Aqueous alteration of silicate glass: state of knowledge and perspectives

Stéphane Gin1,2*, Jean-Marc Delaye1, Frédéric Angeli1 and Sophie Schuller1

The question of silicate glass chemical durability is at the heart of many industrial and environmental issues, with certain glasses, such as bioglasses, needing to transform rapidly, while others, like nuclear glasses, extremely slowly. Due to the wide diversity of the chemical composition for these types of materials and their metastability—no thermodynamic equilibrium can be reached between glass and solution—the evaluation of chemical durability remains a scientific challenge. In this article, we review the current state of knowledge on glass alteration mechanisms and kinetics, and point to some perspectives for glasses for which no direct experimental validation is currently possible. Thanks to the development of novel techniques and international collaborations, a better understanding of the mechanisms involved has been achieved. Mechanistic models have been developed at some specific scales, although holistic models still need further development to link the various scales and perform reliable predictions.

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INTRODUCTION

Silicate glasses are manmade materials with various applications in our daily life, ranging from tableware, labware, windows in construction, insulation, and reinforcement, to high technology fields such as optical lenses, displays, fibers, biomedicine, and nuclear waste disposal, all due to their ability to be customized1-3. Among glass materials' properties, chemical durability is important in many of the above-mentioned applications. In particular, the reaction of a glass in aqueous environments and its alteration behavior is critical in applications such as biomedicine (e.g., bone repair)4, where fast and controllable dissolution is preferred, or in agriculture to supply plants with nutrients5. For nuclear waste disposal, on the contrary, it is essential6 to maintain waste glass long-term stability to prevent the release of radioactive elements into the biosphere for periods of thousands to hundreds of thousands of years. Silicate glasses can also be produced naturally on Earth. Basaltic glasses are volcanic glasses found in different locations, including submarine, subglacial, or lacustrine environments. Understanding their alteration mechanisms and rates is a preliminary step toward the understanding of different natural phenomena such as volcano slope stability7, the chemical mass balance of the oceans8, and the geological history of the planet Mars9.

The term 'silicate glass' refers to any kind of vitreous materials in which SiO2 is the most abundant oxide. Among others, these include the well-known categories of soda-lime glasses, borosilicate glasses, aluminosilicate glasses, etc. The terms 'alteration' or 'corrosion' refer to the transformation of the glass into solid alteration products and aqueous species. The term 'dissolution' is employed when glass transforms into aqueous species only.

The alteration of silicate glasses is a complex phenomenon that depends both on the nature of the glass material, i.e., its chemical composition, structure (short and medium-range order), and bonding, and also on the environmental conditions, such as temperature, pH, Eh, and solution chemistry. The plethora of influencing parameters makes alteration difficult to predict from simple descriptors or parameters arising from structural analysis or tests performed in accelerated conditions. Calculation of a glass alteration rate can be further complicated by the fact that different mechanisms or different coupled mechanisms can control glass alteration during the reaction. This has led scientists to distinguish three main kinetic regimes depending on which mechanism or set of mechanisms control the alteration process: the initial dissolution rate regime (stage I), the residual alteration rate regime (stage II), and (for some glasses) a sudden potential acceleration of the alteration (stage III) (Fig. 1). Stage I takes place in diluted and renewed solutions. A high concentration of dissolved silica which reduces the affinity for matrix dissolution and the formation of a passivating alteration layer on the glass surface allows the establishment of stage II with low residual rates, while massive precipitation of secondary phases can trigger an acceleration of alteration (stage III). In this article, we review the progress made in understanding silicate glass alteration mechanisms (second section), and then focus on the three aforementioned kinetic regimes (third section). Note that most of the knowledge and examples given in this article are based on glass behavior in liquid water. However, a separate section gives information on glass interaction with water vapor, as atmospheric alteration is also a key aspect of glass degradation.

AN INTRODUCTION TO THE MECHANISMS OF GLASS ALTERATION

In contact with water, the glass surface first reaches an electrochemical equilibrium10 before undergoing chemical attack through ion-exchange and hydrolysis reactions of metal-oxygen bonds, driven by the difference between the chemical potential of the solid and that of the fluid11. Because of this, the concentration of glass species in the solution increases, meaning a decrease in the Gibbs free energy of the glass-solution system. The behavior of silicate glass also depends on the fact that the reaction

1CEA, DES, ISEC, DE2D, University of Montpellier, Bagnols sur Cèze, Montpellier, France. *email: stephane.gin@cea.fr
between silica and water (Eq. 1) is reversible\textsuperscript{12}
\begin{equation}
\text{SiO}_2 + 2\text{H}_2\text{O} \rightleftharpoons \text{H}_4\text{SiO}_4
\end{equation}

Therefore, in a solution containing orthosilicic acid, matrix dissolution slows down and above the saturation level, precipitation reactions take place, generally on the glass surface. Furthermore, Si atoms facing water and bound to O atoms of the glass network (Si tetrahedra not fully hydrolyzed) can form silanol groups, which can, in turn, condensate to reform siloxane bonds (Eq. 2).

\begin{equation}
\equiv \text{Si} - \equiv \text{OH} + \equiv \text{OH} \rightleftharpoons \equiv \text{Si} - \equiv \text{O} - \equiv \text{Si} + \text{H}_2\text{O}
\end{equation}

These reactions of precipitation and condensation can involve other elements from the glass or supplied by the solution. They result in the formation of an alteration layer, generally composed of a Si-rich hydrated and porous layer, called a gel, and secondary phases\textsuperscript{13}. In some circumstances, the gel layer can be transport-limiting for aqueous species. It then becomes a passivating layer. Figure 2 summarizes the main features of silicate glass alteration.

The rate of glass alteration can be described as the rate at which the glass hydrates and transforms into new, thermodynamically more stable phases (alteration products) and dissolved ions. It can be determined from a mass balance based on the glass components present in the leachate and those retained in the alteration layer. In practice, one focuses on elements not retained in the alteration layer. Such elements are called alteration tracers.

Difficulties arise when trying to link processes described at atomic scale to the observations made at macroscopic level. Glass displays a disordered structure at medium range (>5 Å) characterized by a broad distribution of bond distances and bond angles. See for instance the case of ISG, a 6-oxide borosilicate glass used as a reference material within the glass corrosion community (Table 1)\textsuperscript{14}, in which Si–O–Si bond angle ranges between 120 and 180°, and Si–O bond distance between 1.5 and 1.75 Å, with significant differences between the bulk and the glass surface\textsuperscript{15}. Therefore, a simple reaction, such as the hydrolysis of Si–O–Si, can be more or less easy depending on its local configuration because of a large distribution of the activation energy\textsuperscript{16}. A mesoscale approach is thus necessary to bridge the gap between the atomic level, where first principles or molecular dynamics (MD) calculations can be performed, and the macroscopic level. Moreover, the reacting surface of the glass is usually covered with an alteration layer, which means that local conditions at the reaction front (pH, concentration of dissolved species) are potentially different from those of the bulk solution\textsuperscript{17}. Another difficulty arises from the chemical complexity of multicomponent glasses, as glass/water interaction cannot be described by a simple solid/liquid thermodynamic equilibrium, such as that applicable for pure amorphous silica\textsuperscript{11}. Although one could theoretically define a state corresponding to an equilibrium between the glass phase and the liquid phase, based on enthalpy and entropy contributions\textsuperscript{18}, this equilibrium is not achievable in practice because secondary phases with low solubility and fast precipitation rates will control the solution chemistry, maintaining the glass-solution system well away from saturation. Because of the high complexity of the system and the difficulties listed above, empirically-inspired approaches are still widely used to assess glass durability. Nonetheless, progress has been achieved in the understanding of glass alteration over recent years, thanks to the development of new simulation methods, analytical techniques, and international efforts\textsuperscript{18,19}.  

Fig. 1 Stages of alteration undergone by a glass altered in static mode. (Reprinted from Vienna et al.\textsuperscript{62} with permission, copyright John Wiley and Sons).

Fig. 2 Illustrations of glass transformation into alteration products. a Representation of a cross section of an altered piece of glass. The alteration layer is generally made up of a hydrated layer (also called ‘interdiffusion’ or ‘leached’ layer), the gel layer (amorphous hydrated polymerized Si-rich material) which can be made of several sub-layers depending on how it forms, and secondary phases (precipitates). b Scanning electron microscopy image of a piece of SON68 glass, a 30 oxide borosilicate glass (Table 1), altered for 2 months at 150 °C in deionized water. The sample was broken and observed from the edge. Secondary phases precipitated on the top of the gel layer consist of poorly crystallized phyllosilicates. c Transmission Electron Microscopy bright field image of a SON68 glass sample (composition given in Table 1) altered for 26 years at 90 °C in granitic water. Pores within the gel can be seen in bright.
the recent literature. Figure 3 provides some examples of the mechanisms both qualitatively and based on atomistic simulation techniques have been dedicated to products through the alteration layer. In recent years, many studies exchange, hydrolysis and bond reformation, Ostwald ripening (gel glass. These mechanisms are related to adsorption/desorption, atomistic level, involving species from the solution and from the consequence of the accumulation of countless mechanisms at the macroscopic behavior of a glass in contact with water is the molecular mechanisms of glass alteration.

The macroscopic behavior of a glass in contact with water is the consequence of the accumulation of countless mechanisms at the atomistic level, involving species from the solution and from the glass. These mechanisms are related to adsorption/desorption, exchange, hydrolysis and bond reformation, Ostwald ripening (gel maturation), precipitation, and the transport of reactants and products through the alteration layer. In recent years, many studies based on atomistic simulation techniques have been dedicated to better understanding these mechanisms both qualitatively and quantitatively. A review of the methods available is proposed in the recent literature. Figure 3 provides some examples of the advanced simulation methods used to understand glass alteration mechanisms. In the following sections, we discuss the outcomes of the most relevant studies on the basic processes of glass alteration: Adsorption, exchange, and hydrolysis. Note that because of the timescale accessible to current atomistic simulations (ps to ns timescale), not all the processes listed above have been studied yet.

