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1 Recent Progress in Low-Carbon Binders

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8 Abstract:

9

10 The development of low-carbon binders has been recognized as a means of reducing the carbon 11 footprint of the Portland cement industry, in response to growing global concerns over CO₂ 12 emissions from the construction sector. This paper reviews recent progress in the three most 13 attractive low-carbon binders: alkali-activated, carbonate, and belite-ye'elimite-based binders. 14 Alkali-activated binders/materials were reviewed at the past two ICCC congresses, so this paper 15 focuses on some key developments of alkali-activated binders/materials since the last keynote 16 paper was published in 2015. Recent progress on carbonate and belite-ye'elimite-based binders 17 are also reviewed and discussed, as they are attracting more and more attention as essential 18 alternative low-carbon cementitious materials. These classes of binders have a clear role to play 19 in providing a sustainable future for global construction, as part of the available toolkit of cements.

20

Keywords: low-carbon cementitious binders; alkali-activated material; carbonated binders;
belite-ye'elimite binders; sustainability

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26 **1. Introduction**

27 Growing concerns over the greenhouse emissions profile of the Portland cement and concrete 28 industry have led to a very high level of recent interest in the development of low-carbon binders 29 as alternatives to Portland-based cements. This paper reviews recent progress in the three most 30 prominent classes of low-carbon binders: alkali-activated or geopolymer, carbonate, and belite-31 ye'elimite-based binders. Alkali-activated binders/materials were reviewed at the past two (13th and 14th) International Congresses on the Chemistry of Cement (ICCC) [1, 2], and in discussing 32 these materials, this paper focuses on some of their key developments since the last keynote paper 33 34 was published in 2015 [2]. These include rheological properties, setting behavior, structural 35 characterization, dimensional stability, durability, and their applications. However, this paper does 36 not aim to provide a detailed overview with respect to all existing insights into these materials, 37 which can be obtained in other references including [3-5], but rather focuses on the most important 38 new information that has been obtained in the past 4 years. Recent progress on carbonate and 39 belite-ye'elimite-based binders will also be reviewed and discussed in detail, as these are 40 attracting more and more attention as essential parts of the 'toolkit' of alternative low-carbon 41 cementitious materials.

42

43 2. Alkali-Activated Binders

44 **2.1. Raw materials**

45 **2.1.1 Activators**

As numerous workers have promoted alkali-activated binders as a potentially low-carbon cementing system during the past decades (see Section 2.11 for more detailed discussion of advances in the environmental assessment of these materials), increased scrutiny has fallen on the selection of the activator for use in these binders. This also has cost implications: the activator is usually the most expensive component of an alkali-activated binder, particularly if it has been produced at high purity for use in other industry sectors (which is the case for most commercial alkali silicate solutions), where such high purity may be less important to its use in alkaliactivation. So, alternative routes to alkali-activation that do not require the use of large volumes of commercial sodium silicate solutions have received serious recent attention. The production of silicate activators from olivine [6] or from waste glass [7-10], and the use of biomass ashes as an alkaline activator [11], have been established with some success.

57

58 The use of near-neutral salts as activators has also seen considerable advances in recent years. 59 This appears to be a pathway that is particularly attractive for production of cements based on 60 ground granulated blast furnace slag, which can be made to react and harden in a useful timeframe 61 by the addition of alkali carbonates or sulfates [12, 13]. In some cases, significant benefits can be gained by using a calcined layered double hydroxide as a carbonate-binding mineral additive [14, 62 63 15], to accelerate the reaction of some slags with a modest magnesia content which would 64 otherwise not react sufficiently rapidly with near-neutral salt activators. The combination of 65 calcium hydroxide and potassium carbonate has also been shown to give very good performance 66 as an activator for kaolinite [16], offering a potentially very low-carbon emissions route to the 67 production of affordable binders without needing a clay calcination step.

68

69 2.1.2 Precursors

70 The selection of precursors available for use in alkali-activation has also broadened significantly 71 in recent years, with particular emphasis being placed upon the use of materials for which there 72 is not strong competition in demand from utilization in blends with Portland cement. For example, 73 calcined non-kaolinitic clays [17-20], palm oil fuel ash [21, 22] or other minerals [23-26], have 74 been shown to yield alkali-activated binder systems with technical properties that are attractive in 75 given applications. Various industrial by-products or wastes without current large-scale utilization 76 as supplementary cementitious materials have been tested and validated for use in alkali-activated 77 binders, including red mud [27-29] and various glassy wastes including slags, some of which can 78 benefit from thermal re-processing or modification to improve their reactivity before use [10, 30-79 35]. Detailed work is also ongoing to better understand the reactivity of fly ashes under alkaliactivation conditions [36-39], and to valorize kaolinitic clay resources which are not of sufficient purity for use in other applications such as ceramic whitewares or coatings [40-42]. Common to many of these types of materials, the role of iron in alkali-activation precursors (and in the resulting binders) is beginning to be understood to some degree [19, 43, 44]. However, a detailed description of its reactivity and the structural implications of its inclusion in the binding gel still remain very much elusive. This is an area in which further advances are expected – and needed – in the coming years.

87

88 Some of the potential precursors described here and in the broader literature are only available in 89 commercially viable quantities in limited locations - but in the locations where they are available, 90 utilization in alkali-activated binders can be extremely attractive as a local solution to the needs 91 of the construction industry. This ability to achieve local specificity in materials design and 92 specification highlights one of the key strengths of alkali-activation, which is its ability to make 93 use of a wide (and ever-growing) range of materials as precursors. However, it also raises 94 challenges in standardization and specification, as it is almost impossible to write a prescriptive 95 recipe-based standard that covers such a broad set of potential material chemistries. This highlights the need for performance-based specification of alkali-activated binders rather than 96 97 relying on a prescriptive approach; discussion will return to this point in Section 2.10 in 98 consideration of durability.

99

100 2.2. Rheology

An understanding of the rheological properties of alkali-activated cement pastes, mortars and concretes is essential to ascertaining their consistency and workability, and consequently their ease of casting or placement. In addition, the rheological properties of alkali-activated cementbased materials have a strong influence on their microstructure, mechanical properties, and durability. The identification and modeling of rheological characteristics of alkali-activated materials (pastes, mortars and concretes) have been intensively studied since the last ICCC, as this has been identified as an area with major practical importance (and challenges) for the fieldapplication of these materials.

109

The rheological behavior of alkali-activated slag (AAS) pastes, activated with NaOH alone or in 110 111 combination with Na₂CO₃, was similar to the rheology observed in Portland cement pastes and 112 could be fitted by the Bingham model. Conversely, the AAS pastes activated with waterglass 113 (AAS-Wg) required description by the Herschel-Bulkley model, as shown in Fig. 1 [45]. Moreover, 114 their rheology depends on both the SiO₂/Na₂O ratio of waterglass and the Na₂O concentration of the activator. The early-age formation of a C-S-H type gel in silicate-activated AAS due to the 115 reaction between silicate species in waterglass and Ca^{2+} as it dissolves from slag particles, and the 116 117 effect of this gel formation on the paste rheology, has been confirmed in several publications [3, 118 4].

119



Figure 1. Shear stress versus shear rate curves showing paste hysteresis cycles: (A) NaOH-activated
paste (AASN); (B) waterglass-activated paste (SiO₂/Na₂O=1.2 in the activator; AAS-Wg).
Reproduced from [45].

124

120

The high viscosity of alkali silicate-activated cements, sometimes also accompanied by a high yield stress, is one of the critical challenges that hinder their wide application. Favier et al. [46] identified that this high viscosity was intrinsically due to the use of a viscous alkaline silicate activating solution, not controlled by interparticle contacts. Yang et al. [47] focused on ameliorating the rheological performance of sodium silicate-activated fly ash/slag pastes using fly ash microspheres as an inorganic dispersing agent. Rheology is also temperature-dependent; Mehdizadeh and Najafi Kani [48] determined an "apparent activation energy" parameter from the temperature dependence of the rheology of alkali-activated phosphorous slag (AAPS) paste, based on the Arrhenius viscosity model, and determined an activation energy of 42 (\pm 3) kJ/mol for the temperature range 10-40°C, which was of a similar magnitude to the equivalent parameter determined for Portland cement pastes.

136

137 The nature of the mixing protocol has been identified to be a key determinant in AAS mortar and 138 concrete rheology. In a series of studies, the Bingham model gave a good fit for all the PC and 139 AAS mortars and concretes tested [49, 50]. A longer time of mixing had an adverse effect on 140 rheology, but gave a slight improvement in hardened performance. In AAS-Wg concrete, the application of a longer mixing time can enhance the rheological behaviour and improve the 141 142 mechanical properties, as the input of mixing energy can partially break down the early-stage 143 microstructure to allow further reaction to continue. A longer mixing time raised the degree of 144 thixotropy in PC and in NaOH-activated slag concretes, which can be attributed to the formation 145 of fine particles induced by over-mixing, but decreased flocculation and lowered the degree of thixotropy in sodium silicate-activated slag concrete. 146

147

148 Alkali-activated materials (AAMs) have also been proven as useful model systems for the testing 149 and validation of mini-slump [51] and creeping sphere [52] rheological measurement methods. 150 Each different approach to rheological testing, including but not limited to these methods, brings 151 access to different shear rate regimes and different shear histories in the sample, which can enrich 152 the understanding of time-dependent rheological processes in a complex fluid. Understanding the 153 time-dependence of the rheology of alkali-activated pastes is not straightforward [53, 54], as these 154 materials tend to be thixotropic in addition to the reversible non-Newtonian aspects of their 155 behavior, but it is often challenging to distinguish true thixotropy from the gel structural evolution 156 that is also taking place during the rheological measurements.

158 2.3. Chemical Admixtures

159 Related to the challenges of rheology control in AAMs, it has been identified that it is critically 160 important to improve the use and applicability of admixtures to improve rheology under alkali-161 activation conditions. Due to the very significant differences in surface chemistry, zeta potential 162 and dissolution mechanisms when comparing alkali-activation to Portland cement hydration [55, 163 56], it is important to design organic molecular architectures that are specifically applicable in 164 alkali-activated binders. Recent efforts [57, 58] have demonstrated important progress in this 165 regard, including demonstration of an allyl ether-based PCE with short side chains that gave 166 extremely effective plasticizing performance in an NaOH-activated slag paste at a dose as low as 167 0.05 wt.% [57]. Keulen et al. [59] also showed that a proprietary PCE admixture could 168 dramatically increase both the slump and the slump retention of alkali-activated fly ash-slag 169 blended concretes, as shown in Fig. 2.

