LABORATORY TESTING OF LITCO GLASSES


Argonne National Laboratory
Chemical Technology Division
9700 South Cass Avenue
Argonne, IL 60439

For presentation at the
American Ceramics Society Meeting
Cincinnati, OH
April 30 - May 4, 1995

This work supported by the U.S. Department of Energy, Lockheed Idaho Technologies Company, under contract DE-AC07-94ID13223.
DISCLAIMER

Portions of this document may be illegible in electronic image products. Images are produced from the best available original document.
LABORATORY TESTING OF LITCO GLASSES

Adam Ellison, S. Wolf, E. Buck, J. S. Luo, N. Dietz, J. K. Bates, W. Ebert
Chemical Technology Division, Argonne National Laboratory, Argonne, IL 60439

INTRODUCTION

The purpose of this program is to measure the intermediate and long-term durability of glasses developed by Lockheed Idaho Technology Co. (LITCO) for the immobilization of calcined radioactive wastes. The objective is to use accelerated corrosion tests as an aid in developing durable waste form compositions. This is a report of tests performed on two LITCO glass compositions, Formula 127 and Formula 532.

The main avenue for release of radionuclides into the environment in a geologic repository is the reaction of a waste glass with ground water, which alters the glass and releases its components into solution. The reaction rate is highest when glass corrosion products in solution are dilute (the forward rate). As the concentrations of corrosion products in solution increase, the chemical potential gradient between glass and water decreases, slowing the reaction (the interim rate). After prolonged contact with water, the concentrations of dissolved glass components increase to a point where secondary crystalline phases precipitate from solution. If these phases remove components from solution faster than they are replenished by the glass, the concentrations of components in solution will remain at or below the saturation value. This may increase the rate of glass corrosion (the long-term rate). Data concerning the forward rate, the interim rate and the long-term rate are all needed to evaluate glass performance.

These stages in glass corrosion are analyzed by using accelerated laboratory tests in which the ratio of sample surface area to solution volume, \( SA/V \), is varied. At low \( SA/V \), the solution concentrations of glass corrosion products remain low and the reaction approaches the forward rate. At higher \( SA/V \) the solution approaches saturation levels for glass corrosion products. At very high \( SA/V \) the solution is rapidly saturated in glass corrosion products and secondary crystalline phases precipitate. Tests at very high \( SA/V \) provide information about the composition of the solution at saturation or, when no solution is recovered, the identities and the order of appearance of secondary crystalline phases.

Tests were applied to Formula 127 and Formula 532 glasses to provide information about the interim and long-term stages in glass corrosion. Product consistency tests method B (PCT-B [1]) at \( SA/V \) of approximately 2,000 and 20,000 m\(^{-1}\) were
performed to measure the interim corrosion rate of both glasses. PCT-B tests highlight the effects of solution chemistry on glass corrosion, and long-term PCT-B tests can also provide information about secondary phase formation and long-term performance of the glass. Argonne National Laboratory vapor hydration (VH [2]) tests were used to determine the identities and order of appearance of secondary crystalline phases, information that is important to the long-term corrosion rate. These tests may indicate whether the presence of secondary crystalline phases increases the glass corrosion rate.

EXPERIMENTAL METHODS

In a PCT-B test a mass of glass of a particular size fraction is placed in contact with deionized water and heated at fixed temperature for various lengths of time. At the end of the test, the leachate solution is filtered to remove suspended glass particles and the composition of the solution is analyzed. The sample surface area is estimated by assuming that the glass grains are spheres with diameters equal to the average mesh size. On this basis, \( SA/V \) is 2000 m\(^{-1}\) for -100+200 mesh grains in contact with 10 times their mass of water and 20,000 m\(^{-1}\) for -100+200 mesh grains in contact with an equal mass of water. The release of components to solution is expressed as normalized release, \( NL_i \):

\[
NL_i = \frac{\text{concentration}}{f_i \cdot SA/V}
\]

where \( f_i \) is the weight fraction of element \( i \) in the glass. The value of \( NL_i \) is an estimate of the mass of glass that reacted to account for the concentration of the element \( i \) in solution. PCT-B tests reported in this study were performed at 90°C in Teflon™ vessels.

In the VH test a monolithic sample is placed in contact with water vapor in a sealed vessel at elevated temperature. At relative humidities above ~90% a thin film of water condenses on the sample. The high \( SA/V \) of this condition rapidly saturates the solution with glass corrosion products, resulting in rapid formation of secondary crystalline phases. Over time, growth of secondary crystalline phases changes the composition of the solution, and the crystalline phases may enter into reactions with one another, producing a more stable phase assemblage in the process. At the end of a test, the glass surface is analyzed using scanning electron microscopy (SEM) and analytical electron microscopy (AEM) or diffraction to identify the secondary crystalline phases that have formed. The solution evaporates after the test is terminated, so solution analyses cannot be performed. The VH tests in this study were performed at 200°C.

