# **Fly ash-Ca(OH)<sup>2</sup> reactivity in hypersaline NaCl and CaCl<sup>2</sup> brines**

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#### 1

### 2 **ABSTRACT**

- 3 The disposal of highly concentrated brines from coal power generation can be effectively
- 4 accomplished by physical solidification and chemical stabilization (S&S) processes that utilize fly
- 5 ashes as a reactant. Herein, pozzolanic fly ashes are typically combined with calcium-based
- 6 additives to achieve S&S. While the reactions of fly ash-(cement)-water systems have been
- 7 extensively studied, the reactivity of fly ashes in hypersaline brines (ionic strength, *I<sup>m</sup>* > 1 mol/L)
- 8 is comparatively less understood. Therefore, the interactions of a Class C (Ca-rich) and a Class F
- 9 (Ca-poor) fly ash were examined in the presence of Ca(OH)2, and their thermodynamic phase-
- 10 equilibria modeled on contact with NaCl or CaCl<sup>2</sup> brines for 0 ≤ *I<sup>m</sup>* ≤ 7.5 mol/L. At low ionic
- 11 strengths (< 0.3 mol/L), reactivity and stable phase-assemblages remain effectively unaltered.
- 12 But, at high(er) ionic strengths (> 0.5 mol/L), the phase assemblage shows a particular
- 13 abundance of Cl-AFm compounds (i.e., Kuzel's and Friedel's salt). Although Kuzel's and Friedel's
- 14 salt formation enhances Class F fly ash reactions in both NaCl and CaCl<sub>2</sub> brines; NaCl brines
- 15 compromise Class C fly ash reactivity substantially, while CaCl<sub>2</sub> results in the reactivity
- 16 remaining essentially unchanged. Thermodynamic modeling that accounts for the fractional,
- 17 and non-congruent dissolution of the fly ashes indicates that their differences in reaction
- 18 behavior are provoked by differences in the prevalent pore solution pH, which affects phase
- 19 stability. The outcomes offer new insights for matching fly ashes, Ca-additives, and brines, and
- 20 accounting for, and controlling fly ash-brine interactions as relevant to optimizing physical
- 21 solidification and chemical stabilization applications.
- 22
- 23 **Keywords:** Fly ash; Brine encapsulation; Solidification/stabilization; Thermodynamic modeling
- 24

#### 25 **INTRODUCTION AND BACKGROUND**

- 26 As industrial sites face increasing challenges in managing wastewater, effective technologies for
- 27 managing such reject streams are needed. A primary concern for many sites is finding a way to
- 28 eliminate the discharge of wastewater as they work towards meeting zero-liquid discharge
- 29 (ZLD) guidelines. One such wastewater is Flue Gas Desulfurization Wastewater (FGDWW) from
- 30 coal-fired electricity generation which contains high levels of dissolved salts (0-5 mol/L of total
- <span id="page-0-0"></span>31 dissolved solids), $1,2$  including alkali cations, halide anions, and heavy metals. A potential
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 solution to safely dispose of such wastewaters, as well as other chemically similar wastewaters (i.e, coal ash landfill leachate, contaminated groundwater, and produced water) is to 34 encapsulate such wastewater in a solid matrix. $3,4$  To reduce disposal volumes, wastewaters are often concentrated resulting in a hypersaline solution (i.e., referred to as a brine hereafter due to their high salt concentrations). In the encapsulation or S&S process, the solution reacts with a binder to form a cemented/cohesive monolith. This monolith retains the dissolved salts 38 originally present in the brine.<sup>3,4</sup> Such salts may be entrapped by precipitation as insoluble 39 species,<sup>5</sup> by sorption on the surfaces of hydrated phases,<sup>6</sup> and/or by incorporation into the 40 Iattice (structure) of the hydrated phases.<sup>7,8</sup> Furthermore, the solution may also be entrapped within the pore spaces of the solidified mass. Many cementitious binders have been examined 42 to obtain a solid that can be safely landfilled. $9,10$  In particular, there is considerable interest in the use of fly ashes; i.e., a co-product of coal combustion on account their ability to motivate so-called pozzolanic reactions upon reaction with water contained in the brine, in the presence of calcium-based additives including quick lime, hydrated lime, and/or ordinary Portland 46 cement (OPC).<sup>3,4,9</sup> Recently, Song et al. suggested that typical ASTM classifications of fly ashes 47 that are used to assess their use in construction applications (e.g., using ASTM C618) may have 48 led to the exclusion of fly ashes whose reactivity was underestimated.<sup>11</sup> Lower-performance requirements (e.g., in terms of strength) of encapsulation materials would imply that a wide range of fly ashes, including current production and harvested and reclaimed ("historical production") fly ashes, may be suitable for use in S&S applications. This is important as the diminishing availability of high-quality fly ashes would not, in the short- to medium-term, affect the use/availability of fly ashes for brine encapsulation. This is significant because fly ashes, e.g., as compared to cement-based materials, provide a lower cost solution, and lower-carbon footprint solution, for S&S applications, wherein the use of cement-based materials is desirable 56 as (minor) additives, but not as the dominant component of the binder composition.<sup>12,13</sup> 57 The performance of a binder for encapsulation applications is ascertained in terms of its ability to immobilize constituents that could potentially contaminate groundwater. The brine's composition (i.e., in the case of coal-power based electricity production) is often dominated by 61 NaCl, CaCl<sub>2</sub> and/or CaSO<sub>4</sub>; although solubility considerations ensure that, typically, the

