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Investigation of Parameters Affecting Gypsum Dewatering Properties in a Wet Flue Gas Desulphurization Pilot Plant

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ABSTRACT: Wet flue gas desulphurization (FGD) plants with forced oxidation, installed at coal and oil fired power plants for removal of SO\(_2\)(g), must produce gypsum of high quality. However, quality issues such as an excessive moisture content, due to poor gypsum dewatering properties, may occur from time to time. In this work, the particle size distribution, morphology, and filtration rate of wet gypsum formed in a pilot-scale experimental setup, operated in forced oxidation mode, have been studied. The influence of holding tank residence time (10–408 h), solids content (30–169 g/L), and the presence of impurities (0.002 M Al\(_2\)F\(_6\); 50 g quartz/L; 0.02 M Al\(^{3+}\); and 0.040 M Mg\(^{2+}\)) were investigated. In addition, slurry from a full-scale wet FGD plant, experiencing formation of flat shaped crystals and poor gypsum dewatering properties, was transferred to the pilot plant to test if the plant would now start to produce low quality gypsum. The crystals formed in the pilot plant, on the basis of the full-scale slurry did, however, show acceptable filtration rates and crystal morphologies closer to the prismatic crystals from after pilot plant experiments with demineralized water. The gypsum slurry filtration rates were generally high, but a shorter residence time (10 h) and gypsum crystals experiencing breakage and/or attrition (408 h) showed slightly lower filtration rates. Both these experiments contained a higher fraction of fines, which may explain the slightly lower filtration rates. Crystals formed at a higher solids concentration and longer residence time (169 g/L and 120 h) showed a higher proportion of flat crystals, and the XRD pattern contained strong peaks at 31.1° (as the full-scale gypsum) and 29.1°, but no change in the filtration rate was obtained. It has not been possible in the pilot plant to form flat gypsum flakes with poor dewatering properties similar to those observed in full-scale plants.

1. INTRODUCTION

The combustion of fossil fuels, such as coal, oil, and natural gas, covers a substantial part of the global energy demand. Fuel resources can be efficiently utilized in combined heat and power plants with flue gas cleaning technologies, such as selective catalytic reduction (SCR), wet flue gas desulphurization (FGD), and electrostatic precipitation, installed to minimize emissions.

The wet flue gas desulphurisation process can remove acidic gases (SO\(_2\), HCl, and HF) in an absorption step by bringing the flue gas into contact with an alkaline slurry, where the solids consist of finely ground limestone (with impurities such as silica, iron, and magnesium) and gypsum particles.\(^1,2\) The HCl and HF absorbed remain in solution, and the concentration of Cl\(^-\) and other impurities are controlled by a purge stream. The SO\(_2\) dissociates to hydrogen sulfite ions (HSO\(_3^-\)), which are again oxidized to sulfate ions (SO\(_4^{2-}\)) by oxygen from air injection. Finally, SO\(_4^{2-}\) ions combine with Ca\(^{2+}\) ions and crystallize as gypsum (CaSO\(_4\cdot2\)H\(_2\)O) according to the following equation

\[
\text{CaCO}_3(s) + \text{SO}_2(g) + 2\text{H}_2\text{O}(l) + \frac{1}{2}\text{O}_2(g) \rightarrow \text{CaSO}_4\cdot2\text{H}_2\text{O}(s) + \text{CO}_2(g)
\]  

(1)

In 1999 the gypsum producing wet scrubber constituted the majority (>60%) of FGD capacity installed at power plants worldwide\(^4\) and is still the dominant FGD technology today producing a significant part of the world’s gypsum. The crystallization process includes the formation (nucleation) and growth of solid crystals from a supersaturated solution (SS). Super saturation is expressed mathematically by

\[
SS = \frac{a_{\text{Ca}^{2+}}a_{\text{SO}_4^{2-}}a_{\text{H}_2\text{O}}^2}{K_{s,\text{CaSO}_4\cdot2\text{H}_2\text{O}}} - 1
= \frac{\gamma_{\text{Ca}^{2+}}\gamma_{\text{SO}_4^{2-}}\gamma_{\text{H}_2\text{O}}^2(\text{H}_4\text{O}_6^{2-})}{K_{s,\text{CaSO}_4\cdot2\text{H}_2\text{O}}} - 1
\]

(2)

