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ABSTRACT: Alkali-activated materials (AAMs) have high potential as an alternative binder to ordinary Portland cement (OPC), because of their high performance and lower CO2 emissions. While there is a general consensus about their strength and durability advantages over OPC, there is a widespread debate regarding their durability. Some groups believe that the availability of wide scientific/technical background, together with the already-known OPC durability problems, are sufficient for their commercialization; others consider the durability of AAMs to be an unresolved issue. This controversy represents one of the limitations facing their bulk applications. The present work provides an overview of the latest developments on the durability of fly ash/slag-based AAMs with the aim to update recent findings regarding their behavior under aggressive conditions (sulfates, freeze–thaw, chloride, carbonation, acid, efflorescence). This review will provide a better understanding of the durability issues of AAMs, which will stimulate further research to develop the appropriate testing methods and help to promote their commercialization.

1. INTRODUCTION

Cement and concrete are central to modern civilization, with its reliance on the built environment to provide a high quality of life. Next to water, concrete is the most consumed substance on the planet and is produced in volumes exceeding 10 billion tons per year worldwide.1−3 Ordinary Portland cement (OPC)-based concrete is still the material of choice and the most widely used in construction and civil engineering, because of the ease of use and generally good performance, including excellent thermal and mechanical properties, besides offering an average life cycle of more than 100 years.

However, with the current focus on environmental issues, the cement industry causes serious destruction of natural quarries through the extraction of raw materials. It also contributes ~8% of global CO2 emissions,4−5 and consumes more than 10% of the total energy used.5,7 The total emissions footprint of ~0.8−0.9 t CO2 per ton of OPC produced are mainly due to the decomposition of limestone (0.53−0.55 t) and the combustion of fossil fuels (0.34−0.39 t) during cement production.3,4

After the 2009 Copenhagen climate summit, where the international community agreed to keep global average temperature increase below 2 °C, a more significant reduction in CO2 emissions (20%−50%) was strictly adopted by several countries/continents to guarantee a “normal” human habitability. In this context, the European Union has made an unilateral commitment for 2020 to reduce overall greenhouse gas emissions from its 28 member states by 20%, compared to 1990 levels. In the same way, China, as the world’s largest emitter of CO2, announced that, by 2020, its carbon dioxide emissions per unit of gross domestic product (GDP) will be reduced by 40%−45% from the 2005 level.11,12 Consequently, the cement industry is under pressure to reduce both energy use and greenhouse gas emissions and is actively seeking alternatives to these serious concerns. The interest in reuse of other industrial byproducts such as fly ash (FA) from coal combustion, blast furnace slag (BFS) from iron production, silica fume (SF) from ferrosilicon production, and rice husk ash (RHA) from burning agricultural rice waste as supplementary cementitious materials (SCMs) has been greatly increased and new eco-efficient binders such as calcium aluminate cements,13−16 calcium sulfoaluminate cements,17−21 super-sulfated cements,22−24 and alkali-activated binders25−41 have been developed. Among them, alkali-activated materials (AAMs), and those classified as geopolymers,42,43 are a high-profile example able to produce cement and concrete with the advantages of OPC but with a large reduction in CO2 emissions and general aspects of good performances (high early strength, good resistance against acid and sulfate attack, etc.). AAMs and GPC have emerged as novel engineering materials with the potential to form a substantial element of an environmentally sustainable construction and building products industry.44−47 AAMs are basically formed by aluminosilicate precursors, which are generally, but not limited to, industrial byproducts such as FA and BFS, aggregates, and activator.

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Figure 1. Fly ash slag specimens aspect after immersion in Na₂SO₄ (left) and MgSO₄ (right) solution for 90 days: (A) w/b 0.40, (B) w/b 0.50, and (C) w/b 0.60. (Reproduced from ref 63. Copyright 2013, Springer, Dordrecht, The Netherlands.)

be drawn between “magnesium sulfate attack” (where both Mg²⁺ and SO₄²⁻ are capable of inducing damage in the structure), and general processes related to the presence of sulfate, accompanied by other, nondamaging cations. Komišenov et al. 66 studied the effects of external sulfate attack (5% Na₂SO₄ solution) on mechanical and micro-structural properties of alkali-activated slag (AAS), compared with portland-slag cement (CEM II/A-S 42.5N). They concluded that, after 90 days, AAS showed significantly higher resistance to sulfate attack, with respect to the benchmark CEM II, and attributed these findings to the absence of portlandite and the unavailability of aluminum, substituted in C-S-H (I) or present in hydrotalcite, for reaction with sulfates. These results are in agreement with those reported previously, 65 66 where a comparative study between AAS and OPC concrete exposed to 5% magnesium sulfate and 5% sodium sulfate solutions during one year showed that the decrease in compressive strength was ~17%–23% for AAS and ~25%–37% for OPC-based concrete. Bakharev 66 investigated the influence of activator (type, concentration, and the type of cation in the sulfate media) on the stability of alkali activated fly-ash-based geopolymers. Two accelerating methods were applied: (i) increased reaction surface (small specimens/large surface areas) and (ii) increased concentrations of aggressive agents. In addition, each of solutions was applied into three different alkali-activation-based specimens and cured at 95 °C for 24 h. The first group was activated with NaOH, the second with Na₂SiO₃, and the third one with a mixture of NaOH and KOH. The best sulfate resistance in different sulfate solutions was observed in NaOH-based geopolymers, because of the stable cross-linked aluminosilicate polymer structure. Moreover, good resistance was linked to a refined pore structure as the average pore size of the tested specimens prepared with NaOH activator was minor. The compressive strength of the slag.
samples was increased by 4%–12% over the exposure time. In
the case of specimens prepared using sodium silicate, the author
attributed the loss of strength to ettringite formation (observed
in the XRD patterns); however, we believe that the formation
probability of this phase in low-calcium systems such as FA-
based concrete is very low, making XRD results of this work
insufficient to confirm the presence of ettringite, taking into
account that, in sodium silicate FA samples, no deep cracks
have been detected in neither needle-shaped particles
(ettringite) were observed from SEM micrographs. The least
changes in specimens were found in the solution of 5% sodium
sulfate + 5% magnesium sulfate. The most significant
microstructural changes and strength fluctuations were
observed when 5% solutions of sodium sulfate and magnesium
sulfate are used. Diffusion of alkali ions into the solution caused
significant stresses and the formation of deep vertical cracks in
the specimens prepared using a mixture of NaOH and KOH.
Sulfate resistance of fly-ash-based geopolymer concrete from
blended ash of pulverized fuel ash and palm oil fuel ash has
been investigated in under 5% sodium sulfate exposure for a
period of 18 months and compared to that of OPC concrete
under the same exposure conditions. Changes in mass,
compressive strength, products of degradation, and micro-
structure have been evaluated in both systems. A better
resistance to sulfate attack in FA concrete, compared to that in
OPC, has been reported, which was attributed to low calcium
phases, low water demand, and compact microstructure in FA-
based concrete. Similar results have been reported in previous
work conducted on heat-cured fly-ash-based geopolymer
concrete after exposure to sodium sulfate solution for up to
one year. However, none of these works explicitly
reported the important role of stable zeolites that are generally
formed in geopolymer systems, which can encapsulate sulfates
by preventing leaching. The formation of gypsum and then
ettringite in OPC concrete caused a reduction in compressive
strength, physical damage at the corners and edges of
specimens, and mass loss as the final deterioration indicator.
No significant changes in appearance have been detected after
18 months of immersion and only 4% of mass loss has been
found in geopolymer concrete. However, in OPC concrete,
more than 20% mass loss was reported, again indicating better
resistance of FA-based geopolymer systems. It has been also
reported that compressive strength of geopolymer concrete
was increased by 7% and that of OPC decreased by 63% after
more than one year of exposure to 5% sodium sulfate solution.
The increase in compressive strength of geopolymer systems
also reported by Ismail et al. and Bascarevic et al. was
attributed to the fact that sodium sulfate is acting as an activator
rather than an aggressive agent, allowing the alkali activation
reaction to continue in the sulfate solution. However, from the
later work conducted in two different FA samples over a period
of 365 days, a reduction in compressive strength of some
mixtures after sulfate treatment was reported. The same
authors also reported that more-porous fly-ash-based geo-
polymer mortars demonstrated an increase in compressive
strength. This was in overall disagreement with the current
observation that a compact and denser microstructure displays
better mechanical, chemical, and physical properties. The
phenomenon was attributed to the prolonged geopolymerization
in sulfate solution favored by the presence of high amount
of alkalis. The authors emphasized that leaching is apparently
less affected by sulfate ions than by the high pH of the sulfate
solution. The initial pH value of Na2SO4 solution was 6, while
the final was reported to be ∼12. Based on the theory, the main
degradation mechanism of aluminosilicate compounds at high
pH is the hydrolysis of the siloxane bonds induced by OH−
attack, which later causes desilication of the geopolymer matrix.
Consequently, Si4+ ions were leached, which was confirmed by
29Si MAS NMR analysis. The decreased Si/Al ratio was one
important indicator of matrix degradation due to leaching of
SiO4. The authors attributed the higher observed leaching to the
higher pH of the sulfate solution.