- 62 abundance of  $SO_4^2$  is often far inferior to Cl anions. The presence of Cl at high concentrations
- 63 can strongly affect the phase assemblage that forms in cementitious systems. For example, Cl-
- 64 species can physisorb onto the C-S-H phases, $14-16$  and they can also occupy the interlayer
- 65 positions in AFm phases (i.e. phases of the general formula  $[Ca_2(A),Fe)(OH)_6] \cdot X \cdot xH_2O$  with  $X =$
- 66 OH,  $SO_4^2$ ,  $CO_3^2$  and/or Cl bto form hydrocalumite (Ca<sub>2</sub>Al(OH)<sub>12</sub>[Cl, CO<sub>3</sub>,OH]<sub>2</sub>·4H<sub>2</sub>O), Kuzel's salt
- 67 (Ca<sub>2</sub>Al(OH)<sub>6</sub>[Cl, SO<sub>4</sub>, OH]·2H<sub>2</sub>O) or Friedel's salt (Ca<sub>2</sub>Al(OH)<sub>6</sub>[Cl, OH]·2H<sub>2</sub>O) ["Cl-AFm phases"
- 68 hereafter].<sup>17-21</sup> On the other hand, in lime-enriched systems, CI species can additionally react
- 69 with portlandite (Ca(OH)<sub>2</sub>) to form calcium oxychloride compounds.<sup>22-24</sup>
- 70
- 71 Thermodynamic modeling based on the minimization of Gibbs free energy is a powerful tool for
- 72 assessing stable phase equilibria and equilibrium compositions of the solid-, liquid (and gas)-
- 73 phases as relevant to encapsulation applications. But, the current implementations of such
- 74 modeling often rely on assumptions including: congruent dissolution of the fly ash,<sup>9</sup> and most
- 75 often, an inability to incorporate the actual degree of fly ash reaction which could be
- 76 particularly affected in hypersaline environments that feature a reduced water activity.<sup>25-30</sup> For
- example, the dissolution rates of crystalline and amorphous materials could either enhance or
- 78 decrease in hypersaline environments in relation to their composition, and concentration.<sup>31-36</sup>
- 79 For example, seawater has been suggested to enhance the fly ash's pozzolanic reactions.<sup>37</sup>
- Since changes in reactivity alter not only the kinetics of reactions, but also the phase
- 81 assemblage that forms, it is important to assess these aspects due to the obvious implications
- 82 on contaminant retention by the solidified waste forms. With this focus in mind, this study
- 83 examines the effect of NaCl or CaCl<sub>2</sub>-based hypersaline brines on the pozzolanic reactions of Ca-
- rich (Class C) and Ca-poor (Class F) fly ashes. Special focus is paid to model, compare, and
- ascertain discrepancies between the results of thermodynamic modeling and experimental
- assessments of stable phase assemblages for mature systems (i.e., aged greater than 10 days at
- 87 50 °C). The outcomes provide new insights to understand how the composition of the brine affects the stable phase assemblage that forms thereby offering guidance to screen and select
- the most compatible combinations of fly ashes, and brines to ensure optimal wastewater
- encapsulation (i.e., physical solidification and chemical stabilization: S&S).
- 

## **MATERIAL AND METHODS**

## *Raw material characterization*

- A Class C and Class F fly ash were considered herein. Their bulk oxide compositions, which
- encompass both crystalline and amorphous compounds as determined using X-ray fluorescence
- (XRF), is shown in [Table 1.](#page-3-0) The crystalline phases present, and the quantity of the amorphous
- phase present, were quantified using Quantitative X-Ray Diffraction (QXRD) and Rietveld
- 98 Fefinement.<sup>38,39</sup> Zincite (ZnO, purity: 99.999%) was used as an internal standard at a mass
- 99 Ioading of 10 mass % in the Rietveld refinement which was carried out using Profex.<sup>40</sup> The
- quantities and types of crystalline phases present is shown in [Table 2.](#page-3-1) The average composition
- 101 of the amorphous phase was calculated by subtracting the quantity of crystalline phases from
- 102 the bulk XRF composition, and is shown i[n Table 3.](#page-4-0)
- 
- Both fly ashes were subjected to 50 min of dissolution in dilute conditions (solid-to-liquid ratio,
- 105 mass basis, s/l = 1:1000) in deionized water (DIW, Milli-Q: 18.2 M $\Omega$ /cm) at 50 °C under agitation
- in high density polyethylene (HDPE) containers. The residual solids were retrieved from the
- 107 container after 50 minutes of dissolution by vacuum-filtration through a 11 µm paper filter,
- 108 dried at 60 °C, and analyzed using XRD to determine which crystalline phases had dissolved.
- Based on this analysis, the crystalline phases were designated as: not soluble (NS), partially
- soluble (PS), or soluble (S). These are designations not so much of "solubility" per se, but rather
- describe the extent to which a compound may dissolve over the period of dissolution, i.e.,
- either remaining undissolved, being partially dissolved, or fully dissolved (see [Table 2\)](#page-3-1).
- 
- 114 Simulated brines were prepared by dissolving NaCl (99% purity) or CaCl<sub>2</sub>·6H<sub>2</sub>O (99% purity) in
- deionized water (DIW) at room temperature to obtain solutions with salt concentrations of 0,
- 116 0.01, 0.1, 0.5, 1, 2 (CaCl<sub>2</sub> and NaCl), 2.5 (CaCl<sub>2</sub> only) and 5 (NaCl only) mol/L. These
- concentrations translate to ionic strengths (*Im*; mol/L) of 0.01, 0.1, 0.4, 1, 2, and 5 mol/L for the
- 118 NaCl solution, and 0.03, 0.3, 1.5, 3, 6, 7.5 mol/L for the CaCl<sub>2</sub> solutions. Note that, for CaCl<sub>2</sub>, the
- highest concentration used was 2.5 mol/L (*I<sup>m</sup>* = 7.5 mol/L), as higher concentrations do not
- 120 allow the formulation of a fluid cementitious paste at the s/l studied. Cementitious
- 121 formulations were composed by combining 55 mass % fly ash, 10 mass % portlandite (Ca(OH)<sub>2</sub>;
- 122 purity > 95%) and 35 mass % brine. The formulation was mixed for 45 s at 270 rpm and 1 min at
- 123 480 rpm at room temperature using a high-shear immersion mixer. The resulting pastes were
- 124 rapidly poured into glass ampoules, hermetically sealed, and placed into a TamAir isothermal
- 125 calorimeter at 50 °C for analysis of their heat release behavior. Heat flow and cumulative heat
- 126 release were measured to assess the rate and the extent of fly ash reaction. A temperature of 127 50 °C was chosen as it is known to be sufficient to accelerate fly ash reactions<sup>#4,41,43</sup> In general,
- 128 the heat release showed little if any change ( $d\dot{Q}/dt$  < 0.1 mW/g/s) after 10 days (see Figure S1),
- 129 following which the ampoules were extracted, and the samples crushed and immersed in
- 130 isopropanol (IPA), to induce solvent-exchange, for 1 week to cease reactions.<sup>44</sup> Thereafter, the
- 131 samples were dried under vacuum for another week, following which they were crushed, milled
- 132 using an agate mortar and pestle and then sieved through a 300 µm sieve prior to any
- 133 additional characterization.
- 134