For growth to take place, SS must be positive. The particle size distribution (PSD) and morphology obtained depend on the growth rate (supersaturation-dependent) as well as the potential adsorption of foreign species onto specific crystal surfaces.\(^3\) The gypsum produced is withdrawn, washed, and dewatered using hydrocyclones, centrifuges, or vacuum belt filters until a commercial product is obtained. Commercial applications include wallboard and cement production and soil conditioning, for which different specifications apply, with respect to purity, moisture content (size and morphology dependent), and the content of different impurities (residual limestone and calcium sulphite, CaSO\(_4\cdot2\)H\(_2\)O). Wet FGD gypsum crystals are usually 5–50 μm, rhomboidal in shape, and of a high purity (>98%),\(^4\) but quality issues, such as a decreased dewatering rate, can occur resulting in expenses for additional
processing or disposal. Decreased dewatering rates may be caused by both increased levels of small particles and by morphology changes, which favor a closer packing of the gypsum crystals (Figure 1). Short residence times, excessive nucleation (high supersaturation) or breakage may decrease the particle size, while the crystal morphology can be affected by temperature, growth rate (supersaturation), and impurities as discussed in further details below.

Commercial gypsum may also be obtained from other sources, for instance mining of deposits from brine evaporation or industrial processes such as phosphoric acid and hydrogen fluoride production. Differences in gypsum properties may result from different growth conditions such as temperature, concentration of salts, and humic compounds (organic constituents of soil). An increasing concentration of deprotonated humic compounds has been reported to favor gypsum twin/rosette aggregate morphologies, while high temperature and salinity (60 °C and 30% NaCl, respectively) yield blocky aggregates of bassanite (2CaSO₄·H₂O). Industrial gypsum may also exhibit different morphologies, probably due to the presence of specific impurities which occupy the most favorable binding sites, thereby affecting the rate of ion incorporation (crystal growth) and the resulting crystal morphology. Differences in the supersaturation of the liquid phase may also affect morphology, with higher degrees of supersaturation favoring longer and thinner needle-like crystals. Hemihydrate and gypsum formed during phosphoric acid production has in the presence of 0.100 mol/kg aluminum fluoride complexes (AlF₆³⁻ to AlF₅²⁻) and elements from the lanthanides series (at a concentration of 0.0003 M) shown inhibited growth of specific crystal faces, changes in crystal morphology (toward flat shaped crystals), filterability, and washability of the filter cake, but it should be remembered that this process takes place at a lower supersaturation and higher temperatures (up to 90 °C) than wet FGD. Iron, manganese, and magnesium have also been reported to negatively affect the dewatering performance of wet FGD gypsum. A contamination with 0.02 M Mg²⁺ or Fe³⁺ was enough to reduce the growth rate and change the crystal morphology toward needle-shaped crystals. In addition, breakage along characteristic mineral planes may also change the crystal morphology obtained—gypsum can exhibit perfect cleavage in one direction because of the water layers separating calcium-sulfate layers in the crystal lattice. Several studies on the theoretical predictions of gypsum equilibrium morphology (prismatic/rhombic) and growth morphology (flat/plate-like, due to water absorption) have furthermore been published.

In this work the dewatering properties of gypsum produced in a wet FGD pilot plant have been investigated and compared to wet FGD gypsum from a full-scale plant (Asnæs Power Plant Unit 2, DK) experiencing dewatering problems. The full-scale wet FGD plant in question is a CT-121 jet bubbling reactor and a dewatering system consisting of hydrocyclone batteries followed by a vacuum belt filter.

2. STRATEGY OF INVESTIGATION

This investigation of wet FGD gypsum dewatering properties is based on a series of pilot-scale experiments with different slurry compositions and residence times. The pilot plant enables a high degree of control of the experimental conditions such as gas phase composition, slurry pH, and residence time. The conditions of the experiments performed have been chosen to investigate the effect of slurry composition (impurities, content of fines and flat crystals), solid concentration, and slurry residence time on gypsum PSD, morphology, and filtration rate.
Aluminum fluoride complexes, Mg\(^{2+}\), Al\(^{3+}\), and full-scale slurry with high concentrations of fines have all been linked to morphology changes (inhibited growth of specific crystal faces) and poor dewatering properties. Long residence times and high solid concentrations can occur at full-scale wet FGD plants during low sulfur coal/biomass combustion at partial power plant load and these conditions create a potential for changes in PSD and morphology through breakage/attrition. Slurry samples were withdrawn and analyzed in terms of filtration time, PSD, morphology, and elemental composition (selected elements only).