WATER ADSORPTION ON GLASS

Recent studies have focused on the chemical reactions occurring at the glass (pure silica, bioactive glass, borosilicate glass)—water interface. Results show that water molecules have a higher affinity for alkali or alkaline earth cations or non-bridging oxygen atoms (NBO) than Si–O–Si linkages. Water dissociates and forms Na–OH, Ca–OH, and NBO–H groups. When a film of water molecules is simulated at the glass surface, collective motions occur, and protons resulting from the dissociation of water molecules jump onto water molecules before being captured by a cation or an NBO at the glass surface. These results have been confirmed by classical molecular dynamics (MD) with shell-model potentials. However, no particular affinity between small Si–O–Si rings (size 2 or 3) and water molecules has been observed, although the large local stresses concentrated on the small rings should favor their hydrolysis.

Mahadevan et al. studied the chemisorption of water on a silica surface by classical MD using a dissociative potential for water specially fitted to represent silica–water interface. As inferred from experiments, a water molecule can form two Si–OH, and NBO–H groups. When a film of water molecules is simulated at the glass surface, collective motions occur, and protons resulting from the dissociation of water molecules jump onto water molecules before being captured by a cation or an NBO at the glass surface. These results have been confirmed by classical molecular dynamics (MD) with shell-model potentials. However, no particular affinity between small Si–O–Si rings (size 2 or 3) and water molecules has been observed, although the large local stresses concentrated on the small rings should favor their hydrolysis.

For the sake of comparison, \( r_0 \) has been given at 90 °C pH 9. For each study (T, pH) conditions of \( r_0 \) measurement are given. If they differed from the selected conditions, \( r_0 \) was calculated at 90 °C, pH 9 with Eq. (4). If \( E_a \) and \( \eta \) were not available, \( r_{0,\text{calc}} \) was determined with \( E_a = 60 \text{ kJ} \cdot \text{mol}^{-1} \) and \( \eta = 0.4 \).

\[ r_0 = 8.2 \text{ g} \cdot \text{m}^{-2} \cdot \text{d}^{-1} \text{ at } 90 \text{ °C, pH 9} \]

Table 1. Glass composition and initial dissolution rate, \( r_0 \), of some common silicate glasses.

| Glass composition in mol% | ISG | SON68 | Albite glass | Basaltic glass | Obsidian | Roman glass | Soda-lime | Pyrex | Bioglass 45S5 | E |
|---------------------------|-----|-------|--------------|----------------|----------|-------------|-----------|-------|----------------|---|
| SiO2                      | 60.1| 52.7  | 75           | 61.6           | 83       | 72.4        | 72.9      | 81.0  | 46.1           | 57.0 |
| B2O3                      | 16.0| 14.0  |              |                |          |             | 13.0      |       | 6.1            |    |
| Al2O3                     | 3.8 | 3.4   | 12.5         | 10.9           | 8.2      | 1.1         | 1.0       | 2.0   | 8.8            |    |
| P2O5                      | 0.1 |       |              |                |          |             |           |       |                | 2.6 |
| Fe2O3                     | 3.0 |       |              |                | 5.1      | 0.6         | 0.1       |       |                |    |
| Na2O                      | 12.6| 11.4  | 12.5         | 3.2            | 3.9      | 19.4        | 12.3      | 4.0   | 24.4           | 0.6 |
| Li2O                      | 4.6 |       |              | 2.5            |          |             |           |       |                |    |
| K2O                       | 2.9 |       |              |                |          |             |           |       |                |    |
| CaO                       | 5.7 | 5.0   | 14.6         | 1.0            | 5.4      | 7.2         | 26.9      | 19.6  |                |    |
| MgO                       | 0.3 |       |              |                | 6.6      | 0.4         | 6.5       |       |                |    |
| BaO                       | 0.3 |       |              |                |          |             |           |       |                |    |
| SrO                       | 0.2 |       |              |                |          |             |           |       |                |    |
| ZrO2                      | 1.7 | 1.5   |              |                |          |             |           |       |                |    |
| MoO3                      | 0.9 |       |              |                |          |             |           |       |                |    |
| TiO2                      |     | 0.1   |              |                | 0.4      |             |           |       |                |    |
| Σ2RE2O3                   | 0.9 |       |              |                |          |             |           |       |                |    |
| Other                     | 2.0 | 2.0   | 1.5          |                | <1       | <1          |           |       |                |    |
| Initial dissolution rate, \( r_0 \), in g·m⁻²·d⁻¹ in the (T, pH) conditions given in the reference (Ref.) |
| T (°C), pH | 90 | 90 | 90 | 90 | 157 | 90 | 87 | 90 | 37 | 90 | 90 |
| r₀ | 8.2 | 0.3 | 0.26 | 0.8 | 6 × 10⁻³ | 5.9 | 7.7 | 0.5 | 45 | 4.6 | 9 |
| Ref. | 60 | 166 | 256 | 222 | 257 | 245 | 257 | 257 | 258 | np | |
| r₀ | 8.2 | 0.3 | 0.26 | 1.4 | 2 × 10⁻³ | 2.5 | 23 | 3 | >8000 | 4.6 | |

For the sake of comparison, \( r_0 \) has been given at 90 °C pH 9. For each study (T, pH) conditions of \( r_0 \) measurement are given. If they differed from the selected conditions, \( r_0 \) was calculated at 90 °C, pH 9 with Eq. (4). If \( E_a \) and \( \eta \) were not available, \( r_{0,\text{calc}} \) was determined with \( E_a = 60 \text{ kJ} \cdot \text{mol}^{-1} \) and \( \eta = 0.4 \).

\[ r_0 = 8.2 \text{ g} \cdot \text{m}^{-2} \cdot \text{d}^{-1} \text{ at } 90 \text{ °C, pH 9} \]

Initial dissolution rate \( r_0 \) recalculated at 90 °C pH 9 from the measured value

\[ r_{0,\text{calc}} \text{ at } 90 \text{ °C, pH 9} = 8.2 \text{ g} \cdot \text{m}^{-2} \cdot \text{d}^{-1} \text{ at } 90 \text{ °C, pH 9} \]

Initial dissolution rate \( r_0 \) in g·m⁻²·d⁻¹.
zone of a silicate glass is not a surface but a volume, whose size depends on both the connectivity of the silicate network and the pH.

To go deeper in our understanding of the chemical processes at work, first principle calculations using the DFT method have been performed simulating an interface between a SiO₂–B₂O₃–Na₂O glass and water. Both a single water molecule and a film of water molecules were simulated. Adsorption energy calculations showed that three coordinated B atoms (IIIIB) have a lower affinity for water molecules than four-coordinated B atoms (IVB). However, IIIB atoms are more accessible to water molecules than IVB atoms, which explains why it has been observed experimentally that IIIB atoms are preferentially dissolved rather than IVB. This relationship between the quantity of accessible sites and the alteration rate has also been pointed out in a SiO₂–CaO system.

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WATER DIFFUSION/IION EXCHANGE AT THE GLASS–WATER INTERFACE

Beyond adsorption, hydration, and exchanges between water and mobile glass cations (alkaline, alkaline earth elements) have been studied experimentally and by using different molecular simulation techniques. The importance of the glass topology, in particular the ring size distribution, was underlined decades ago by Bunker. Water molecules are able to penetrate the glass by diffusion only if voids with diameters larger than ~0.7 nm are available (the water molecule diameter is equal to 0.28 nm).

Otherwise, water penetration occurs mainly through hydrolysis reactions. This threshold at around 0.7 nm is important, as it separates two diffusion regimes: liquid-state diffusion above the threshold with $D_{\text{app}}$ of the order of $10^{-9}$ m$^2$.s$^{-1}$ and solid-state diffusion below with $D_{\text{app}} < 10^{-15}$ m$^2$.s$^{-1}$. Note that recent MD simulations have tended to lower the threshold below 0.5 nm in pure silica. Interestingly, it was shown recently that, owing to complexity of the topology and the existence of multiple pathways for water molecules, actual gels cannot be characterized by a unique $D_{\text{app}}$. It is thus better to consider both open channels and closed pores. In simplified systems, MD techniques provided a more detailed description of water diffusion mechanisms involving alkali–H$_2$O exchanges, proton diffusion, and a possible braking role of the glass surface on the water diffusion. In porous silica, a proxy for the gel of multi-component glasses, the diffusion of water molecules is impacted by the topology of the porous network. In particular, water diffusion slows down in low connected pores. Ohkubo et al. also observed a large decrease in the water diffusion compared to diffusion in bulk water in simulated gels prepared from an alumino-borosilicate glass after removing the soluble species (B and Na). A water diffusion coefficient of $8 \times 10^{-16}$ m$^2$.s$^{-1}$ was calculated. The interactions between the water molecules and the atoms located at the interface between the reticulated network and the pores are responsible for this low diffusivity compared to water self-diffusion ($\sim 2 \times 10^{-9}$ m$^2$.s$^{-1}$ at room temperature). Similar observations were obtained with a cylindrical pore dug out in a silica glass containing pure water or alkalis in water. The water diffusion coefficient decreases as the ionic radius of the alkali increases.

In a system simulated by ReaxFF and consisting of three layers (pure silica, an intermediate gel layer, water), water molecules diffused significantly more slowly in the gel than in bulk water. At ambient temperature, the decrease factor is roughly equal to 30 compared to the diffusion coefficient of bulk water. Moreover, as the gel–water interactions proceed, the gel network becomes more reticulated because of the hydrolysis of the weakest Q$_n$ species (Q$_n$ stands for the reticulation of the silicate network; $n$ is the number of bridging O linking a Si atom to the network, $n$ can vary from 0 to 4), and because of the cross-linking siloxane bonds. Interestingly, the gel repolymerization could be
linked to the drop in the alteration rate, as pores formed by the release of mobile species tend to close\textsuperscript{30,36,37}. When a system with a SiO\textsubscript{2}–Na\textsubscript{2}O glass in interaction with water was simulated with the same ReaxFF potentials used in Rimsza and Du's work\textsuperscript{38}, different reactions occurred depending on the depth. Near the surface, H\textsubscript{2}O dissociated to form a Si–OH group accompanied by a Na\textsuperscript{+}/H\textsuperscript{+} exchange. Deeper below the surface, H\textsubscript{2}O dissociation mechanisms always occurred in conjunction with H\textsuperscript{+} jumps from NBO to NBO. Finally, below the subsurface region, only H\textsuperscript{+} jumps were observed. These mechanisms have been confirmed in a recent simulation by Mahadevan on aluminosilicate glasses with varying Na contents\textsuperscript{39}. When the Na\textsubscript{2}O content increases, water penetrates faster and silanol groups form more deeply. In addition, some dissolved Na atoms remain close to the glass surface, surrounded by water molecules.