<span id="page-3-0"></span>

<span id="page-3-1"></span>**Table 2.** The crystalline phases present in the fly ashes as determined using QXRD (mass %). Here,  $^{NS}$  indicates the phases that were determined to be not-soluble,  $^{PS}$  indicate the phases that are partially soluble with the percentage dissolved recorded in brackets, and <sup>s</sup> indicates the phases that are completely dissolved after 50 minutes of fly ash dissolution in dilute  $conditions (s/l = 1:1000)$ 

| 50.15151515777<br>-------     |                   |                     |
|-------------------------------|-------------------|---------------------|
|                               | <b>Class C</b>    | <b>Class F</b>      |
| $Quartz - SiO2$               | 5.0 <sup>NS</sup> | 6.0 <sup>NS</sup>   |
| Periclase - MgO               | $3.3$ NS          | $\qquad \qquad$     |
| Free lime - CaO               | 1.4 <sup>5</sup>  | 0.4 <sup>5</sup>    |
| Anhydrite – CaSO <sub>4</sub> | 17S               | 0.6 $^{\mathrm{s}}$ |
|                               |                   |                     |

<span id="page-3-2"></span> $*$  Assuming an apparent activation energy of the pozzolanic fly ash reactions to be on the order of 50 kJ/mol<sup>41,42</sup> implies that curing at 50 °C for 10 days equates to around 50 days of curing at ambient conditions (25 °C).



<span id="page-4-0"></span>

137

138 *Material characterization*

139 *Thermogravimetric analysis*: Thermogravimetric analysis (TGA) was performed using a Perkin

140 Elmer STA 6000 under a flow of ultra-high purity (UHP)  $N_2$  in aluminum oxide crucibles. A

141 heating rate of 10 °C min<sup>-1</sup> was used between 35 and 950 °C, after 5 min equilibration at 35 °C.

142 The mass loss (TG) and the derivative mass loss (DTG) were used to characterize and quantify

143 several hydrate phases, including portlandite, CI-AFm, etc.<sup>46</sup> The analysis was performed on

144 powdered samples that had undergone solvent exchange using IPA. While this prevents

145 quantification of the free water content, it does not affect the remainder of the solid phase-

- 146 specific mass determinations.<sup>46</sup>
- 147

148 Quantitative information regarding the Cl-AFm phases, i.e., Friedel's salt and Kuzel's salt, can be

149 obtained using TGA.<sup>18,46,47</sup> These hydrates display two significant mass losses in the

150 temperature range of ~100-180 °C and ~180-450 °C. The first mass loss is attributed to

151 interlayer water removal, while the second results from removal of the main-layer water.<sup>46</sup> In

152 our system, the first mass loss occurred coincidentally with that of water removal from C-S-H,

- 153 ettringite and/or monosulfoaluminate. Consequently, this peak could not be used for Cl-AFm
- 154 quantification. The second mass loss, however, occurs in a temperature range where no water
- 155 removal from the other phases is observed. But, since the mass loss of Friedel's salt and Kuzel's
- 156 are superimposed with each other, independent quantifications of each phase was not possible
- 157 herein. Nevertheless, comparing the mass loss measured between ≈250-to-430 °C helps to
- 158 assess the total quantity of Cl-AFm phases formed in each system.
- 159

160 *Infrared spectroscopy:* Solid-state attenuated total reflection Fourier-transform infrared

161 spectroscopy (ATR-FTIR) was performed using a Spectrum Two FT-IR Spectrometer (Perkin 162 Elmer). The powdered samples were pressed using around 90 N of force onto a diamond/ZnSe

- 163 composite crystal to ensure good contact and generate total internal reflection. The spectra 164 reported herein were obtained by averaging 4 scans over the wavenumber range of 4000-to-
- 165  $400 \text{ cm}^{-1}$  at a resolution of 1 cm<sup>-1</sup>.
- 166

167 *X-Ray diffraction:* XRD analysis was performed using a PANalytical X'PertPro diffractometer (θ-θ 168 configuration, Cu-Kα radiation,  $\alpha$  = 1.54 Å) on powdered samples. The scans were acquired 169 using a rotating sample stage between 5° and 70° with a step size of 0.02° using a X'Celerator 2

170 detector. In general, powdered samples were placed in the sample holder and their surfaces 171 gently textured to minimize the potential for preferred orientation related errors.

172

173 *Thermodynamic modeling*: Thermodynamic modeling was carried out using GEM‐Selektor v.3.6 174 (GEMS)<sup>48,49</sup> which incorporates the slop98.dat and Cemdata18 thermodynamic databases.<sup>50–54</sup>

175 To represent the non-ideality of the solutions, the activity coefficients were calculated using the

176 Truesdell-Jones extension to the Debye-Hückel equation that is applicable for  $I_m \approx 2$  mol/L:<sup>55</sup>

$$
\log_{10} \gamma_i = \frac{-A_{\gamma} z_i^2 \sqrt{I}}{1 + aB_{\gamma} \sqrt{I}} + b_{\gamma} I + \log_{10} \frac{X_{jw}}{X_w}
$$
 Equation (1)