From these results, it can be concluded that the relationship
between strength and sulfate treatment is still unclear and
requires deeper investigation in order to identify whether sulfates improve or reduce the mechanical properties of these
binders. We believe that the curing conditions of reference
samples (laboratory conditions, fog room, immersed in tap
water, etc.) may have significant impact on their mechanical
properties and could explain the divergence trends reported on
compressive strength. In the case that sulfates positively affect
the durability of AAMs (by improving strength), should sodium
sulfate be considered as an aggressive agent in future
formulations of appropriate standards for durability testing of
these systems?

Through the review of different studies conducted in sulfate
resistance of slag and/or fly-ash-based AAM, it can be concluded that this phenomenon is mainly affected by the type of cation accompanying the sulfate, although durability can also be influenced by other factors, such as precursors chemical composition, calcium/magnesium content, the exposure conditions, and even sample preparation and curing con-
ditions. Glukhovsky and Krivenko investigated the influence of
different alkaline activator solutions on freeze–thaw resistance.
It has been reported that sodium silicate-activated slag concrete
exhibited the greatest resistance, because of its less porous
structure. AAS concretes can bear 300–1300 cycles of repeated
freezing–thawing cycles, compared to the <300 cycles in the
case of OPC concrete. The influence of slag basicity on the
frost resistance has been also investigated, and it has been
found that the worst performance corresponds to acidic slag.

The freeze–thaw resistance of AAS-based concrete has been
investigated by Fu et al., where an accelerated freeze–thaw
cycling method was applied. The dynamic elasticity modulus
was measured every 25 cycles, up to 300 cycles. The authors
established two freeze–thaw cycle damage models, based on
experimental results: dynamic elasticity modulus attenuation
model and an accumulative freeze–thaw damage model. Each
one was designed with respect to damage variables. Two
functions were used to describe these damage models: power
function and exponential function. Functional equations were
established based on a dynamic elasticity modulus attenuation
model in the literature, and power function models were

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The interpreted results showed significant freeze-thaw resistance of AAS concrete (frost-curing grade above F300 and coefficient freeze-thaw resistance of ~90%), making it a suitable candidate to be used in cold areas. Properties such as low Ca/Si ratio, short original three-dimensional (3D) structure, and better cushion power than that of C-S-H in OPC concrete are observed, which benefits the corrosion resistance of materials. The close-grained hydration products make the structure of AAS concrete more compact, which makes it difficult for water to penetrate and difficult for concrete to become frozen and saturated, which makes AAS concrete more durable.

Cai et al. used response surface methodology (RSM) to analyze the effects and interaction among three relevant parameters on the freeze-thaw resistance of AAS concrete: solution/slag ratio, slag content, and sand content. The most important factor influencing the freeze-thaw resistance of AAS concrete is the air-void structure. The freeze-thaw resistance has a tendency to improve with a smaller air bubble space coefficient and a bigger specific surface area. It has been also found that the lower solution/slag ratios and higher slag content in the mixture design exhibit the best freeze-thaw resistance, while the sand content was found to have a minor effect on the durability.

Škára et al. investigated the resistance of fly-ash-based fly-ash-based geopolymers in geopolymer specimens to alternating freezing and defrosting cycles, according to Czech Standard ČSN 72 2452. Samples with dimensions of 40 mm × 40 mm × 160 mm were cured under laboratory conditions at ambient temperature and 40% relative humidity (RH) for 28 days before freeze-thaw testing. Seven mixture designs were made, adding various materials, such as OPC, limestone (in ground or aggregate form), and three different air entraining agents. Geopolymer samples were kept under water for 150 freezing and defrosting cycles. Samples were broken after 28 days, 6 months, and 1 year, respectively. No bulk disintegration was observed, and no mass loss of samples was observed. The compressive strength values of samples decreased slightly after the defrosting cycles, compared to the values for the samples without exposure to frost tests. Still, no physical deterioration by means of damages or deformations was noticed, indicating that fly-ash-based geopolymers were considerably resistant to frost exposure. Similar conclusions have been reported for fly ash mortars, in comparison to OPC-based specimens.

