Corrosion of ISG by Mg-Si precipitation in presence of Ankerite

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ABSTRACT

Contact with groundwater in the disposal geological site will induce the creation of an amorphous corrosion layer on the high-level radioactive glass. This is connected to silicate saturation conditions in the surrounding medium, and it is influenced significantly by geochemical processes in the near-field minerals at that depth. The international simple glass is a six-oxide borosilicate glass that is commonly used in nuclear interest. It is a simple glass generated from its composition to be an international benchmark glass. The results of the standard materials characterization center leaching tests in double deionized water at 90°C and an initial pH value of 6.3 showed that it reacts with Ankerite for a short period of time. The effect of Ankerite on borosilicate glass durability through magnesium-silicate precipitation has been investigated and confirmed in this study.

KEYWORDS

international simple glass, Mg-Si precipitation, high-level radioactive waste, Ankerite, R7T7 glass

1. INTRODUCTION

The nuclear fuel generated is classified as High-Level radioactive Waste (HLW). The HLW is vitrified and is then discharged into stainless steel canisters as a method of pre-processing, preceding the final disposal of a geologic forming repository deeply during a long-term storage [1–4]. In Hungary, the Boda Claystone at Mecsek site is the candidate host rock of high level radioactive waste [5]. Borosilicate glass, a chemical-durable glass for stabilizing high-level waste by leaching tests for nuclear waste glass corrosion in aquatic environments, is stable against many corrosive parameters, but its durability can be influenced over time by critical extrinsic parameters such as temperature, chemicals, and solvents [6].

Flooding in nuclear waste packs occurs as a result of water diffusion via capillary absorption, which is also a process that influences water diffusion across different rock densities, surface tension, and the angle of contact [7]. Additional criteria should be explored in order to forecast water absorption levels, including the porosity, rhyolitic tuff, and the density of stones [8].

The hydration is due to water diffusion across the glass network and the ion exchange between the positive protons found in the aqueous and the glass alkaline metals, frequently leading to the formation of the surface of an amorphous corrosion layer. In a process defined as a hydrolysis, Ionic-covalent bonds of the most soluble elements in the aqueous solution attack the glass network and a reverse mechanism controlled by pH and temperature will be established if the silica easily dissolved in the aqueous solution can condense as a protective gel layer in the outer shell of the glass [9–11].

The protective gel layer should produce a balancing phase, reducing contact between the glass and the aqueous environment [12–14]. That balance leads to silicate saturation in a solution in which the ultimate silicic acid saturation (H₄SiO₄) occurs. Finally, if the
interventions by iron minerals i.e., Ankerite or other geological factors are not carried out [15–21], the corrosion rate of HLW glass should reach a steady state.

2. MATERIALS

2.1. Ankerite

A natural Ankerite-rich geological sample was employed in this investigation. The sample was crushed into a powder using an electrical mortar grinder of tungsten carbide. The X-ray diffraction measurement proved that the sample is a natural Ankerite including a minor amount of Siderite (Fig. 1).

The iron-rich Dolomites or the iron Dolomites are the most so-called Ankerites. Dolomite-Ankerite is still in a constant transition according to the Fe/Mg ratio. Fe > Mg and Fe > Mn must be present at Ankerite (application of the rule of the predominance cation with predominance valence in each position). That implies that the host rock of the natural Ankerite can be mostly Dolomite [22, 23].

The powder was then sieved using a 100+120 mesh fraction vibratory strainer and collected the 75–125 μm part of the grain size. The Specific Surface Area (SSA) was measured at 225 cm².g⁻¹ using the BET-N₂ approach. A portion with grain size 90 percent below 149.01 μm with an average diameter of 88.91 μm was shown by the Particle Size Analyzer (PSA). For further analytical tests this powdered sample was dried overnight at 105 °C. Inductively Coupled Plasma - Optical Emission Spectrometry (ICP-OES) was used for the elemental analysis of this sample. Thermal analysis has been used to assess the total carbon content. Table 1 shows the elemental composition of Ankerite.