177 where,  $\gamma_i$  is the activity coefficient and  $z_i$  the charge of the *i*<sup>th</sup> aqueous species,  $A_\gamma$  and  $B_\gamma$  are 178 temperature and pressure dependent coefficients,  $X_{iw}$  is the molar quantity of water,  $X_w$  is the 179 total molar amount of the aqueous phase, and  $I$  is the molal ionic strength. A common ion size 180 parameter ( $a = 3.72$  Å) and a short-range interaction parameter ( $b_v = 0.64$  kg/mol) were 181 used, treating NaCl as the background electrolyte.<sup>55,56</sup> Water activity is generally defined as the 182 deviation of the chemical potential of liquid water from its pure state (where water activity = 1) 183 to a 'non-pure' state (where water activity < 1) due to the presence of a solute. To a first 184 approximation, water activity in GEMS is calculated from the osmotic coefficient when the 185 extended Debye-Hückel activity model for aqueous species is used, following Helgeson et 186 al.:<sup>48,55</sup>

187

$$
\ln a_{water} = -\frac{\phi m_{\Sigma}}{55.508435}
$$
 Equation (2)

188

189 where,  $m_{\overline{y}}$  is the sum of all species molalities and  $\phi$  is the osmotic coefficient, both of which 190 are calculated within GEMS as following Helgeson et al.<sup>55</sup>

191

 The system modelled (55 g of FA, 10 g of portlandite, and 35 g of brine) is equivalent to that studied experimentally. Here, brine concentrations of 0, 0.01, 0.1, 0.5, 1, and 2 (NaCl only) 194 mol/L and up to 1 mol/L of CaCl<sub>2</sub> were considered to conform to the limits of applicability of Equation (1). Of course, as water is consumed over the course of reaction, the ionic strength exceeds the 2 mol/L limit. Although GEMS offers an option to use Pitzer's model which can 197 accurately describe ion activities in highly-concentrated solutions  $(I_m < 6 \text{ mol/L})$ , <sup>57</sup> this module was not used due to a lack of ion-specific parameters to consider all the species that are present herein. Nonetheless, it should be noted that previous comparisons of ion activities determined using Pitzer's equations and the parameterization of the Truesdell-Jones ion- activity model for simple Ca, Na, and Cl containing systems yield results within ±30% for *I<sup>m</sup>* <  $\frac{4 \text{ mol/L}}{9.57,58}$  Given that  $I_m < 3.5$  mol/L for all solution compositions considered herein, while the absolute quantitative accuracy of the predictions of phase equilibria would degrade for *I<sup>m</sup>* > 2 mol/L, qualitative indicators (e.g., the types, and relative abundance of phases formed) would nevertheless continue to be relevant to the systems studied. As such, some uncertainty is expected in the quantitative (although not qualitative) outcomes of the simulations performed 207 at higher ionic strength.<sup>48,58</sup> In general, portlandite is considered to be fully consumed over the course of reactions, while the fly ashes are considered to show fractional reactivity based on: (a) the near-inert nature of the insoluble crystalline phases (e.g., quartz), (b) incomplete reaction of some partially soluble crystalline phases, (c) complete consumption of the highly-211 reactive crystalline phases (e.g., CaSO<sub>4</sub>), and (d) the congruent dissolution of the amorphous phase in accordance with its average composition [\(Table 3\)](#page-4-0).

## **RESULTS AND DISCUSSION**

 *Hydrate formation:* The formation of crystalline and amorphous reaction products was 216 examined in the Class C and Class F fly ashes after 10 days of reaction at 50 °C. Among the crystalline phases, ettringite (PDF #04-013-3691), monosulfoaluminate (AMCSD #0014757), a magnesium-aluminum hydrotalcite-like phase (referred to as 'hydrotalcite' hereafter, PDF #00- 014-0525), katoite (AMCSD #0006980), Kuzel's salt (PDF #00-019-0203), Friedel's salt (AMCSD #0014832), and strätlingite (AMCSD #0006404) were observed. Expectedly, the phase assemblages formed differ considerably across Class C and Class F fly ashes (see [Figure 1\)](#page-7-0). For example, upon reaction in DI-water, the Class F fly ash system shows the presence of ettringite (identified by its main peak at 9.08° 2θ), which in time, transforms into monosulfoaluminate (9.93° 2θ). In contrast, the Class C fly ash shows the presence of monosulfoaluminate, katoite (32.61° 2θ), hydrotalcite (11.63° 2θ) and traces of strätlingite (7.00° 2θ). The introduction of low concentrations of NaCl or CaCl<sup>2</sup> (*I<sup>m</sup>* < 0.3 mol/L) does not have significant effects on the phase assemblages formed across both classes of fly ashes. However, higher abundances lead to 228 substantial modifications, as observed elsewhere.<sup>3,59</sup> More specifically, for the Class C fly ash in NaCl system, ionic strengths above 0.1 mol/L inhibit monosulfoaluminate, katoite, strätlingite and hydrotalcite formation (see [Figure 1a](#page-7-0)). These phases are replaced by Cl-AFm phases. 231 Friedel's salt forms for ionic strengths greater than 0.5 mol/L in NaCl system. For CaCl<sub>2</sub>, ionic strengths above 0.3 mol/L similarly inhibit monosulfoaluminate, katoite and hydrotalcite formation [\(Figure 1b](#page-7-0)). The co-formation/-persistence of Kuzel's salt and Friedel's salt is observed in the ionic strength range of 1.5-3 mol/L, while Friedel's salt is the main hydrate 235 formed, alongside a minority of ettringite for ionic strengths greater than 3 mol/L (CaCl<sub>2</sub>).

236 Although no crystalline C-S-H phases were observed, the presence of amorphous C-S-H (i.e.,

237 generally with Ca/Si < 1.45)<sup>60–62</sup> is expected because of the pozzolanic reaction between the fly 238 ashes and portlandite.

239



<span id="page-7-0"></span>Ms = monosulfoaluminate, FrS = Friedel's salt, KzS = Kuzel's salt, HT = hydrotalcite, St = strätlingite and K = katoite  $(Ca_3Al_2(SiO_4)_{3-x}(OH)_{4x}$  with x = 1.5-3). For the unreacted Class C fly ash, the peaks at  $31.25^\circ$  and  $32.6^\circ$  20 are attributed to CaO and MgCO<sub>3</sub>, respectively.