Shu reported that AAS concrete showed excellent freeze-thaw resistance, thus fitting their application in tunnel lining in extremely cold regions. It was also shown that, using a blend of fly ash and slag in concrete synthesis, the relative dynamic modulus was ~90% after 300 freeze-thaw cycles with a slight mass loss and physical deterioration due to thin surface layers that favorably restrain freeze-thaw damage of concrete. Bilek et al. have reported worse frost resistance in AAMs than in OPC, which was attributed to the presence of a higher amount of free water available for freezing in the structure of alkali-activated concrete. Few results on the effect of FA/BFS on scaling durability of concrete have been reported and discussed by Pigeon et al.; however, these results were related to OPC-based concrete in which a small amount of FA/BFS (up to 20%) has been used as supplementary cementing materials.

From the reported study, no clear conclusions were drawn when OPC was partially replaced by BFS.

Based on the given literature review, evaluation of a suitable method for testing in a laboratory environment, replicating in-service conditions, remains to be performed. Determination of parameters related to mix design and sample conditioning is needed in order to define general recommendations for the synthesis of geopolymers resistant to freeze-thaw in cold climate zones.

Another interesting point that was not deeply investigated through the literature and therefore requires special attention in future works is the relationship between pore structure and frost resistance in AAMs. The nature and particle size of the precursor used can strongly affect the pore structure, and consequently, the frost resistance of concrete.

2.3. Chloride Resistance. The resistance to chloride attack has been recognized as one of the most concerned aspects of concrete durability. Chloride attack could destroy the passivated layers on the surface of reinforced steel bars, initiate the electrochemical corrosion of steel reinforcement, and consequently reduce the structural capability of concrete elements. Chlorides do not usually harm the concrete matrix as much as its induced corrosion of reinforced steel bars, because the degree of structural change caused by chloride is generally low in Portland cement binders and even lower in AAMs. Friedel’s salt is formed through the conversion of layered calcium sulfoaluminate hydrates into chloroaluminate in OPC mortar after the ponding test of chloride, but the chloride in alkali-activated binders does not involve the formation of crystal phases of new chloroaluminate. The resistance to the transport of chloride ions to the surface of reinforced elements comes to be the first priority in the study of chloride attack in AAMs.

The rapid chloride permeability test (RCPT), conforming to ASTM Standard C1202, is widely accepted in practice. The RCPT test, in fact, measures the conductivity of the material, rather than the actual permeability. The effects of conductivity of pore solution, dependency on all the mobile ions rather than the ions of interest, and joule heating on the experimental results make RCPT inadequate, especially when comparing across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries. Moreover, the high voltage (typically 60 V) applied across different types of concretes with different pore solution chemistries.
To date, there are a limited number of studies on the chloride ingress of AAMs, compared to OPC-based ones, and most of these studies were based on experimental investigation either in the laboratory or in the field. Miranda et al.88 reported that the chloride-free activated fly ash mortars were also able to passivate steel reinforcement as quickly and effectively as OPC mortars, but an addition of 2% (by binder weight) of chloride during the mixing operation would multiply the corrosion rate by a factor of ~100. According to Kannapiiran et al.,89 fly-ash-based reinforced geopolymer concrete beams with heat curing for 24 h revealed no visible traces of corrosion after 180 days of immersion in 5% HCl solution, and the penetration depth of chlorides into concrete was <4 mm, which was much lower than the cover to steel. Kupwade-Patil et al.90 studied the durability of steel-reinforced concrete made from alkali-activated fly ash and OPC in a cyclic wet–dry chloride environment over a period of 12 months. The authors found that geopolymer concrete specimens where only microcracks indications of corrosion products were observed via SEM at the matrix/rebar interface exhibited lower average diffusion coefficients, lower chloride contents, and lower porosity, compared with their OPC counterparts where multiple gross corrosion products were observed. Zhu et al.91 found that (i) the unsaturated alkali-activated fly ash paste and mortar may have higher chloride penetration rate than portland cement paste and mortar, and (ii) only the mortars with a liquid/solid ratio of 0.6 and with 40% slag substitution exhibit a similar chloride penetration rate as the portland cement mortar at w/c = 0.5. The RCPT and NSSM tests have been used to evaluate the chloride transport behavior of alkali-silicate-powder-activated slag concrete.85 The results showed that the RCPT values of the AAS concrete were generally lower than those of OPC concrete and water-glass-activated slag concretes, while the NSSM coefficients were similar to those of OPC and water-glass-activated slag concretes. According to Ismail et al.,84 the chloride migration testing by the NordTest NT Build 492 standard accelerated method and by chloride ponding showed that the diffusion of chloride in AAS-based binders is much lower than that observed in OPC ones (Figure 2), but the volume of permeable voids and sorptivity testing showed that AAS mortar and concrete exhibit higher water absorption than OPC-based samples. The authors attributed this divergence to the specified standard preconditioning (drying) of the samples prior to water absorption testing as drying is likely to induce desiccation and consequent microcracking of slag-based binding gels.

The chemistry of reaction gels and pore structure significantly controls the transport properties of chloride ions in the AAMs in the chloride-containing environment. Bernal et al.81 reviewed the influences of the nature of the binding gel on the chloride permeability in alkali-activated slag/fly ash systems; they concluded that highly durable binders can be designed through the analysis and tailoring of binder gel chemistry. Ismail et al.84 mentioned that a denser Al-substituted calcium silicate hydrate (C-A-S-H) gel in AAS concrete contributed to a higher durability under chloride exposure, while the inclusion of fly ash (BFS:FA = 100:0, 75:25, 50:50) promoted the formation of more porous sodium aluminosilicate (N-A-S-H) type gels, reducing the resistance to chloride ingress.91 Yang et al.87 reported that the incorporation of slag as a secondary precursor in fly-ash-based geopolymers contributed to the refinement of pore structure and thus restricted the transportation of chloride ions in the paste. Zhu et al.92 showed that lower porosity and higher tortuosity are both helpful toward decreasing the chloride penetration rate in alkali-activated fly-ash concrete. Ravikumar et al.93 investigated the effect of sodium silicate modulus on the chloride penetration of AAS. It has been reported that increasing the Na2O/slag ratio reduced RCPT and NSSM coefficients of solid sodium silicate-activated slag concretes, and increasing the SiO2/Na2O ratio of the activator benefically influenced the transport parameters of liquid sodium silicate-activated slag concretes. The authors mentioned that both the 28-day and 56-day RCPT and NSSM values of AAS concrete decrease as the silica content increases, with a limiting value beyond which the chloride transport parameters are insensitive to the content of silica. They also reported that the powder alkali silicate-activated slag mixtures consistently show better resistance to chloride ion transport than the water-glass-activated mixtures, which can be attributed to a more-refined pore structure indicated by a smaller critical pore size. The same authors reported in one of their recent works83 that the critical pore size is more influential than porosity in determining the chloride transport properties of AAS concrete.

