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# Impact of freeze–thaw weathering on integrity, internal structure and particle release from micro- and nanostructured cement composites†

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Nowadays, both micro- and nanomaterials are processed in cement to improve application properties or reduce cement consumption in concrete. The present study deals with the impact of nanostructured SiO<sub>2</sub>, TiO<sub>2</sub> and milled slags in cement composites on material properties, aging resilience and release behaviour into the environment. In this context, hydrated and hardened cement pastes were weathered artificially by means of the standardised capillary suction of de-icing solution and repeated freeze–thaw cycle test method. Mineralogical, physical, mechanical and chemical properties as well as hydration products were analysed by electron microscopy, mercury intrusion porosimetry, thermogravimetry as well as differential scanning calorimetry and ultrasonic propagation analysis. Release characterisation was performed concerning weathering-induced fragment release into the liquid phase by gravimetry and sanding-induced particle release into the air by gravimetry, differential electrical mobility analyses, time of flight spectrometry, condensation nuclei counting and electron microscopy. Freeze–thaw weathering led for all cement composites to considerable but different progressed deteriorations. The worst material performance was observed for pure cement and the cement composite with state of the art ground granulated slag. The cement composites with nanostructured SiO<sub>2</sub> and activated ground granulated slag showed a higher durability and less deteriorations. These effects could be explained by structure density, porosimetry and the formed hydration products. Furthermore, it was shown that weathering-induced changes in the material properties lead to significant changes concerning sanding-induced particle release into air. In summary, this study investigated the consequences of (nano)-materials, which modulate the recrystallization during hardening of cement pastes to a solid, internally nanostructured material, on multi-structural release. In this context, critical stress scenarios, which are encountered during intended use, were simulated. The transforming synthesis differentiates this case study from almost all other release studies for nanomaterials, which typically assess polymer nanocomposites, where nanomaterials act as additives, pigments or fillers in an otherwise unchanged matrix.

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## Environmental significance

Environmental impacts can change substantially the material properties of nanoenabled products and thus the release behaviour into the environment. In this context, the study addresses cements composites that have the potential to improve sustainably the construction material concrete. Results of a comprehensive testing program provide a mechanistic insight concerning the impact of freeze–thaw weathering on integrity, internal structure and particle release behaviour of micro- and nanostructured cement composites. Since nanomaterial use in cements is accompanied by a transforming synthesis (recrystallization during hardening), the consequence of nanomaterial use in cements was investigated that differentiate this study from almost all others release studies. It is shown that nanomaterial additives have the potential to modulate the release behaviour over the lifecycle of cement composites.

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## 1. Introduction

Based on the advantageous mechanical properties and excellent processability, Portland cement concrete is one of the most important construction materials all over the world. In most cases, cement is the basic component of concrete and serves therein after the addition of mixing water as binder material due to its chemical transformation into a solid continuous



phase. The high demand on concrete requires also huge amounts of cement that lead to a current worldwide cement production volume of around 4 billion tonnes per year.<sup>1</sup>

Beside the benefits of concrete, one global challenge is the considerable carbon dioxide emission (approx. 5 wt-% of global anthropogenic CO<sub>2</sub> emissions)<sup>1</sup> accompanying the production of cement (e.g. 0.562 t<sub>CO<sub>2</sub></sub>/t<sub>cement</sub> in Germany in 2015);<sup>2</sup> especially due to limestone decalcification and clinker burning. Approaches to reduce emissions deal with improved cement compositions or the development of advanced concretes. One possibility is the substitution by suitable latent hydraulic materials (i.e., activator-based material that cured chemically with water in water) like fly ash or ground granulated blastfurnace slag (GGBS).<sup>3</sup> Another way is the use of high or ultrahigh performance concrete, which contain specific fillers or functional additives like hydraulic micrometre-sized silica,<sup>4</sup> nanostructured silica,<sup>4–11</sup> carbon nanotubes<sup>12,13</sup> or carbon fibres.<sup>14</sup>

During the lifetime of concrete, the material ages due to temperature changes and attack of aggressive media like acid rain, saline solutions from de-icing salt, sewage or carbon dioxide and sulphur oxide as gaseous pollutants. Deterioration of concrete increases the probability of fragmentation, and the probability of molecular or particulate additives to leach into the surrounding environment. The processes and consequences of such deterioration have to be evaluated and quantified. Especially nanomaterials are associated with a higher uncertainty of environment, health and safety considerations in comparison to more conventional materials, the increasing application of nanomaterials as filler or functional additive in cements and concretes demands also an appropriate risk assessment. The nanoform of a substance may have a lower threshold of adverse effects than the non-nanoform of the same substance, especially due to the increased specific surface area, which in turn is a prerequisite for the nanomaterial's performance as a binder.