241 For the Class F fly ash in NaCl system, high ionic strengths (*I<sup>m</sup>* > 0.1 mol/L) suppress

 monosulfoaluminate formation, while the formation of ettringite too is noted to be suppressed at ionic strengths of 5 mol/L. The formation of Friedel's salt (11.20° 2θ) is favored as sulfate is 244 replaced by chloride as the interstitial anion within the AFm interlayer. In CaCl<sub>2</sub> system, ionic 245 strengths above 0.03 mol/L prevent monosulfoaluminate formation, and ettringite's formation is inhibited at ionic strengths above 3 mol/L. Kuzel's salt formation (9.89° 2θ) is observed between 0.03 mol/L < *I<sup>m</sup>* < 0.3 mol/L, while only Friedel's salt formation is observed at ionic 248 strengths higher than 0.3 mol/L due to higher quantity of solubilized Ca<sup>2+</sup> and Cl<sup>-</sup> available in solution.

250

251 ATR-FTIR was used to ascertain the presence of amorphous hydrated phases. The presence of 252 amorphous C-S-H was detected across both fly ash compositions after 10 days (peak between 253 950 and 965 cm<sup>-1</sup>, [Figure 2a](#page-8-0) and Figure S2). The position and shape of the peak is similar to that 254 of Al-containing C-S-H as previously identified by Kapeluszna, et *al*.<sup>61</sup> While it is possible that the 255 amorphous C-S-H formed may transform into variants of increasing crystallinity – in time, 256 particularly at somewhat elevated curing temperatures – no crystalline C-S-H phases were 257 detected over the course of the examinations carried out herein. In general, the C-S-H formed 258 in the presence of NaCl brines shows a peak position and a peak intensity that is unaffected by 259 NaCl concentration [\(Figure 2b](#page-8-0) and c). This suggests, to the first order, that there is no structural 260 incorporation of Na<sup>+</sup> or Cl<sup>-</sup> into the C-S-H, and that, rather, Na<sup>+</sup> and Cl<sup>-</sup> interact with the C-S-H 261 via physical (sorption) processes. $14-16$  In contrast, the C-S-H peak intensity decreases with 262 increasing CaCl<sub>2</sub> concentrations, while the peak position shifts to higher wavenumbers (for both 263 Class C and Class F fly ashes), with the emergence of a second peak around 1015 cm<sup>-1</sup> (Class F 264 fly ash only). Both Ca/Si and Al/Si ratios of the C-(A)-S-H phases can influence the infrared peak 265 position. In particular, an increasing Ca/Si ratio and decreasing Al/Si ratio of C-(A)-S-H have both

- 266 been shown to shift the peak position to higher wavenumbers.<sup>61</sup> Both of these compositional 267 alterations can also result in the formation of a second peak around 1000 cm<sup>-1 61</sup> As such, first, 268 it is indicated that the shift in the peak position in the CaCl<sub>2</sub> system is linked to enhanced Cl-269 AFm phase formation that consumes mobile Al present in the system; resulting in a decrease of 270 the Al/Si ratio. Second, an increasing CaCl<sub>2</sub> concentration provides additional calcium, which 271 likely results in an increase of the Ca/Si ratio. The indications suggest that both an increase in 272 Ca/Si and decrease in Al/Si ratios play a role in altering the peak position in the CaCl<sub>2</sub>-brine 273 systems. Additionally, increasing CI-AFm formation in the CaCl<sub>2</sub> system likely consumes the C-274 (A)-S-H, as attested by the peak intensity decrease that is observed experimentally. Herein, Cl-
- 275 AFm formation results in the decomposition of C-(A)-S-H into portlandite and amorphous silica, 276 the former of which serves as a reactant in CI-AFm formation.
- 277



<span id="page-8-0"></span>**Figure 2.** IR spectra of **(a)** unreacted and reacted (10 days hydration at 50 °C in DIW) Class C and Class F fly ash. **(b)** A zoomed-in view of the peak attributed to C-S-H for the Class C and **(c)** the Class F fly ash reacted in NaCl or CaCl<sub>2</sub> brines (the full FTIR spectra are shown in Figure S2).

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279 *Heat release and Cl-AFm formation:* The degree of fly ash reaction is difficult to establish 280 quantitatively. While several methods exist to assess the degree of reaction of fly ash, including 281 selective dissolution,  $63-65$  SEM-image analysis,  $26,64,66$  etc., these methods present considerable 282 uncertainty and/or are time-consuming. $64$  First, it is instructive to assess how the cumulative 283 heat release measured until 10 days is affected by brine composition (see [Figure 3a](#page-9-0)). In general, 284 a substantial effect of NaCl and CaCl<sub>2</sub> on fly ash reactivity is noted. The effect of both salts is 285 minimal for ionic strengths below 0.3 mol/L across both fly ash types, which is consistent with 286 the XRD data that showed little-to-no changes in phase assemblages in this ionic strength 287 range. In contrast, for ionic strengths in excess of 0.3 mol/L, NaCl strongly suppresses the 288 reactivity of Class C fly ash, while the reactivity of Class F fly is even somewhat enhanced; the 289 latter, particularly for ionic strength in excess of 1 mol/L. On the other hand, while CaCl<sub>2</sub> slightly 290 elevates Class F fly ash reactivity, the Class C fly ash's reactivity is initially suppressed and 291 recovers for ionic strengths only in excess of 1.5 mol/L. This may indicate that reductions in 292 water activity resulting from the presence of salts prominently affect Ca-based compounds 293 which are more prevalent in the Class C fly ash as compared to Class F variants. However, the 294 non-monotonous trend in heat release wherein the reactivity of Class C (and to a smaller

 extent, Class F) fly ash appears to elevate for ionic strengths greater than 1 mol/L (Class C) and greater than 0.3 mol/L (Class F) remains unclear currently.