Activator dosage in terms of SiO2 or Na2O (by binder weight) can significantly affect the resistance to chloride penetration. Chindaprasit et al.35 reported that the chloride penetration and corrosion of embedded steel in the high-

**Figure 2.** Boundary of chloride penetration in concretes cured for 28 days at the end of the NordTest procedure, as a function of the slag/fly ash ratio: (A) 100 wt % slag, (B) 75 wt % slag/25 wt % fly ash, (C) 50 wt % slag/50 wt % fly ash, and (D) OPC. (Reproduced with permission from ref 84. Copyright 2013, Elsevier BV, Amsterdam.)
calcium fly-ash-based GPC decreased as the NaOH concentration increased, after 3 years of exposure under the marine site, which was attributed to the refinement of the pore structures, as a result of better polycondensation reaction in the relatively high concentration of NaOH. Ma et al. showed that increasing both the silica and alkali content of the activator results in a lower total porosity and a finer pore system, hence reducing the permeability of alkali-activated fly-ash paste. Law et al. assessed the durability of AAS concrete with a range of sodium oxide dosages and activator modulus ($M_a$) values. They found that between $M_a = 1$ and $M_a = 1.25$, the increased reaction of the slag grains is offset by an excess of activator, resulting in fewer, but larger, microcracks. Through the investigation of pore solution and alkali diffusion in fly ash (class F, class C) and slag-based AAM, Lloyd et al. reported that these systems are prone to alkali leaching, because the alkalics are highly mobile in the pore system, making their retention difficult. The authors suggested the use of a Ca-rich precursor as an effective way for reducing alkali mobility and consequently to keep the pore solution alkalinity at an adequate level, which may avoid steel corrosion and improve the resistance to chloride attack.

Thus far, there have been some aforementioned studies on the chloride permeability of AAMs from the point of view of either scientific or technical research, but it is far enough as many challenges are still confronted ahead. The factors affecting the binding capacity of chloride in portland cement-based materials have been studied extensively, but they are not yet studied comprehensively in AAMs. Many researchers have investigated the chloride permeability of AAMs in comparison with portland cement-based materials, but there is no explicit and uniform criterion for the selection of its counterpart to compare. Many experimental studies have been conducted on the chloride permeability, but the database of chloride migration coefficients is not sufficient for AAMs. Consequently, another challenge is presented, in that modeling chloride transport in AAMs is scarce, because of insufficient information describing chloride penetration with time. There are some proposed hypotheses in the research of chloride ingress, but they are not yet studied in detail (for example, the sorption of chlorides on the secondary layered double hydroxide phases forming in AAMs). In portland cement and its blended cement concretes, the relationships between chloride penetration and other performance properties have been well-documented, but these relationships are not established in AAMs. All of these challenges provide significant indications for further research to contribute to the study of chloride attack in AAMs.

2.4. Carbonation Resistance. Carbonation is the chemical reaction taking place between the reaction products of a cementitious matrix and the CO$_2$ in the atmosphere, promoting the formation of carbonates. In OPC concretes, the pH is mainly controlled by the amount of portlandite (Ca(OH)$_2$); however, in the case of GPC and AAMs, the pH of the system is mainly controlled by the pore solution. Thus, the carbonation mechanism in these binders occurs in two stages: first, carbonation of the pore solution, leading to a reduction on pH and precipitation of Na-rich carbonates; then, decalcification of the gel and structure deterioration. The process of carbonation is mainly controlled by reactivity and diffusivity of CO$_2$. Diffusivity of CO$_2$ is more related to the interconnectivity of the pore structure and the carbonation exposure conditions while reactivity of CO$_2$ is related to its concentration, type of binder, the gel maturity and chemistry of pore solution. It is evident that concentration of CO$_2$ plays an important role for both reactivity and diffusivity, affecting the carbonation rate during material exposure.

Applying more than 1% CO$_2$ under laboratory conditions during accelerated carbonation significantly underestimates the service life of alkali-activated binders, as reported by Bernal et al. According to the authors, during natural carbonation, evolution of the binder structure requires a long period of time (years), while accelerated tests are applied on relatively young concretes for a short period of time (weeks). Thus, the influence of gel maturity is essential when analyzing the results of accelerated carbonation exposure. Based on thermodynamic calculations and XRD results, they established a relationship between alkalinity and CO$_2$/$\text{HCO}_3^-$ ratios in simulated pore solutions with different concentrations of NaOH added in the activator, for natural (0.04% CO$_2$) and accelerated (4% CO$_2$) carbonation. The pH reduction in pore solution, because of accelerated carbonation, was 2 orders of magnitude lower, in comparison with natural carbonation exposure. The pH has a tendency to increase as the NaOH concentration increases up to 0.5 mol/kg; after that, the composition of pore solution does not change significantly. It was noted that if the accelerated carbonation is applied to induce steel corrosion of alkali-activated binder, the results will not be representative of in-service performance under natural conditions, because of the rapid decrease in pH in the pore solution and extensive degradation at high CO$_2$ concentration during testing.

The effect of exposure conditions (i.e., concentration of CO$_2$, temperature, and relative humidity in accelerated carbonation testing of AAMs) has been investigated by Byfors et al. and recently by Bernal et al. The lower carbonation rates at higher relative humidity values (RH > 80%) were attributed to the fact that the pores are waterlogged and almost fully saturated with moisture, so the diffusivity of CO$_2$ is significantly reduced, even when a higher concentration of CO$_2$ is applied, as stated by Houst et al. Similar conclusions have been reported where testing samples with low water absorption (i.e., initially highly saturated and refined pore networks) at high RH gives a very low carbonation rate in the early stages of the test. On the other hand, at lower relative humidity values (RH < 50%), the diffusivity of CO$_2$ also decreases, because of unsaturated conditions in the pore structure of concrete, hindering the solvation and hydration of the CO$_2$ to form carbon acid. The level of relative humidity widely used in accelerated carbonation testing of OPC concrete is RH = 50%−70%, providing the best results, which almost reproduced real in-service conditions of concrete carbonation.

The highest carbonation rates reported in alkali-activated BFS−metakaolin systems were observed in specimens exposed at RH = 65% ± 5%, where a partially saturated moisture condition produced the highest volume of permeable pores and accelerated the carbonation reaction process. Although in this work, the authors make a clear correlation between relative humidity and carbonation rate, the disagreement between residual compressive strength and relative humidity trends was not well-documented and still unclear. Another interesting point that was raised by the same authors and, consequently, requires further investigation was the limitation of the phenolphthalein method mostly at higher CO$_2$ concentrations (3% in their study).

Deja studied the carbonation depths in AAS mortars and concretes, in comparison with OPC-based specimens. Com...
pressurized strength was increased at longer exposure times to carbonation in both binders, because of the precipitation of carbonates, which resulted in a refined pore structure of specimens. In this study, the relative humidity used (RH = 90%) and CO₂ concentration (100%) values are significantly higher than those commonly used, which make it difficult to formulate any relationship or comparison with other results.