170



171

172 Figure 2. Slump and slump retention of alkali-activated concretes (binder 73.7% fly ash, 25% BFS,

176 There has also been important work aimed at improving the open working time of alkali-activated

^{173 1.3%} Na₂SiO₃·5H₂O powder; activator 3 M NaOH) as a function of PCE admixture dose.
174 Reproduced from [59].

177 concretes and grouts, which is problematic in some applications due to the relatively rapid 178 workability loss that is shown by some alkali-activated mixes (including in cases where 179 workability can be lost even though setting is not unduly rapid). This will be discussed further in 180 Section 2.4. Many chemical retarders for PC are not compatible with AAMs [60], but citric acid 181 [61], d-gluconic acid [62], borate and phosphate [63] have all been described to give useful 182 retardation in specific cases. However, the appropriate selection of a retarder depends critically 183 on the role and content of calcium within the alkali-activation process; high-calcium mixes tend to be more effectively retarded by small organics that can complex Ca²⁺ as it is released from the 184 solid precursor and thus delay the precipitation of C-A-S-H type gels, whereas low-calcium mixes 185 186 appear to be more amenable to the use of inorganic retarders. Although care is required to select admixtures that can give retardation without loss of some percentage of the final strength 187 188 development, it is not always straightforward.

189

2.4. Setting time

191 The setting time of an alkali-activated material (AAM) is a critical performance parameter that 192 affects its practical application. The practical ability to control setting time of alkali-activated 193 binders can determine the time window available for mixing, transportation, and casting of 194 concrete. However, fast setting is a feature of AAS-Wg based materials; the setting time of this 195 type of binder is often less than 30 min [64]. Setting behavior is affected by many factors, such as 196 raw materials, specimen preparation and process conditions, as well as the use of additives as 197 discussed in section 2.3. Lower-calcium alkali-activated binders have been identified to set 198 through a gel percolation process [65], distinct from the localized precipitation of C-A-S-H type 199 gels that characterizes the early stages of slag alkali-activation. This distinction brings important 200 new abilities to understand and manipulate the setting processes of each binder type.

201

Li et al. [66, 67] found that the setting times of alkali-activated slag-fly ash and alkali-activated slag-metakaolin pastes were prolonged with increased fly ash or metakaolin contents. Li et al. [64]

204 summarized the setting times of slag-based AAMs designed in the SiO₂-Al₂O₃-CaO system with 205 a single activator (sodium silicate of modulus 1.4), as shown in Fig. 3. They found a general trend 206 that the setting time decreased with increasing CaO content in the SiO₂-Al₂O₃-CaO system. The 207 role of Ca (and correspondingly also Mg) as a network modifier in the slag glass leads to more 208 rapid dissolution of these precursors, whereas fly ash does not show the same degree of reactivity 209 as it is a more chemically durable glass. However, when comparing blast furnace slags of different 210 chemistry, the ability to use a single indexing parameter to predict reaction kinetics (as measured 211 by isothermal calorimetry) remains elusive [68, 69], as it is becoming clear that many parameters 212 beyond simple glass chemistry play important roles in determining the rate of slag reaction under 213 alkali-activation. An indexing approach that also includes particle size distribution parameters has 214 been proposed for fly ash activation [37], but this will also require further development before it 215 could be considered in any way broadly applicable.







Figure 3 Setting times of AAMs (color scale, in minutes) in the SiO₂-Al₂O₃-CaO solid precursor
 system [64]

Careful selection and/or combination of activators is another way to manipulate the setting times of AAMs. Shi & Day [70] and Bernal et al. [13, 71] have illustrated that a prolonged induction period, which can take up to 3-5 days in some slag-based AAM systems, can be obtained when sodium carbonate is used as the activator. They showed that the Na₂CO₃ promotes the formation

of calcite and mixed sodium-calcium carbonates prior to C-(A)-S-H; and that the fairly modest pH (between 11 and 12) generated from this activator gives a low rate of initial dissolution of the slag [70]. However, high-magnesia blast furnace slags do react relatively rapidly with Na_2CO_3 as an activator [14], and can under some conditions give higher early strength with this activator than with sodium silicate, when using a sufficiently high-MgO slag [72].

230

231 Li et al. [66, 67] further investigated the setting time of alkali-activated slags, and found that the 232 setting time was prolonged with an increase in the dose of Na₂CO₃ as activator, especially when 233 considering the final setting time and a slag of moderate MgO content, as shown in Fig. 4. The 234 initial setting time of Na₂CO₃-AAS in that study was more than 300 min, while the final setting 235 time was about 6 days. A wide range of setting time results were also shown in the work of 236 Fernández-Jiménez & Puertas [73] (mortar setting time above 3 d), Kovtun et al. [74] (concrete 237 setting time about 8 h), and Atis et al. [75] (paste setting time about 3 h). However, a well-balanced 238 mixture of Na₂CO₃, Na₂O·*r*SiO₂·*n*H₂O, and/or NaOH can take advantage of the influence of each 239 of the respective anion groups present, leading to the ability to control and tailor setting time, 240 workability and mechanical properties [71, 76].

241



Figure 4. Contours of setting time (min) of alkali-activated slag with Na₂CO₃-NaOH-Na₂SiO₃

activators: (a) Initial setting time, (b) Final setting time [66, 67]

245

244

Garg and White [77] investigated how nano-ZnO retards the setting of alkali-activated materials, by pair distribution function (PDF) analysis and isothermal calorimetry. Fig. 5 illustrates the reaction between tetrahydroxozincate ions (Zn(OH)₄²⁻) and calcium ions (Ca²⁺) in the retardation stage, that can affect the nucleation/growth of the C-(A)-S-H type gel because Ca²⁺ is bound into a calcium zincate phase. In the other hand, nano-ZnO does not significantly influence the alkaliactivation reaction of metakaolin-based binders [77], as interactions involving calcium play a pivotal role in dictating the effectiveness of nano-ZnO in retardation of AAMs.





Figure 5. A schematic outline of the retardation process caused by adding nano-ZnO to alkaliactivation of slag, sketched based on the discussions in Reference [77]

257

254

258 **2.5. Structural characterization**

259 2.5.1 Experimental approaches

In alkali-activated binder systems, as in Portland-based cements, the disordered, complex and multiphase nature of the reaction products that contribute to the main strength gain means that it is difficult to gain a full understanding of the binder characteristics from any small subset of 263 common materials science techniques [2]. Therefore, cementitious materials (and AAMs in 264 particular) provide fertile ground for the development and implementation of new approaches to 265 the experimental characterization of complex materials, and the continued structural 266 characterization of alkali-activated materials is still a key field of investigation. The C-S-H type 267 gel (often represented as C-(N)-A-S-H) and layered double hydroxide (LDH) secondary phases 268 that dominate higher-calcium AAMs, and the alkali-aluminosilicate (N-A-S-H) gel formed in 269 lower-calcium AAMs, are all highly complex in chemistry, but are becoming better understood 270 through the application of both conventional and advanced experimental analytical tools. The use 271 of new analytical techniques as well as novel research routes during the past years has provided 272 valuable insight into the structure of alkali-activated binders, and there is no evidence to suggest 273 that major advances in this area are likely to end any time soon.

274

275 There has been much recent focus on the factors affecting aluminum and alkali uptake into the C-276 S-H structure by analysis of synthetic gels. The full details of this work are beyond the scope of 277 the current review, and an excellent overview was provided in a paper presented at the 2015 ICCC 278 [78]. A low Ca/Si ratio favors the incorporation of Al(IV) into C-S-H gel, whereas a high Ca/Si ratio results in more octahedrally coordinated Al(VI) [79, 80] that is predominately present in 279 280 "third aluminate hydrate" (TAH) and in AFm phases [81]. High Al and alkali content, as in the 281 case of many AAS binders, leads to co-existence of C-A-S-H and N-A-S-H type products, 282 although these two gels can be very difficult to distinguish and isolate from each other unless 283 detailed structural models are applied to aid in the interpretation of spectroscopic data [82]. Cross-284 linking within the C-A-S-H type structure has also been identified as playing a key role in the 285 structural description and understanding of the binding phases formed in high-Ca AAMs [82-84]

286

Nuclear magnetic resonance (NMR) spectroscopy is now well known as a key technique to track the information on the local bonding environments of silicon and aluminum atoms. The structure of C-(N)-A-S-H gel has been described by various groups [83, 85-88]. However, NMR has a limited capacity to assess medium range ordering. X-ray and neutron pair distribution function 291 (PDF) analysis are now well established as techniques for understanding the gel structure formed 292 in alkali-activated binders, due to the ability of the PDF technique to probe the local atomic 293 structure of disordered materials [89], although the analysis of multi-phase materials by this 294 technique remains very challenging. Gong & White [90] used X-ray total scatting and PDF 295 analysis to study the impact of chemical variability on phase formation in alkali-activated 296 granulated blast-furnace slag (GGBS). The primary reaction product after alkali-activation was 297 C-(N)-A-S-H gel with a highly disordered structure, and PDF analysis revealed that atomic arrangements, as shown in Fig. 6 for the short-range ordering < 5 Å, in the C-(N)-A-S-H gel 298 299 differed depending on the chemical composition, especially the calcium content, of the raw 300 material. The gel connectivity of C-(N)-A-S-H gel increased with decreased calcium content, as 301 shown in Fig. 6, represented to a first approximation by the intensity of the peak just above 3 Å 302 assigned to Si-Si/Al and Mg-Al correlations. Magnesium in alkali-activated GGBS pastes is 303 primarily incorporated in the secondary reaction product, which is a hydrotalcite-like phase, as 304 will be discussed in more detail below. In-situ X-ray total scattering measurements and PDF 305 analysis are also starting to be widely used in the study of alkali-activated binders, as one of few 306 techniques that can give time-resolved information about gel local structure during setting and hardening [91-93]. 307





Figure 6. X-ray PDFs of pastes produced from hydroxide-activated granulated blast-furnace slags
(slags from different sources as noted in the legend), showing the short-range ordering (< 5 Å) [90].