 The mass loss from the Cl-AFm phases derived from thermal analysis (i.e., as measured between 250-and-430 °C, see Figure S3) shows that, expectedly, the amount of Cl-AFm formation increases with the Cl-content of the brine (see [Figure 3b](#page-9-0)). The formation of Cl-AFm phases is particularly pronounced for ionic strengths greater than 0.1 mol/L. Across both fly 302 ashes, a higher CI-AFm content is observed in the case of CaCI<sub>2</sub>-based brines, as compared to their NaCl counterparts on account of the provision of mobile Ca-species in the former. Interestingly, when the mass loss from thermal analysis corresponding to the Cl-AFm compounds is mapped as a function of the cumulative heat release, a "V-shape" – like the heat release behavior (se[e Figure 3a](#page-9-0)) – is observed in the case of the Class C fly ash [\(Figure 3c](#page-9-0)), while a linear trend emerges in the case of the Class F fly ash [\(Figure 3d](#page-9-0)). The correlation between the heat release and Cl-AFm phase abundance indicates that, broadly speaking, the formation of (one or more) Cl-AFm phases is a dominant reaction in fly ash-brine systems. Cl-AFm formation is beneficial to Class F fly ash reactivity, as it occurs in addition (i.e., not at the expense of) to ettringite formation. In contrast, Cl-AFm formation is detrimental to Class C fly ash reactivity, since it occurs at the expense of strätlingite, monosulfoaluminate and katoite; which are 313 observed to be destabilized in the NaCl  $(I_m > 0.5 \text{ mol/L})$  and CaCl<sub>2</sub>  $(I_m > 0.3 \text{ mol/L})$  containing systems.



<span id="page-9-0"></span>**Figure 3. (a)** Cumulative heat release after 10 days of hydration at 50 °C as a function of the ionic strength. **(b)** Mass loss from Cl-AFm phases – measured between ~250 and ~430 °C – as a function of the ionic strength. The cumulative heat release after 10 days as a function of the mass loss from Cl-AFm phases for the: **(c)** Class C fly ash, and **(d)** Class F fly ash.

*Thermodynamic modeling:* Thermodynamic modeling based on the minimization of Gibbs free

 energies was used to estimate the stable phase assemblages that exist as a function of fly ash reactivity. Herein, the heterogeneity of the fly ash is taken into account, unlike most studies

that considered the congruent dissolution of fly ash content based on its average composition

321 as determined from XRF.<sup>9</sup> Thus, the crystalline phases that feature widely varying reactivities

are programmed to dissolve following the observations of the short-term dissolution analysis.

Based on this analysis, while quartz for example is assessed to be non-reactive, anhydrite

- 324 (CaSO<sub>4</sub>, i.e., present in both Class C and Class F fly ash) is assumed to rapidly react and be
- 325 consumed. The amorphous phase dissolves congruently in relation to its average composition
- 326 (shown in [Table 3\)](#page-4-0) following a linear dissolution expression from 0 to 100 mass %.
- 327
- 328 A key question here is related to estimating the degree of fly ash reaction following 10 days of
- 329 hydration at 50 °C. The degree of fly ash reaction was inferred by analyzing when the ratio of
- 330 masses of a single well-characterized crystalline phase, e.g., portlandite (CH: Ca(OH)<sub>2</sub>) is equal
- 331 to unity; i.e., when the modeled quantity of a given phase is equivalent to its content
- 332 established by experimental, i.e., TGA and/or XRD assessments (e.g., when  $CH_m/CH_e \approx 1$ , where
- 333 the subscripts 'm' and 'e' indicate modeled and experimental assessments). While it does not
- 334 have any implications on the assessed degree of fly ash reaction, it should be noted that,
- 335 expectedly, the quantity of amorphous C-S-H formed could not be determined experimentally
- 336 and was thus considered part of the amorphous compounds as calculated by XRD analysis. For 337 example, [Figure 4\(](#page-10-0)a-b) shows the modelled phase assemblage obtained in DIW for the Class C
- 338 and the Class F fly ash, respectively, as a function of the fly ash degree of reaction. The red
- 
- 339 dashed lines indicate the degree of fly ash reaction for which the condition CH<sub>m</sub>/CH<sub>e</sub>  $\approx$  1 is 340 satisfied for the crystalline phase of interest; herein, portlandite.
- 341



<span id="page-10-0"></span>**Figure 4.** Thermodynamic modelling of **(a)** a Class C and **(b)** a Class F fly ash hydration in DIW in presence of  $Ca(OH)_2$ . Dashed red lines indicate the degree of reaction for which the best agreement is found between simulations and experiments phase quantity (on a mass bases). CH = portlandite, Ms = monosulfoaluminate,  $C_2AH_6$  = katoite, CNASH = sodium substituted calcium aluminum silicate gel, MAH = hydroxyl hydrotalcite. **(c)** Fly ash degree of reaction as a function of ionic strength. **(d)** Comparison between the fly ash degree of reaction modelled and the experimental cumulative heat release.

- 342
- 343 Interestingly, it is noted that while the modeled phase assemblage of the Class F fly ash in DIW
- 344 shows very good agreement with experimental data, comparatively, the Class C fly ash is not as
- 345 well modeled. Particularly, in the case of the modeled Class C fly ash system (in DIW), the
- 346 katoite content is underestimated, and the monosulfoaluminate content overestimated. This
- 347 may suggest that the dissolution of the amorphous phase in the Class C fly ash may be
- 348 incongruent, a not unexpected consideration, given the substantial Ca-content therein. In spite
- of these differences, in general, reasonable agreement in the modeled and experimental phase
- relation is observed for both Class C and Class F fly ashes up to *I<sup>m</sup>* = 2 mol/L (NaCl) and 3 mol/L
- (CaCl<sub>2</sub>). The degree of reaction of the Class F fly ash increases with increasing ionic strength
- regardless of the type of salt considered [\(Figure 4c](#page-10-0)). On the other hand, the degree of reaction
- of the Class C fly ash decreases at *I<sup>m</sup>* = 0.5 mol/L for NaCl and remains unchanged with increasing ionic strength. In contrast, the degree of reaction of the Class C fly ash decreases at
- 355  $I_m = 1.5$  mol/L for CaCl<sub>2</sub> brines and increases thereafter with at higher ionic strengths. These
- observations are consistent with the cumulative heat release, and a good correlation is
- observed between the inferred degree of fly ash reaction and the cumulative heat release (see [Figure 4d](#page-10-0)).
- 