The possibility for a proton to jump from site to site has also been evidenced in a work using DFT calculations dedicated to the hydration of aluminosilicate glasses\textsuperscript{40}. After the bond dissociation, Si–OH and Al–OH groups formed. During the remaining simulations, different reactions of proton exchanges around the Si or Al sites were observed.

Using Car Parrinello Molecular Dynamics, Tilocco et al. showed that in dry bioactive glasses with quite a low SiO\textsubscript{2} content, Na ions can migrate by a jump process with the initial and final polyhedra around the Na sharing common O atoms\textsuperscript{41}. It is also possible for a Na ion to pass temporarily through a Ca site during its migration, due to the considerable flexibility of the glass network. When the glass surface is in contact with water, Na ions can exchange with H\textsubscript{2}O molecules inducing water dissociation\textsuperscript{42}.

Coupling experiments and classical MD simulations, the blocking role of Ca in relation to the water diffusion has been evidenced\textsuperscript{43}. In particular, in the composition domain investigated in the study, an increasing content of CaO in the glass resulted in a decrease in the amount of bottlenecks larger than that of the water molecule, which in turn led to a drop in water diffusivity. In borosilicate glasses, the mixing of Si\textsuperscript{4} and Si\textsuperscript{6} complexes the mechanisms\textsuperscript{16}. When a Na\textsuperscript{+} near a BO\textsubscript{4}\textsuperscript{−} is exchanged with a hydronium ion, one B–O–Si bond breaks, and the Si\textsuperscript{5} is converted into a Si\textsuperscript{6}. In parallel, a silanol group forms and the hydronium dissociates.

The examples given here emphasize the complexity of the ion-exchange processes taking place in silicate glasses. They account for the experimental observations related to water labeled with isotopes such as \textsuperscript{2}H = D or \textsuperscript{18}O, where, even in simple soda-lime glasses, the exchange ratio D/Na can vary from 1 to 3\textsuperscript{44,45}. Furthermore, they explain how ion-exchange and water ingress in the solid work together, as experimentally evidenced by Rutherford backscattering (RBS) and Resonant nuclear reaction analysis (RNRA) techniques applied to various samples of natural and nuclear glasses altered between 60 and 200 °C\textsuperscript{46}. Regarding the time scales accessible to the simulation techniques, it is still not possible to numerically investigate low apparent diffusivity determined experimentally in a passivating gel, where values lower than 10\textsuperscript{−20} m\textsuperscript{2}/s have been reported\textsuperscript{46,47,48}. This would however be of great interest in order to explain the cause and the dynamics of passivation.

**HYDROLYSIS**

The investigation of hydrolysis mechanisms by first principles calculations began several decades ago by focusing on F\textsubscript{1}–O–F\textsubscript{2} bridges (F\textsubscript{1,2} = Si or Al) in reduced clusters, Q\textsubscript{n} entities, or small –O–Si–O– rings\textsuperscript{49–52}. It appeared that the results depended on several factors, namely the cluster size, the cluster termination types, and the number of water molecules around the cluster\textsuperscript{50,53}. For instance, Pelmschensikov demonstrated that the activation energy for hydrolysis, \( E_{\text{hydrolysis}} \), i.e., energy barrier necessary to form the activated complex, increases significantly when a Si–O–Si linkage is embedded in a crystalline structure; the larger the connections with the structure, the higher the \( E_{\text{hydrolysis}} \). From Pelmschensikov’s results, it can be inferred that the breaking of the last bond around a SiO\textsubscript{4} tetrahedron in pure silica will be the easiest, energetically. This would confirm a hypothesis made in the 1980s, leading to the widely used macroscopic first order law (Eq. (3), Table 3) to calculate a glass dissolution rate as a function of the concentration of dissolved silica\textsuperscript{54}.

Despite the above-mentioned limitations, calculations with small clusters are of interest and enable the topological characterization of the different steps leading to the bond breaking. In particular, it has been shown that during the Si–O– Si breakage, an intermediate five-coordinated Si atom is formed conjointly with the H\textsubscript{2}O dissociation, before the equilibration of the final configuration formed with two separated SiO\textsubscript{4} entities containing Si–OH groups\textsuperscript{55}.

Work by Kagan et al. has proposed a statistical analysis of different sites at the surface of a silica–water system, using the potential mean force (PMF) method to determine the activation energies for hydrolysis of various linkages\textsuperscript{56}. The force fields used in this study were those developed specially to simulate hydrated silica systems by classical molecular dynamics\textsuperscript{24,36}. The Q\textsubscript{3} to Q\textsubscript{2} and Q\textsubscript{2} to Q\textsubscript{1} conversions have \( E_{\text{hydrolysis}} \) equal to 58.6 kJ/mol. The Q\textsubscript{3} to Q\textsubscript{2} conversion has a lower activation energy because of local strains around the Q\textsubscript{2} entities on the silica surface. \( E_{\text{hydrolysis}} \) associated with the Q\textsubscript{1} to Q\textsubscript{0} conversion is equal to 52.7 kJ/mol. The fact that the hydrolysis of the last bond around a Q\textsubscript{0} entity corresponds to the lowest energy barrier is in agreement with Pelmschensikov’s work\textsuperscript{57}. It was also shown that \( E_{\text{hydrolysis}} \) of a –Si(OH)\textsubscript{3} group is only slightly dependent on the nature of the connected Si\textsuperscript{58}. This conclusion was derived from calculations on a surface of quartz, where –Si(OH)\textsubscript{3} groups were connected to either an edge site, a kink site, or a terrace site.

To our knowledge, Zapol et al.’s work is the only study dedicated to hydrolysis mechanisms in borosilicate glasses using ab initio methods\textsuperscript{59}. The authors showed that \( E_{\text{hydrolysis}} \) necessary for the dissociation of B–O–B and B–O–Si linkages are lower in acid conditions compared to neutral or basic conditions, which is in fair agreement with experimental observations pointing out that the preferential dissolution of B compared to Si increases with decreasing pH\textsuperscript{60,61}. Moreover, the \( E_{\text{hydrolysis}} \) for B–O–B and B–O–Si linkages dissociation are much lower in acid and neutral conditions compared to the \( E_{\text{hydrolysis}} \) for Si–O–Si dissociation, but the gap decreases in basic conditions.

As a whole, despite progress in this area, there is still a lack of statistics in the determination of the activation energies for the dissociation of the various linkages in silicate glasses to account for structural disorder in these materials. This limits the use of these data to investigate macroscopic systems.

**MECHANISMS RESPONSIBLE FOR THE INITIAL DISSOLUTION RATE**

At the macroscopic level, the initial dissolution rate, \( r_0 \), depends on both the glass composition and environmental factors such as pH, temperature, and the solution composition\textsuperscript{52,62}. From a mechanistic standpoint, \( r_0 \) results from the following processes described above: adsorption, exchange, and hydrolysis reactions. We remind the reader that in this regime, the solution remains undersaturated, with no secondary phases able to form in the conditions tested. From the state of knowledge previously discussed, we can make several observations: (1) The mechanisms at molecular level are extremely complicated, as they involve collective effects of water molecules and concern not only a linkage but the whole local environment around a given atom (at least the second shell of cations undergoing hydrolysis). (2) The release of a glass cation by exchange or hydrolysis involves local structural reconfigurations after its release, such as the possible
change in coordination of the surrounding cations, the formation of silanol groups, or the diffusion of protons or water molecules. To date, scientists have not yet investigated all the linkages and configurations even within simple silicate glasses by classical MD or ab initio techniques. Therefore, some fundamental parameters of glass dissolution are missing. (3) Ion-exchange and hydration processes are intimately coupled with hydrolysis of covalent bonds forming the silicate network. The way the first mechanism impacts dissolution of Si from the glass is not yet understood. The main theory for glass dissolution relies on that developed for silicate minerals in the 1980s. This approach, inherited from the transition state theory (TST), supposes that dissolution of the entire mineral or glass is controlled by an elementary reaction. The theory was later improved by Oelkers, who considered a series of elementary reactions as the reaction progresses. However, glasses differ from minerals in their structural disorder and in the possibility of water diffusion within the structure after exchanges with lattice modifier cations. It has been shown experimentally that the degree of disorder within the glassy structure, generated by fine-tuning the cooling rate of the molten glass, significantly impacts .

MECHANISMS RESPONSIBLE FOR THE RESIDUAL RATE

It has taken many years for the glass corrosion community to understand the fundamental reasons for the drop in the alteration rate, but it seems well established now that a thermodynamic effect and a transport-limiting effect work together. The thermodynamic effect can be seen as a decrease in the chemical affinity for the hydrolysis of the silicate network (Si–O–M linkages, M = glass formers). The transport-limiting effect is related to the formation of a Si-rich gel layer, whose structure reorganizes and becomes a diffusion barrier for aqueous species. In a transient period between Stages I and II, the concentrations of glass network formers in solution increase and eventually reach steady states. These are interpreted in a first approximation as the saturation of the fluid with respect to the alteration layer. Since the alteration layer is generally made up of several phases, several competing equilibria must be considered. This makes the system impossible to describe by a single equation, unless it can be demonstrated that a single reaction controls the whole system. Such a demonstration applied to multicomponent glasses has not yet clearly established. Moreover, even if only a gel forms on the glass surface, it is difficult to consider that a single thermodynamic equilibrium can describe the system because the gel does not necessarily form by precipitation of aqueous species, and does not necessarily have a constant composition and structure in volume and time. A time and depth-dependent composition and structure would suppose that the equilibrium continuously changes. In that case, a solid solution model would then be preferable to accommodate these dynamics.