313 A method to investigate the chemistry of aluminosilicate-based cementitious binders by alkali-314 activation of high-purity synthetic amorphous aluminosilicate powder has been proposed by 315 Walkley et al. [94]. The phase evolution and nanostructure development of these materials have 316 been examined after activation [95, 96]. Using this information, a new structural model of alkali 317 aluminosilicate gel (N-A-S-H) gel frameworks has been proposed based on data from solid-state nuclear magnetic resonance spectroscopy (¹⁷O, ²³Na, and ²⁷Al) [97]. Fig. 7 shows part of the 318 319 proposed conceptual model for this gel structure, with charge-balancing of the partial negative 320 charges on bridging oxygen associated with tetrahedral Al provided by sodium and by extra-321 framework Al (Al_{EF}). The alkali aluminosilicate gel predominantly comprises fully coordinated $(Q^4 \text{ or } q^4)$ Si and Al units. A considerable proportion of Al³⁺ in tetrahedral coordination exists in 322

323 sites of lower symmetry, where some of the charge-balancing capacity in the gel is provided by 324 extra-framework Al species which have not previously been observed in these materials. Greiser 325 et al. [98] conducted advanced multi-dimensional NMR analysis of N-A-S-H gels derived from 326 various amorphous silica sources and also identified extra-framework Al sites, although 327 contended that their results were consistent with tetrahedral Al_{EF} sites, as also noted by Brus et al. 328 [99] rather than the octahedral Al_{EF} identified by Walkley et al. [97]. There is evidently important 329 further work required to unravel the nanostructural details of the N-A-S-H gel structure, and multi-330 nuclear and multi-dimensional NMR techniques are expected to underpin and inform these efforts 331 in coming years.

332



Figure 7. ²⁷Al and ²³Na MAS and 3QMAS NMR analysis for the gel of alkali-activated synthetic precursor (Si/Al=2), and a 3D representation of a polymerized section of the N-A-S-H gel showing various constituent environments as marked, adapted with permission from [97]. Copyright American Chemical Society.

340 Scanning electron microscopy (SEM), together with energy-dispersive X-ray spectroscopy (EDX), 341 has been widely used as a powerful tool by different scholars [91, 100, 101] to track 342 microstructural evolution in alkali-activated binders, including some important work on model 343 systems where "microreactors" were ion-milled into slag grains and the morphology of the reaction products formed by their reaction with different alkaline solutions monitored [102]. Fig. 344 345 8 shows examples of the data that were obtained using this approach, where both the concentration and the nature of the alkaline solution led to remarkable differences in the appearance of the 346 347 reaction products formed. The large platelets of hydrotalcite-like minerals and the globular C-S-348 H type gel are particularly evident at the highest concentration of KOH tested, but there is a clear 349 progress of the alkali-activation reaction under all conditions depicted.

350



352 Figure 8 (a) A microreactor ion-milled into a GGBS grain; and microreactors exposed for 2 days to:

353 (b) 0.1 M KOH; (c) 10 M KOH; (d) 1 M NaOH. Reproduced from [102]

354

355 The use of SEM as an analytical technique is very well established in the field of cementitious 356 materials, both for imaging and for determination of elemental compositions. Particularly 357 important information has recently been obtained regarding the Mg-Al layered double hydroxide 358 (hydrotalcite-like) phase in alkali-activated slags. Richardson & Li [103] used SEM-EDX to 359 determine the Mg/Al ratio in KOH-activated blast furnace slag paste after 18 years of curing; a 360 ratio of 2.6 was determined, in excellent agreement with calculations based on XRD data for that 361 binder. Ke et al. determined Mg/Al ratios closer to 2.0 for the corresponding Mg-Al LDH phases 362 in Na₂CO₃-activated slag binders [14] and in Na₂SiO₃-activated slag binders [104], also using 363 SEM-EDX, and consistent with older literature including [105] and others. This phase is often 364 described in the cements literature as being simply "hydrotalcite", but in a mineralogical sense, 365 true hydrotalcite has Mg/Al = 3.0 and contains carbonate in its interlayer [106], whereas the LDH 366 phases formed in many alkali-activated binders will be carbonate-free. The carbonate-containing 367 member of the hydrotalcite family with Mg/Al = 2 is correctly called meixnerite, whereas the carbonate-free "M₄AH₁₃" hydrotalcite-group composition, which is probably the most relevant to 368 369 most alkali-activated binding systems, does not have a formal mineral name. So, the continued 370 description of the Mg-Al LDH phase formed in alkali-activated binders as "hydrotalcite-like" seems satisfactory, but it does need to be clearly identified that this is not true hydrotalcite in the 371 372 mineralogical sense.

373

Transmission electron microscopy (TEM) has also been applied to the analysis of alkali-activated binders with some success [103], but sample preparation for this technique remains challenging and the samples are prone to beam damage. Helium ion microscopy (HIM), a technique with some resemblances to SEM but potentially offering higher spatial resolution, was proposed by Morandeau et al. [107] for studying the nanoscale structure of alkali-activated materials. HIM involves imaging of sample surfaces by detecting secondary electrons that are excited from the 380 sample surface by bombardment with helium ions. Spatial resolution is high due to the specific 381 nature of the beam-sample interaction [108]. Hence, in comparison with SEM, this method is 382 well-suited for resolving nanometer-scale surface morphologies and porosity present in rough and 383 irregular fractured samples, including alkali-activated binders. Fig. 9 shows a selection of HIM 384 images of AAS obtained by Morandeau et al. [107]. Finer heterogeneous morphological details 385 have been captured. Two types of C-(N)-A-S-H gel have been identified, with the 'inner' gel 386 showing a foil-like morphology while the 'outer' gel appearing more globular. The use of HIM 387 with EDX analysis can provide new insight into the structure of alkali-activated materials as well 388 as other binder systems, and although the HIM instruments are still expensive, this technique is 389 becoming more widely available in the international community.

390



391 392

Figure 9. HIM images (a) GGBS particle covered by 'foil-like' C-(N)-A-S-H gel; (b) the surface of
the concave spherical void: C-(N)-A-S-H gel. Reproduced from [107].

395

Scattering and diffraction-based techniques have been used to provide insight into AAMs during the initial [109-111], medium-term [112], and later-age [113] evolution of gel, crystallite, and pore structure in AAMs. The combination of neutron and X-ray scattering, applied in parallel to provide different aspects of the required information, has also given important new insight into the highlyconnected nature of the pore structure in metakaolin-based AAMs [114]. The marked differences 401 in pore geometry and its evolution with curing as a function of the nature of the alkali cation 402 present have also been examined using the combination of PDF, small-angle X-ray scattering and 403 electrical impedance methods [115]: the structure-forming (kosmotropic) Na⁺ leads to a pore 404 network structure that evolves considerably over a 5-year curing duration, while the pore networks 405 generated through the use of chaotropic (K⁺ or Cs⁺) alkali cations are much more stable during 406 extended curing periods [115].

407

408 Another essential aspect of alkali-activated binder chemistry that has received attention - and 409 answers to some key outstanding questions - in recent years is the exact chemical nature of the 410 green coloration in alkali-activated slag binders (and similarly in high volume Portland-slag 411 blends). There has been much speculation in the past that this coloration is due to the presence of 412 polysulfide species (resulting from the release of sulfide by slag dissolution), but the previous 413 spectroscopic evidence was far from sufficient to fully substantiate this argument. However, 414 Chaouche et al. [118] have recently used synchrotron-based X-ray absorption near edge 415 spectroscopy (XANES) to demonstrate that the blue/green regions of alkali-activated (and 416 Portland cement-blended) slag binders contain the characteristic spectroscopic features of the 417 trisulfur (thiozonide) radical anion S_3^- , the same species that gives color to ultramarine pigments. 418 Consistent with this, Le Cornec et al. [119] have also recently applied vibrational spectroscopy to 419 the analysis of greening effects in 70% GGBS-30% PC blended cements, and identified 420 spectroscopic features consistent with the confinement of various sulfur radical ions (S_2^- , S_3^- and S₄) within the interlayer space of LDH phases, proposed to be of the AFm family. They claimed 421 422 that all three of these polysulfide species were present at similar ratios in the hydration products 423 of various slags tested, which modifies the characteristically blue pigmentation of the S_3^- radical 424 anion (which is the most prominent species in their spectra also), to instead give a green color 425 [119]. This is clearly an important step forward in understanding the fundamental science of 426 AAMs, but also has implications for understanding the role of binder redox chemistry in 427 controlling steel corrosion processes.

429 **2.5.2 Modelling approaches**

430 There have also been notable recent advances made in the modelling of AAM binder chemistry 431 by a variety of modelling approaches at different length scales. At an atomistic level, a number of 432 molecular dynamics (MD) studies have generated model structures claiming to represent N-A-S-433 H gels. However, the majority of these studies have not included water in a realistic or reasonable 434 manner, and so have generated structures of anhydrous or partially-hydrated glasses rather than 435 anything representative of a N-A-S-H structure that could form by precipitation from an aqueous solution. Features such as edge-sharing tetrahedra and extensive Al^{IV}-O-Al^{IV} bonding, which are 436 437 not observed to any significant degree in hydrous aluminosilicate minerals, can be viewed as 438 indicators of such an unrealistic structure. An exception to this trend is the work of Lolli et al. 439 [120], who used MD to generate correctly hydrated N-A-S-H gel structures based on three approaches: a "crystalline" structure based on adjusting the sodalite framework to the desired N-440 441 A-S-H stoichiometry; a "defective" structure generated by introducing defects into the sodalite 442 framework and allowing this to relax (in the presence of water) using MD, and an "amorphous" 443 structure based on SiO₂ glass adjusted to the desired N-A-S-H stoichiometry. Among these three 444 models, the "defective" structure (Fig. 10) gave the best match to experimental PDF data, and also yielded predictions of nanoscale mechanical properties and porosity that are consistent with the 445 available literature, while complying with the requirements for predominantly Q^4 bonding, the 446 447 absence of edge-sharing tetrahedra, and agreement with Loewenstein's principle of Al-O-Al 448 avoidance [121]. This can therefore be considered to be a reasonably representative structural 449 model for N-A-S-H gel at this length scale, and is consistent at a chemical level with the schematic 450 description of the potential site types that was shown in Figure 7.