 *Effect of brine concentration on hydrate formation:* An important experimental observation in 361 the case of the NaCl and CaCl<sub>2</sub> brines is that, despite presenting similar initial Cl<sup>-</sup> concentrations, the phase assemblages formed vary considerably, even for a single fly ash across the two brine compositions (see [Figure 1\)](#page-7-0). This implicates the role of the counterion. Interestingly, the GEMS modeling indicates that the pore solution pH observed in the NaCl brine is consistently higher 365 than that of the CaCl<sub>2</sub> brine, for both Class C and Class F fly ashes [\(Figure 5\)](#page-13-0); in some cases, up 366 to 1 pH unit. Of note, while Cl<sup>-</sup> species are consumed from both brines to form the Cl-AFm 367 compounds, little if any Na<sup>+</sup> is taken up into the solids, other than Na<sup>+</sup> that may be sorbed by, 368 or incorporated into C(-N)-A-S-H.<sup>67,68</sup> Expectedly, the consumed Cl<sup>-</sup> ions are compensated by 369 OH<sup>-</sup> that forms, e.g., via the dissociation of water to ensure charge neutrality, resulting in a pH 370 increase as has been observed previously in seawater systems.<sup>59</sup> In contrast, Ca<sup>2+</sup> is consumed from the solution to form C(-N)-A-S-H, AFm phases, and conceivably portlandite; whereby the 372 consumption of Ca<sup>2+</sup> is compensated by an increase in the acidity ([H<sup>+</sup>] abundance) of the solution resulting in a decrease of pH.

375 Since the stability of numerous cementitious phases is sensitively linked to the solution pH (e.g., ettringite and AFm), the solution pH variations begin to explain why different phase

377 assemblages are observed (see Supplementary Information: Figure S4),<sup>69–72</sup> in NaCl and CaCl<sub>2</sub>-

based brines. It is furthermore noted that the simulated pH for each of the brine systems varies

- considerably across different ionic strengths [\(Figure 5\)](#page-13-0). Although the pore solution chemistry
- and pH were not specifically analyzed herein, previous studies have shown favorable
- 381 agreement between measured and modeled pore solution attributes.<sup>30,73,74</sup> As a result of the variation in pH, for Class F fly ashes, the differences in pH alter the quantity of ettringite
- formed, as in Cl-containing solutions, ettringite stability decreases with increasing pH, and no
- ettringite formation is observed above a pH of 12.5 (see Supplementary Information: Figure S4).
- 385 This is, however, only valid for Cl- concentrations greater than 0.1 M as ettringite is known to be
- 386 stable at  $pH > 13$  in DIW and in dilute CI solutions. In the case of the Class C fly ash in the CaCl<sub>2</sub>
- system, the pH remains in a range that allows ettringite formation. In the NaCl system however,
- the pH approaches the limits of stability for Kuzel's and Friedel's salts since for pH > 13.5, both
- Cl-AFms are replaced by katoite (see Supplementary Information: Figure S4) although the
- kinetics of this transformation are likely affected by the particular chemistry of the pore
- solution.

393 All systems modelled here show differences in Cl-binding efficiency (see Supplementary 394 Information: Figure S5), resulting from the differences in the reactant (i.e., brine and fly ash) 395 compositions which dictate the amount and stability of phases formed. Importantly, since the 396 relationship between the chloride concentration of the brine and the amount of Cl-AFm formed 397 is not fully linear [\(Figure 3b](#page-9-0)); the Cl-binding (i.e., ability to retain Cl species within hydrated 398 phases) varies accordingly (see Figure S5). Class F fly ash systems, in particular, cannot retain 399 more than 40% of the mobile CI<sup>-</sup> content, regardless of the type (Na, Ca) of brine. On the other 400 hand, while Class C fly ash shows a reduced efficiency to retain mobile Cl<sup>-</sup> at high ionic strengths 401 it can achieve monotonically increasing Cl-binding with CaCl<sub>2</sub> up to 88%; even for the highest 402 ionic strengths considered. These results highlight the need for incorporating cost-effective 403 additives that offer mobile Ca, and Al (i.e., to favor CI-AFm formation) to maximize Cl-404 encapsulation across a diversity of brine and fly ash compositions*.*

405

406 *Additional considerations:* The results herein indicate that there are two key attributes that are 407 vital for accurate thermodynamic modeling of brine encapsulation/S&S including:

408 • Compositional heterogeneity of the fly ash, and,

409 • The effects of brine composition and concentration (ionic strength) on fly ash reactivity. 410 Foremost, it is inaccurate to consider congruent dissolution of the bulk fly ashes. Rather, it is 411 necessary to consider the distinguished reactivities of the crystalline and amorphous phases 412 and make use of dissolution assessments to establish which phases may be considered readily 413 reactive (i.e., similar or higher reactivity as the fly ash's glass), or not. SEM analysis of Class F fly 414 ashes has previously showed that the amorphous content is relatively homogenous,  $25,26,28,75$ 415 and that the dissolution rate of its main components are similar.<sup>26</sup> This implies that assuming 416 congruent dissolution of its amorphous phase is reasonable. Surprisingly, thermodynamic 417 modeling of a Class C fly ash was unable to replicate experimental phase balances as accurately 418 when its amorphous phase was assumed to react (dissolve) congruently. This may indicate that 419 it is necessary to consider the dissolution rates of its major glass components discretely, or 420 establish an incongruent dissolution expression for the average glass composition.<sup>26,29</sup> It is 421 further observed that the brine composition markedly affects fly ash reactivity. Some changes 422 in fly ash reactivity, particularly of Class C fly ashes, are indeed attributable to water activity 423 reductions. 44,76,77 Other aspects, perhaps including the stability of the different Cl-AFm phases, 424 and the implications of the persisting solution pH and composition, are required to understand 425 the nature and kinetics of phase transformations that may result in Ca- and Na-brine systems as 426 a function of time, ambient temperature, and moisture conditions (i.e., relative humidity, RH). 427 428 Coming back to considerations of temperature, and reaction consideration, Han, et al.<sup>41</sup> studied 429 the pozzolanic reactions of Class F fly ashes in water at various temperatures and suggested an 430 apparent activation energy of fly ash reactions in the range of 43-50 kJ/mol. This is in 431 agreement with the data of Bentz<sup>42</sup> across a variety of systems (35-50 kJ/mol). These values 432 are, however, much lower than that expected for CAS  $[CaO-AI<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub>]$  glass dissolution (70-