In recent years, several experimental studies concerning release into air or water from solid nanocomposites, especially for the plastic, paint and coating industry, were performed,<sup>15–17</sup> whereof some dealt also with the impact of the materials life cycle,<sup>18–25</sup> i.e., the effect on release by the change of material properties due to aging. An increasing number of studies concerning the release or exposure characterisation emerges for concretes<sup>26–32</sup> or hardened cement pastes,<sup>19,33–35</sup> but rarely differentiate the influence of nanomaterial additives.

The effect of artificially weathering on the release behaviour into air from cement nanocomposites (containing 2 wt-% of carbon nanotubes resp. 1 wt-% nanosized calcium silicate hydrates) was previously analysed by simulated weathering based on a method that was standardised for plastics/coatings (ISO 4892-2:2006 (ref. 36)).<sup>19</sup> Neither the surface characterisation concerning morphology and chemistry nor the performed release analyses for weak (abrasion) and moderate (sanding) mechanical stress showed significant differences between the aged samples and their unaged

counterparts. However, the aging by UV irradiation with or without rain may not have tested the most relevant mechanism of degradation for cementitious systems. Hirth *et al.* analysed in detail sanding-induced fragments from CNT-containing cements concerning CNT protrusions.<sup>33</sup> Further release analyses by mechanical shear (sonication) during immersion in water showed a high fragment release rate that was slightly reduced by embedded CNT fillers,<sup>34</sup> and where recently followed up by high-throughput ecotoxicity screening of the hazards of released fragments.<sup>37</sup> A high fragment release was also observed during leaching (7 d, ultrapure water, stirring) of photocatalytic cement nanocomposites (approx. 2.85 wt-% photocatalytic TiO<sub>2</sub>).<sup>35</sup> Bossa *et al.* showed furthermore that the release quantities increase with increasing surface porosity, which accompany with higher water to cement ratios (*w/c*) or longer leaching time.

The present work, which deals with the effects on the binder matrix of cementitious materials (i.e., hardened cement pastes), advances from less intense processes such as leaching or UV irradiation to a more aggressive weathering that was especially designed for concrete – the standardised capillary suction of de-icing solution and freeze thaw test (CDF) according to CEN/TS 12390-9:2016.<sup>38</sup> To characterise the impact of this weathering procedure on integrity, internal structure and particle release from micro- and nanostructured cement composites, a comprehensive test program was performed before, during and after CDF. In this context, the modulation of fragmentation and release by different materials, which are used to optimize the commercial value of concrete by reducing embodied energy, is considered comparatively.

## 2. Experimental details

### 2.1 Samples systems and sample preparation

Overall five types of cement composites (Table 1) based on high purity Portland cement with low iron content (CEM I 52.5R, HeidelbergCement AG, Heidelberg, Germany) and four different additives were analysed in this study. The solely hydrated cement (CEM\_hyd) served therefore as reference material to study the impact of the additives. In order to compare also the release behaviour of the cement composites with the ones of concrete, analyses were also performed for a high performance concrete (FuturBeton), which contains beside CEM and HKP-GGBS as clinker substitute also sand and coarse grain in different sizes.

The additives GGBS and HKP-GGBS (High Kinetic Processing-GGBS) were produced based on processing of a commercial granulated blastfurnace slag (GBS, ThyssenKrupp Steel Europe AG, Duisburg, Germany). Therefore, 5 kg of GBS were dried at 60 °C and ball milled (drum mill Comb03-A03, Zoz GmbH, Wenden, Germany) with 44 kg corundum balls (ball diameter between 5/4" and 7/8", total filling level of 66.7 vol-%) at 50 rpm for 4 h to obtain GGBS. Afterwards, GGBS was processed to HKP-GGBS by high energy milling (Simoloyer® CM08-8lm, Zoz GmbH, Wenden, Germany). More details



**Table 1** Composite IDs, additive IDs, formulation for preparation of cement/concrete composite samples and dry composite densities