452

Figure 10. Representation of part of the N-A-S-H gel structure generated by application of molecular
dynamics to relax a defective sodalite framework. Reproduced with permission from [120].
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456

457 At the mesoscale, Valentini [92] adapted an established code designed for Portland cement 458 hydration simulations to describe the activation of metakaolin by different alkaline solutions, 459 while Yang & White [122] advanced the use of on-lattice coarse-grained Monte Carlo simulations 460 to describe activation of different aluminosilicate precursors. Modeling approaches such as these 461 are computationally intensive and need care in parameterization and specification to ensure that 462 atomic-scale interactions are replicated as accurately as possible on the mesoscale, but also 463 provide unparalleled access to mechanistic and microstructural information on a length scale of 464 up to hundreds of nanometers, which is very difficult to access experimentally in real-time.

465

Thermodynamic modeling of phase assemblages in AAMs has been an area of particularly important recent developments, where the application of a detailed ideal solid solution model for the C-(N,K)-A-S-H system [123] supported by the availability of improved solubility data [124] has significantly moved forward the state of the art. This has enabled advances in phase assemblage predictions for alkali-activation of blast furnace slag [125-127], including prediction 471 of phase diagrams for a wide range of slag and activator compositions [128]. Solubility 472 information for N-A-S-H gels [129], and improved information for aqueous species in the N-K-473 A-S-H-Cl system [130], are bringing the opportunity for modelling of some lower-calcium binder 474 systems [131], although much work is still required to develop and validate the necessary range 475 of model constituents to enable full description of the phase assemblages in these binders. In 476 particular, the database of zeolite phases available for inclusion in models of the (N,K)-A-S-H 477 system requires expansion, as there are significant gaps in the literature here, although constrained 478 to some degree by issues of metastability and difficulties in actually defining "solubility" in many 479 instances.

480

481 **2.6. Microstructure and mechanical properties**

482 The development of a detailed understanding of the microstructure and mechanical properties of 483 AAMs is obviously key to the application of these materials in civil and infrastructure applications. 484 In particular, it is essential to understand whether the engineering design equations that have been 485 established for conventional concretes are also broadly applicable to AAM concretes. A detailed 486 review of the mechanical properties of AAM concretes has been provided recently by Ding et al. 487 [132], and the full scope of that review will not be repeated here. However, it should be noted that 488 in many cases, the general functional forms of relationships that work well in describing the 489 characteristics of Portland cement-based concretes also appear valid for AAM concretes, but some 490 re-fitting of parameters seems necessary.

491

492 Analysis of the stress-strain characteristics of AAM concretes has tended to show that these 493 materials show a higher tensile strength, lower modulus of elasticity, and lower Poisson's ratio 494 than conventional Portland cement concretes, as reported by e.g. [133-135] and many others. 495 Thomas and Peethamparan [136] also showed that the specimen size effect in compressive 496 strength testing of AAM concrete cylinders was well described by the established models for 497 Portland cement in the case of AAMs based on GGBS, but observed an unexpectedly strong size 498 effect in AAMs based on fly ash. This was attributed to microcracking effects in the fly ash-based 499 binders; microcracking of AAMs is certainly an area requiring more detailed analysis, and will be 500 revisited below (section 2.8) in the discussion of dimensional stability. The creep of AAM 501 concretes also requires further attention. There are indications that although the early-age creep 502 of these materials resembles that of Portland cement-based concretes, the deceleration of creep 503 over extended timeframes may be less dramatic in AAM concretes, meaning that longer-term 504 creep processes must be taken into account in structural design procedures [137]. AAMs have also 505 been observed to have a higher fracture energy [138, 139] and a more compact interfacial 506 transition zone [138, 140] than comparable Portland cement-based materials, and undergo a more 507 localized cracking process [141]. The strong aggregate-paste bond also gives relatively high 508 fatigue resistance [142, 143].

509

510 An important finding underpinning much of the analysis of AAM property-microstructure 511 relationships was the identification by Winnefeld et al. [68] that the degree of reaction of the blast 512 furnace slag precursor appears to be a characteristic parameter which controls strength, across a 513 range of slag sources and activators. It is quite probable that this relationship is critically 514 dependent on the pore structure of the AAM binder; Ranjbar et al. [144] obtained strengths of 515 over 130 MPa by hot-pressing fly ash-based AAMs to reduce porosity, while Rouver et al. [145] 516 showed a clear relationship between Young's modulus and pore volume of a range of metakaolin-517 based AAMs. Blyth et al. [146] also showed orders-of-magnitude differences in intrinsic 518 permeability between hydroxide-activated and silicate-activated slag binders, but without a 519 corresponding difference in the Young's modulus, which was attributed to a very marked 520 reduction in the characteristic pore diameter upon silicate activation. Bernal et al. [147] also 521 reported a surprisingly low dependence of mechanical properties on total pore volume in alkali-522 activated slag mortars, where water/binder ratios of 0.40 and 0.44 gave similar 28-day strengths 523 and identical 56-day strengths. Babaee and Castel [148] used water vapor sorption as a sensitive 524 probe of pore structure in blended fly ash-slag AAMs, distinguishing the very fine pores which 525 dominate slag-rich pastes from the mesoporous nature of fly ash-rich binders. Hu et al. [149] 526 reported that the compressive strength of alkali-activated slag/fly ash mortar is mainly affected by 527 total porosity and porosity of capillary pores, with sizes ranging from 10 to 10⁴ nm. The change 528 of pore structure in the mortar was strongly influenced by the activator silicate modulus, alkali 529 dosage and fly ash content.

530

531 **2.7. One-part alkali-activated binders**

532 One-part ('just add water') alkali-activated materials can be treated as an important step towards 533 to the commercial-scale development of these low-carbon binders. A new review on one-part 534 AAMs has been published [150] which included systematic analysis of the available literature, so 535 the current paper will not attempt to repeat the full scope of that review, which covered one-part 536 AAMs in respect of raw materials, admixtures, optimum calcination, composition, curing 537 conditions, and mechanical strength. Fig. 11 illustrates the general procedure to prepare one-part 538 AAMs by adding water to a dry mixture of solid alkali-activator and a solid aluminosilicate 539 precursor, including a calcination step if necessary. This technology has been considered as a 540 method to face some technical challenges related to conventional (two-part) AAMs, in particular 541 the question of how to handle large amounts of activator solutions which may be viscous, 542 corrosive, and/or hazardous, on site in a construction application. It should be noted that the pH 543 of most alkali-silicate activators is actually similar to that of fresh Portland cement paste, but the 544 fact that these may need to be stored and handled in large quantities by personnel who are not 545 specialized in chemical handling is nonetheless an important reason to drive forward the 546 development of one-part AAMs.



5. Solid alkali activator 🚺

549 Figure 11. The general procedure of one-part AAM (geopolymer) preparation. From [150].
550

551 In one-part alkali-activated binders, any substance that dissolves sufficiently rapidly, and offers 552 alkali cations and provides a high pH environment to facilitate dissolution of the aluminosilicate 553 precursor, can in principle be used as an activator [151]. Sodium metasilicate powders 554 $(Na_2SiO_3 \cdot xH_2O, 0 \le x \le 5)$ has been studied as a solid activator in one-part alkali-activated binders 555 [152-154]. Anhydrous sodium metasilicate was reported to contribute to higher compressive 556 strength and better workability than its hydrous counterparts when used to activate fly ash and 557 blast furnace slag-based binders, and is available commercially in the form of spray-dried powders 558 that appear quite amenable to use in alkali-activation processes at an acceptable cost.

559

560 Hybrid alkaline cement, where Portland cement and an alkaline activator are added in parallel, 561 can also be regarded as a type of one-part AAM binder. The nanostructural evolution of these 562 cements has been presented by García-Lodeiro et al. [155] via a descriptive model. Fernández-563 Jiménez et al. [156] investigated the hydration mechanisms of fly ash-based alkaline hybrid 564 cement as a function of the nature of the activator as it was supplied in different forms (solid and 565 liquid). When solid sodium sulfate was used as the activator in hybrid binder systems, slightly 566 higher mechanical strength and less AFt and AFm phases were obtained than when it was added 567 as a liquid. Both early age reaction kinetics and the nature of reaction products were influenced 568 by the form of addition of the activator.

569

570 Various different precursors, and combinations of precursors, have been validated for use in one-571 part AAM binders. In addition to the more commonly used GGBS [157, 158], fly ash [159, 160] 572 and calcined clays [161], there has also been meaningful work dedicated to the development of 573 one-part AAMs from industrial waste silicas and NaAlO₂ [162][163], and red mud [164, 165]. In 574 many of these binder systems, a careful balance needs to be drawn between adding sufficient 575 alkalis to enable rapid strength development, and avoiding the excessive alkali levels that may 576 lead to efflorescence. Ongoing work to understand the causes and implications of efflorescence 577 in both one-part and two-part AAM binder systems [166-168] is certainly necessary to underpin 578 the development and deployment of one-part AAMs.

579

580 Qu et al. [169] produced a pre-industrial hybrid alkaline cement, manufactured in a Latin 581 American plant on a scale of around 20 tons. The proportions used were 30 % Portland clinker + 32.5 % blast furnace slag + 32.5% fly ash + 5% solid activator (the main salt is Na₂SO₄). This 582 583 hybrid alkaline cement was showed to react with water at ambient temperature and reached a 584 compressive strength around 32 MPa at 28 days, with acceptable setting time and early strength. Further, the cement paste was tested at up to 1000°C, and showed better high-temperature 585 586 resistance than Portland cement due to the recrystallization of new poorly hydraulic phases, 587 mainly in gehlenite- and rankinite-type phases [170, 171]. Velandia et al. [172] also demonstrated 588 the production of concretes with good performance using a hybrid fly ash-Portland-Na₂SO₄ binder, 589 and provided extensive data about the durability performance of these concretes, including 590 correlations between key durability parameters and compressive strength at ages of up to 1 year.

591

592 **2.8. Dimensional stability**

593 The dimensional stability of any cementitious binder is a critical factor in determining its use in 594 engineering applications, as concretes are required to neither shrink nor expand excessively in 595 service. The relatively low level of bound water present in AAMs, particularly those with low 596 calcium content, has been observed to cause some susceptibility to drying shrinkage, as has 597 recently been reviewed by Mastali et al. [173]. This area has attracted a high degree of attention 598 in the literature in recent years, as the importance of understanding and controlling shrinkage has 599 become more and more evident. Low-calcium AAMs do not self-desiccate during curing in the 600 same manner as does Portland cement during its hydration [174], as there is not such a strong 601 chemical driving force withdrawing water from the pore fluid into solid phases. However, they do 602 undergo autogenous shrinkage which can lead to early-age cracking if not appropriately controlled 603 [175].