A combination of simulation and analytical techniques, including MD simulation, atom-probe tomography (APT), time of flight secondary ion mass spectrometry depth profiling (ToF-SIMS), transmission electron microscopy (TEM), spectroscopic ellipsometry (SE), infrared spectroscopy (IR), nuclear magnetic resonance spectroscopy (NMR), and and dyes diffusivity measurement, have been deployed on ISG glass. The goal was to better understand the mechanisms of gel formation and maturation. Using different Si isotopic ratios in the leaching solution and in the glass, and post mortem characterizations, it was demonstrated that the gels formed in circum-neutral and slightly alkaline pHs resulted from an in situ reorganization of the silicate network after partial detachment of Si enabled the release of the soluble elements. This reorganization implies that condensation reactions of siloxane bonds occur before all the bonds linking one Si tetrahedron to another are hydrolyzed. In these cases, only the outermost gel layer is in equilibrium with the bulk solution. Another qualitative proof of in situ replacement was supplied a long time before the collaborative work on ISG by post mortem characterization of highly irradiated glasses which were embodied in the gel part of the fission tracks inherited from the parent glass. A gel formed by the precipitation of dissolved species would not have kept such features. Although the interfacial congruent dissolution/precipitation (ICDP) model is valid for most silicate minerals, it is now clear that the formation of gel layers on glass can proceed in different ways. The weight of the local mechanism compared to the complete dissolution/precipitation process depends on the glass composition and the pH. This point of view remains widely debated in the literature, as certain authors claim that the ICDP model can be generalized to any kind of silicate material. Several promising ways to settle this controversy and develop a general theory for silicates would be to:

(1) Investigate the behavior of various polymorphs to understand how the structure of the dissolving material impacts the characteristics of the alteration layer (see for instance the work comparing albite glasses and crystals).
(2) Explore a broad compositional domain from highly depolymerized to highly polymerized glasses under both acid and basic pH conditions.
(3) Probe the leaching materials at atomic and nanoscopic scales into both the structure of the materials and the reactivity of the surfaces by spiking solutions or solids with isotopes and monitoring their behavior with advanced analytical techniques, or by using in operando techniques such as in situ TEM or Raman.

The existence of a residual alteration rate was postulated for the first time in 1984 by Grambow, who argued that no thermodynamic equilibrium between the glass surface and the solution could take place, and that hydration of the glass would never stop. It was then experimentally verified in 1990. Beyond Grambow’s hypothesis, several other possible individual or coupled rate-controlling mechanisms have been evaluated, such as ion-exchange, precipitation of secondary phases, gel dissolution, and diffusion of water, silicon, or tracking elements through the gel, but no definitive demonstration has been made. Consequently, no model is currently comprehensive enough to accurately predict the residual rate of a given glass altered in a given set of experimental conditions (kinetic models are discussed in section “Holistic models for glass alteration”). Several factors account for this profusion of hypotheses: (1) many intrinsic and extrinsic factors affect glass behavior (composition and structure of the glass, temperature, pH, solution composition, and radioactivity), making the development of a unified approach challenging. (2) Residual rates are generally extremely low (3 to 5 orders of magnitude lower than ), resulting in small amounts of alteration products, which in turn make detailed characterizations difficult. Last but not least, (3) the discovery of a correlation does not mean that the root causes have been identified. There is still some work to be done to develop a truly comprehensive understanding of the residual rate. It could start from the observation that the interfacial dissolution/precipitation mechanism and the leaching mechanism constitute two extreme cases, and that all the intermediate situations are possible and can be explored by a glass/solution system, even during an experiment.

EFFECTS OF SOLUTION COMPOSITION

Both the pH and the ions in solution can affect the mechanisms of silicate glass alteration. The pH affects the molecular mechanisms of bond breaking and bond reforming, as most of the chemical reactions involve H+ or OH−. The pH also affects the speciation of elements in water, and thus the equilibria between solid phases and the aqueous species. Siaq diminishes the rate at
which Si\textsubscript{glass} detaches from the surface. This effect, related to the reversibility of the dissolution of SiO\textsubscript{2} in water (Eq. 1), impacts the whole alteration process as, except for highly depolymerized glasses which undergo ion-exchange, the other glass constituents cannot be released if Si–O–Si bonds are not broken. The first order rate laws were derived on the basis of this observation (see section “Continuous scale models”)): 

\[
r = r_0 \left(1 - \frac{[H_4SiO_4]}{K}\right)
\]

(3)

where \( r \) is the glass corrosion rate, \( r_0 \) is the initial (or forward) dissolution rate, \([H_4SiO_4]\) is the activity of orthosilicic acid at the glass surface, and \( K \) is the equilibrium constant (which equals the activity of orthosilicic acid at saturation).

It is worth noting that certain silicate glasses are more soluble than amorphous silica (SiO\textsubscript{2}am). This is the case, for instance, for Al-free high B, Na content glasses\(^{106}\). In other words, the saturation of the solution with respect to SiO\textsubscript{2}am, the glass still dissolves and releases orthosilicic acid. Consequently, the solution becomes supersaturated until it reaches the threshold where SiO\textsubscript{2} particles precipitate. The additional amount of Al to these glasses decreases their apparent solubility, which dramatically affects the mechanisms of gel formation once silica saturation conditions are reached\(^{69}\).

Certain elements other than Si\textsubscript{glass} in solution can also affect the mechanisms of bond breaking\(^{69}\) or the diffusivity of the passivating layer\(^{94}\), but the greatest influence of exogenous elements is on the nature of secondary phases formed during the alteration process\(^{86,92,95–101}\). There has been a general observation that the elements such as Fe, Mg, or Ni precipitating with Si to form silicate minerals will increase glass alteration rate. To form these precipitates, Si can be supplied either by the solution or by the gel layer. In the first case, the drop in the concentration of Si triggers the dissolution of the gel, and thus sustains glass alteration. The other reason is that secondary silicate phases do not passivate the glass surface, at least in short-term experiments. If the gel is passivating, the consumption of Si makes the gel more porous, less dense, and this accelerates the transport of reactants and products. Some examples are given in the following sections. There is a wide variety of silicate minerals which can form during glass alteration: amorphous silicates, phyllosilicates, calcium silicate hydrates, zeolites... In a first approximation, the consequence on glass durability depends on the type and rate of secondary phases formed\(^{91}\).

### EFFECTS OF MICROORGANISMS

Although most of the studies on glass alteration published in the literature to date have focused on abiotic processes, it has also been established that microorganisms can interact with glass, leading to potential synergistic effects\(^{102}\). Microorganisms (bacteria, fungi) have been shown to impact glass alteration by a variety of mechanisms, especially through a direct impact of bacteria attached to the glass surface\(^{103}\) and also by the possible effect of microbial metabolites excreted by the cells\(^{104}\). The formation of a biofilm on the glass surface is not systematic. It depends on physical and chemical factors, which may be influenced by the glass composition and particularly by the presence of nutrients in the glass (Fe, Mn, P...), surface (substrate and organism) hydrophobicity, charge distribution, solution chemistry (i.e., pH and ionic strength), contact angle, and roughness\(^{104}\). Biofilm formation on the glass surface can act as a diffusion barrier for aqueous species, induce mechanical stress, or favor the formation of secondary phases (biomineralization)\(^{102}\). Depending on which mechanism dominates in the conditions of interest, the effects of microorganisms can either enhance or reduce glass alteration\(^{105,106}\). For nuclear waste glasses under geological storage conditions, the direct effect of microorganisms on glass alteration should be neglected, but redox reactions involving multivalent elements such as Fe or S could impact the local chemical conditions that could indirectly impact the glass. These potential indirect effects require further investigations.

### EFFECTS OF IRRADIATION

Nuclear waste glasses contain fission products (mostly \(\beta\) and \(\gamma\) emitters) and minor actinides (mostly \(\alpha\) emitters). Radioactivity can affect glass durability either through structural damage within the glass structure or through the radiolysis of the interfacial water, modifying the chemical environment in the zone under reaction. Most of the structural damage within a glass is due to recoil nuclei associated with \(\alpha\) decay\(^{107}\). Defects in the solid are usually in the form of dangling bonds (non-bridging oxygen atoms and peroxo bridges), oxygen-deficient centers, molecular oxygen, and coordination changes of certain network formers such as boron\(^{108,109}\). In borosilicate glasses, ballistic damage causes an increase in tri-coordinated B species at the expense of four-coordinated B species, with a detrimental consequence on the fraction of Na modifier\(^{110}\). The impact of irradiation on the medium-range order results in an increase in the disorder\(^{111,112}\). MD simulations reveal an increase in the mean ring size in the glass, which can increase water diffusion in the material\(^{113,114}\). Healing processes have been observed, which can mitigate the detrimental effects mentioned above\(^{111,115}\).

Alpha-emitters are generally less soluble elements which are highly retained in the gel and secondary phases\(^{107}\). The gel also undergoes radiation damages that could impact its thermodynamic and transport properties. The consequences on the glass of such changes require further investigation\(^{116}\).

Together, the different types of structural change can impact Stages I and II of glass alteration owing to a significantly higher reactivity of the reacting interface as compared to a non-irradiated glass sample\(^{109}\). The differences in behavior between alpha-irradiated and non-irradiated specimen depend on both the glass composition and the dose of irradiation, until a plateau is reached corresponding to a complete irradiation of the whole sample\(^{116,117}\). In general, simple glasses and high doses mean a greater impact. Regarding the mechanisms involved, the potential effects of irradiation on gel properties, the speciation of certain redox-sensitive species, and the nature of secondary phases remain subjects of investigation, especially when considering the complex scenarios expected for nuclear waste glass in a geological repository.

### EFFECT OF SURROUNDING REACTIVE SOLIDS

The presence of other solids near glass can change the driving forces for glass dissolution and thus the overall dynamics of the system. The effects on glass durability resulting from interactions with other solids chiefly depend on the transport of chemical species in the fluid between the dissimilar solids. If transport is slow or the distance is large, the effect will be small, and vice versa. We detail two emblematic examples: nuclear waste glass in interaction with a metallic canister, and bioactive glass interacting with bone.