In a recent study of Bernal et al., the AAS concrete specimens were exposed to the natural carbonation for 7 years. It was observed that different activation conditions lead to different carbonation rates. It was also shown that the detected carbonation depth under accelerated carbonation testing was relatively higher than that measured under natural conditions. This was attributed to the thermodynamic differences between accelerated and natural carbonation conditions, which considerably affects the degradation mechanism under these conditions. Similar results have been reported on AAS concrete. The microcracking in partially carbonated AAS concretes was also observed, which can be associated with the autogenous shrinkage along the duration of reaction. The same assumption was made in other works. The authors concluded that AAS binders can be more susceptible to shrinkage-related processes than portland-cement-based binders. This was even emphasized in the early stage of curing. In addition, the extent of shrinkage is considered to be largely affected by the composition and concentration of the alkaline activator solution.

The effect of the binder composition and curing conditions on carbonation rate was extensively studied. Bernal et al. found that the higher amount of paste in AAS concrete reduced carbonation depth. The authors also reported that activation conditions—mainly, the concentration of the activator—greatly influenced the carbonation depth.

The influence of slag chemistry on the kinetics of reaction and the structural evolution of the solid phases forming AAS binders has been investigated. It was found that increasing the MgO content induced a significant reduction in the carbonation rate (Figure 3). This phenomenon has been attributed to the formation of layered double hydroxides with a hydrotalcite-type structure as a secondary reaction product in systems with high content of MgO (>5 wt %). The advantage of layered double hydroxides is its ability to absorb CO₂, thus improving the performance of AAS binders when exposed to carbonation.

Puertas et al. studied the carbonation of water glass or NaOH-activated slag pastes and mortars under saturated CO₂ conditions. It has been found that, regardless of the type of activator used, carbonation occurs on and decalciﬁes the C-S-H gel. The mechanical properties of carbonated mortars are dependent on the nature of the alkali activator. Using water glass, the decalciﬁcation of the C-S-H gel leads to a loss of cohesion in the matrix, an increase in porosity, and a decline in mechanical strength. When NaOH is used, carbonation enhanced mortar cohesion, because of the precipitation of greater amounts of calcium carbonate in the pores, causing a decline in total porosity and average pore size, and consequently, an increase in mechanical strength. In the water-glass-activated slag systems, no signiﬁcant effect, either on their behavior after carbonation or on the nature of the reaction products, has been found after the inclusion of organic admixtures. More intense and deeper carbonation was found in AAS than in portland cement mortars. Based on their experimental results on paste specimens, the author proposed two different carbonation mechanisms in OPC and AAS systems. In OPC, carbonation occurs in both the portlandite and the C-S-H gel, whereas in AAS, carbonation occurs directly in the C-S-H gel, probably through the reaction between H₂CO₃ dissolved in the aqueous phase and the Ca²⁺ ions in the interlayer zone, between the silicate chains.

Criado et al. studied the effect of curing conditions on the carbonation of the reaction products of alkali-activated FA systems. It was found that a quick carbonation process could be favored if the curing conditions are not suitable. The initial carbonation of the system involves the reduction of the pH levels; therefore, the ash activation rate and the mechanical strength developments are notably slowed. The carbonated phases have been identiﬁed in samples that were exposed directly in the oven alongside a porcelain capsule containing water, regardless of the duration of thermal treatment and the type of activator used. The atmospheric CO₂ reacts with the sodium present in the system, producing sodium bicarbonates instead, which reduces the amount of sodium available for the formation of N-A-S-H gel. On the other hand, the authors reported that the carbonation does not interrupt it, because of the amount of soluble sodium, which decreased at longer curing times.

The carbonation of reinforced geopolymer concretes produced from three alkali-activated fly ashes with different CaO content (1.97, 5.00, and 12.93 wt %, respectively) has been recently studied. Specimens were subjected to...
accelerated carbonation at 5% CO₂ for a period of 450 days. Changes of concrete structures were determined and electrochemical measurements were performed over the time of testing to evaluate the effects of carbonation on the corrosion of the steel rebar. Accelerated carbonation induced a considerable reduction in the pH, a reduction in the compressive strength, and a consequent increase in the total porosity in all examined concrete specimens. The reduction in pH is mainly linked to the carbonation of the pore solution according to the fact that sodium carbonates are identified as the main reaction products produced in carbonated specimens. Greater changes in porosity were observed in the case of high-Ca fly ash-based concretes. It was also concluded that low-Ca class F fly ashes reduced the risk of corrosion in the steel reinforcement to a greater extent than those containing higher levels of Ca, with regard to the chemistry and physical properties of these materials.

The carbonation resistance of alkali-activated fly ash, slag, or their blend is mainly influenced by preconditioning of the specimens prior to carbonation testing, carbonation shrinkage induced by decalciﬁcation of the binding products, and the chemistry of the pore solution. The standard methodology, which is similar to that which exists for the OPC system, is needed in alkali-activated binders, speciﬁcally development of a technique that will imply accelerated carbonation testing.

Determination of appropriate exposure conditions, such as the concentration of CO₂, the relative humidity (expressed as a percentage), and the time of exposure needed to replicate a real in-service environment must be postulated. This represents another challenge as the performances of AAMs under accelerated carbonation and natural carbonation have showed some divergences. Shi et al. reported a natural carbonation rate of existing aged structures in service, based on AAS concrete, of <1 mm/yr; however, under accelerated carbonation exposures (7% CO₂) for 240 h, a very high carbonation rate (between 13 and 25 mm) was reported.

2.5. Acid Resistance. Concrete structures can suffer serious damage when exposed to acidic media such as hydrochloric, acetic, nitric, and sulfuric acids. The most important cause of acid-induced damage to infrastructure elements is biogenic sulfuric acid corrosion, which often occurs in sewer systems. Several studies showed that AAMs, including those derived from high-Ca precursors, exhibit better acid corrosion resistance than Portland cement, because of the differences in the nature of their hydration products. Bernal et al. concluded that AAS exhibits better acid resistance than Portland cement, retaining 75% of their original strength after 150 days of exposure to acetic acid. Higher stability of AAS binder under acetic acid attack is attributed to lower initial permeability, higher alkalinity of the pore solution, and low CaO/SiO₂ ratio in the AAS system. Decalciﬁcation of the AAS binder through formation of calcium acetate leaves a residual aluminosilicate-type gel in the corroded area, which is more soluble and more mechanically sound than the silicate gel formed in Portland cement binders, thus contributing to the higher acid resistance of AAS binder.