604

605 Higher-calcium AAMs such as sodium-silicate activated GGBS can self-desiccate, in part due to 606 the formation of hydration products, and also because the high ionic strength of their pore solution 607 reduces water activity to draw the relative humidity below 100% [176]; this latter effect is likely 608 to be the cause of any observed self-desiccation effects in lower-calcium binders that do not 609 chemically incorporate water of hydration. Ye & Radlińska [177] proposed that the drying 610 shrinkage of alkali-activated GGBS involves densification of the C-(N)-A-S-H as its structure is 611 damaged by reductions in relative humidity, as neither the moisture loss nor the drying shrinkage were reversible upon soaking of dried specimens. Shrinkage mitigation strategies similar to those 612 613 that are implemented in conventional Portland cement have been evaluated recently for 614 application in AAS {Ye, 2017 #741}. Chemical shrinkage of alkali-activated GGBS and GGBS-615 rich blends has been studied experimentally [178] and also identified through thermodynamic 616 modeling [125, 127]. Thomas et al. [179] identified a beneficial role for heat curing in reduction 617 of drying shrinkage, while Gao et al. [180] applied a particle packing model at paste scale to 618 optimize blends of GGBS and fly ash for minimum porosity and shrinkage. The shrinkage 619 properties of alkali-activated binders based on different blended precursors have also been 620 reported [181, 182]. Shrinkage-reducing admixtures [183-186] and super-absorbent polymers 621 [187-189] have also shown some effectiveness in reducing drying shrinkage, as has the tailored 622 design of blended activators [190]. While shrinkage control in AAMs does remain an area of open 623 research, with many questions yet to be answered, the fact that this broad range of approaches have all shown some potential for success is a strong indication that this is not an intractable issue.

626 **2.9. Durability**

627 The durability of alkali-activated binders, and concretes produced from them, has been reviewed 628 in various publications [151, 191-193]. In most cases, AAMs have been tested according to 629 methods devised and validated for the testing of Portland cement-based binders; there are ongoing 630 discussions around whether this is entirely appropriate, including through the work of a RILEM 631 Technical Committee [194, 195], and it appears that in the majority of cases there are details of 632 the standard testing methodologies that will require modification if they are to give truly 633 meaningful results for AAMs. Sample preconditioning has been highlighted as an area requiring 634 particular care when designing tests for AAMs, as some of these materials can be damaged by the 635 preconditioning regimes that are often applied to Portland cement-based materials before testing, 636 particularly when very vigorous drying is applied [93, 196-198]. A performance-based 637 specification designed specifically for application to AAMs has been released in the UK, based 638 on minimal adaptations to established Portland cement testing methodologies [199], and efforts 639 are also ongoing in other countries and through multinational collaborative programs; it is 640 expected that this will be an area of rapid development in the coming years.

641

642 It has long been identified that binder carbonation under exposure to CO_2 is an area of durability 643 that requires careful consideration when designing and specifying AAMs. Early accelerated 644 testing at high CO₂ partial pressures appeared to show that alkali-activated binders would be very 645 susceptible to carbonation, but this was not directly matched by observations under natural 646 conditions. The reasons for the sometimes very poor performance of AAMs under accelerated 647 carbonation exposure is now understood to be related to specific changes in the carbonate-648 bicarbonate equilibrium of the AAM pore solution at elevated CO₂ partial pressures [200]. This 649 can give an unrepresentative reduction in pH compared to natural carbonation exposure, which 650 has been shown to give a much less marked reductions in pH [201, 202]. Relative humidity control during carbonation testing has also been shown to be highly influential in determining theobserved rate of carbonation [203].

653

654 The carbonation of AAMs has been shown to be strongly dependent on binder microstructure, and 655 particularly the degree of microstructural evolution and pore network refinement achieved prior 656 to the start of carbonation exposure, which can be influenced by curing, as well as various mix 657 design parameters such as the activator dose [204-207]. The presence of hydrotalcite-type LDH 658 phases has been identified as being particularly crucial in enabling carbonation resistance, whether 659 these are produced directly as a result of activation of an Mg-containing precursor, or due to the 660 addition of a supplemental Mg source (or calcined LDH as a seeding/templating agent) [208] [15, 661 209, 210]. The mechanisms of carbonation shrinkage in alkali-activated slag binders have also 662 been identified [211]. Together, these new aspects of insight provide essential steps toward 663 designing AAMs that can appropriately resist carbonation in service, and also in understanding 664 the connections between accelerated and natural carbonation mechanisms to enable the design of appropriate laboratory tests for the prediction of field performance. Electrochemical examinations 665 666 of carbonated AAM concrete showed that the binders have been capable of keeping the 667 reinforcement in a passive condition even with the lowered pH caused by the accelerated 668 carbonation [212, 213], but this does necessitate further investigation.

669

670 In many steel-reinforced concrete applications, the service life of a structure or element is 671 governed by the ability of the concrete to protect the steel from chloride-induced corrosion. The 672 rate and mechanisms of chloride transport in AAMs have been reviewed in detail by Osio-673 Norgaard et al. [214]. Thomas et al. [215, 216] have provided a comparison of chloride test 674 methods as applied to these materials. Hu et al. [144] found that some alkali could leach out during 675 specimen saturation before the electrically accelerated chloride transport test, and that the water-676 to-specimen ratio could have a critical effect on the passed charges, but not on the chloride 677 migration coefficient of the specimens. This is an active area of work in international organizations including RILEM and the European Federation for Corrosion, who have established 678

679 working groups to investigate chloride transport and corrosion initiation in AAMs, with a 680 particular view toward the development of more appropriate testing methods. Noushini and Castel 681 [217] have recently discussed the development of performance-based criteria for AAM concretes 682 based on chloride ponding and electrochemically accelerated tests, which is an essential step 683 toward performance-based standardization of AAMs. For electrochemical testing, it also appears 684 likely that the proportionality constants applied in the relationships that are commonly used to 685 obtain material parameters from polarization curves (Tafel slopes) for Portland cement will need 686 to be re-assessed for AAMs, as there appear to be significant deviations from the classical 687 electrochemistry of Portland cement when considering the particular pore fluid chemistry of 688 AAMs [218].

689

690 Chloride binding, particularly by hydrotalcite-type LDH phases, has been identified by some 691 authors to be very influential in determining chloride transport through AAMs [219, 220], 692 although other authors did not identify strong evidence for chloride binding in alkali-activated 693 slag concretes [215, 221]. There is a clear need for further developments to resolve this open 694 question, which is of fundamental importance to service life prediction for reinforced AAMs 695 under chloride exposure.

696

697 Ma et al. [222] linked chloride diffusivity, electrical resistivity, and corrosion testing of reinforced 698 alkali-activated concretes, and highlighted the importance of sulfide (provided by blast furnace 699 slag when used as a precursor in AAMs) in defining the corrosion rate post-initiation. The role of 700 sulfide has also been identified in studies of steel corrosion in simulated alkali-activated slag pore 701 solutions [223-225] [226], and in various types of mortar specimens [227-229]. The very high 702 pore solution pH of some AAM binders has also been shown to generate unconventional 703 threshold-like relationships in chloride initiation, and also to give chemical protection of steel 704 reinforcement even at high chloride concentrations [230-232]. Mundra et al. [226] also developed 705 a classification scheme for alkali-activated and slag-blended binders as shown in Fig. 12.



70(

Figure 12. Overview of the classification of cements, particularly of AAMs, based on internal
redox conditions, and the parameters influencing the onset of steel pitting and the servicelife of these binders. AAFA = alkali-activated fly ashes, AAMK = alkali-activated metakaolin.
Adapted from [226], under Creative Commons license conditions.

711

712 Questions around alkali-silica reactions, analogous to those which can lead to damaging 713 expansion in Portland cement binders with reactive aggregates, also arise regularly in discussions 714 of AAMs because of the high levels of alkali present in these binders. However, the results of 715 testing with a broad range of binder-aggregate combinations have shown that alkali-silica 716 reactions do not appear to be particularly problematic in AAMs with aggregates of 'normal' 717 reactivity [233]. It is possible to induce alkali-silica reaction expansion under accelerated 718 conditions and with the use of a reactive aggregate [234, 235], but in the majority of cases, AAM 719 mortars show less expansion than plain PC mortars with the same reactive aggregates [234, 236-720 238]. The relatively high Al concentration in the pore solution of AAMs, and in some cases also the lack or near-absence of dissolved Ca, appears to be important in restricting alkali-silicadamage [234, 237, 239].

723

724 The freeze-thaw and frost-salt resistance of AAMs do appear to merit further attention, as the 725 literature on this topic contains many case studies but lacks consistent or systematic analysis 726 across the class of materials in general. Sulfate attack on AAMs has also been studied according 727 to a number of test methods, largely because such testing is often requested by specifiers or end-728 users as sulfate attack can be problematic for Portland cement-based materials. However, the 729 fundamental mechanism of Portland cement sulfate attack, with expansive processes involving 730 the monosulfate-AFm phase, is not possible in most AAMs as this phase is absent from the hydrate 731 products. Sulfuric acid attack on AAMs is, however, a relevant mechanism related to use in sewer 732 infrastructure and other highly aggressive environments [240, 241], and the performance of 733 AAMs (particularly those with low Ca content [242, 243]) under such conditions has been 734 observed to significantly exceed that of most other cementitious binders [242, 244]. Organic acid 735 resistance has also been reported to be a strength of low-calcium AAMs, as small organic acids damage calcium-rich binders through complexation and removal of Ca²⁺ ions, but this mechanism 736 is much less significant for AAMs that do not rely on calcium as a key binder constituent [245, 737 738 246].

739

740 Testing of AAM durability in the field has generally shown results that are consistent with 741 laboratory trials under non-accelerated or minimally-accelerated conditions; the materials that 742 have been put into service under varying conditions have in many cases served very well, 743 including concretes dating back to the 1950s [151, 247, 248], and more recent demonstration or 744 full-scale infrastructure projects [249-252] including an airport in Australia that was constructed 745 largely from alkali-activated concretes [253]. Such projects are essential in building stakeholder 746 acceptance of AAM technology, and in using the experience gained to guide standards development, to ensure that the materials selected, specified and used are fully fit for purpose 747 748 [254].