### NUCLEAR WASTE GLASS

Materials in contact with or near a borosilicate nuclear glass may include carbon steel\(^{118–125}\), stainless steel\(^{126–130}\), and their corrosion products, including magnetite (Fe\textsubscript{3}O\textsubscript{4})\(^{131–136}\), goethite (FeOOH)\(^{134}\), iron hydroxide (Fe(OH))\(_\textsubscript{2}\)\(^{120}\), siderite (FeCO\textsubscript{3})\(^{123}\), or various types of cementitious materials\(^{137,138}\). Several mechanisms have been identified: (1) precipitation of iron-silicates, which suppresses the saturation of Si in solution, and continuously drives...
glass alteration at a higher rate than that of glass altered without the presence of metals and their corrosion products118,121–125,139. (2) sorption120,128,131,133,134 and precipitation132 of silica onto the surface of metallic iron and Fe-containing corrosion products. (3) adverse effects of Fe2+ and Fe3+ cations140,141, which can strongly acidify the solution. (4) incorporation of Fe into the glass alteration layer, potentially deteriorating the passivating properties of the gel121–125. It is generally believed that like precipitation in the bulk solution, the precipitation of iron-silicates in porous gels may act as a sink for silicates, because these precipitates remove Si from the gel and hinder condensation reactions and gel maturation125,136,142. As a result, the gel is destabilized and the alteration rate of glass increases. In addition, the formation of Fe- and Si-containing phyllosilicates consumes hydroxyl ions and induces a pH decrease, resulting in the enhancement of glass alteration via the ion-exchange mechanism142. Finally, (5) enhanced alteration driven by the localized corrosion of metals129,143 (Fig. 4). More details regarding the corrosion interactions between different materials can be found in a recent review article143. The presence of cement near the glass can have opposite effects (increase or decrease glass alteration) depending on the pH137,144. In this case, the role of Ca is key.

BIOACTIVE GLASS

Bioactive silicate glasses are used in medicine to stimulate bone regeneration4. The four-component glass 45S5 (Table 1) was discovered 50 years ago by Larry Hench, and since then more than a million patients have undergone surgery where this glass is used to repair bone defects in the jaw and in orthopedics5. For such applications, glass particles are pressed into the bone defect. The bioactivity of the glass is intimately linked to its alteration properties. Bone bonding is attributed to the precipitation of a hydroxyapatite (HCA) layer on the altered glass surface, which interacts with collagen fibrils of damaged bone to form a bond. Furthermore, the release of Si and Ca from the glass stimulates osteogenesis, and HCA provides a surface suitable for osteogenic cell attachment and proliferation. The mechanism of HCA precipitation is well understood. First, the glass undergoes ion-exchange, leading to a rapid release of Na and an increase in the pH near the glass surface. High local pH promotes hydrolysis and a condensation reaction of the remaining silica layer. The migration of Ca and P through the silica layer eventually leads to the formation of an amorphous film of Ca–P, which reacts with carbonate ions and hydroxyl groups to crystallize HCA4. The high pH near the glass surface is also known for its bactericidal properties145. The biological interactions of the HCA layer are complicated and less understood. They involve protein actions, incorporation of collagen fibrils, attachment of bone progenitor cells, cell differentiation, the excretion of bone extracellular matrix, and eventually its mineralization146. There are many other applications of bioglasses and many other compositions than that of glass 45S5, including glasses from other families (borate, phosphate glasses)147.

ATMOSPHERIC ALTERATION

The literature on glass alteration by water vapor is quite limited compared to that in liquid water. A recent review by Majérus et al.148 concluded that the alteration of silicate glass in relative humidity (RH) <100% differs from that in liquid water at high S/V because the hydration process does not involve element release into the fluid, but rather a redistribution within the alteration layer, except for the elements which could be volatilized149. Therefore, the flux of water molecules, the local chemistry near the glass surface, and the porosity of the hydrated layer differ from the liquid medium. This results from the difference in the mechanisms between the two media, so that glasses that are classified as the most durable in aqueous phase can be the least durable in vapor phase150,151. In other words, to date, good knowledge of how a glass behaves in liquid water cannot be used to infer its behavior in vapor phase.
The RH plays a key role, as it determines the number of layers of water molecules that build on the glass surface\(^{152}\). In general, the hydration rate (in this situation, the authors refer to hydration rate instead of alteration, corrosion, or leaching rate) increases with RH and temperature\(^{153}\). Depending on the glass composition, temperature, and RH, the hydration rate can be constant or decreasing following a square root of time dependency, potentially related to a diffusion-controlled behavior\(^{154,155}\). Even in a square root of time behavior, the apparent activation energy is high (>60 kJ.mol\(^{-1}\)), suggesting that coupled processes of transport of reactive species and bond breaking operate simultaneously. A recent study showed that some elements such as B or I can be volatilized during atmospheric alteration at 90 °C\(^{156}\). This makes the hydrated layer more porous and reactive than expected.

Secondary phases form more rapidly in atmospheric conditions because of the confined conditions. This could be explained by the local conditions at the glass surface that differ from that in liquid water. Bates et al. estimated that the \(S/V\)—where \(V\) is the volume occupied by the water molecules adsorbed on the glass surface—\(S\)—is 10\(^3\) times greater in vapor phase than in liquid phase\(^{157}\). Abrajano et al. estimated a pH of about 10–12 in the film of water formed in vapor phase for an initial pH of 6–7\(^{151}\). Various salts can precipitate with alcalis and volatile species such as chlorine, sulfate, carbon dioxide... Moreover, silicate minerals such as calcium silicate hydrates, zeolites, or clay-type minerals can also form. The precipitation of secondary phases gives direct effects, such as a modification of the visual appearance of the material along with changes to the chemical conditions at the glass surface, promoting or reducing further alteration\(^{148}\).

The main lesson from the recent work in this field is that the current understanding of glass alteration in atmospheric conditions cannot be deduced from knowledge obtained in liquid water. Although the processes at a molecular level are the same, the interplays between the mechanisms differ and this has an impact on the macroscopic behavior of the material. Therefore, there is a need to connect the various scales in order to decipher the rate-limiting steps depending on all the intrinsic and extrinsic variables.

**KINETICS OF GLASS ALTERATION**

In this paper, we have followed the widely used description of glass alteration in three kinetic regimes (Fig. 1)\(^{62}\). Stage I for ion-exchange and matrix dissolution, with no backward reaction of condensation and precipitation capable of impacting the solution chemistry or the properties and reactivity of the glass surface; Stage II for the residual rate regime associated with a Si-saturated solution, a passivating gel, and potential secondary phases; and Stage III for a potential acceleration of glass alteration due to the partial or total loss of the protective layer of the gel.

**STAGE I**

Among the three kinetic regimes listed above, Stage I may be seen as the most simple, as the glass undergoes only dissolution and the solution remains far from saturation, meaning that no backward reaction can occur. However, owing to the complexity of the glass structure, predicting \(r_0\) remains challenging. To date, no model has been developed which is capable of predicting, a priori and without resorting to empirical parameters, the initial dissolution rate of a silicate glass in a given environment. In this regime, the most common practice is to calculate the glass dissolution rate from the release of Si into the solution from static (batch) or dynamic (flow through) leaching experiments. This release is generally linear with time and congruent with the other glass elements. This is also true for glass modifiers sensitive to ion-exchange, but only if the glass is highly polymerized and the pH not too acidic. Otherwise, glass modifiers are released incongruently and faster than Si due to a fast \(\text{H}^+ / \text{Na}^+\) exchange. As an example, the dealkalized layer is about 100 nm thick in champagne bottles which have held wine for many years\(^{158}\). In lead-crystal bottles, unlike Pb, Na and K also undergo preferential leaching in acetic acid solutions simulating beverages\(^{159}\). Pb, which is a major health hazard, is strongly retained in the alteration layer because the silicate network repolymerizes after the release of alkali.

Figure 5 shows the typical evolution of the normalized mass loss of Si and Na under Stage I glass corrosion. Na can be preferentially released depending on its content in the glass and the pH. The stars correspond to the time necessary to reach a steady state (release rate of Na = release rate of Si).

\[
NL(i)(g \cdot m^{-2}) = \frac{C(i) \cdot V}{S \cdot x_i} + \sum NL(i)_{\text{samplings}} (4)
\]

where \(C(i)\) is the concentration of an element \(i\) in the solution, \(V\) the volume of solution, \(S\) the glass surface area, \(x_i\) the mass fraction of \(i\) in the glass, and \(\sum NL(i)_{\text{samplings}}\) represents the contribution of the various samples taken in the leachate. The glass composition is known (nominal or analyzed composition), concentrations of \(i\) and solution volume are monitored over time, and \(S\) is treated with simple considerations. Depending on the authors, either the specific surface area determined from gas adsorption methods or the geometric surface using spheres as proxies for glass particles have been considered. Furthermore, a glass surface area can be considered as constant or decreasing with time\(^{160}\). For more details, see Fournier et al.\(^{161}\). In most kinetic studies, \(S\) is treated with simple considerations which do not take into account what actually happens at small scale. From a general standpoint, this issue must be considered cautiously, as several studies have reported significant changes to the reacting surface area during the reaction due to roughening\(^{48}\), pitting\(^{162-164}\), or cracking\(^{165}\).

\[
r_0(g \cdot m^{-2} \cdot d^{-1}) = \frac{d(NL(Si))}{dt} (5)
\]

\[
r_0(\mu m \cdot d^{-1}) = \frac{r_0(g, m^{-2} d^{-1})}{\rho} (6)
\]

where \(\rho\) is the glass density.
Another accurate way to calculate \( r_0 \) is to monitor the retreat of the glass surface as a function of time. This can be performed by vertical scanning interferometry (VSI), after coating a small portion of the surface to serve as a reference\textsuperscript{155,164}. But, if a gel forms, this technique will underestimate \( r_0 \).