ID of composite <sup>a</sup> [-]	ID of additive [-]	Formulation for preparation additive/water/superplasticizer/others [wt-% related to cement]	Dry composite density [g cm <sup>-3</sup> ]
CEM_hyd	—	0.00/40.0/0.0/0.0	1.82
CEM_GGBS	GGBS	42.9/57.1/0.0/0.0	1.70
CEM_HKP-GGBS	HKP-GGBS	42.9/57.1/0.0/0.0	1.81
CEM_SiO <sub>2</sub>	SiO <sub>2</sub> _nano	1.0/40.0/0.6/0.0	1.65
CEM_TiO <sub>2</sub>	TiO <sub>2</sub> _nano	1.0/40.0/0.6/0.0	1.81
FuturBeton	HKP-GGBS	42.9/43.0/0.52 <sup>b</sup> /643.1	2.34

<sup>a</sup> Note: IDs are supplemented by a star, when weathered samples are discussed. <sup>b</sup> Glenium® ACE 431, BASF Construction Solutions GmbH, Trostberg, Germany: PCE-content: 30% (only for these concrete samples used).

concerning the production of HKP-GGBS are provided by Funk *et al.*<sup>39</sup> Beside the two modifications of blastfurnace slag, two commercial nanomaterials, *i.e.*, synthetic amorphous silica (SiO<sub>2</sub>\_nano, CAS 112945-52-5, primary particle diameter of 26 nm, corresponds to NM203 of the JRC repository)<sup>40</sup> and titanium dioxide (TiO<sub>2</sub>\_nano, CAS 13463-67-7, primary particle diameter of 21 nm, corresponds to NM105 of the JRC repository)<sup>41</sup> were chosen for cement composite preparation. Data on the elemental composition and the mass-specific surface area for each of the processed raw materials are provided in Table S1.†

As it can be deduced from the formulation data in Table 1, the cement pastes preparation was performed with technical relevant amounts of additives and a constant water to cement ratio (*w/c*) of 0.4 according to DIN 1045-2:2008 (ref. 42) to i) ensure complete hydration and hardening and ii) prevent incomplete binding (as typical for *w/c* > 0.4). In order to receive a suitable processability, the components of each composite were stirred with mixing water at room temperature for at least 2 min. In the case of the blast furnace slag composites (CEM\_GGBS, CEM\_HKP-GGBS), the slag additive and the cement were mixed in advance by a continuous ploughshare mixer (Ploughshare® Food Mixer, Gebrüder Lödige Maschinenbau GmbH, Paderborn, Germany). For the preparation of the nanomaterial-containing cements pastes (CEM\_SiO<sub>2</sub>, CEM\_TiO<sub>2</sub>), SiO<sub>2</sub>\_nano and TiO<sub>2</sub>\_nano respectively were dispersed homogeneously in the mixing water with the addition of a superplasticizer (MasterGlenium® SKY 592, BASF Construction Solutions GmbH, Trostberg, Germany) by means of a dissolver prior the addition of cement.

Three different types of test specimens were casted based on the prepared cement/concrete pastes: i) 100 × 100 × 100 mm<sup>3</sup> cubes according to EN 12390-2:2009 (ref. 43) for the standardised artificial freeze–thaw weathering, ii) non-standardised discs with a nominal diameter of 110 mm (bottom side diameter of 115.36 mm, bevelling of 15°, height of 10 mm) for release experiments on unaged samples and iii) standardised (mortar standards) 40 × 40 × 160 mm<sup>3</sup> prisms. To achieve comparable curing conditions, all test specimens were stored according to DIN EN 196-1:2016 (ref. 44) for at least 28 days under wet conditions at room temperature prior to any analysis.

## 2.2 Sample conditioning by artificial freeze–thaw weathering (CDF)

The cubes of the cement composites CEM\_hyd, CEM\_GGBS, CEM\_HKP-GGBS and CEM\_SiO<sub>2</sub> were artificially weathered by means of the CDF method according to CEN/TS 12390-9:2016 (ref. 38) that was developed to simulate aggressive environmental conditions for geographic regions where both freeze–thaw-cycles and seawater or de-icing salt does matter. Therefore, 5 afore casted and defined stored (*i.e.*, 1 d in covered casting mould, 6 d in water bath, 20 d in climate chamber) test specimens per cement composite were sealed on their side facades and partly submerged (approx. 5 mm) in a 3 wt-% aqueous NaCl solution for 7 d (within a test container with (10 ± 1) mm solution level and (5 ± 1) mm sample distance pieces). After gravimetrical determination of the amount of penetrated NaCl solution, the still submerged samples underwent 28 repeated freeze–thaw cycles. Each freeze–thaw cycle (FTC) based on 4 h successive cooling from 20 °C to –20 °C, resting at –20 °C for 3 h, successive reheating from –20 to 20 °C for 4 h and resting at 20 °C for 1 h.