SAMPLE CHARACTERISTICS

The compositions of Formula 127 and Formula 532 glasses are shown in Table 1. The composition of Formula 127 was taken from a report by Staples et al. [3]. Formula 127 is a translucent green glass containing 5-10 vol % crystalline CaF\(_2\).
Most fluorite crystals are 2 to 5 μm spherulites distributed 5 to 15 μm from one another. Some are in the form of skeletal masses up to 5 mm in diameter (Fig. 1). Semi-quantitative energy dispersive x-ray spectroscopy (EDS) indicates that the interstitial glass (the dark areas in Fig. 1) is depleted in Ca and F relative to the bulk.

The nominal composition of Formula 532 is shown in Table 1. It is a transparent, deep-blue glass with swirls of much darker glass in places and relatively large (1-5 μm) blebs of light-colored, undissolved or partially-reacted material. EDS analyses of the light-colored material indicate that its composition is similar to that of the bulk glass though significantly depleted in sodium. Scanning electron microscopy (SEM) and electron diffraction reveal that 5-10% of the bulk material is the lithium aluminosilicate mineral virgilite, LiAlSiO₄. The virgilite crystals have highly irregular faces, indicating that they are fragments of still larger (> 100 μm) crystals. These larger crystals may have been excluded by crushing the samples for PCT-B tests.

Table 1. Compositions of Formula 127 and Formula 532 Glasses.

<table>
<thead>
<tr>
<th>Element</th>
<th>Formula 127 at %</th>
<th>Formula 127 wt %</th>
<th>Formula 532 at %</th>
<th>Formula 532 wt %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>4.1</td>
<td>5.1</td>
<td>8.0</td>
<td>11.6</td>
</tr>
<tr>
<td>B</td>
<td>1.3</td>
<td>0.6</td>
<td>5.3</td>
<td>3.1</td>
</tr>
<tr>
<td>Ca</td>
<td>5.2</td>
<td>9.6</td>
<td>9.6</td>
<td>14.3</td>
</tr>
<tr>
<td>Cu</td>
<td>0.6</td>
<td>1.8</td>
<td>0.5</td>
<td>1.6</td>
</tr>
<tr>
<td>Li</td>
<td>3.6</td>
<td>1.1</td>
<td>8.8</td>
<td>3.3</td>
</tr>
<tr>
<td>Na</td>
<td>4.1</td>
<td>4.3</td>
<td>4.2</td>
<td>5.3</td>
</tr>
<tr>
<td>Si</td>
<td>18.2</td>
<td>23.5</td>
<td>14.3</td>
<td>21.8</td>
</tr>
<tr>
<td>Ti</td>
<td></td>
<td>1.2</td>
<td>5.3</td>
<td>3.1</td>
</tr>
<tr>
<td>Zr</td>
<td>1.8</td>
<td>7.5</td>
<td>1.1</td>
<td>3.3</td>
</tr>
<tr>
<td>F</td>
<td>11.6</td>
<td>10.1</td>
<td>11.6</td>
<td>10.1</td>
</tr>
<tr>
<td>O</td>
<td>49.5</td>
<td>36.4</td>
<td>57.8</td>
<td>50.2</td>
</tr>
</tbody>
</table>

**Fig. 1** Electron backscatter image of a polished surface of Formula 127 glass.

**FORMULA 127 TEST RESULTS**

Figure 2 is a graph of the normalized releases of B, Li, Na and Si from Formula 127 glass as a function of the duration of the PCT-B tests. Normalized releases for each element were greater for tests performed at 2,000 m⁻¹ than for tests performed 20,000 m⁻¹ because relatively little glass needs to dissolve at the higher SA/V to raise the concentrations of components in solution to near their saturation levels. Normalized releases decreased in the order Li, B > Na > F > Si > Ca >> Al, Cu, Zr. SEM analysis of the PCT-B samples indicated that exposed fluorite crystals
grow during the course of the tests (Fig. 3). Analytical electron microscopy (AEM) analysis showed that the glass immediately around the fluorite grains was extensively reacted, whereas the bulk glass showed little evidence of reaction and no reaction layer.

After 98 days at 20,000 m⁻¹, calcium silicate crystals started to form, and after 182 days, isolated crystals of a Si-O-F phase (possibly containing Li or B) also formed. No other secondary crystalline phases were identified in solids analyses of PCT-B tests. The rate of release of glass components in 20,000 m⁻¹ tests (Fig. 2) may increase between 98 and 189 days: one-year tests to be terminated later this year may shed light on this issue. If this small increase is real, then the formation of secondary phases has a minor impact on the rate of corrosion of Formula 127 glass.