Among the various parameters impacting the magnitude of \( r_0 \), we find the temperature and the pH yield effects, which are well understood, while the effects of glass composition\textsuperscript{167} and solution composition are less understood. The effect of temperature and pH can be modeled with an Arrhenius law and a power law, respectively:

\[
r_0 (g \cdot m^{-2} \cdot d^{-1}) = k \cdot e^{\Delta H / RT} \cdot 10^{-n \cdot pH}
\]

(7)

where \( k \) is a kinetic constant, \( E_r \) an apparent activation energy, and \( n \) a coefficient for the pH dependency. Similarly to crystals, \( t_0 \) against pH displays a V-shape curve with a minimum at a pH corresponding to the zero net proton charge (pH\text{\textsubscript{net}}), and an increase in both acid and basic regions for highly polymerized glasses\textsuperscript{159,168–170}, or a continuously decreasing trend for low polymerized glasses\textsuperscript{171}, as reported for silicate minerals. There has been no systematic study on the effect of solution chemistry on \( r_0 \). Several potential mechanisms have been identified, such as the formation of outer-sphere or inner-sphere complexes between interactions in solution and surface sites, the modification of the electric potential of the interfacial double layer, the nucleation of secondary phases, or the modification of the interfacial pH\textsuperscript{17,92,144,172,173}. These effects can account for a supplementary term in Eq. (7), as suggested by previous work on silicate minerals\textsuperscript{174}.

A challenging task for the glass community is to understand the effect of glass composition on \( r_0 \). Table 1 summarizes the values of \( r_0 \) determined or recalculated at 90°C and pH\textsuperscript{90°C} 9 for some nuclear, bioactive, natural, and industrial common glasses. It can be seen that \( r_0 \) varies widely depending on the glass composition. At the first order, the Si content plays a major role—glass durability increases with increasing Si content—but the other glass constituents can dramatically and non-linearly affect \( r_0 \). The case of Al is particularly interesting. Depending on the authors, the addition of Al to silicate glasses increases\textsuperscript{175} or decreases\textsuperscript{176} their chemical durability. This apparent contradiction can be explained if one considers the various effects of Al on a glass structure. On the one hand, except in peraluminous glasses, Al decreases the amount of sodium ions bonded to NBOs by increasing sodium ions for charge compensation of [AlO\textsubscript{4}]\textsuperscript{−} tetrahedral units\textsuperscript{177}. It also increases the glass network connectivity and results in a decrease in water diffusivity, and when present in a glass, stabilizes four-fold coordinated Mg and Zn, which then act as glass formers\textsuperscript{173}. All these effects increase glass durability. But on the other hand, first principle calculations have demonstrated that Q\textsubscript{4}\textsuperscript{Al–Q\textsubscript{4}\textsuperscript{Si}} linkages are easier to hydrolyze than Q\textsubscript{4}\textsuperscript{Al–Q\textsubscript{4}\textsuperscript{Al}}. This result can explain why it is that in highly polymerized glasses with the same fraction of NBO, a higher Al content yields higher \( r_0 \). The example of aluminum highlights the complexity of the mechanisms involved, and the difficulty of attributing a single effect of a given element on the durability of silicate glasses.

The first approaches attempting to develop predictive models for \( r_0 \) relied on thermodynamic models\textsuperscript{176,179}. Reasonable trends were obtained between \( r_0 \) and the free energy of hydration of glasses, although the hypothesis that glass can be seen as the sum of pure oxides is questionable given that the mixing enthalpy and the entropy tied to the structural disorder were not considered. Later, statistical approaches\textsuperscript{180} or Monte Carlo simulations with probabilities of dissolution of glass network formers fitted from experimental data\textsuperscript{161,181,182} were developed. However, the incomplete set of mechanisms incorporated in these models have prevented accurate predictions from being made. Recently, advanced techniques have been developed, such as science informed machine learning (ML) which overcome the intrinsic limitations of classical ML approaches (violation of the laws of physics, or unreliable extrapolations)\textsuperscript{183} or quantitative structure properties relationship (QSPR) approaches. These link structural descriptors to various glass properties, including \( r_0 \)\textsuperscript{184–187}. The efficiency of these approaches is partly due to the recent improvements made in modeling glass structures by MD. The development of new interatomic potentials enables simulated structures to be in fair agreement with actual ones, even for borosilicate glasses which were previously poorly modeled because of the various coordinations of B atoms\textsuperscript{186}. With these structural models, it has become possible to determine the number of constraints per atom, \( n_c \), and apply the topological constraints theory (TCT)\textsuperscript{189,190} or determine a mean bond strength, \( F_{\text{net}} \), that can be correlated to \( r_0 \). Suitable linear correlations between \( n_c \) (or \( F_{\text{net}} \)) and log(\( r_0 \)) were obtained with these models, if one considers restricted compositional domains\textsuperscript{191–193}. However, several limitations have been pointed out. These models require calibration, and the dissolution rate of an unknown glass can be calculated only if one knows the dissolution rate of a reference glass or a set of references glasses (training set). With a broad compositional domain, the uncertainties for the predictions remain much larger than for the measured rates. An uncertainty with a factor of \( \sim 7 \) was calculated for a compositional domain with \( r_0 \) spanning 5.5 orders of magnitude\textsuperscript{61}, compared to a relative uncertainty of just 30% for a measured value of \( r_0 \)\textsuperscript{161}. These limitations mean more effort is necessary in order to better refine the structural descriptors. For example, in the TCT approach no difference is made between Si and Al, as both have 4 bond stretching constraints and 5 bond bending constraints per atom, whereas as discussed above, Al impacts glass durability when substituted for Si\textsuperscript{176}. In the \( F_{\text{net}} \) approach, instead of taking the M–O bond strength, first principles calculations could be performed to estimate the energy barrier for hydrolysis of the various M–O–M bonds\textsuperscript{194}.

**STAGE II**

As indicated above, Stage II of glass corrosion begins when the concentrations of low soluble glass network formers (Si, Al, Zr,…) in solution have reached steady states. The corresponding alteration rate is called the residual rate, as the affinity for matrix dissolution is minimum. This residual rate, \( r_r \), is highly dependent on the glass composition—several studies have reported wide variations of \( r_r \) for only slight glass compositional variations\textsuperscript{98,194,195}—but not in the same way as for the initial dissolution rate\textsuperscript{180,196}. In other words, knowledge of \( r_0 \) cannot be used to predict \( r_r \). This is because the rate-controlling mechanisms are not the same.

Various experimental protocols have been used to measure \( r_r \). In general, crushed glass is placed in contact with a given volume of deionized water in a Teflon or stainless steel vessel, and placed in an oven at the target temperature\textsuperscript{180,194,195} and pH\textsuperscript{180,195}. From that point on, two strategies are possible. Either a series of small vessels is prepared and each vessel is used for a particular duration, or a single vessel is used and the solution is sampled at various time durations. Samples are analyzed by classical solution analysis methods, such as ICP-OES or ICP-MS. The residual rate is derived from \( N(t) \) calculated for the tracer elements (Eq. 4). In that case, alkalis or boron are used as glass dissolution tracers, instead of silicon. The criteria taken into account for selecting a tracer include high solubility in the test conditions and a low retention in the alteration layer. In most studies on nuclear glass, B has been considered as an ideal tracer\textsuperscript{198}. Recently, it has been shown that when ISG is corroded in Stage II, some B is retained within an inner layer close to the pristine glass surface, called the reactive zone as it is also richer in hydrogenated species\textsuperscript{58,61}. In general, the...
question of the tracer must always remain central when studying the glass dissolution.

During experiments, the volume of sampled solution can be replaced by fresh water (pseudo-dynamic test) or not (static test). To better monitor the glass response for a given solution composition, experiments can be conducted in dynamic conditions. The use of glass powder ensures a high S/V ratio and thus increases the accuracy of the rate determination from solution analysis. An alternative approach consists in altering the geometry of the reacting glass and opens up the possibility of running depth profiling analysis.

For glasses that can form a passivating gel, $r$, usually starts diminishing proportionally to $t^{-1/2}$ (the concentration of tracer elements released into the solution increases proportionally to $t^{1/2}$). The rest of the curve depends on both the glass composition and the alteration conditions ($pH$, solution composition): it can be continuously decreasing, constant, or fluctuating. There is no general practice for calculating $r$. In some studies, the authors considered the long-term near constant rate when calculating $r$, while in others the authors used the value corresponding to the most advanced stage of reaction in the tested conditions but without checking that the rate was at its minimum. This can make inter-study comparisons difficult. Table 2 summarizes values of $r$, for SON68 glass, the reference non-radioactive glass which simulates the R7T7 glass produced at La Hague. Data came from various experiments conducted at 90°C in deionized water, in Si-rich solution, in synthetic clayey groundwater, or at various pHs imposed by a strong acid or base. All the protocols described above have little impact on the $r$. This reveals that in this regime, glass behavior is not sensitive to S/V. Only the experiment in groundwater had a significant impact on $r$. This can be explained by both a pH effect and the precipitation of Mg-silicate at the expense of the passivating gel. The impact of pH on $r$ is not clear. On the one hand, the pH seemed to have a minor effect on $r$; on the other hand, it significantly affected the residual rate of ISG. Note that in Gin et al. the pH of the pH 8 and pH 7 experiments fluctuated, despite numerous manual corrections. The mechanisms governing the residual rate also depend on T. It has been found that the apparent activation energy measured in the Stage II regime was close to that determined when glass undergoes only dissolution (stage I).

### STAGE III

Stage III of glass corrosion is characterized by a sudden increase of the alteration rate. The period of slow corrosion in Stage II before the alteration takes off is very sensitive to both glass composition and environmental conditions. The resumption can happen after a minute, a few decades, or perhaps never. This issue is particularly serious for nuclear glasses, because it must be demonstrated that Stage III is unlikely if one wants to develop a performance model based on Stage II. As recalled in a recent review paper, Stage III has been widely investigated in the past. It has been established that when alteration resumes, the corresponding rate, $r_{res}$, can be close to but is always less than the initial dissolution rate measured in the same (T, pH) conditions. Interestingly, it has been found that the lower the pH, the lower the ratio $r_{res}/r_{o}$. This means that for a given glass, hyperalkaline media favor Stage III with rates close to $r_{o}$, whereas circum-neutral pH media, such as those of most ground waters, favor alteration rates close to $r_{res}$. In his review, Fournier concluded that the Stage III is mostly associated with the precipitation of zeolites and, to a lesser extent, calcium-silicate-hydrates (CSH), although other causes such as the formation of cracks within the passivating layer or an increase in the reacting surface area have been reported. The zeolites and CSH precipitate at the expense of the passivating gel, and do not form a diffusion barrier for aqueous species, at least in the experimental test conditions. At low temperatures in natural environments, it is not known if zeolitization of basaltic glass is associated with fast alteration rates. Jantzen et al. developed a powerful approach based on observations and thermodynamic considerations to predict if a glass is likely to reach Stage III, depending on its composition. The authors found that the Si/Al ratio of the gel is the key point. If Si/Al <1, glasses tend to remain in Stage II even if clay minerals precipitate, whereas if Si/Al >1, the presence of alkali or OH− in the leachate (strong bases) generate zeolites accompanied by a marked increase of the glass dissolution rate. However, this approach does not tell us when alteration will resume, and at what rate. To date, mostly thermodynamic models have been tested, showing that zeolite precipitation can occur and control glass dissolution depending on the glass and the solution composition.