2.2. International simple glass

International Simple Glass (ISG) is a borosilicate six oxide glass with the same elemental ratios as SON68 (the inactive reference glass for the French R7T7 glass). It is commonly used to investigate the HLW glass’s durability. The ISG ingot was cut into 10 × 10 × 2 mm plates (coupons) with a diameter of 12.7 cm. The polishing machines using oil-based diamond suspension spray have been used to polish the ISG coupons. Each coupon’s Surface Area (SA) was geometrically measured with a digital caliper and found to be 2.4 cm² but was adjusted to 3.1 cm² after application of a surface roughness factor equal to 1.5 which was calculated using BET-N₂ to account for the additional surfaces. The Inductive Coupling Plasma-Mass Spectrometry (ICP-MS) techniques were used to detect the ISG’s composition in percentage ratios. The mass fraction of each element/oxide in the ISG reacted is shown in Table 2.

Table 1. Oxide ratios in natural Ankerite, derived from ICP-OES and thermal analysis

| Method       | Element | Mass (%) | Element | Mass (%) |
|--------------|---------|----------|---------|----------|
| ICP-OES     | SiO₂    | 0.56     | CaO     | 11.7     |
|              | K₂O     | 0.55     | Li₂O    | 0.03     |
|              | Na₂O    | 0.88     | Fe₂O₃   | 37.6     |
|              | Al₂O₃   | 0.30     | MgO     | 11.9     |
|              | MnO     | 1.18     | SO₃      | 1.50     |
|              | CuO     | 0.53     | ZnO      | 0.03     |

| Method       | Element | Mass (%) |
|--------------|---------|----------|
| Thermal analysis | CO₂      | 35.7     |

TOTAL (%) 102.5

Table 2. Results of the common oxide ratios and mass percentage ratios in ISG

| Method       | Oxide | Mass (%) | Element | Mole (%) |
|--------------|-------|----------|---------|----------|
| ICP-OES     | SiO₂  | 55.6     | Si      | 26.0     |
|              | B₂O₃  | 16.0     | B       | 4.61     |
|              | Na₂O  | 13.2     | Na      | 4.91     |
|              | Al₂O₃ | 5.50     | Al      | 1.45     |
|              | CaO   | 5.53     | Ca      | 3.95     |
|              | ZrO₂  | 2.98     | Zr      | 2.21     |
| ICP-MS      | Others | <0.79 | Others | <0.25 |

TOTAL (%) ~99.63 ~43.38

The commonly performed static leach tests include the standard Materials Characterization Center (MCC) leach test for nuclear waste forms has been established by the U.S. Department of Energy at the Pacific Northwest Laboratory (PNL) [24]. It provides a comparison of the durability of candidate waste forms developed for the stabilization of high-level nuclear wastes [25]. The MCC-1 test has already been standardized by the American Society for Testing and Materials (ASTM) through committee C-26 as Standard C1220-92 [26].

Fig. 1. The XRD pattern of natural Ankerite sample, collected from the mining area in Rudabánya, Hungary
The very low Surface area to Volume (S/V) used in the MCC-1 test represents the improbable event of the headspace of a waste canister being instantaneously filled with groundwater and provides a simple method of comparing the relative durability of different homogeneous waste glass. The MCC-1 test is commonly used for low S/V (i.e., ~10 m⁻¹) glass at 90°C. Teflon vessels and ASTM deionized water are required for this test [24].

In the current experiment, the MCC-1 test was employed with PolyTetraFluoroEthylene (PTFE) vessels of a 30 ml capacity each. The leaching was conducted at a temperature of 90°C. The test samples were fabricated by adding specific (~300 mg) ISG coupons and (~240 mg) Ankerite powder to the containers, pouring 24 mL of ASTM-I water (having a conductivity and resistivity of 0.055 μS.cm⁻¹, 18.2 MΩ.cm) in each container, then deoxygenating the samples by Argon gas. The pH of the ASTM-I water was adjusted to 6.3 at 25°C by adding 10 μL of 0.08 g.L⁻¹ NaOH.

Throughout this study, three experimental systems were set up and they were admitted in the oven for (3, 7, 14, 28, 90 days) reacting durations at 90°C:
- System (MCC-1): PTFE vessels were assembled using ISG coupons, each coupon was put into a separate vessel;
- System (reference Ankerite): PTFE vessels were assembled using Ankerite powder. This system was used to facilitate Normalized mass Loss (NL) calculation of dissolved elements, resulting solely from the corrosion product of ISG;
- System (MCC-1 + Ankerite): PTFE vessels were assembled using ISG coupons and Ankerite powder.