### 3. EXPERIMENTAL SETUP AND PROCEDURE

**Description of Setup.** The wet FGD pilot plant, outlined in Figure 2, simulates a single vertical channel in a full-scale cocurrent flow wet FGD grid absorber. A 110 kW natural gas burner generates the flue gas and SO\(_2\) (AGA Gas AB) is subsequently added before it is brought into contact with a

![Outline of wet FGD pilot plant used in the experiments. The absorber is based on the falling film principle, and the gas and liquid phases flow cocurrently. Reprinted with permission from the work of Hansen et al. Copyright 2011 American Chemical Society.](image)

**Table 1. Overview of Experiments Performed and Operating Conditions**

<table>
<thead>
<tr>
<th>experiment</th>
<th>addition feed tank</th>
<th>addition holding tank</th>
<th>slurry (\tau) (h)</th>
<th>flue gas (N m(^{-3})/h)</th>
<th>(\text{SO}_2) ppm(v)</th>
<th>duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a: ASV2 slurry</td>
<td>ASV liquid phase(^b)</td>
<td>ASV slurry</td>
<td>30</td>
<td>18</td>
<td>990</td>
<td>(\sim 1\tau)</td>
</tr>
<tr>
<td>1b: base case</td>
<td></td>
<td></td>
<td>30</td>
<td>17</td>
<td>950</td>
<td>(\sim 2^{1/2}\tau)</td>
</tr>
<tr>
<td>2: seed crystals</td>
<td>ASV process water</td>
<td>AVV gypsum(^d)</td>
<td>28</td>
<td>17</td>
<td>1030</td>
<td>(\sim 2\tau)</td>
</tr>
<tr>
<td>3: Al(_2)F(_6)</td>
<td>0.002 M Al(_2)F(_6)</td>
<td>0.002 M Al(_2)F(_6)</td>
<td>25</td>
<td>17</td>
<td>1000</td>
<td>(\sim 2r)</td>
</tr>
<tr>
<td>4: quartz(^c)</td>
<td>50 g quartz/l</td>
<td>50 g quartz/l</td>
<td>30</td>
<td>17</td>
<td>1030</td>
<td>(\sim 5r)</td>
</tr>
<tr>
<td>5a: Mg(^{2+})</td>
<td>0.04 M Mg</td>
<td>0.04 M Mg</td>
<td>35</td>
<td>17</td>
<td>1010</td>
<td>(\sim 3r)</td>
</tr>
<tr>
<td>5b: Al(^{3+})</td>
<td>0.02 M Al</td>
<td>0.02 M Al</td>
<td>27</td>
<td>17</td>
<td>1010</td>
<td>(\sim 2r)</td>
</tr>
<tr>
<td>5c: base case</td>
<td></td>
<td></td>
<td>30</td>
<td>17</td>
<td>980</td>
<td>(\sim 1^{1/2}\tau)</td>
</tr>
<tr>
<td>6: (\tau = 10) h</td>
<td></td>
<td></td>
<td>10</td>
<td>18</td>
<td>980</td>
<td>(\sim 5r)</td>
</tr>
<tr>
<td>7: (\tau = 80) h</td>
<td></td>
<td></td>
<td>83</td>
<td>16</td>
<td>270</td>
<td>(\sim 2^{1/2}\tau)</td>
</tr>
<tr>
<td>8a: (\tau = 120) h high density</td>
<td></td>
<td></td>
<td>122</td>
<td>17</td>
<td>510</td>
<td>(\sim 1^{1/2}\tau)</td>
</tr>
</tbody>
</table>

\(\tau = \) residence time. \(^{b}\)Coarse gypsum particles separated by sedimentation and liquid phase including fines used for limestone feed slurry. \(^{c}\)Quartz PSD (20\% < 5 \(\mu\)m). \(^{d}\)Gypsum from Avedøre Power Station (high filtration rate).
slurry based on Faxe Bryozo limestone, with a purity above 97 wt %, in the 7 m absorber. Multiple sampling sites along the absorber enable measurements of gas phase composition profiles and slurry pH. The slurry is collected in a 110 L holding tank, where air injection, reactant addition (for maintaining a constant holding tank pH), and slurry removal (for maintaining a constant slurry level) take place. From the holding tank, the slurry is recycled to the absorber, ensuring a liquid/gas ratio of 16–18 L slurry/m³ flue gas (STP).