Allalverdi et al. investigated the acid resistance of alkali-activated fly ash and slag mixtures exposed to sulfuric acid solution with different concentrations. The corrosion mechanism of hardened paste at relatively high concentrations of sulfuric acid (pH ~1) consists of two steps. The first step is an ion exchange reaction between the charge-compensating cations of the framework (Na⁺, Ca⁴⁺) and H⁺ or H₂O⁺ ions from the solution, along with an electrophilic attack by acid protons on polymeric Si–O–Al bonds. The electrophilic attack of acid protons results in the ejection of tetrahedral aluminum from the aluminosilicate framework. In the second step, the exchanged Ca ions diffusing toward the acid solution react with counter-diffusing sulfate anions, resulting in the formation and deposition of gypsum crystals inside a corroding layer, which provides a protective effect inhibiting the total process of deterioration. At mild concentrations of sulfuric acid (pH ~2), the first step of the total corrosion process continues until results in the formation of shrinkage cracks. When shrinkage cracks become wide enough, sulfate anions diffuse into the cracks, and react with the counter-diffusing Ca ions, resulting in the formation and deposition of gypsum crystals. At relatively low concentrations of sulfuric acid (pH ~3) and for limited periods of exposure time (~90 days), the corrosion mechanism was described as the leaching of charge-compensating cations and the ejection of tetrahedral aluminum with no gypsum deposition, which is identical to that of pH 3 nitric acid.

The durability of geopolymer concrete prepared using blended ash of pulverized fuel ash and palm oil fuel ash when exposed to 2% sulfuric acid solution for 18 months has been investigated by Ariffin et al. and compared to OPC performances under the same conditions. From the evaluation of mass and strength loss, it has been reported that geopolymer concretes exhibit better acid resistance than OPC, based on their lower mass and strength loss. While the average mass and strength loss for geopolymer were 8% and 35%, these parameters are considerably high for OPC, reaching values of 20% and 68%, respectively. One of the possible reported explanations to OPC degradation was related to the reaction between acid and Portlandite, which could induce tensile stress, resulting in cracking and scaling of concrete. However, no additional information concerning the origin of stress, such as the nature/type of products formed during the acid–base reaction and how their formation could affect the microstructure integrity, was provided. The decrease in geopolymer strength was not well documented, and only the breakage of aluminosilicates bonding was postulated as the main cause of geopolymer degradation. One important factor that was not discussed in the reported results and could be responsible for the decrease on mass and strength is the initial porosity of geopolymers. This parameter has been deeply analyzed by Bakharev during a comparative acid resistance study of fly ash, fly ash+OPC, and OPC specimens. It has been concluded from this study that the three systems had very similar porosity, as measured by Brunauer–Emmett–Teller (BET) surface area analysis (17.5%, 18%, and 16.6%, respectively) but their performances in durability tests were very different; such behavior was attributed to the difference in their average pore diameter, where fly ash geopolymer specimens with a pore diameter of ~45 Å were the most durable, while OPC specimens having a pore diameter of ~100 Å were the least durable in the acidic environment.

On the other hand, the examination of XRD data before and after immersion in sulfuric acid solution showed an increase of zeolite phases (sodalite, natrolite) formed after acid attack; however, no comment on their effect on the durability of the geopolymer matrix was reported. Contrarily, Fernandez-Jimenez et al. reported the destruction of zeolite phases and dealumination of N-A-S-H gel after immersion in HCl solution; however, Bakharev attributed the loss of strength in fly-ash-based geopolymer (8FAK) exposed to sulfuric acid to the increased formation of Na–P1 zeolite (gismondine).
While the loss in both strength and mass has been widely accepted and used to assess the acid resistance of cement-based materials including AAM, the suitability of these parameters has been subjected to recent debates. Provis et al. reported that the use of compressive strength loss, as a measure of degradation during an accelerated test with a duration of weeks or months, can be complicated by the increase in strength of the undamaged binder regions during the test, which, to some extent, can counteract the strength losses in the degraded binder. Another drawback reported by the same authors was related to the percentage mass loss measured at a given corrosion depth, which is closely linked to the sample geometry; therefore, a larger sample will lose less strength at the same corrosion depth as a smaller sample, leading to severe difficulties in comparing results between investigations. The corroded depth was proposed to be a better and more suitable parameter than the loss of mass or compressive strength able to be measured with higher accuracy, and reproducibility of the acid resistance of AAM.133,135

Recently, Tahri et al. investigated the resistance to chemical attack (by sulfuric and nitric acid) of fly ash geopolymers for coating portland cement concrete. It has been found that the fly ash geopolymeric mortar shows a good performance for sulfuric acid concentrations of both 10% and 20%. Even for a sulfuric acid concentration of 30%, this mortar shows good acid resistance for immersion for 14 days. After nitric acid attack, fly-ash-based mortars show a relatively low weight loss (∼2%), even after 56 days of immersion, and these materials showed a clear vulnerability to acid attack only for a very high concentration (30%), in comparison with other polymeric resins based on epoxy or acrylics. No explicit explanation for the important decrease on fly ash geopolymer mass loss (20%–25% after 14–56 days of exposure to acid attack) was mentioned in their discussion. This decrease in performance could be associated with either the relatively high permeability of fly-ash-based geopolymers, together with their lower rate strength development, or the increase in calcium content in the mixture (as they used calcium hydroxide as partial replacement (10%) for fly ash in order to improve mechanical properties). These findings indicated that the role of calcium in alkali-activated material performances remains poorly defined and still need more investigation to draw suitable correlations between calcium content and the engineering and durability properties of the final product.

2.6. Efflorescence. Efflorescence is defined as the formation of a surface deposit that occurs in both natural and artificial environments. It is considered to be a typical surface defect commonly observed on brick, mortar, and concrete facades. This defect is associated with the growth of salt crystals on a surface, originating from the material itself or the surrounding environment and caused by chemical reaction and/or evaporation of the salt solution. In conventional cement-based materials, the most common efflorescence is the formation of calcium carbonate via the following equation:

\[
\text{CO}_2(\text{g}) + \text{Ca}^{2+} + 2\text{OH}^- \rightarrow \text{CaCO}_3(\text{solid}) + 2\text{H}_2\text{O}
\]

This process was described to occur in six steps: dissolving of \(\text{CO}_2(\text{g})\) in \(\text{H}_2\text{O}\) at the air/water interface on the surface of products, conversion of \(\text{CO}_2\) to aqueous species, release of alkalis, dissolution of \(\text{Ca(OH})_2\), diffusion of reactants through solution, and precipitation of calcium carbonate.137