749 **2.10.** Applications of alkali-activated materials

Alkali-activated binders are finding rapidly increasing uptake in a growing range of applications, and this section will provide a very brief overview of some of these. The primary route to market for these materials in the short term will almost certainly be as a binder in concretes, and increasingly sophisticated approaches to the design and use of AAM concretes (rather than just directly applying protocols used for Portland cement concretes) are being published for concretes based on alkali-activated GGBS [60, 255-257], fly ash [258-260], metakaolin [261], and various blends of these materials [259].

757

758 AAMs, including particularly the lower-calcium "geopolymer"-type materials, are also attracting 759 attention as matrices for the conditioning and immobilization of radioactive wastes; the ability of 760 these materials to host, and bind, radioisotopes of cesium and strontium has been demonstrated 761 and analyzed in some detail [262-264]. The effective immobilization of cesium in Portland 762 cement-based matrices is well known to be challenging, and so the availability of a cementing 763 system that can restrict its movement is highly desirable. There have been important recent investigations of the potential for compatibility of AAM matrices with complex waste streams 764 765 containing multiple radioisotopes [265, 266], with oily wastes [267, 268], with ion exchange media [269, 270], and with graphitic or metallic wastes [271-273]. AAM matrices have also been 766 767 demonstrated to show generally good stability under irradiation [274-276], and a hydrogen 768 radiolytic yield that depends on water content and pore structure [274].

769

The ability to produce lightweight AAMs has been investigated by numerous groups, as reviewed recently by Bai & Colombo [277] and by Zhang et al. [278]. Successful approaches have included various types of templating by organic foams or emulsions [160, 267, 268, 279-281], foaming by peroxide addition [282] or by metal powders [283] and the use of lightweight aggregates [284]

AAMs have also been tested – and in some cases validated – in a broad range of 'niche'
applications in recent years; a non-exhaustive selection of these includes:

- Well cementing in hydrocarbon and geothermal industries [285-287]
- Chromatographic substrates [288]
- Manufacture of composites [289, 290], including strain-hardening and/or ductile
 "engineered composite" materials [291-293]
- 781 Repair mortars [294]
- Materials for additive manufacturing or "3D printing" through various extrusion-based
 and powder bed processes (Fig.13 [295]) [296-298]
- Moderate-temperature refractories or fire-resistant construction materials [299-302]
- 785



- Figure 13. Additive manufacture of a vase from a metakaolin-based AAM, by extrusion. Reproduced
 from [295].
- 789

786

790 **2.11. LCA and environmental aspects**

When considering any type of cement as a potentially "eco-friendly" or "low-carbon" alternative to established technologies, it is essential that the actual environmental footprint of both the conventional and innovative materials are sufficiently well understood and quantified, to enable a fair comparison to be made. However, this is an area in which most current research publication practice in the field of alternative cements falls well short of providing the information needed for informed decision-making. This is potentially in part because of the trend for technical authors to justify the importance of their work (to funders, editors or other stakeholders) by 'advertising' the class of materials they prefer to study, and in part because the rigorous comparative environmental assessment of two construction materials is actually a highly specialized and challenging research task in itself. In the specific context of AAMs, Habert & Ouellet-Plamondon [303] have provided some very insightful discussion and assessment of data sources, and highlighted in particular the importance of understanding and controlling the environmental footprint of the alkali activator when designing and specifying an AAM mix design.

804

805 Another critical aspect of the assessment of sustainability is the need to conduct a locally-specific 806 determination of energy supply and transport options, and their costs and environmental footprints, 807 as these will differ very strongly between locations worldwide. This means that it is impossible to 808 conduct a valid, generic assessment of the emissions footprint of an AAM at a useful level of 809 precision, without knowing where in the world it will be used. The emissions attributed to 810 electricity generation differ widely from (e.g.) hydroelectric to nuclear to coal sources, and AAMs are much more dependent on electrical energy in production than is Portland cement. This opens 811 812 some attractive possibilities when considering binder production using a decarbonized electricity supply [304], which may be a strong point in favor of the use of AAMs in regions where low-813 814 carbon electricity is available.

815

816 The trend in the academic literature recently has therefore been toward regionally-specific (or 817 very localized) assessments of AAMs for use in particular applications or concrete/mortar 818 products in the Americas [305-307], Europe [308-310], and Australasia [311, 312]. The general 819 trend observed in these studies is that AAMs offer greenhouse emissions savings compared to a 820 Portland cement baseline, and usually on the order of 40-60%, but somewhat increased the 821 environmental impact in other non-greenhouse categories, such as abiotic depletion, ozone layer 822 depletion, fresh and marine water ecotoxicity, and human toxicity, that are considered in the life-823 cycle assessment process. It has also been identified that there is a strong need for more refined 824 and updated life-cycle inventory data for activator constituents including sodium silicate [303,
825 305, 313], as the currently available international databases do not reflect current production 826 practice or processes. The assessment of recyclability [314, 315] and release of potentially 827 problematic elements (toxic or naturally occurring radioactive materials) [316-318] from AAMs 828 in service has also received some attention as an essential constituent of a full cradle-to-grave or 829 cradle-to-cradle environmental assessment. This will doubtless gain further importance as non-830 carbon emissions become more of a focus in material and product assessments worldwide, while 831 the sources and characteristics of waste materials used in AAM production become ever more 832 diverse.

833

834 The other critical aspect that needs to be considered in environmental analysis of AAMs is 835 durability (at both material and element/structure scale), and this was discussed in Section 2.9 836 above. Considering all of these aspects together, and to conclude the discussion of AAMs, it 837 should be identified that AAMs are becoming a mature class of materials whose nature and 838 properties are increasingly well understood, and which offer numerous attractive opportunities to 839 exercise their desirable technical and environmental characteristics for the benefit of society. They 840 should not in any way be viewed as a panacea for all problems in the construction materials sector, 841 and nor are they likely to be universally suitable as a replacement for Portland cement-based 842 binders across the full range of applications in which cements are used, for both technical and 843 logistical (materials-supply) reasons. However, as a constituent of the future toolkit of cements, 844 AAMs do bring very significant value.

845

846 In the following sections, the focus of this review will turn to some other types of cementing 847 systems, which are not yet as widely deployed as AAMs, but which can also form valuable 848 components of the cements toolkit.

849

850 **3. Carbonate Binders**

851 The concept of carbonatable binders is based on the fact that some raw materials can harden

through carbonation. Carbonated cementitious binders have attracted wide attention in recent years, attributed to their rapid strength gain and the sequestration of CO₂ when exposed to a CO₂ rich environment [319, 320]. This section will discuss carbonated calcium silicate, carbonated Portland cement, MgO-based cement and carbonated waste-derived binders. These four binder types react with water and CO₂, yielding strong cementing materials.

857 **3.1. Carbonated calcium silicate binders**

Tricalcium silicate (C₃S), β-dicalcium silicate (β-C₂S), γ-dicalcium silicate (γ-C₂S), tricalcium disilicate (C₃S₂) and monocalcium silicate (CS) can react with CO₂ and form strong monolithic matrices [321-323]. Ashraf & Olek [324] reported that the carbonation of pure calcium silicates consists of two distinct processes: an initial phase-boundary controlled process, and then a subsequent product layer diffusion controlled process. The reaction rate constant was found to vary based on the calcium silicate phases; β-C₂S has the highest reaction rate, followed by C₃S, γ -C₂S, C₃S₂ and finally CS.

865

The carbonation products of pure calcium silicate are calcium carbonate and Ca-modified silica 866 867 gel or silica gel [323, 325, 326]. The calcium carbonate crystals resulting from carbonation of C_3S , 868 C₂S, C₃S₂ and CS include the polymorphs calcite, aragonite, and vaterite [326, 327]. The presence of a ¹³C CP/MAS NMR signal in carbonated C₃S, C₂S, and C₃S₂ phases can be attributed to the 869 870 additional formation of amorphous calcium carbonate (ACC), as shown schematically in Fig. 14. 871 However, ACC is not formed in carbonated CS under the same environmental conditions, Fig.14 872 [326]. The presence of poorly crystallized forms of CaCO₃ tends to increase the strength of the 873 carbonated calcium silicate matrices [328]. Furthermore, the values of the elastic modulus of 874 CaCO₃-rich binders can vary over a relatively wide range due to the presence of different 875 polymorphs of CaCO₃ crystals [327].





879 Figure 14. Proposed distribution of the pores (smaller than 10 nm) in carbonated calcium silicate
880 matrixes a) without the presence of amorphous calcium carbonate (ACC) and b) in the presence of

amorphous calcium carbonate (ACC), adapted from [319].

882

As shown in Fig. 15, the degree of polymerization of Ca-modified silica gel (which is essentially defined by the inverse of the Ca/Si ratio) is nearly the same for all the carbonated calcium silicates discussed, except for the carbonated C_3S , which has a slightly lower degree [326]. However, all the calcium silicates reach a similar overall carbonation level due to the retardation of the carbonation reaction after formation of nearly the same amounts of CaCO₃ (Fig. 16), indicating that effect is related to blockage of the surfaces of potentially reactive particles by the precipitated carbonates.

890



891

892Figure 15. Average Ca/Si atomic ratios of Ca-modified silica gel phase formed during the893carbonation reaction of the calcium silicate samples [321].



Figure 16. CaCO₃ contents (by mass, %) as a function of carbonation duration for different calcium silicate phases [326].

Among these calcium silicates, carbonation of C₂S has attracted more attention due to the different crystal polymorphs that it can take, particularly β -C₂S and γ -C₂S, as γ -C₂S can be produced at much lower synthesis temperatures than the conventional hydraulic calcium silicate phases. Chang et al. found β -C₂S and γ -C₂S to absorb 9.2% and 18.3% of their theoretical levels of CO₂ after 2 h of carbonation, respectively [328], but β -C₂S showed the twice the compressive strength compared to γ -C₂S. A similar result was reported by Guan et al., who found that the compressive strength of carbonated y-C₂S was 52.4 MPa after 2 h carbonation [329]. Calcite and aragonite are the main crystals formed by carbonation of γ -C₂S, and amorphous Ca-modified silica gel lacking long-range order was also formed [329-331]. Mu et al. [327] proposed a conceptual model of the carbonation process of a γ -C₂S particle, as shown in Fig. 17.