## 2.3 Material characterisation

The cement composites were characterised before, during (*i.e.*, between single freeze–thaw cycles) and after freeze–thaw weathering concerning surface structure and hydration products by scanning electron microscopy (SEM), water absorption by gravimetry (GM), water content by simultaneous thermal analyses (STA), porosity by mercury intrusion porosimetry (MIP) and the relative dynamic *E*-modulus (RDM) by ultrasonic propagation analyses. In the cases of SEM, STA and MIP, analyses were performed on the base of fragments taken from the inner side of the cubes to avoid artefacts, which can originate from mechanical-induced densification of outer regions.

Surface structure analyses, especially for the evaluation of the hydration progress and for the identification of hydration products, were performed on the base of a gentle beam SEM (JSM-7500F, JEOL Ltd., Tokyo, Japan).

Water absorption analyses by gravimetry were performed before, during and after the 7 d lasting storage in the NaCl solution (*i.e.*, prior the freeze–thaw weathering procedure). For this purpose, the samples were removed from the test container and placed on an analytical balance.



Water content analyses were conducted within a nitrogen atmosphere by simultaneous thermal analyses (STA 449 F1 Jupiter® with platinum furnace, NETZSCH-Gerätebau GmbH, Selb, Germany), which comprises both thermogravimetry (TG) and differential scanning calorimetry (DSC). Prior to the analyses, all samples were dried at 35 °C to constant weight and grinded by pestle and mortar. The employed crucibles and sample holders were made of a platinum–rhodium alloy. The temperature gradient was chosen to be 10 K min<sup>−1</sup> within a temperature range of 30–1000 °C, while measurement data were evaluated with commercial analytical software (Proteus®, NETZSCH-Gerätebau GmbH, Selb, Germany).

Pore size analyses were realised by MIP (Autopore 9600, Micromeritics®, Aachen, Germany) over a pressure range between 0.10 psia and 61 000 psia (sample quantity 2.5 g ± 0.2 g), while data evaluation was performed with commercial software (MicroActive, Micromeritics®, Aachen, Germany).

Ultrasonic propagation analyses, *i.e.*, transit time analyses, for the determination of the RDMs were performed for five times (*i.e.*, for the FTCs 0, 4, 8, 16, 28). Analyses were performed after each of the mentioned FTCs in the wake of fragment release characterisation (see chapter 2.4) and sample mass characterisation by placing the samples between ultrasonic transmitter and receiver.

## 2.4 Release characterisation

Release characterisation was performed for weathering-induced fragment release into water and sanding-induced release into air prior and after freeze–thaw weathering.

Weathering-induced fragment release into water was characterised according to the non-size-selective CDF method of CEN/TS 12390-9:2016,<sup>38</sup> which probes rather the coarse particle fraction, for five times (*i.e.*, for the FTCs 0, 4, 8, 16, 28). After each of the mentioned FTCs, the test container, which contained sample and NaCl solution, was transferred to an ultrasonic bath and treated for 3 minutes. Afterwards, the NaCl solution was filtrated (folded filter, particle retention 8–10 µm, Sartorius™ FT-4-303-270, Sartorius AG, Göttingen, Germany) and the release quantities determined gravimetrically. According to CEN/TS 12390-9:2016,<sup>38</sup> the results of the first FTC were discarded.

Sanding-induced particle release into air was simulated according to ISO/DIS 21683:2018(E)<sup>45</sup> by means of a sanding apparatus that was design to mimic professional sanding processes (similar to other release processes accompanied with moderate mechanical stress on surfaces like driving on highways) on the base of single short-term testing (*i.e.*, <10 s) under clean room conditions.<sup>46</sup> The sanding apparatus contains a vertical aligned revolution-controlled rotatory sander (Model Dremel 400 Series Digital Dremel Europe, Breda, The Netherlands), which interacts due to gravity with the sample surface that is supplied horizontally beneath the sanding tool. Sanding procedures were performed in analogy to Göhler *et al.*<sup>21</sup> with P600 sanding paper (WPF P600, Mirka