In AAM, mostly those having a porous and open microstructure (generally with, but not limited to, a low calcium content such as fly ash), efflorescence is caused by excess alkali oxide (sodium, potassium, etc.) remaining unreacted in the material, because of the relative mobility of the alkali cations within the aluminosilicate framework, particularly when the material is exposed to cycles of wetting/drying or moisture transfer. A hydrous alkali carbonate, \(\text{Na}_2\text{CO}_3\cdot7\text{H}_2\text{O}\), has been identified by XRD analysis as the main efflorescence product of sodium silicate activated fly ash.136 These results are in agreement with the mechanism of the efflorescence formation in geopolymers described through the following equations:

\[
\text{CO}_2(\text{g}) + 2\text{OH}^- \rightarrow \text{CO}_3^{2-} + \text{H}_2\text{O}
\]

(2)

\[
\text{CO}_3^{2-} + 2\text{Na}^+ + 7\text{H}_2\text{O} \rightarrow \text{Na}_2\text{CO}_3\cdot7\text{H}_2\text{O}(s)
\]

(3)

AAM are generally more prone to efflorescence formation, compared to portland cement. This behavior could be related to their higher porosity, to the high alkali concentration in the pore solution, and to the weak binding property of alkali cations in the aluminosilicate framework.133,135,140,141,146

Efflorescence phenomena have been scarcely investigated, compared to other AAM issues. In a recently published work, it has been reported that only ~10 works related to efflorescence could be found in the literature and only few of them are directly focused on efflorescence. One of the most important reasons that could be related to the lack of research on this topic is whether efflorescence is considered to be a durability issue or only a simple aesthetic problem. In some works, it has been reported that the efflorescence products are whitish and structurally harmless but aesthetically undesirable. The same authors consider them to be unsightly, and so it is desirable to avoid them; however, they are rarely harmful to the performance of the material. According to other studies, the efflorescence process was considered only to be unsightly, but not expensive.124,137 Alexander et al. emphasized that, depending on the environmental conditions, efflorescence can be observed on the surface of the concrete, or subflourescence could occur below the surface, or both. They also highlighted that efflorescence is not aesthetically pleasing and can often be easily eliminated by washing the surface, while subflourescence is more likely to generate stresses that will deteriorate the structure. Recently, Yao et al. investigated the effect of efflorescence on the compressive strength and shrinkage property of alkali-activated fly ash and slag blends (BFS:FA = 970:100:0, 75:25, 50:50, 25:75, 0:100). Figure 4 shows efflorescence on the surface of investigated specimens at different aging regimes. It has been found that efflorescence led to the strength loss associated with the formation of carbonate crystals in the binder and the damage of the microstructure. They also concluded that shrinkage behavior was linked to efflorescence, in addition to the relative humidity. These latest results clearly showed that efflorescence is a durability issue rather than aesthetic problem, since it produced a reduction in the alkalinity of the binder; the loss of alkalinity may damage the protective layer around the steel reinforcement, resulting in its accelerated corrosion and, consequently, the concrete service life reduction. On the other hand, the efflorescence products deposited in the binder, mainly alkali carbonates such as 84

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Cured at room temperature than in samples cured at 80 °C under these activation conditions. From the study of hybrid alkaline cements based on fly ash (32%–60%) and portland cement (30%–40%), it has been concluded that no efflorescence was detected in any of the investigated mixtures. The absence of efflorescence was attributed to the low amount of (sodium/potassium), as well as to the fact that some sodium/potassium replaced Ca2+ in C-S-H hydration products. Another factor that influences efflorescence is the pore volume and permeability. In portland cement, the water participates in the hydration products; therefore, as the hydration progresses, the pore volume decreases over time. While the first reason seems reasonable as the amount of Na/K hydroxide used does not exceed 5%, the second argument is not clear, since no information about the porosity of different investigated mixtures was reported. In addition, this explanation takes into account only the portland cement content: it does not consider the big variation in the amount of fly ash in each mixture (from 32% to 60%) and its effect on total binder permeability.

The use of potassium hydroxide, instead of sodium hydroxide, in the activator was reported to be beneficial in controlling the efflorescence issue, because the K+ cations are strongly bound in the nanostructure of the gel phases in the binder. However, Skvra et al. later reported that both sodium and potassium are weakly bound in the nanostructure of (N,K)-A-S-H gel, and they determined that the alkalis (Na, K) can be almost completely leached from the investigated alkali-activated binder without compromising compressive strength. These findings open new debates, first on the role of alkalis in the stability of gel nanostructure and then the efflorescence effect on mechanical properties. A recently published work showed that sodium can partially form Na-O-Al(Si) in the gel structure, in which the Na–O bond is relatively intense, and partially form Na(H2O)6+ in Na+(H2O)6+, in which Na is weakly associated with water molecules. The authors were skeptical with the Skvara hypothesis (weakly bonded alkalis in the form of Na(H2O)6+) and stated another hypothesis, where alkalis have different states (more than the two discussed above), and each state has its own leaching rate. Concerning the efflorescence effect on compressive strength, most results found in the literature confirmed the negative influence of this process. However, other aspects, such as the relationship between efflorescence and microstructure and how this phenomenon will affect the performances of the final product, are still unclear and need further investigation.

Another aspect of controversy which is still subject to debate and is still unsolved is the differentiation between the process of efflorescence and natural/atmospheric carbonation. While some authors claimed that efflorescence is distinct from the process of natural carbonation as carbonation usually results in binder degradation, pH reduction and the deposition of carbonate reaction products in the bulk of the sample, which may or may not be visible to the naked eye, whereas efflorescence causes the formation of visible surface deposits. However, recent studies showed that efflorescence produced a reduction in alkalis, which could reduce pH and accelerate the corrosion process, and the deposition of alkali salts, which may affect some properties of the AAM. Moreover, the reduced alkali concentration in the matrix due to diffusion toward the surface will affect or suppress the later activation of residual precursors. In addition, the crystallization pressure due to the precipitation of alkali carbonates in the mixtures is hindering the growth of the gel network, which is crucial for the development of physical and mechanical properties.

To reduce efflorescence in AAM, different solutions have been proposed. Allahverdi et al. summarized efflorescence control in three methods: (i) adjustment of chemical formulation of the used alkali activator, (ii) using different curing conditions, and (iii) using different admixtures to enhance microstructure densification of AAM matrix. Naja Kani suggested that efflorescence can be reduced either by the addition of alumina-rich admixtures such as metakaolin, ground granulated blast-furnace slag, andd calcium aluminate cements or by hydrothermal curing. The additional alumina supplied by the high alumina cement admixtures leads to an increased extent of cross-linking in the geopolymer binder, reduces the mobility of alkalis (which is the key cause of efflorescence in these materials), and also generates a hardened geopolymer binder product with markedly improved mechanical properties, compared to the systems with no admixtures. It has been also concluded that hydrothermal curing at temperatures of 65 °C or higher provides a significant effect in efflorescence reduction, as well as slight strength improvements. Similar results have been reported by Zhang et al. where a decrease in the efflorescence rate due to the local reorganization and crystallization of N-A-S-H gels has been observed after hydrothermal curing. In addition, the study of fly ash- based geopolymers efflorescence behavior was found to be strongly dependent on the type of alkali activator solution, the curing temperature, and the slag addition; at the same alkali content (in terms of Na2O to solid precursor mass ratio), soluble silica present in the activator restricts the early age efflorescence of geopolymers cured at room temperature, but promotes early age efflorescence at 80 °C. This is mainly because of the finer pore size distribution developed in samples.