912

Figure 17. A conceptual carbonation model diagram of γ -C₂S particle. From [327]

913

914 **3.2. Carbonated Portland cement (PC) binders**

915 It is well known that accelerated early-age carbonation of Portland cement binders can result in 916 rapid strength gain and lower permeability. This is because the carbonation reaction leads to a 917 reduction in total porosity through the formation of carbonation products that occupy more space 918 than the portlandite that they are replacing. Shi et al. [332] proposed the use of pre-conditioning 919 to improve the accelerated carbonation of PC binders; the compressive strength of carbonated 920 concrete after proper pre-conditioning then 2 h of CO₂ exposure is similar to that of the concrete 921 after 24 h of steam curing. Furthermore, the carbonated concrete exhibits a similar compressive 922 strength to that of steam-cured concrete during winter weathering exposure [333]. Shi et al. [334, 923 335] suggested that pre-conditioning environments have the most crucial effect on the 924 effectiveness of CO₂ curing. Additionally, the temperature of the samples rises very quickly once 925 the samples are exposed to CO_2 ; this could reach a peak value of 70 °C during the first 15-20 min, 926 then goes down gradually with time.

927

828 Kenward et al. [336] studied hydration of an oil-well cement in the presence and the absence of 929 pure CO_2 gas. The carbonate formed was initially amorphous calcium carbonate that was not 930 detectable by XRD, but this changed to crystalline calcite detectable by XRD within 24 h. The 931 addition of carbon dioxide did result in performance benefits.

933 Shah et al. [331] have suggested that the precipitation of the three polymorphs of calcium 934 carbonate (calcite, vaterite, and aragonite) takes place in carbonated PC, while Castellote et al. 935 [333] only detected calcite in carbonated PC. CaCO₃ polymorphism is sensitive to pore fluid 936 chemistry, carbonation conditions and duration, and may also involve amorphous phases as noted 937 in Section 3.1, so this is a complex area requiring further analysis. Based on thermogravimetric 938 analysis, the main mass loss from carbonated PC pastes takes place between 600°C and 950°C. 939 The poorly crystalline carbonates, preferentially associated with C-S-H carbonation, decomposed at a temperature below 600°C, while the decomposition temperature of well crystallized CaCO₃ 940 941 is above 600 °C [320, 337, 338]. The carbonated PC was strongly decalcified to form these CaCO₃ phases along with a Ca-modified silica gel, identified via the decrease of the Q^1 and Q^2 sites, and 942 increase in Q³ and Q⁴ sites, according to ²⁹Si MAS NMR analysis. ²⁷Al MAS NMR spectroscopy 943 944 confirmed that the aluminum-bearing phases, containing mainly AlO₆ (ettringite and AFm) and 945 AlO₄ sites (C-A-S-H) were dissolved to form an alumino-silicate amorphous gel (with Al as AlO₄), 946 characterized by a broad resonance that was always positioned at the same chemical shift (55 ppm) 947 [339].

948

949 **3.3. Magnesium-based cement (MC) binders**

950 In recent decades, the use of reactive magnesium oxide (MgO) in PC has received more and more 951 attention. Carbonation of magnesium-rich cements improves the compressive strength of these 952 cementitious materials, which is attributed to the densification of materials caused by the formation of nesquehonite (MgCO₃·3H₂O), dypingite (Mg₅(CO₃)₄(OH)₂·5H₂O) and artinite 953 954 (Mg₂(OH)₂CO₃·3H₂O) [340, 341]. Mo et al. [342] suggested that a large amount of calcite and a 955 relatively smaller amount of aragonite are the calcium carbonates formed, while magnesian calcite is formed due to the incorporation of Mg^{2+} in the carbonated phase, and nesquehonite is formed 956 957 only in pastes containing at least 40% reactive MgO. Nesquehonite has been identified as the key 958 binding phase in other potential carbonated magnesia-based binders [343].

960 Figure 18 shows the morphology of carbonated magnesia cement [344]. The needle-like 961 nesquehonite and disk/rose-like hydromagnesite/dypingite, which are the main sources of strength 962 cement formulations, The development in these are observed. disk/rose-like hydromagnesite/dypingite crystals that formed due to the carbonation process could be 963 964 distinguished from the hydromagnesite seeds included within the initial mix, which possesses a 965 ground ball-like morphology.

966





H₂O·S1.0. From [344].

968 Figure 18. SEM images of H₂O samples after carbonation: (a) H₂O·S0, (b) H₂O·S0.5 and (c)

969

967

970

The area of magnesia-based cements is very diverse, and includes cements which harden and gain strength by various combinations of carbonation and other chemical reactions, as reviewed in detail recently by Walling & Provis [345]. These cements are proposed for use in many applications ranging from large-scale construction to nuclear waste immobilization, and in some cases offer the possibility for notable CO₂ emissions savings compared to conventional Portland cement. The supply of MgO is constrained in some areas due to cost or resource availability, but this is not universally the case, and materials of very good technical and environmental performance can certainly be produced using this chemistry.

979

980 **3.4. Carbonated waste-derived binders**

981 Steel slag is a broad classification for several types of industrial by-products produced during the 982 steel making process, which may be regarded as a waste-derived binder precursor [346-348]. 983 Generally, the components of steel slag include hydraulic calcium silicates (C_3S , β - C_2S), non-984 hydraulic calcium silicates (e.g. γ -C₂S, CS), and free CaO, each of which can react with CO₂. 985 Formation of calcium carbonate in the form of calcite and aragonite in the carbonated steel slag 986 binders causes microstructural densification associated with a reduction in the total porosity, and 987 hence improves the compressive strength. A carbonated steel slag binder was observed to show a 988 shift in its dominant pore diameter from 0.3-3 μ m before carbonation, to <0.1 μ m in the carbonated 989 paste [349]. The free CaO in the steel slag is partially or completely consumed due to the reaction 990 with CO₂, which improves the volume stability of the binder [350, 351]. Calcium carbonate (as 991 calcite and aragonite) is the main carbonate product formed, and portlandite and calcium silicate 992 seem to be more carbonation-reactive than the Fe-bearing phases that are also present [349]. 993 Monkman et al. [352] reported the possibility of using a carbonated ladle slag as a fine aggregate. 994 After carbonation, calcium carbonates and spurrite were detected as new phases by XRD, together 995 with the consumption of hydrogarnet and calcium hydroxide. Mortars made with the slag sand 996 demonstrated strengths comparable to mortars made with conventional river sand.

997

High calcium fly ashes have an attractive capacity to be used for mineral sequestration of CO_2 under controlled conditions [353, 354]. A recent study show that Ca-rich fly ashes react readily with gas-phase CO_2 to produce robustly cemented solids which can achieve a compressive strength of around 35 MPa and take up 9% CO_2 under optimized conditions [355]. Mahoutian and Shao [356] implemented a low temperature process to produce a binder material from blends of fly ash and ladle slag. CO_2 gas (99.5% purity) was used for carbonation of the synthesized cement for 2 hours and showed that the early age carbonation curing increased the subsequent hydration strength.

1006

1007 4. Belite-Ye'elimite Binders

1008 Ye'elimite, or calcium sulfoaluminate (Ca₄Al₆O₁₂SO₄), is the main mineral in CSA cement clinker, 1009 and has a crystallographic structure belonging to the sodalite family [357]. CSA cements have 1010 been developed on a commercial basis and used in real applications, primarily in China, since the 1011 1970s [358]. These cements are normally used as components in specialty applications because 1012 of their higher price compared to Portland cement. However, as low-carbon binders, the interest 1013 in these binders from the cement industry continues to increase because it is closer to the objective 1014 of 'eco-friendly' than many of the other low-carbon binder systems that are still under R&D [359, 1015 360]. The highly innovative production of ye'elimite-containing clinkers burning waste elemental 1016 sulfur as fuel, meeting both energy and materials supply demand in a single step, has also been 1017 demonstrated in a full-scale kiln [361], with the potential for further scale-up.

1018

1019 Ye'elimite reacts very quickly with water and contributes to the development of early strength of 1020 this binder, forming monosulfate, ettringite, and amorphous aluminum hydroxide as major 1021 hydrates. Various other reaction products can be obtained, such as strätlingite, 1022 monocarboaluminate, and gibbsite depending on the minor phases in the CSA cement [362, 363]. 1023 Normally calcium sulfates are used to adjust the binder hydration reactions, and to promote the 1024 formation of ettringite rather than monosulfate [364, 365]. Dicalcium silicate (belite) and ferrites 1025 are present as additional main mineralogical components of CSA cements. In this paper, the 1026 authors will focus on some belite-ye'elimite binders containing more belite than ye'elimite, which 1027 is different from CSA cements that contain more ye'elimite than belite. Both belite and ye'elimite 1028 are lower energy minerals compared with tricalcium silicate (alite), which are suitable for low 1029 energy and low-carbon clinker cement production and applications.

1030 **4.1. Belite-ye'elimite-ferrite binders**

1031 Belite-ye'elimite-ferrite (BYF) binders, also known as calcium sulfoaluminate or sulfobelite 1032 cement [359, 360], have belite as the main phase (45-75%), and ye'elimite as a second component 1033 (20-45%). This approach to manufacturing BYF binders allows the use of less expensive Al-rich 1034 raw materials, due to the lower ye'elimite content in the clinker compared to 'conventional' CSA 1035 cements which require a higher-purity Al source. The recent main research interest in this type of 1036 binder is related to the understanding of ye'elimite hydration, that should be carefully controlled 1037 to achieve desired rheology and setting time, and also on achieving more reactivity of the belite 1038 component that contributes to the later growth in strength. BYF binders are not yet in large-scale 1039 industrial production, but have been developed to pilot scale by some cement companies under 1040 certain national and multi-national projects.

1041

A study by Cuesta et al. [366] on the early hydration mechanisms of synthetic ye'elimite revealed that the polymorphism of ye'elimite (orthorhombic stoichiometric and pseudo-cubic solidsolution ye'elimite) influenced the hydration kinetics, together with the w/c ratio and the solubility of the additional sulfate sources.

1046

1047 Recently, new data on the hydration of BYF cements have been published by Álvarez-Pinazo et 1048 al. [367]. 'Non-active' clinker (containing β -belite and orthorhombic ye'elimite) and 'active 1049 clinker' (containing α'_{H} -belite and pseudo-cubic ye'elimite) have been studied with different 1050 calcium sulfate sources. The findings of this study showed that the active-clinker mortar 1051 developed higher compressive strengths than non-active-clinker mortars, independent of the 1052 choice of sulfate source, and it formed higher quantities of ettringite during hydration and less 1053 AFm compared to non-active cements. Another interesting finding that should be mentioned is 1054 that the paste with basanite (CaSO₄ \cdot 0.5H₂O) as the sulfate source showed the highest viscosity 1055 values and a hysteresis cycle attributed to fast setting, more so than gypsum- and anhydrite1056 containing pastes; this behavior can be adjusted by adding a small amount of polycarboxylate1057 based superplasticizer (SP) (0.05 wt%) without changing the phase assemblage [368, 369] (Fig.
1058 19). This offers a possibility to add a superplasticizer normally used in Portland cement binders
1059 to control the rheological behaviour of BYF cements.