- 433 90 kJ/mol for nuclear glasses,<sup>78</sup> 102 kJ/mol for two class F fly ashes in hyperalkaline solutions<sup>79</sup>).
- 434 This may suggest that, in brine encapsulation applications, glass dissolution is transport-
- 435 controlled due to the fast early-precipitation of the hydrated phases. It should be noted that, in
- brine encapsulation, aging occurs over a period of several months. As such, the systems
- considered herein may still be somewhat "immature" and may be expected to show ongoing
- evolutions in phase balances, in time. Unsurprisingly, despite the presence of sufficient
- Ca(OH)2, Class F fly ash reactivity remains low (<15 mass % after 10 days at 50 °C). In contrast,
- Class C fly ash reacts more substantially (up to 50 mass % after 10 days at 50 °C), but its
- reactivity is strongly inhibited at high-NaCl concentrations likely due to water activity
- 442 considerations although much less so in CaCl<sub>2</sub>-based brines. It should also be noted that while
- 443 the GEMS simulations indicate that the pozzolanic reaction of Class C and Class F fly ashes with
- portlandite results in the formation of C-(N)-A-S-H compounds, this may not be the main driving reaction. In fact, both our isothermal calorimetry and TGA data suggest that the formation of
- Cl-AFm phases is the major contributor to Cl-uptake (and chemical reaction) in these systems.
- 



<span id="page-13-0"></span>**Figure 5.** A comparison of the (pore) solution pH after 10 days of hydration at 50 °C (i.e. for the degree of fly ash reaction shown in [Figure 4c](#page-10-0)) as a function of the solvent chemistry (DIW, CaCl<sub>2</sub> or NaCl) for Class C and Class F fly ash systems as calculated by GEMS modeling.

Nevertheless, taken together, the approach shown here demonstrates how GEMS-type

- geochemical simulations can be used to rapidly design and screen formulations for S&S applications, with due considerations of chemical attributes, e.g., pH, brine composition, fly ash
- composition, etc. and the implications on wastewater, and ion-encapsulation consumption. This
- is important to identify the optimal combinations of brine and fly ash, and to also assess if
- additives may be needed, e.g., cement, lime, etc. to enhance S&S performance. This work also
- shows a means to assess the differences in pore solution pH that may result in different
- formulations, since it is sought to maintain a pH > 13 to ensure that heavy metals precipitate as
- 457 insoluble species.<sup>5,6</sup> While this work has focused on Na, Ca-based chloride brine compositions,
- in reality, the range of possibly compositions is much broader. For example, not only can brines
- contain species other than Na, Ca, and Cl, but more broadly, water treatment systems may also
- produce sulfate-rich brines including those based on magnesium compositions. While
- consideration of each of these aspects is beyond the scope of this paper, the general approach
- elaborated herein can be used to study the interactions and compatibility of diverse brine and fly ash types and compositions. This is significant as opportunities exist for knowledge of the
- nature developed herein to influence the selection of upstream treatment technology options
- such as reverse osmosis or thermal evaporation such that an optimal brine concentration can
- be achieved and matched to the coal-combustion residuals produced over the course of electricity production operations.
- 

### **SUMMARY AND CONCLUSIONS**

- This paper has shown that the brine composition (NaCl or CaCl2) and its ionic strength (0 ≤ *I<sup>m</sup>* ≤
- 471 7.5 mol/L) strongly affect: (1) the phase assemblage formed, and (2) the fly ash degree of
- 472 reaction. In general, hypersaline NaCl and CaCl<sub>2</sub> brines strongly promote the formation of Cl-
- AFm phases including: Friedel's and/or Kuzel's salts. The formation of the Cl-AFm phases,
- however, is counterbalanced by the destabilization of other phases that are otherwise stable in
- DI-water (e.g. katoite, strätlingite, monosulfoaluminate, etc.) with increasing chloride
- abundance. Thermodynamic modeling indicates that such phase stability is related to
- 477 differences in the prevailing pH: e.g., a strong increase in pH is observed in NaCl systems with
- 478 increasing CI<sup>-</sup> consumption. In contrast, Ca<sup>2+</sup> is consumed alongside CI<sup>-</sup> to form hydrated phases 479 in a CaCl<sub>2</sub> system, and the pH consequently decreases. These differences in pH, and phase
- stability as a function of brine composition and ionic strength also result in differences in the
- reactivity of Class C and Class F fly ashes, wherein the reactivity of the Class C fly ashes is
- furthermore affected by reductions in the water activity with increasing brine salinity. Taken
- together, the outcomes of this work illustrate the need to select the appropriate combination
- of fly ash type and additive to match brine composition, to ensure effective S&S. This
- information is important as it enhances our ability, *a priori*, to rapidly screen S&S compositions
- as a function of practical considerations including: (desired) extent of water volume reduction,
- 487 Cl<sup>-</sup>, and/or SO<sub>4</sub><sup>2-</sup> consumption, the prevailing pH of the pore solution, etc. Amongst others, such
- information forms a critical input to reactive-transport calculations which seek to assess the
- performance and contaminant retention ability of encapsulation materials and S&S operations.
- 

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- 

## **SUPPORTING INFORMATION**

- Additional isothermal calorimetry data (Figure S1), IR spectroscopy data (Figure S2), TGA data (Figure S3), and equilibrium diagram of ettringite and monosulfoaluminate as a function of pH and Cl concentration (Figure S4).
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#### **SYNOPSIS**

- This work increases our understanding of the interactions between hypersaline brines and the
- resulting cementitious matrix formed during solidification and stabilization processes.