GmbH, Sulzbach, Germany), a contact force of 0.5 N and a cutting velocity of 1.8 m s<sup>−1</sup> that results in a cutting power of 1.3 W. Released airborne particles were characterised by time of flight spectrometry (TOF, Model APS 3321, TSI Inc. Shoreview, USA), differential electrical mobility analyses (DEMA, Model EEPs 3090, TSI Inc., Shoreview, USA), condensation nuclei counting (CNC, Model CPC 3775 and Model CPC 3022, TSI Inc. Shoreview, USA) and by SEM (Gemini 982, Carl Zeiss AG, Jena, Germany) after electrostatic deposition (electrostatic precipitator according to Dixkens & Fissan,<sup>47</sup> Prototype of Model NAS 3089, TSI-Inc., Shoreview, USA). Area specific particle release numbers were calculated according to Göhler *et al.*<sup>21</sup> on the base of measured particle size distributions (total size range 5.6 nm–20 µm) and particle number concentrations as well as realised experimental conditions. To enable sanding with the given apparatus, the sample specimens were converted at first into practicable forms. The dish-shaped samples of the non-weathered cement composites were cut by a table saw into ashlar-formed subsamples (20 mm × 70 mm × 10 mm), while adequate fragments of the destructed cuboids were fixated within a silicon elastomer to receive a plane-parallel sample surface for testing. Furthermore, the at least 28 d cured cement/concrete samples were prior the analyses thermal equilibrated, *i.e.*, the samples were dried for ≥24 h at 45 °C and subsequently cooled to room temperature within desiccators.

## 3. Results and discussion

As mentioned before, this study deals with the effects on the binder matrix of cementitious materials. Accordingly, most of the following results apply solely for hardened cement pastes (solely cement, water and additives) that must not be confused with mortar (contains additionally sand) or concrete (contains mortar and gravel).

### 3.1 Material properties

**3.1.1 Phenomenological condition after artificial freeze–thaw weathering.** The artificially freeze–thaw weathering caused considerable but different progressed deteriorations of each composite, as exemplarily shown in Fig. 1 by two of five weathered specimen each. The worst visual deteriorations were observed for CEM\_GGBS\* followed by CEM\_hyd\* and CEM\_SiO<sub>2</sub>\*, while CEM\_HKP-GGBS\* showed the lowest ones. CEM\_GGBS\* and CEM\_hyd\* are clearly cracked and spalled, whereas CEM\_SiO<sub>2</sub>\* and CEM\_HKP-GGBS\* show a lower internal damage and less crack formation.

Crack formation and fragment spalling are mainly attributed to the volume expansion of water from residual moisture during freezing that pushes the cement matrix apart if there is lack for compensation due to low porosity. Likewise, a crack formation is also possible due to shrinkage during hydration. The availability of water within the samples based typically on residual moisture from sample preparation (*w/z* ratio of 0.4) prior freeze–thaw weathering and water







Fig. 1 Test specimens of a) CEM\_hyd\*, b) CEM\_GGBS\*, c) CEM\_HKP-GGBS\* and d) CEM\_SiO<sub>2</sub>\* after artificial freeze-thaw weathering.

absorption by capillary suction after sample curing during freeze-thaw weathering.

It is believed that CEM\_GGBS contained the highest residual moisture due to its composition, which causes a high internal pressure during freezing and lead to high deteriorations. The water absorption by capillary suction of CEM\_GGBS should be lower than for CEM\_hyd since it is assumed that CEM\_GGBS has a denser surface due to filler presence. The reduced deterioration of the samples CEM\_SiO<sub>2</sub>\* and CEM\_HKP-GGBS\* is attributed to their filler-induced high surface density that should go along with a reduced water absorption by capillary suction.

Beside the volume expansion of water from residual moisture during freezing, hydration products (see below) can also contribute to crack formation, for example due to the formation of voluminous ettringite structures (sulphate attack) that can be attributed to increased proportions of tricalcium aluminate in the processed cement.

**3.1.2 Surface structure and hydration products (SEM image analysis).** The hydration of cement, which contains tricalcium silicate (Ca<sub>3</sub>SiO<sub>5</sub>, alite, C<sub>3</sub>S), dicalcium silicate (Ca<sub>2</sub>SiO<sub>4</sub>, belite, C<sub>2</sub>S), calcium aluminoferrite (Ca<sub>2</sub>(Al,Fe)<sub>2</sub>O<sub>5</sub>, brownmillerite, C<sub>4</sub>AF) and tricalcium aluminate (Ca<sub>3</sub>Al<sub>2</sub>O<sub>6</sub>, C<sub>3</sub>A) as well as calcium sulphate (CaSO<sub>4</sub>, C<sub>5</sub>), lead to the formation of different hydration products like calcium silicate hydrates (CSH, C-S-H: flakes, dendrites, gel), calcium hydroxide (Ca(OH)<sub>2</sub>, portlandite, CH: plates) and hydrous calcium aluminium sulphate (Ca<sub>6</sub>Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>(OH)<sub>12</sub>·26H<sub>2</sub>O, ettringite, C<sub>3</sub>A·3C<sub>5</sub>·H<sub>32</sub>, Aft: acicular, striated prismatic). The mentioned hydration products were observed for both the non-weathered and the artificially weathered cement/concrete samples.