Additional details concerning the pilot plant can be found in previous publications.20,21 Experimental Procedure. The experimental series were initiated by at least 2 days of desulphurisation of a 1000 ppm(v) SO₂ (g) flue gas stream using a feed stream containing 7.1 wt % Faxe Bryozo limestone. This allowed the system to approach steady state operation in terms of gypsum content, residual limestone content, and desulphurisation degree. The different operating conditions in Table 1 were then introduced one at a time and operation continued for at least one residence time (usually more). If an experiment is performed with the slurry from the preceding experiment, this is indicated by the use of the same “exp number” but with a letter (e.g., “a”, “b”) to distinguish experiments. Quartz was added in exp 4 to investigate the influence of fines, which has been linked to changes in morphology.21 A diluted feed stream was used to generate a short residence time in exp 6, and a concentrated feed stream was used to increase the slurry density (higher gypsum concentration) in exp 8a. Experiment 8b investigates the influence of long-term breakage/attrition on crystal size and morphology while no crystal growth takes place, simulating wet FGD operation during low sulfur coal/biomass combustion at partial power plant load. Chemicals used include: MgSO₄·7H₂O (Fluka 99%), AlCl₃·6H₂O (Sigma Aldrich 99%), NaF (Sigma Aldrich 99%), and quartz powder (Ormslev kvartsværk “Kvartsmel” d₂₀% < 5 μm).

The limestone consumption (change in weight of feed tank), the flue gas flow rate (Flemco flowmeter), the holding tank pH (Yokogawa FU20), and the SO₂ content (Rosemount NGA 2000) were monitored throughout the experiments. Each experiment was concluded by withdrawing two slurry samples from the holding tank for analysis of particle size distribution, crystal morphology, and filtration rate. The slurry samples for PSD analysis were stored for up to a few days at room temperature before analysis. The limited solubility of gypsum (about 2 g/L), compared to the slurry concentration (>100 g/L), and its invariability with temperature22 ensured that no major changes in PSD took place during storage, as has been previously verified.23 Two individual PSD analyses were performed by laser diffraction (Malvern Mastersizer S longbed) on an ethanol/slurry dispersion of each sample, all analyses representing the average of 5 measurements. The filtration rate was obtained by measuring the time it took the 80 mL slurry sample to pass through a 0.45 μm filter in a Büchner funnel. The collected crystals were subsequently dried at 40 °C and used to study the morphology and crystallographic structure by scanning electron microscopy (FEI Inspect S) and X-ray powder diffraction (PANalytical X’pert PRO Diffractometer). Information on the crystallographic structure/crystal lattice planar distance (d) is obtained by applying Bragg’s law (nλ = 2d sin θ) for a range of incident angles (θ). A few samples were furthermore screened in terms of the elemental composition of the liquid and solid phase, and the concentration of selected elements (Al, Na, Cl, F) was determined. The samples for analysis of the ionic liquid composition were passed through a 0.45 μm filter and diluted 1:2 with laboratory grade pure water to prevent further precipitation. The solid crystals (collected from the filter) were either dissolved in a warm mixture of acids (HF and HNO₃), neutralized by H₂BO₃, and analyzed by inductively coupled plasma optical emission spectrometry (Perkin-Elmer Optima 3000 ICP-OES) for cation detection or dissolved in excess laboratory grade pure water and analyzed by ICP-OES (Spectro Ciros-CCD) for anion detection.

4. RESULTS AND DISCUSSION

The experimental work includes 12 wet FGD experiments using different slurry compositions (1a–5c), solids content (7–8b), and residence times (5c–8b) to investigate their impact on gypsum particle size distribution, morphology, and filtration rate. Details of the experiments are given in Table 1.

Effect of Experimental Conditions on Filtration Rate. The filtration times and slurry solids concentrations of the experiments performed are shown in Table 2. The diluted limestone feed stream used in the experiment with a short residence time (τ = 10 h) also causes a lower solids concentration. The solids concentrations in the base case and the Al³⁺ experiments are also below the average, due to a minor water leak from an external water tank entering the system through the air injection line. Experiments 5a (Mg²⁺), 5b (Al³⁺), and 5c (base case), with very similar particle size distributions and morphologies (will be illustrated in the next sections) yield reasonably consistent filtration times (75, 55, and 65 s, respectively).