At the end of the designated leaching period, the hot samples were cooled to room temperature and then centrifuged at 4,500 rpm for 10 min. The pH value of the leaching solution was measured immediately for all samples at 25°C. pH records are plotted in Fig. 2. The leaching solution was then filtered through a Millipore filter (0.45 μm pore size) and 15 milliliters of the filtrate were taken for ICP-MS analysis.

4. RESULTS

4.1. Acidity evolution

In the extracted solutions, pH values were shown (Fig. 2) in two ranges:
(i) pH (7.7–8.3);
(ii) pH (6.2–9.2).

Samples include solely Ankerite powder at 90°C, and samples including Ankerite powder and ISG coupons at 90°C indicate a pH of a range (i). At 90°C the pH values were shown in range (ii) on the reference samples of the glass coupon. Dissolution of Ankerite was found for pH improvements over the first 28 days, which accelerated the ISG corrosion rate [20]. As iron minerals may accelerate their hydrolysis predominantly [27], which will lead to hydroxyl ions being released and H⁺ intake from the solution being enhanced [28].

On day 90, and due to the continuous silicate dissolution in the MCC-1 system, an elevation occurred in the pH value, which increased the silicate’s solubility while the system was attempting toward saturation. Consequently, the higher pH values on day 90 were eventually attributed to glass corrosion [29]. At the same time, the “MCC-1 + Ankerite” system maintained at steady-state pH after 28 days reaction due to the influence of Ankerite, which regulated the pH and clogged the ISG alteration layer’s porosity. This represented a lower rate of ISG corrosion versus MCC-1 as described in Fig. 4.

4.2. Leaching solution analysis

Iron (Fe) concentrations in the ISG systems were lower than the detection limit in the present experiment, which might be owing to its rapid integration into the Si network during the first corrosion phase [16]. Magnesium solubility was observed in various systems. However, lower concentrations in the ISG coupon system were attributable to Mg-Si precipitation, as indicated by the extra silicic acid created as a consequence of increased glass corrosion rates owing to the presence of adequate magnesium in the solution [29].

Due to the development and precipitation of Mg-Si, which may be Sepiolite [30], the main silicic acid concentrations were depleted, and the deficit will be composed of silicic acid provided by glass corrosion. Mg concentrations in the reference Ankerite system achieved a high value on day 90 of this experiment, but the opposite occurred in the “MCC-1 + Ankerite” system. This was due to the initiation of Mg-Si precipitations, which increased silicate solubility as a consequence of the higher pH. The reduced Mg amounts measured at day 90 support this theory (Fig. 3).

The normalized loss (NLₐ) was calculated using Eq. (1) [7] for the main elements resulting from the dissolution of
the glass samples (Na, B, Si, Ca, Al); concentrations of Zr were not sufficient for said calculations,

\[ NL_i = \frac{m_i}{f_i \cdot SA} \]

where \( m_i \) is the mass of element \( i \) in the leachate (g); \( f_i \) is the mass fraction of element \( i \) in the pristine solid (unit less) and \( SA \) is the sample geometric surface area (m\(^2\)).

Figure 4 depicts the effect of Ankerite by displaying total values of normalized mass loss. The MCC-1 method, however, produced better outcomes. Clogging the porosity of gel [31] was probably mainly due to dissolving products of Ankerite [32]. The normalized weight transfer by a layer that evolves as the glass erodes and/or a rise in glass corrosion product concentrations in the solution; Fig. 4 shows the inverse proportional link between leaching duration and ISG corrosion rate. This reduces the driving force of mass transfer, which slows down the several processes that release these components. The corrosion rate \( (r) \) for all leaching methods is shown in Fig. 4 and has been computed by using the following Eqs (2) and (3), used by Neill et al. [15] for ISG:

\[ r = \frac{d(E_{th(B)})}{dt}, \quad (2) \]

\[ E_{th(B)} = \frac{NL_B}{\rho_{ISG}}, \quad (3) \]

where \( r \) denotes the glass corrosion rate (nm.d\(^{-1}\)); \( E_{th(B)} \) is the equivalent thickness of Boron (nm) calculated from the following equation; \( NL_B \) is the normalized mass loss for B (g.m\(^{-2}\)) and \( \rho_{ISG} \) is the density of ISG (2.500 g.cm\(^{-3}\)).

