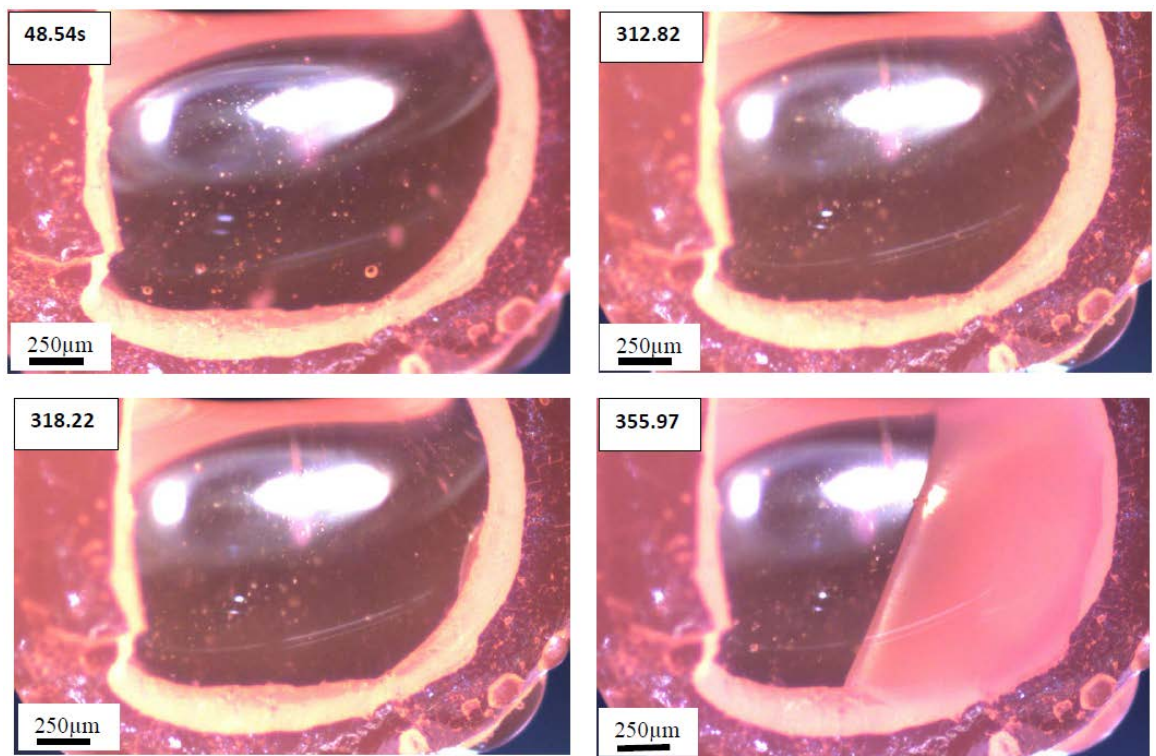


Figure 5: Crystallization of a diopside glass sample at 1100°C

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