The filtration time of the full-scale slurry (exp 1a, 1200 s) differs significantly from the other experiments (generally about 60 s), including an experiment with seed crystals from Avedøre power plant (exp 2). However, gypsum crystals formed in the pilot plant in the presence of ASV2 liquid phase show a highly reduced filtration time (about 180 s). The gypsum produced at short residence times and after 408 h attrition/breakage both showed slightly increased filtration times (140–170 s). The filtration rate may be affected by mechanisms changing the morphology or the PSD, this will be discussed in further detail in the following sections.
Effect of Process Parameters on Gypsum Particle Size Distribution. Figures 3 and 4 and Table 3 show the particle size distribution and the volumetric mean particle diameter, respectively. The base case particle size distribution and mean particle size corresponds well with previously published data from the pilot plant ($d_{50} = 42.8 \mu m$)\(^5\) while it was slightly lower in fines than previously investigated gypsum from full-scale plants ($d_{50} = 34.9−41.3 \mu m$).\(^5\) A high mean particle size and good filtration properties were obtained for the base case experiment as well as experiments with Al\(^{3+}\) and Mg\(^{2+}\) addition. A lower mean particle size is obtained in the experiments with quartz addition and long residence times, but short filtration times are still obtained. The particle size distributions in these experiments are shifted toward smaller particles, but without any significant changes in the level of particles smaller than 10 \(\mu m\). The experiments with full-scale slurry, short residence times, and 408 h breakage (1a, 1b, 6, and 8b) yield a lower mean particle size, higher levels of particles below 10 \(\mu m\) (least pronounced in exp 8b), and longer filtration times (especially for the full-scale slurry). A short residence time means less growth time for the crystals, thereby yielding smaller crystals, while the experiments with longer residence times (7, 8a, and 8b) may facilitate breakage/attrition and a slower crystal growth rate (a lower SO\(_2\) concentration was used yielding a lower supersaturation). Fine particles may furthermore be introduced into a full-scale plant via dust in the flue gas, impurities in the process water and excessive nucleation.

Effect of Experimental Conditions on Crystal Morphology. The crystals formed during the base case experiment were columnar shaped (Figure 5), corresponding well with the morphologies reported in literature at lower degrees of saturation ($S \leq 1.2$).\(^7,8\) A similar morphology was obtained in experiments with Al\(_2\)F\(_6\) (exp 3), quartz (exp 4), Mg\(^{2+}\) (exp 5a), Al\(^{3+}\) (exp 5b), and a prolonged residence time (exp 7), the crystals in the latter experiment were, however, a bit more compact. The strongest X-ray diffraction peak of these gypsum samples (Figure 6) was at 11.6° (020 face), with strong and moderate peaks at 20.7° (−1−21 face), 23.4° (040 face), 29.1° (−1−41 face), and an additional moderate peak at 31.1° (121 face) for the gypsum with quartz addition and the gypsum formed at a prolonged residence time. The peaks at 20.7° and 29.1° were more distinct in the base case gypsum than in the other samples. The shorter residence time, lower gypsum
concentration, and, thereby, potentially higher supersaturation (not measured) in experiment 6 formed more elongated crystals (Figure 7) compared to the other experiments, also in agreement with other investigations.\(^8,9\) The strongest X-ray diffraction peak was still at 11.6\(^\circ\) (020 face), with moderate peaks at 20.7\(^\circ\), 23.4\(^\circ\), and 29.1\(^\circ\). The thin gypsum flakes from the full-scale wet FGD plant differed significantly from the crystals from all other experiments. The crystal aspect ratio did, however, decrease, and the filtration rate increase, in the pilot plant during growth in the presence of ASV liquid phase (experiment 1a, see Figure 8). The strongest X-ray diffraction peak was at 31.1\(^\circ\) with moderate peaks at 11.6, 20.7\(^\circ\), 29.1\(^\circ\), 33.3\(^\circ\) (051 face), and 35.9\(^\circ\) (200 face). The intensity of the 20.7\(^\circ\) and 29.1\(^\circ\) peaks increased significantly during experiment 1a. Crystals formed at higher solids concentrations, and longer residence times (exp 8a) showed a higher proportion of flat crystals (as in Figure 8) and strong XRD peaks at 20.7\(^\circ\) (strongest), 29.1\(^\circ\), and 31.1\(^\circ\), while crystals exposed to long-term attrition (exp 8b) showed similar XRD peaks (29.1\(^\circ\) and 31.1\(^\circ\) were slightly more powerful) and more rounded shapes (Figure 9), similar to gypsum from full-scale wet FGD plants.\(^5\)

The flatter crystals formed at higher solids concentrations and longer residence times (exp 8a) also possessed the strong XRD peak at 31.1\(^\circ\) seen for the ASV2 gypsum, but also a much stronger 29.1\(^\circ\) peak, and the morphology change did not affect the filtration rate. In summary, none of the pilot plant experiments performed could form the flat ASV2 gypsum crystals associated with low filtration rates.