### Table 2. Summary of $r$, determination for SON68 glass altered at 90°C.

| Onset solution | Mode | S/V (m−1) | pH$^{90°C}$ | Δt or t (days) | Log($r_r$) | Ref. |
|---------------|------|-----------|-------------|----------------|------------|------|
| DIW           | S    | 1200      | 9.7         | 365            | −3.4       | 194  |
| DIW           | S    | 1200      | 9.7         | 1825           | −3.3       |      |
| DIW           | S    | 8000      | 9.2         | 365            | −3.5       |      |
| DIW           | S    | 8000      | 9.1         | 1825           | −4         |      |
| DIW           | S    | 200,000   | 9.2         | 365            | −4         |      |
| DIW           | S    | 200,000   | 9.4         | 1825           | −3.5       |      |
| DIW           | S    | 8000      | 9.3         | 4015           | −3.8       |      |
| C(Si) = 100 mg.L$^{-1}$, pH 9 | S    | 8000      | 9.1         | 365            | −3.7       | 194  |
| C(Si) = 250 mg.L$^{-1}$, pH 9.3 | CF   | 14,000    | 9.3(c)      | 200            | −3.3       | 199  |
| DIW           | S    | 1200      | 8.9(c)      | 500−4500       | −3.9       | 98   |
| DIW + KOH     | S    | 1500      | 7, 8, 9, 10, 10.5 | 594 | [−4; −3.5] | 206  |
| Cox groundwater | S    | 1000      | 6.1         | 273−673        | −2.7       | 92   |

DIW stands for deionized water. Cox stands for Callovo-Oxfordian claystone. S, CF stand for static, or continuous flow, respectively. The pH is given at 90°C (c) = calculated or estimated from data obtained at room temperature and corresponds to a mean value at the time when $r_r$ was calculated. $r_r$ is given in g.m$^{-2}$.d$^{-1}$. Uncertainty on $r_r$ is estimated at 0.2 log units.
HOLISTIC MODELS FOR GLASS ALTERATION

Here we discuss the models able to calculate the glass dissolution rate and the formation of alteration products, compared to time and environmental conditions.

Two kinds of models have been developed: continuous scale models including a single equation or a set of equations, and probabilistic models based on Monte Carlo (MC) algorithms.

CONTINUOUS SCALE MODELS

The most popular continuous scale models are summarized in Table 3. The questions discussed here focus on whether these models can simulate the transition between Stages I and II and the residual rate regime, and if they are able to assess the risk of Stage III. We will not discuss the theoretical bases of these rate laws, but only their application to account for experimental observations.

Some models simulate the decrease in the rate with an affinity term only54,65,220,221, while others couple an affinity term and the transport of aqueous species through the passivating gel and/or the transport of water in the pristine glass91,222–226. The affinity term (1-IAP/IAPsat) describes the degree of saturation of the solution with respect to the dissolving phase in terms of an adequate ion activity product IAP, with the index “sat” denoting saturation. The affinity-based rate laws suppose that either a thermodynamic equilibrium between the outer surface of the reacting material and the solution can stop further glass dissolution56,220,221, or that the thermodynamic equilibrium applies between the buried glass surface and the interfacial solution224. The predictions of these affinity-type models are at odds with observations on borosilicate glass, where alkalis and boron continue to dissolve even when concentrations of low soluble glass network formers (elements forming the outer surface) have reached saturation87. For this type of glass, it is thought that alteration proceeds beyond saturation of the fluid through the dissolution of B, because (1) B–O–Si linkages break faster than Si–O–Si linkages, and (2) dissolution of B is irreversible (B–O–Si bonds cannot reform), therefore B cannot contribute to the polymerization of the passivating layer. But this is not so clear-cut, since a recent study on ISG has shown that the activation energies for the hydrolysis of the two kinds of bonds are very close and that a high concentration of B and Ca in solution seems to limit further B release61. Therefore, at least for this type of glass, the dissolution of B and the transport of B(OH)3 and B(OH)4− through the gel could be rate-limiting in Stage II. For basaltic glass, the rate law proposed by Daux et al. predicts that the rate is zero when a solution is saturated with the leached outer layer made up of Si, Al, and Fe221. Techer et al. proposed an affinity term only based on Si, applied at the pristine glass surface, and added a diffusion term of Si through the gel layer222. The model predicts that the rate is zero when the concentration of Si at the glass surface reaches saturation. However, when there was a rapid reaction progress (long duration, high S/V, static conditions), an extremely slow but not zero residual rate was measured197. Therefore, the two models developed for basaltic glass (Table 3) cannot account for this low rres.

For B-free glasses with no or a low amount of non-bridging oxygens (NBO), such as obsidian, albite, or nepheline glass, the existence of a residual rate has not been demonstrated. At the first order, it depends on whether the glass structure allows water molecules to diffuse in the pristine material. If water can diffuse in the solid, alteration could proceed through ion exchange, hydrolysis reactions, and local rearrangements16,197,222. This can be considered as a residual alteration even if no glass components are released into the solution. Another possible cause of a residual alteration could be the difference in chemical potential between the altered glass surface in equilibrium with the bulk solution and the pristine glass surface. A significant difference would account for the maintenance of a residual transformation of the pristine glass into alteration products, with a potential supplemental effect if alteration products are transport-limiting for water molecules. A recent MD study to compare the behavior of vitreous albite (NaAlSi3O8) with its crystalline polymorph demonstrated that water can diffuse in the glassy structure but not in the crystal87. This structural difference was assumed to impact the formation mechanism of the passivating layer. It remains to be shown whether the glassy form exhibits a non-zero residual rate.

Direct evidence using isotopic (29Si, 18O) markers has been obtained showing that a dense gel layer can be transport-limiting for water molecules and some dissolved glass species30,61. The kinetic models coupling affinity and diffusive transport are therefore a priori better able to predict the residual rate of these glasses. However, it remains unclear which chemical species are involved in the diffusion-limiting mechanism. It is likely that several of them can play a role, depending on the gel characteristics (porosity, surface charge) and the environmental conditions (T, pH…). Progress here could come from the identification of the chemical species diffusing in the gel and the direct measurement of their transport properties.

### Table 3. Selection of models developed to account for the drop in the alteration rate of silicate glass.

| Reference | Glass | Affinity term | Thermodynamic equilibrium | Elements taken into account | Transport term |
|-----------|-------|---------------|---------------------------|----------------------------|----------------|
| 222 Basaltic | Glass surface/interfacial solution | Si | Siaq | SiO2 |
| 221 Basaltic | Leached glass/bulk solution | Si, Al, Fe | No | No |
| 54 Nuclear glass | Glass surface/bulk solution | Si | SiO2 | SiO2 |
| 224 Nuclear glass | Glass surface/interfacial solution | Si | SiO2 | SiO2 |
| 91 Nuclear glass | Gel/bulk solution | Si | All dissolved glass cations(1) | H2O |
| 223 Nuclear glass | Gel/bulk solution | Si, Al | All dissolved glass cation(1) | H2O |
| 220 Nuclear glass | Gel/bulk solution | All gel-forming elements | No | No |
| 225 Silicate glass | Gel/bulk solution | Si | Baq, Naaq, H2O | H2O |
| 65 Silicate glass | Si gel/bulk solution | Si(2) | No | No |

The table lists the main mechanisms included in the models described in the literature. The affinity term accounts for the dissolution of the glass or the gel, which can be considered as a diffusion barrier or not. The transport term—generally a Fick’s equation with a constant diffusion coefficient—can account for the gel formation or the passivation effect of the gel. (1) In the GRAAL91,223, a single apparent diffusion coefficient is considered for all the diffusing species, including water in the glass. (2) Oelkers’ model considers several chemical reactions taking place in series56. The last reaction (the most likely for modeling the residual rate) corresponds to the dissolution of the remnant silicate network. Bourcier’s model considers the formation of a diffusion layer formed by ion-exchange but the key mechanisms to account for the long-term dissolution rate is the equilibrium between gel and solution220.

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of the diffusion coefficients of these species. Another important aspect concerns the formation dynamics of the passivating layer. Recent experimental studies have shown that the passivating gel undergoes local reorganization, leading to pore ripening, and a decrease in the reactivity with water\textsuperscript{30,228}. Except for the model developed by Ma et al.\textsuperscript{225}, none of the models listed above take into account the effect of maturation.

Overall, when considering the glass/gel/solution system, there is still some progress to be made in identifying the rate-limiting mechanisms and their inclusion in a reactive transport model in order to be able to make accurate predictions. This will benefit from the progress made in the development of advanced analytical techniques\textsuperscript{17,209,229}, along with simulation techniques at molecular and mesoscale levels\textsuperscript{230}.

The precipitation of secondary phases is a key phenomenon for Stage II. In most of the experimental studies, secondary phases form at the expense of the passivating gel (Fig. 7). When secondary phases precipitate, they change the solution chemistry, which in turn can change the rate of formation or dissolution of the passivating gel. From a modeling standpoint, the data required to take secondary phases into account include, in order of priority, the solubility product, and the kinetics of precipitation. The rate laws listed above are implemented in reactive transport codes, which use thermodynamic databases with most of the secondary phases which can form.