In this context, Fig. 2 shows selected SEM-images regarding surface structure and hydration products on the example of the weathered cement composites. In contrast to

CEM\_hyd\*, SEM-images show that CEM\_GGBS\*, CEM\_HKP-GGBS\*, CEM\_SiO<sub>2</sub>\* and CEM\_TiO<sub>2</sub> have a denser cement stone, which is attributed to different phenomena. The activation of HKP-GGBS increases significantly its reactivity that lead to a denser matrix.<sup>48</sup> A similar if not the same effect can be attributed to the increased pozzolanic reaction in the case of CEM\_SiO<sub>2</sub>, where silica reacts with portlandite to additional CSH structures. Furthermore, slag-containing cements form denser phases at the end of strength development than Portland cements.<sup>48</sup> Moreover, the denser cement stone is also attributed to the physical filler effect, whereof fine particles can fill the voids between coarser particles.

Furthermore, SEM-images based on artificially weathered cement composites showed also the formation of unwanted calcium carbonate structures (CaCO<sub>3</sub>, calcite as well as vaterite), which grow during the weathering procedure due to the contact of portlandite with carbon dioxide (CO<sub>2</sub>) from the environmental atmosphere. The later ones could be observed especially for CEM\_HKP-GGBS\* and CEM\_SiO<sub>2</sub>\* and are attributed to the high surface area reactivity of the processed additives. For CEM\_SiO<sub>2</sub>\*, this is due to the pozzolanic reaction of silica. In the case of CEM\_HKP-GGBS\* it is caused by mechanochemically modified phases of GGBS and the resulting changed reactivity. Since different phases are formed in CEM\_GGBS\* and CEM\_HKP-GGBS\*, the changed reactivity of HKP-GGBS by tribochemistry is obvious. Additionally, voluminous halite (NaCl) structures, whose formation is attributed to the de-icing saline solution and which could lead to a higher porosity by matrix displacement could be observed for CEM\_SiO<sub>2</sub>\*.

Beside hydration products, SEM-images of artificially weathered samples showed also the presence of micro-cracks, whose formation is already discussed in the previous subchapter.