1060



1061

Figure 19. Flow curves of different BYF clinkers with different additional sulphate sources (redrawn based on [367])

1064

Morin et al. [370] studied five different BYF cements by experimental analysis and thermodynamic modeling to track the hydration kinetics and phase assemblage, which were influenced by the quantity of anhydrite, the w/c ratio, and the clinker fineness. The results indicated that with increasing addition of anhydrite, belite hydration was delayed, which contributed to the formation of a strength plateau between early ye'elimite hydration and later belite and ferrite hydration. Also, a higher fineness of cement together with increased w/c ratio leads to higher belite hydration kinetics. Those authors proposed that the question of whether there
is enough water provided to form all hydration products strongly influences the hydration of belite
and ferrite phases.

1074

1075 **4.2. Belite-alite-ye'elimite binder**

1076 Because some belite-ye'elimite-ferrite cements present quite low mechanical strength due to their 1077 high content of belite with slow reactivity, methods to improve early strength have been sought. 1078 One such approach is a clinkering method aiming to introduce a reactive alite phase into the belite-1079 ye'elimite binder system, as an alternative way to improve the mechanical strength. However, the temperature incompatibility between alite formation (above ~1300 °C) and ve'elimite 1080 1081 decomposition (between 1300 and 1350 °C) brings difficulty in achieving the coexistence of alite 1082 and ye'elimte phases in clinkering processes. However, this problem can be solved and controlled 1083 by addition of minor quantities of CaF₂ [371] or other oxides, such as ZnO, B₂O₃, or Na₂O, in the 1084 raw meal [372].

1085

1086 Chitvoranund et al. [373] prepared a clinker by firing limestone, tuff, gypsum and calcium fluoride 1087 (used as mineraliser) in a laboratory furnace at 1300 °C for 45 min, which requires a synthesis 1088 temperature 150-200°C lower than traditional PC clinker. The minerals present in the clinker 1089 included alite (48.3%), belite (1.5% α '-C₂S + 10.3 % β -C₂S + 2.2% γ -C₂S), ye'elimite (9.6%), 1090 and ferrite (12.9%). Later, the ground clinker was mixed with 5% anhydrite to make a so-called 1091 alite-calcium sulfoaluminate cement. The hydration products were mainly C-S-H, ettringite, 1092 monosulfate, and portlandite, and hydration rates are rapid. Thermodynamic modeling revealed 1093 that the cement reacted strongly within the first 10 days of hydration, then the reaction process 1094 slowed down and was almost completed by 100 days. Ferrite exhibited reactivity in the presence 1095 of C₃S, and was consumed to give monosulfoaluminate and katoite. The compressive strength of 1096 mortars developed quite rapidly, from 10 MPa at 1 day to 35 MPa at 28 days. The release of CO₂ 1097 from this approach to clinker production is estimated at about 11-12 % less than conventional 1098 Portland cement without the consideration of other factors.

1099

In another study by Londono-Zuluaga et al. [374], a novel clinkering process to prepare belite-1100 alite-ye'elimite (BAY) binders has been optimized (900 °C/30 min - 1300 °C/15 min) and scaled-1101 1102 up to 2 kg scale. This process involved as raw meal a mix of natural limestone, sand, iron oxide 1103 (a byproduct of the sulfuric acid industry), gypsum, and kaolin. The main mineralogical 1104 composition of their final scaled-up BAY clinker was 60.6 % belite, 14.3 % of alite and 10.4% 1105 ye'elimite, on a mass basis. BAY cements were prepared by mixing the scaled-up clinker with 12 1106 wt.% anhydrite. The analysis of hydration highlighted that the main reaction products are 1107 ettringite, AFm phases (monosulfoaluminate and strätlingite), katoite, and C-S-H. Ye'elimite 1108 reacted with anhydrite to be completely consumed within 1 day, alite and ferrite almost fully 1109 reacted after 7 days, and belite showed a typical slower hydration behavior. Portlandite was not 1110 detected in the pastes at testing ages of 1, 7 and 28 days; it was speculated to be consumed to form 1111 katoite, AFt phases or monosulfoaluminate. The compressive strength of BAY mortars was recorded to be higher than that of a BYF binder prepared by the same group, at any testing age, 1112 1113 most likely due to the presence of alite. The influence of fly ash blending in BAY cements has 1114 been also reported by the same authors [375]: with the addition of fly ash, the compressive 1115 strengths of mortars increased to 68, 73 and 82 MPa, for mortars with 0, 15 and 30 wt.% 1116 replacement of BAY cement by fly ash respectively, at 180 days. The main hydration products 1117 were AFt, AFm phases, katoite, and C-S-H for all systems studied. Utilizing a small amount of superplasticizer makes it possible to prepare BAF pastes with low viscosity values. The reactivity 1118 1119 of belite appeared to have been inhibited by the high addition of fly ash, and other than the strength 1120 increase, no clear evidence of pozzolanic chemical reaction with fly ash in BAY systems was 1121 obtained. These do appear to be a promising class of cements for future large-scale utilization. 1122 Zhou et al. [376] investigated the influence of the ferrite phase in a similar binder system on its 1123 hydration and mechanical properties.

1125 **4.3. Belite-ye'elimite-ternesite binder**

Belite-ye'elimite-ternesite binders have been treated as another interesting alternative material for reducing CO₂ emissions. The HeidelbergCement Technology Center (HTC) has worked for several years to develop new and innovative techniques of production approaches for this binder, and have published some patents related to this type of cement in recent years.

1130

Ternesite ($C_5S_2\hat{S}$) was first found in Germany as a natural mineral in the 1990s. It is also found in 1131 the crust covering the areas of Portland cement kilns where the temperature is lower than 1250 °C. 1132 1133 The advantages of ternesite-containing clinkers are quite clear due to the lower clinkering 1134 temperature. For a long time, this phase has been regarded as a non-hydraulic material, until it 1135 was recently found to be reactive with aluminum hydroxide. According to Ben Haha et al. [377], 1136 aluminum hydroxide can be used to activate ternesite to form ettringite, strätlingite and C-S-H in 1137 different proportions, depending on the reactivity and reaction degree. Work by Montes et al. [378], 1138 focusing on how other calcium aluminates activate the hydration of ternesite, has also been published recently. Synthetic C₃A, C₁₂A₇, CA and C₄A₃ \hat{S} (ye'elimite) phases were blended with 1139 1140 ternesite separately, then the hydration reactions of the blends were studied through various 1141 techniques. Ternesite was activated in all the blends with aluminates, with descending effectiveness order $C_{12}A_7 \approx CA > C_3A >>> C_4A_3\hat{S}$. Also, the presence of ternesite changes the 1142 hydration products of these aluminates. However, ternesite was less consumed in the samples 1143 1144 mixed with ye'elimite due to the sulfate common ion effect. Even though in this study some 1145 calorimetric evidence of an activating effect was recorded, ternesite could not be regarded as 1146 having been activated by ye'elimite as no strätlingite was detected. The characteristics of ternesite 1147 as a component of belite-ye'elimite (sulfobelitic) binders was later discussed by Blanco and 1148 Carmona [379] who noted that ye'elimite and ternesite can co-exist in the CaO-SiO₂-Al₂O₃-1149 CaSO₄ system.

1150

A single-stage process to produce ternesite-containing clinkers (belite and ternesite-rich calcium
sulphoaluminate) has been proposed by Hanein et al. [380], based on some important new work

in determining and defining the high-temperature thermodynamics of sulfur-containing clinkerization processes [381]. The clinker was obtained in a pilot plant where the partial pressure of O_2 and SO_2 was controlled in the kiln. The target operation temperature was set up to 1260 °C in order to make the process to reach steady state rapidly, corresponding with their thermodynamic calculations that the upper limit stability temperature for ternesite is ~1290 °C. The results also clearly confirmed that ternesite can be synthesized in a dry atmosphere.

1159

1160 **5. Conclusions and Final Remarks**

1161 The development and use of low-carbon binders as an alternative to Portland cement-based 1162 materials, aiming to reduce the carbon footprint associated with construction and other 1163 applications, has made notable progress in recent years. In summary:

1164

1165 a) Alkali-activated binders are very important and high-potential alternative materials, which 1166 are now deployed on a commercial scale in several nations in the world. Recently the 1167 development of understanding on the rheological behavior, setting properties and structural 1168 characterization of alkali-activated binders has advanced rapidly. Progress in formulation of 1169 one-part alkali-activated binders has further approached large-scale production and application. However, the development and optimization of mix designs based on different 1170 1171 raw materials and activators has not yet been systematically understood. Durability 1172 performance appears very good in most areas but needs more detailed work on test method 1173 validation and standardization. Environmental assessment of these materials should also be 1174 improved.

b) Carbonatable binders, regarding as a new approach to address concerns over CO₂ emissions,
still are in a development route. The technology has been advanced recently, especially in the
understanding of accelerating and controlling the carbonation hardening process. The
limitations for these binders in application are also becoming clear, for instance the CO₂-rich
atmospheres required for curing, and the pH reduction that means that use in reinforced

elements will be challenging. However, these binders which may offer very high CO₂ savings
if a circular CO₂ economy develops [360] still deserve serious attention as alternative lowcarbon materials.

Belite-ye'elimite binders: this is a relatively new approach to produce alternative 1183 c) 1184 cementitious materials compared to the conventional CSA cements, targeting a high belite 1185 content in the clinkers. Although belite-ye'elimite-based binders are still under development and have not reached the full scale-up stage, the clinkering process, understanding of 1186 1187 hydration, and the formulation of binders has developed greatly, not only in the scientific 1188 community but also in the cement industry. Good mechanical strength was obtained by hydrating this type of binder. The control of the rheological behavior and setting time have 1189 1190 also been investigated. However, this binder system and technology is not yet commercialized 1191 or standardized. The clinkering process, which depends on different raw materials, should be 1192 optimized for large-scale production.

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