The first phases to appear in VH tests were calcium silicates, one with a Ca:Si ratio near 1:1 and another with a Ca:Si ratio > 2:1, and a crystal rich in Si, O and F (Li or B may also have been present). Irregularly-shaped masses of a phase containing Ca and O also formed, which may be precipitates of Ca(OH)₂. After 14 d the calcium-rich silicate disappears, and the other calcium silicate was in a reaction relationship with CaCO₃, calcite. After 28 d the main calcium-bearing phase was calcite, which grew to 50-100 μm in diameter after 56 d. After 28 d zirconium also appeared in secondary crystalline phases, a Ca-Zr-F phase (possibly CaZrFₓ) and a Ca-Zr-Si-O-F phase. The latter was the only significant Si-bearing phase formed in VH tests. Even after 56 d, the glass itself did not appear to undergo very extensive corrosion. Thus, a relatively small amount of glass was reacting to form the phases observed in the VH tests. This supports the conclusion from the PCT-B test data that secondary phases do not have a large impact on the long-term rate of corrosion of Formula 127 glass: the maximum rate of release at 20,000 m⁻¹ from 28 to 98 d is similar to the long-term rate measured from at 20,000 m⁻¹ from 98 to 182 d.

Nevertheless, Formula 127 appears to absorb or incorporate water during the course of accelerated corrosion tests. Cross-sectioning of grains from PCT-B tests resulted in fragmentation around the borders of the grains, and the depth of fragmentation varied directly with the duration of the experiment. Cross-sectioning grains of unreacted glass did not produce fragmentation, indicating that the near-surface regions of glass grains from PCT-B tests are more fragile than unreacted glass. Cross-sectioning VH test samples also produced fragmentation of the material in contact with water vapor. Furthermore, the cross-sectioned VH test samples pulled back from the epoxy over time, leaving a gap around the periphery of the glass. The width of the gap increased with the duration of the hydration test. The simplest explanation for this is that Formula 127 glass incorporated water during VH tests, and slowly released it after the tests were completed. Thus, while Formula 127 does not dissolve rapidly in water in accelerated tests, water is incorporated into a layer near the glass surface.
RESULTS

Figure 4 shows normalized releases of Al, B, Li, Na and Si from Formula 532 glass as a function of the duration of 2,000 m$^{-1}$ and 20,000 m$^{-1}$ PCT-B tests. Normalized releases decreased in the order B > Li > Na > Al > Si: in particular, $N_{rAl}$ was 20 times greater than for comparable tests performed on Formula 127 glass. The rate of glass dissolution as measured by $N_{rNa}$ increased after 48 d at 2,000 m$^{-1}$ and stayed at this high rate through at least 182 d. By contrast, $N_{rNa}$ and $N_{rSi}$ decreased after 48 d at 2,000 m$^{-1}$ and showed a consistent decrease over the course of the 20,000 m$^{-1}$ tests. After 49 d at 2,000 m$^{-1}$ the zeolite analcime and small crystals with Ti:Al atomic ratio of 1:2 (probably TiAl$_2$O$_5$) formed in Formula 532 PCT-B test samples. These and other secondary phases were present in all PCT-B tests conducted at 20,000 m$^{-1}$. At least five different secondary phases have been identified in long-term (98 d) PCT-B tests, including what appears to be LiOH$\cdot$2H$_2$O, and small irregular particles with EDS spectra similar to that of virgilite (these may be a lithium-bearing zeolite, see below). Given the steady decreases in $N_{rSi}$ and $N_{rNa}$, analcime and other secondary crystalline phases appear to control the rate of glass dissolution after PCT-B tests of relatively short duration.

The VH tests of Formula 532 glass showed that after 7 d an alteration layer formed with an average thickness of 30 μm and pockets as deep as 50 μm. The layer was filled with trapezoidal 1-4 μm crystals that are tentatively identified as a lithium-bearing zeolite bikitaite, Li$_2$Al$_2$Si$_4$O$_{12}$·2H$_2$O [4]. The layer was penetrated by deep
pits that revealed a clay-rich layer below the surface, or in some cases, a second altered layer filled with trapezoidal crystals. Isolated analcime crystals were also found. After 14 d in a VH test, the surface of Formula 532 was extensively altered, as shown in Fig. 5. The clustered, bladed crystals are the lithium bearing zeolite, and the large faceted crystal in the center of the image is analcime. The noodle-like mat layer underlying all the crystals is a clay. No new phases appeared after 49 d at 200°C, but the thickness of the alteration layer increased to approximately 300 μm with pockets as deep as 400 μm (Fig. 6). The layer consists of alternating layers of lithium-bearing zeolite and sodium-rich, aluminum-depleted material. After 56 d no unaltered glass remains.