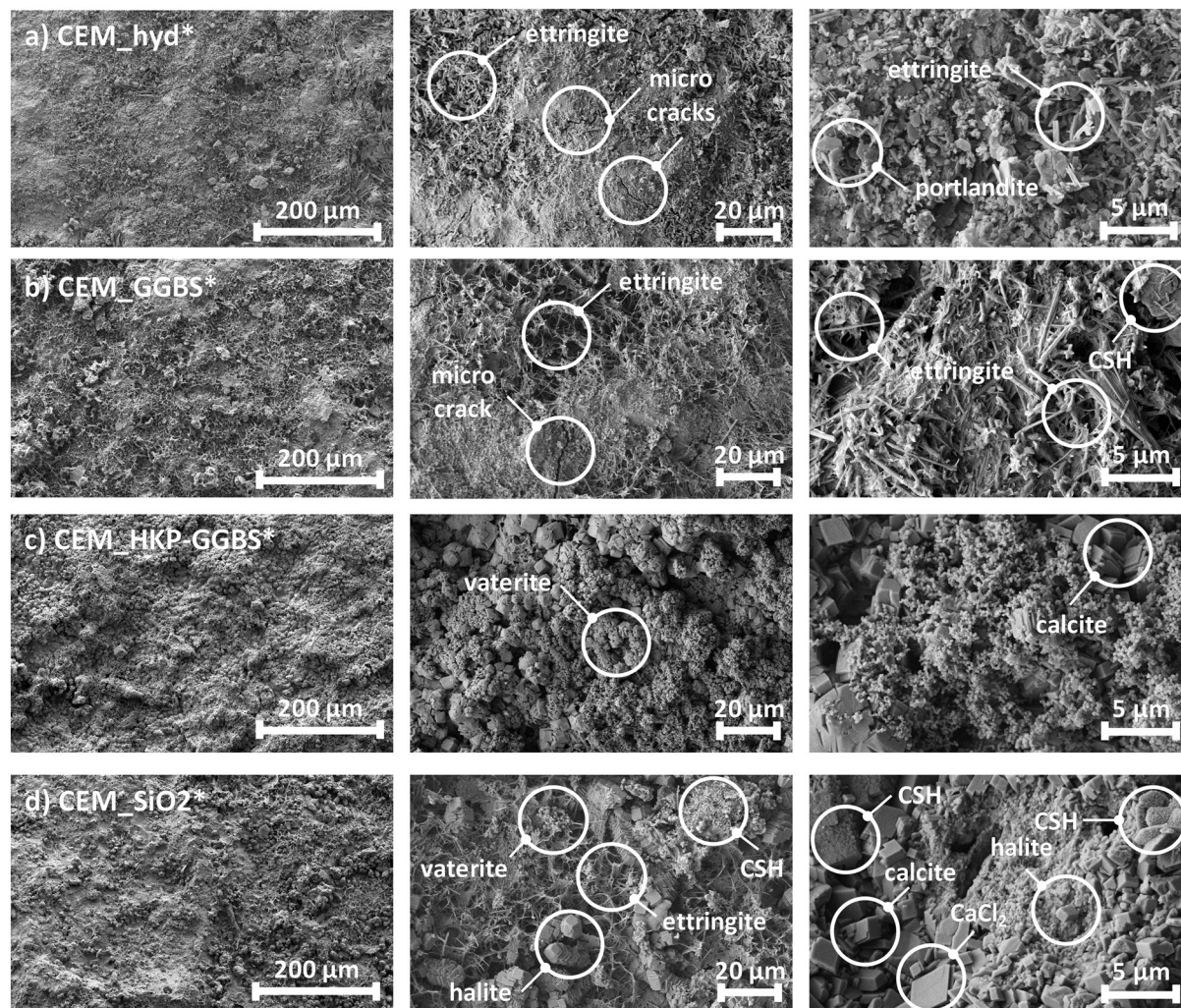


Fig. 2 SEM-images of surface structure with identified hydration products after artificial freeze-thaw weathering based on three different magnifications (300 $\times$ , 1500 $\times$ , 7500 $\times$ ) for a) CEM\_hyd\*, b) CEM\_GGBS\*, c) CEM\_HKP-GGBS\* and d) CEM\_SiO<sub>2</sub>\*.

**3.1.3 Porosity and water absorption behaviour.** The pore size distributions of the artificially weathered cement composites are provided in Fig. 3a), whereof also integral pore volumina of the three pore types (*i.e.*, air pores, capillary pores and gel pores) according to Smolczyk and Romberg<sup>49</sup> were determined (see also Table S2<sup>†</sup>). Furthermore, Fig. 3b) shows the corresponding porosity depended water absorption behaviour of these samples. According to Fig. 3a), the pore size distributions of all weathered samples vary considerably. While all cement composites show nearly the same low content of coarse air pores ( $x_p > 10 \mu\text{m}$ ), more significant differences could be observed concerning capillary pores ( $0.03 \mu\text{m} \leq x_p \leq 10 \mu\text{m}$ ) and gel pores ( $x_p \leq 0.03 \mu\text{m}$ ).

The lowest total pore volume was determined for CEM\_hyd\* ( $81.6 \mu\text{L g}^{-1}$ ), whereas slightly higher values result for CEM\_HKP-GGBS\* ( $82.9 \mu\text{L g}^{-1}$ ), CEM\_GGBS\* ( $83.3 \mu\text{L g}^{-1}$ ) and CEM\_SiO<sub>2</sub>\* ( $83.9 \mu\text{L g}^{-1}$ ). CEM\_GGBS\* showed the highest content of air pores (9.9 vol-%), followed by CEM\_HKP-GGBS\* (8.0 vol-%), CEM\_hyd\* (7.6 vol-%) and CEM\_SiO<sub>2</sub>\* (7.0 vol-%). The highest amount of capillary

pores was determined for CEM\_hyd\* (34.4 vol-%), followed by CEM\_HKP-GGBS\* (29.1 vol-%) and CEM\_SiO<sub>2</sub>\* (27.7 vol-%). Despite a high content of  $1.0 \mu\text{m}$  pores, CEM\_GGBS\* showed the lowest amount (12.3 vol-%) of capillary pores. In contrast, CEM\_GGBS\* showed the highest gel pore volume (77.8 vol-%), followed by CEM\_SiO<sub>2</sub>\* (65.3 vol-%), CEM\_HKP-GGBS\* (62.9 vol-%) and CEM\_hyd\* (58.0 vol-%).