Figure 4. Efflorescence of the alkali-activated fly ash and slag blends with different aging regimes: (a) 7–7bot, (b) 7–28bot, (c) 7–60bot, (d) 7–60con, and (e) 60–60bot. The numbers 2–1 to 2–5 mean the contents of slag, varying from 0% to 100 wt %. Reproduced with permission from ref 143. Copyright 2015, Springer, Berlin, Heidelberg, Germany.)

985 Na2CO3·nH2O, NaHCO3, K2CO3, and KHCO3, could generate some internal stress leading to volume expansion and/or cracking of the binder.
986 To reduce efflorescence in AAM, different solutions have 987 been proposed. Allahverdi et al. summarized efflorescence 988 control in three methods: (i) adjustment of chemical 989 formulation of the used alkali activator, (ii) using different 990 curing conditions, and (iii) using different admixtures to 991 enhance microstructure densification of AAM matrix. Naja 992 Kani suggested that efflorescence can be reduced either by the 993 addition of alumina-rich admixtures such as metakaolin, ground 994 granulated blast-furnace slag, andd calcium aluminate cements 995 or by hydrothermal curing. The additional alumina supplied 996 by the high alumina cement admixtures leads to an increased 997 extent of cross-linking in the geopolymer binder, reduces the 998 mobility of alkalis (which is the key cause of efflorescence in 999 these materials), and also generates a hardened geopolymer 1000 binder product with markedly improved mechanical properties, 1001 compared to the systems with no admixtures. It has been also 1002 concluded that hydrothermal curing at temperatures of 65 °C 1003 or higher provides a significant effect in efflorescence reduction, 1004 as well as slight strength improvements. Similar results have 1005 been reported by Zhang et al. where a decrease in the 1006 efflorescence rate due to the local reorganization and 1007 crystallization of N-A-S-H gels has been observed after 1008 hydrothermal curing. In addition, the study of fly ash- 1009 based geopolymers efflorescence behavior was found to be 1010 strongly dependent on the type of alkali activator solution, the 1011 curing temperature, and the slag addition; at the same alkali 1012 content (in terms of Na2O to solid precursor mass ratio), the 1013 soluble silica present in the activator restricts the early age 1014 efflorescence of geopolymers cured at room temperature, but 1015 promotes early age efflorescence at 80 °C. This is mainly 1016 because of the finer pore size distribution developed in samples

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The most common durability aspects of AAMs based on fly ash and/or slag have been reviewed in this work. The main findings on this topic, as well as the experimental conditions deduced from some articles referenced in this review, are summarized in Table S1 in the Supporting Information. It has been proved that these materials exhibit, when adequately formulated, similar or even better durability performances than OPC-based ones. However, a comparison between different results on AAFA and/or AAS seems more difficult, because of the big divergence in experimental conditions and testing methods adopted by each group.

The versatility of alkaline activation technology, together with the availability of a large number of precursors, may allow the design of several types of cements/concretes with the required durability performances of the area/region where they will be used. In the binary slag-fly ash system, these durability requirements could be achieved by tuning the fly ash/slag ratio and adjusting other parameters such as the mix design, the activator (type, dosage), curing conditions, admixtures, etc.

The interest in geopolymer cement/concrete and other AAMs has increased rapidly in recent years, which has opened new perspectives for using these materials in many applications such as construction, infrastructure, nuclear waste immobilization, etc. However, the large-scale production and commercialization of AAMs and also their use in bulk construction are still locked despite the enormous progresses either in developing new precursors or processing methods and characterization techniques. The lack of a detailed description of their durability, together with the lack of appropriate standards and specifications, are the main issues facing the entrance and consideration of alkali-activated systems in the future construction industry.

3. CONCLUDING REMARKS

From this review on the long-term performance of fly ash/slag-based AAMs during the past decade, the following conclusions can be drawn:

- Most published results have demonstrated that AAMs have generally good performances in terms of durability (mostly their resistance to chemical attacks (acids, sulfates, etc.)).
- The durability of AAMs is mainly controlled by the chemistry of the pore solution, the microstructure of the reaction products, and the chemistry of the precursors.
- Most of the testing methods used to investigate long-term performances of AAMs derived from fly ash, slag, or their blends are developed the same for checking the durability of OPC-based systems; however, the difference on chemistry and structure of these binders, with respect to OPC, could induce inaccurate results, when adopting these standards.
- Some limitations of common testing methods have also been identified, such as repeatability and reproducibility of accelerated carbonation or the determination of the critical chloride threshold level during chloride penetration test. Many factors, such as the pH of the concrete pore solution, the dosage of admixtures, and the detection method, could be related to these issues.
- The lack of appropriate protocols, specifications, and standards for a uniform durability testing method to investigate the durability of fly ash/slag-based AAMs induced some divergences and even contradictions of some published results, making any eventual comparison between them more difficult.

- The scientific community and cement industry should provide the necessary support to technical committees actually working to develop basic recommendations and standards. This would help not only AAMs gain acceptance in the marketplace but also to validate and compare results between different research groups. RILEM Committees—especially TC 247—DTA—and the ASTM C01 Committee (Subcommittee C01.13) have been working the past few years to assess the suitability of OPC standards for testing the durability of AAMs and to build the first standards to achieve this goal.

- New advanced characterization techniques either for precursors or reaction products are needed. Small-angle neutron and X-ray scattering would reveal the detailed microstructure that will help to evaluate the microstructural growth with time as in situ experiments can be performed using these techniques. Raman spectroscopy, confocal microscopy, X-ray photoelectron spectroscopy, etc. could also provide important information.

- An accurate description and prediction of thermodynamic processes involved during gel formation mostly in low-Ca AAMs requires a developed database to provide all input parameters needed for this purpose.

- Since durability is directly related to transport phenomena, a deep understanding of the transport mechanism in geopolymer systems and AAMs is still another big challenge.

- Despite the increasing interest in AAMs, which has provided wide scientific knowledge in this area, some aspects such as degradation mechanisms under simultaneous deterioration processes still must be investigated deeply for an accurate evaluation of their durability. This will be very helpful to design the suitable concrete with the desired performances to specific conditions under which it will be applied.

ASSOCIATED CONTENT

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