Formation of these phases resulted in complete alteration of Formula 532 glass in VH tests. The same phases also formed in PCT-B tests, and their appearance coincided with an increase in the glass corrosion rate. These results suggest that Formula 532 moves into the long-term (secondary-phase dominated) stage of glass dissolution very early compared to Formula 127 glass.

SUMMARY

Formula 127 showed low normalized releases of glass components in PCT-B tests, and VH test data indicated that the secondary crystalline phases did not strongly impact the rate of glass corrosion. The VH test results suggest that a complex, fluoride-rich phase assemblage will eventually form in PCT-B tests. It may be that the main role of secondary crystalline phases is to accommodate fluoride, in which case these phases may not impact the rate of glass dissolution. Fluorite grains on the glass surface appear to control the solution concentrations of Ca and F. In this
sense, fluorite behaves both as a primary phase and as a secondary phase, though the fluorite crystals do not appear to increase the rate of corrosion of Formula 127.

![Cross-section of a monolith of Formula 532 glass after a 49 d vapor hydration test at 200°C unreacted glass is to the left.](image)

**Fig. 6** Cross-section of a monolith of Formula 532 glass after a 49 d vapor hydration test at 200°C unreacted glass is to the left.

Formula 532 showed significantly greater normalized releases for all elements, particularly for aluminum. It is not possible from the solution data to discriminate between Li, Al and Si released from the glass and that released from virgulite, but the integrated effect was to rapidly saturate the solution with Si (and possibly Al) and drive the formation of secondary crystalline phases. Secondary crystalline phases such as analcime and a lithium-bearing zeolite precipitated relatively early in PCT-B tests and appeared to accelerate the corrosion of Formula 532 glass. VH test results indicate that the lithium-bearing zeolite and analcime are the most important secondary corrosion products for Formula 532, and that a large fraction of the total mass of the glass is converted to these phases as corrosion proceeds.

While secondary crystalline phases are thermodynamically more stable than the glass, the glass must break down to form them, and any components that are not soluble in the secondary phases can potentially be released to solution. This is particularly important in the case of radionuclides: the breakdown of a waste glass into secondary crystalline phases that do not incorporate radionuclides will release them into the environment. The results of this study show the importance of identifying primary and secondary crystalline phases and performing long-term, high-SA/V accelerated tests to analyse their effect on the rate of glass corrosion.

These results also illustrate the value of using a suite of tests with different SA/V to analyze the course of glass corrosion. For example, at a fixed point in time Formula 127 glass is at a very different point in its corrosion path than Formula 532 (e.g., Figs. 2 and 4). Comparing these glasses by using a test with fixed SA/V
run for a fixed time period only provides information about the integrated effects of 
corrosion up to that point in time, and these effects clearly differ from glass to 
glass. Thus in 7 d PCT-B tests at 2000 m\(^1\), NL\(_B\) values for Formula 127 and 
Formula 532 differed by a factor of 2.5, but after 182 d at 20,000 m\(^1\), NL\(_B\) values 
differed by more than a factor of 6. The feedback between the evolving 
composition of a solution, the glass, and secondary crystalline phases is very 
complex, so one must exercise caution in extrapolating durability rankings obtained 
in short-term leach tests to geologic time scales. Performing different tests over a 
range of temperatures and SA/V is an effective means of evaluating all three stages 
in glass corrosion and thus, the course of glass corrosion over time.

REFERENCES

Glasses: The Product Consistency Test (PCT) Standard C1285-94, ASTM, 

2. W. L. Ebert, J. K. Bates and W. L. Bourcier, "The Hydration of Borosilicate 
Waste Glass in Liquid Water and Steam at 200\(^\circ\)C," Waste Management 11, 

(1983).

4. E. M. Flanigen, "Crystal Structure and Chemistry of Natural Zeolites," in 

DISCLAIMER

This report was prepared as an account of work sponsored by an agency of the United States 
Government. Neither the United States Government nor any agency thereof, nor any of their 
employees, makes any warranty, express or implied, or assumes any legal liability or responsi-
bility for the accuracy, completeness, or usefulness of any information, apparatus, product, or 
process disclosed, or represents that its use would not infringe privately owned rights. Refer-
ence herein to any specific commercial product, process, or service by trade name, trademark, 
manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recom-
mandation, or favoring by the United States Government or any agency thereof. The views 
and opinions of authors expressed herein do not necessarily state or reflect those of the 
United States Government or any agency thereof.