Boron levels were observed to be substantially greater constantly than silicone levels, a typical characteristic concerning corrosion of borosilicate glass. B is, therefore, better suited to assess glass corrosion rate [29, 33]. Simultaneously, the concentrations of silicon in this study do not apply to the monitoring of glass corrosion because of its constant kinetics and the dissolution/evolution of porous texture in the gel layer [34].

In all systems, the ISG corrosion rate decreased with time (Fig. 4) due to the solution becoming more occupied with glass corrosion products [29]. For the first 14 days of the reaction, the MCC-1 system exhibited the highest corrosion rate and total ISG corrosion products when compared to the “MCC-1 + Ankerite” system. However, on day 90, the ”MCC-1 + Ankerite” system began generating Mg-Si precipitations as the silicate solubility increased with increasing the pH.

5. DISCUSSION

In this investigation, iron corrosion products were represented by Ankerite (CaFe(CO\(_3\))\(_2\)), a common iron carbonate mineral found in soils in various clay matrices [35].
The ISG reference system reached higher corrosion rates and lower pH values during the first 14 days of leaching compared with the Ankerite Mixed System (Fig. 4). At the same time, lower levels of dissolved glass per unit area were exhibited; this is because glass corrosion products are highly concentrated in a solution. ISG glass corrosion rates are reduced by a saturation of the surrounding solution with amorphous silicate (SiO₂). The higher the pH, the higher the silicate in a solution, the higher the pH, the more the pH control was achieved by providing the ideal medium for increased silicate solubility. The gel layer formation was not enough as long as it did not show a substantial reduction in the ISG corrosion rate because of insufficient silicic acid concentration [29].

When Ankerite is present, fewer quantities of boron are released from the glass. Simultaneously, silicon concentrations in MCC-1 systems were much higher. On day 90, they released a high of (85 mg.L⁻¹) compared to “MCC-1 + Ankerite” (19 mg.L⁻¹) at the same time. As already stated, any silicate consumption in the solution will result in further supplies of silicate from the glass. Sorption and precipitation by Ankerite took place, but the impact of precipitation was nevertheless more substantial since sorption is typically more intense at higher concentrations of silicon acid [36]. Due to the low surface area of ISG to leachate volume, there were not adequate concentrations of silicon acid. The principal concentration of silicic acid is conceivable when the amorphous layer, SiO₂, begins to disintegrate and dissolve after 90 days of reaction.

In the experiment conducted, NL is a focal factor in the calculation of the glass corrosion rate and is aimed illustrating a better understanding of the normalized elementary loss of the glass components owing to the probable effects of Ankerite on the pH levels. Investigations and analysis of the ICP-MS data show that the released Mg and other corrosion products i.e., Carbonates from Ankerite increased the pH of the leachate in the initial phase of leaching, thereby increasing silicate solubility and initiating early Mg-Si precipitation. However, the presence of Ankerite during the 90 days of reaction was sufficiently effective to reduce ISG’s corrosion rates by (i) regulating the pH to not approach free pH value (≈ 9); and (ii) its products of dissolution were able to clog the newly produced gel layer which decreased the exchange with the external area.

6. CONCLUSION

This paper illustrated the results of our research in which the influence of Ankerite, Mg-Si precipitation, on ISG durability during the 90 days of disposal was studied. The dissociation of Silicic acid increased as pH increased, requiring more glass to be dissolved to achieve saturation in solution. In comparison to the pure glass water system, adding Ankerite regulated the pH. Thus, throughout the entire period, Ankerite’s pH influence on glass corrosion may explain its impact on glass corrosion. In contrast, the Mg-Si precipitation in Ankerite + ISG system was not aggressive until day 90 of reaction due to the clogging effect, and the system was attempting toward saturation. Under the present experimental conditions, the pH-profile of Mg-Si precipitation over longer durations should be investigated further.

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