A kinetic model was used by Fournier et al. to account for Stage III glass corrosion. The model was applied to ISG altered in NaOH solutions\textsuperscript{231}. The authors used the GRAAL model associated with additional equations for nucleation and the growth of zeolites. Despite the use of fitting parameters, the modeling exercise was helpful and showed that, subject to knowing the composition and the solubility of the gel as a function of pH, it is possible to predict when and at what rate the alteration will resume. It is worth noting that, at laboratory scale, the nature and both the solubility and passivating properties of gels able to form in various environments need to be understood and calculated before a mechanistic model can predict the behavior of glass in both Stages II and III. At a longer timescale, the modeling of nuclear glasses in disposal environments requires an integrated approach and the use of a model relying on the relevant mechanisms. More complicated, slow processes that could impact the durability of glass packages, such as the sealing of cracks by secondary phases formed in Stages II or III, should be considered, especially in a best estimate approach. An illuminating example was given by the alteration of a fractured archeological glass block from the Roman period, altered for 1800 years in seawater. It was demonstrated that the precipitation of clay minerals and calcite within the cracks dramatically diminished the contribution of the internal surfaces to the global alteration of the block. After a few centuries, their contribution became negligible even though their surface was initially 100 times greater than the outer surface\textsuperscript{232}. A simple model based on a constant contribution of all the surfaces would therefore be extremely over-conservative.

**PROBABILISTIC MODELS**

Probabilistic models based on the MC algorithm constitute another kind of kinetic models. They link the elementary mechanisms at the atomistic scale to the dissolution rates at the macroscopic level. This approach has been widely applied to both minerals\textsuperscript{233–235} and glasses\textsuperscript{13,182,236,237}. The parameters are generally fitted from ab initio calculations\textsuperscript{233} or empirically, to reproduce certain experimental behaviors\textsuperscript{238}. Despite some limitations discussed below, it is worth emphasizing the use of these models, as interesting results have already been obtained and the technique is still in development. Let us discuss a few examples.

To simulate glasses within the compositional ranges (61 – \(x\))SiO\(_2\)-17B\(_2\)O\(_3\)-18Na\(_2\)O-4CaO-xZrO\(_2\)\textsuperscript{236} or (70 – 2\(x\))SiO\(_2\)-xAl\(_2\)O\(_3\)-15B\(_2\)O\(_3\)-(15 + x)Na\(_2\)O\textsuperscript{237} glass cations were placed at the nodes of an ordered cubic network (each atom is surrounded by six neighbors). Then some bonds were broken, depending on the coordination number of the cations in the glass: zero for \(\text{VIZr}\), two for \(\text{IVSi}\) or \(\text{IIIAl}\), and three for \(\text{IBB}\). O atoms were not explicitly displayed, and Na and Ca atoms were not bound to the network. A 2D or 3D glass–water interface with thousands to millions of atoms and water molecules was prepared, and probabilities were implemented for bond breaking (Si–Si, Si–Al, Al–Al) and dissolved cation (Si, Al) redeposition on the glass surface. Note that a single set of probabilities was assigned for a given series of glasses. A simulation consists in accumulating thousands to millions of events (dissolution–redeposition) to monitor the glass behaviors.

Step by step, the soluble elements in contact with water were dissolved in solution, leading to the formation of a gel-like altered layer composed of low or non-soluble elements (Si, Al, and Zr in this example). The alteration rate decreased with time because redeposition prevented the accessibility of water to the soluble elements, but once the Si and Al saturation was reached, the rate dropped sharply to zero, as no diffusion mechanisms were implemented, except for one attempt to introduce ion diffusion in the liquid phase\textsuperscript{239}. Owing to the limitations preventing the calculation of \(r_g\), simulations have mostly focused on the effects of soluble or hardener elements on \(r_g\), the transition between Stages I and II, and the characteristics of the alteration layer (thickness, interfacial roughness, pore volume, pore size). For instance, it has been seen that \(r_g\) decreases with increasing Zr content in the glass, because Zr cannot dissolve. However, at high reaction progress rates, high Zr content glasses are much less durable than low Zr content glasses, as Zr prevents gel maturation and pore closing\textsuperscript{236,238,240}. This nonlinear effect of Zr has been confirmed experimentally by porosity investigation using small-angle X-ray and neutron scattering (SAXS, SANS) and the diffusion of dyes monitored by time-of-flight secondary ion mass spectrometry (ToF-SIMS) techniques\textsuperscript{25,240} (Fig. 6).

The impact of NBO, the Si/B ratio, boroxol rings, or Al\(_2\)O\(_3\) content has been investigated in alumino-borosilicate glasses\textsuperscript{181,237}. The quantity of NBO is a key factor that increases the glass dissolution rate. The impact of NBO concentration on glass dissolution rate is nonlinear: at low concentration, the glass dissolution rate is proportional to Q\(_3\), whereas at high concentration, glass clusters surrounded by Q\(_3\) and Q\(_4\) species are released to solution without being dissolved. Furthermore, when the B atoms are clustered within boroxol rings, the glass dissolution rate increases. A clear synergy between the Al and B has been shown. The B release changes in a nonlinear manner when the Al\(_2\)O\(_3\) content increases, with a maximum B release observed for an intermediate Al\(_2\)O\(_3\) content\textsuperscript{237}. This behavior relies on the dynamics of gel reorganization.

MC modeling has also been used to investigate radiation-induced structural damage in glass and its consequences on chemical durability\textsuperscript{173}. Modifications of the local structural characteristics induced by ballistic effects (depolymerization, change in the local angles) were not sufficient to account for the experimental observations on \(r_g\). For that, it is necessary to consider other mechanisms not implemented in the current Monte Carlo approach, such as the acceleration of water diffusion in the irradiated structure.

Despite the interesting results summarized above, the simulation of the glass structure by an ordered lattice constitutes only a rough approximation. The consequences of this oversimplification have been evaluated by comparing the \(r_g\) of an ordered structure to that of a disordered structure prepared by classical MD\textsuperscript{241}. It appears important to represent the surface roughness accurately, because of the strong correlation between this parameter and \(r_g\) (the number of sites accessible to water increases with increasing...
14

the frontier between a passivating gel and secondary phases is
corrosion result from these competitions. It is worth noting that
at the expense of the passivating layer. Stages II and III of glass
layer and have poor diffusion properties. Therefore, they can form
thermodynamically more stable than the amorphous passivation
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exists between the ICDP model and the leaching/in situ
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For applications requiring very resistant glasses, secondary
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and c, gels form by in situ reorganization of the silicate network
after preferential dissolution of alkalis and boron, whereas in Fig.
7b, a series of amorphous layers forms by congruent dissolution
of the glass and reprecipitation. Note that, those solid phases can act
as a sink for hazardous elements contained in the glass. This
can dramatically reduce the source term of vitrified wastes. Furthermore, in some cases, precipitation of secondary
phases can reduce the renewal of fresh water on the glass surface,
resulting in increased durability of the glass.

For some applications, the formation of an alteration layer on
glass, whatever its composition, is detrimental. For instance, this is
the case for (1) tableware, which can lose its transparency after a
few cleaning cycles in a dishwasher; (2) commercial window
glass during the storage period, which can undergo atmospheric
alteration; (3) stained-glass windows, which can lose shine and
luminosity after decades or centuries of exposure to acid rain and
pollutant deposits; and (4) ampoules, vials, or cartridges for
pharmaceutical use, as alteration products can delaminate and
contaminate the active products. For all these situations, a
deeper understanding of the mechanisms involved can help to
find solutions to delay or limit the detrimental effects of aging.
As seen above the case of bioactive glass is the opposite. The
formation of secondary phases is wanted, and must occur quickly,
as that is the way bioactive glasses alter to form HCA to repair
damaged bones in the human body. Unmet challenges always
require more research. In particular, the role of cations released
by glass on cell stimulation has to be better understood.

CONCLUDING REMARKS

Significant progress has been made over recent years in
deciphering glass behaviors in natural, industrial, or living
environments. Progress has come from both understanding the
fundamental processes at molecular level and integrating that
knowledge in models. Zooming in at the molecular level reveals
extremely complex and dynamic processes. The challenge now is
to identify the relevant mechanisms to be implemented in
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must always be kept in mind. Experiments in laboratories as well
as observations of natural systems have demonstrated that silicate
glasses can self-passivate by forming amorphous gels on their
surfaces, but the fate of the protected glass strongly depends on
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FUTURE CHALLENGES AND PERSPECTIVES

The previous sections have shown that there are still some
unresolved issues regarding the mechanisms controlling the three
stages of glass corrosion. These fundamental limitations prevent
the development of predictive models without the use of
empirical convoluted parameters. Having predictive models
available, with calculated or measured fundamental parameters


Fig. 6  Role of Zr on the alteration of borosilicate glasses. Glasses ((61−x)SiO2−xZrO2−17B2O3−18Na2O−4CaO) were studied both by a MC
model and experimentally. a On the left, cross section of altered glass specimen in Stage I and corresponding values of T0 measured at 90 °C,
neutral pH. On the right, cross section of the same glasses at a faster reaction progress. This example highlights the nonlinear effect of Zr,
which can be explained when considering its role in Si dissolution and gel maturation. b TEM image of the gel formed on the Zr-free glass of
the series (x = 0) and the corresponding simulated gel obtained by MC. Both display the same features with the closing of the porosity near
the outer surface. c Comparison of the gel morphologies based on the exponent of the SAXS power law as a function of the ZrO2 content in
glass composition after 106 computation steps for the simulated glasses and after 39 days of corrosion for the real glasses. Data adapted from
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disorder in a glass structure yields isotropic behaviors, mainly in the way of forming alteration products. In that sense, there is still some work to be done to develop a universal theory for silicates. Both experimental and digital tools have made progress, and it is now possible to simulate the dissolution rate of a silicate glass with only parameters calculated at the atomic level. Such a result would open the gate to fully predictive models able to design materials with tailored properties, or to calculate their durability in conditions where experiments are currently impossible or costly.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon request.

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S.G. supervised the work and wrote most of the paper. J.M.D. wrote the section on molecular mechanisms of glass alteration. F.A. and S.C. contributed to the discussion and deeply reviewed the manuscript.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Correspondence and requests for materials should be addressed to S.G.

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