**Gypsum Slurry Impurities.** The X-ray diffraction analysis identified gypsum as the major compound in all samples, but
changes in crystals morphology are often associated with adsorption of small amounts impurities onto specific crystal faces, thereby affecting the rate of ion incorporation (crystal growth) and the resulting morphology. Table 4 shows the concentration of selected impurities (Al, Na, Cl, F, and P) for six full-scale and pilot-scale samples (three with solid gypsum and three with liquid phase). The impurities initially present in the ASV2 gypsum gradually disappear in the pilot plant (exp 1a), and the crystal thickness and filtration rate increase, despite the use of the liquid phase from the full-scale slurry. Traces of iron, potassium, magnesium, silicon, and titanium were furthermore detected in addition to high levels of calcium (from gypsum). Pilot scale experiments with Al3+, Mg2+, and F− added showed no effect of these species on gypsum morphology. The elements present in wet FGD systems depend on fly ash escaping the particulate filter, gaseous elements absorbed, and impurities from limestone and process water. The elements measured in this work correspond well with other investigations, but additional species such as zinc, copper, manganese, molybdenum, and vanadium may also be encountered.

## CONCLUSION

A systematic pilot-scale investigation of wet FGD gypsum dewatering properties has demonstrated that high quality gypsum with favorable dewatering properties can be produced in a pilot scale setup over a wide range of operating conditions. A decreased holding tank solids residence time (from 30 to 10 h) did, however, yield elongated crystals, an XRD pattern dominated by a strong peak at 11.6° (peaks at 20.7° and 29.1° were less powerful than in the base case), a higher fraction of fines, and a slightly lower filtration rate, possibly due to higher degrees of supersaturation. Long-term attrition (408 h) also shifted the PSD toward smaller particles and decreased the filtration rate. Gypsum from a full-scale plant experiencing dewatering problems contained an increased fraction of fines and had a significantly different morphology (thin gypsum flakes) and XRD pattern (dominated by a strong peak at 31.1°). The properties of these crystals did, however, improve (and the impurity content decreased) during subsequent growth in the pilot plant both in the presence of liquid phase from the full-scale plant and demineralised water. A shift toward flatter crystals was obtained at higher solids concentrations and longer residence times (169 g/L and 120 h), and the XRD pattern contained strong peaks at 31.1° (as the full-scale gypsum) and 29.1°, but the morphology change did not affect the filtration rate.

An increased processing time may to some extent overcome poor dewatering properties, but efforts to change the PSD or morphology should also be considered. The presence of growth retarding species may significantly influence the gypsum PSD and morphology but no specific compounds were identified to be of importance in the present study. This effect may be reduced by an increased wastewater withdrawal, optimization of the preceeding particulate control device, a change in the limestone source, and process water pretreatment. High degrees of supersaturation may furthermore increase nucleation rate and favor needle-like crystals. The supersaturation in a wet FGD plant is influenced by the sulfur content of the fuel used, the desulphurisation degree as well as the total crystal area available for growth. Finally, an increased slurry volume or solids concentration can provide both potentially positive (longer residence time) and negative effects (increased exposure to breakage or attrition and an increased surface area due to lower growth rate).

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**Notes**

The authors declare no competing financial interest.

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### NOMENCLATURE

\[ a = \text{activity [M]} \]

\[ c = \text{concentration [M]} \]

\[ d = \text{crystal lattice planar distance} \]

\[ D_{20\%} = \text{particle size under which 20% of the total volume is located [μm]} \]

\[ D_{50\%} = \text{volumetric mean diameter [μm]} \]

\[ F_s = \text{cumulative volumetric distribution [-]} \]

\[ l = \text{particle size [μm]} \]

\[ n = \text{order of reflection} \]

\[ SS = \text{degree of super saturation [-]} \]

\[ γ = \text{activity coefficient [-]} \]

\[ Θ = \text{incident X-ray angle} \]

\[ λ = \text{X-ray wavelength} \]

### REFERENCES


(13) Blyth, G. M. Results of Full-Scale Utility FGD SO2 removal upgrade testing. Tenth Annual Coal Preparation, Utilization, and Environmental Control Contractors Conference, Pittsburgh, July 18−21, 1994.