The results of both porosity and water absorption (Fig. 3) are in line with the presumptions based on the phenomenological observations (Fig. 1) for the cement composites after artificial freeze-thaw weathering. CEM\_GGBS\* showed the highest deteriorations of all samples, which are attributed mainly to the volume expansion of freezing residual water and low compensation volume. This is consistent with the findings that CEM\_GGBS\* contains in comparison to the other samples the lowest content of capillary pores (as well as the lowest water absorption behaviour) and the highest content of gel pores. The structural different and less pronounced deteriorations identified for CEM\_hyd\*, CEM\_SiO<sub>2</sub>\* and CEM\_HKP-GGBS\* were attributed to be caused mainly



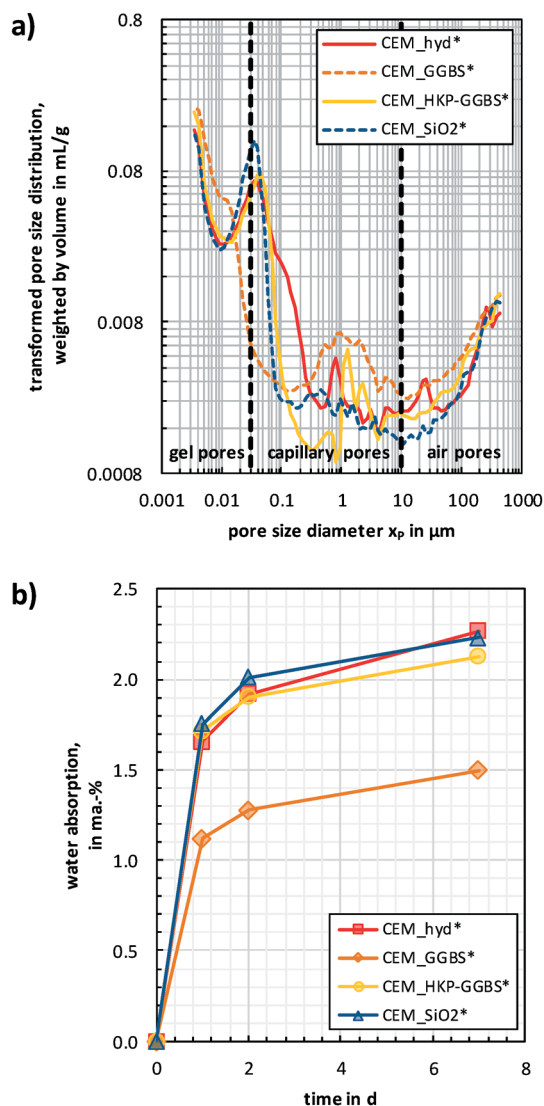


Fig. 3 Pore size distributions (a) and water absorption behaviour (b) of the artificially freeze-thaw weathered cement composites.

by freezing-induced volume expansion of water absorbed by capillary suction during artificial weathering. CEM\_hyd\* contained the highest amount of capillary pores that explains also its more progressed deterioration in comparison to CEM\_SiO<sub>2</sub>\* and CEM\_HKP-GGBS\*. Despite a slightly lower capillary pore volume, the weathering-induced deteriorations of CEM\_SiO<sub>2</sub>\* were more progressed than for CEM\_HKP-GGBS\*. This can maybe attributed to the slightly higher content of gel pores in CEM\_SiO<sub>2</sub>\* that accompany with a superposition of residual water based deteriorations.

**3.1.4 Thermal behaviour and water content.** The thermal behaviour of the artificially weathered cement composites (CEM\_i\*), which comprises also information concerning the content of both free and chemically bound water and thus on specific hydration products, is shown in the STA-thermogram (TG-DSC) of Fig. 4, while integral values of the relative mass change for certain temperature ranges are provided in the supplement (Table S3†).

The results obtained by STA are consistent with the ones of third parties.<sup>50–52</sup> Within the analysed temperature range (30–1000 °C), the DSC data show three characteristic peaks (100–200 °C, 450–550 °C, 650–750 °C), which correlate to significant gradations of the TG-curves. The first DSC-peak (100–200 °C) is based on the dehydration of both free water and chemically bound water from CSH. The second DSC-peak (450–550 °C) originates from the dehydroxylation of chemically bound water of portlandite (Ca(OH)<sub>2</sub>), while the third DSC-peak (650–750 °C) is based on the decarbonation of calcium carbonate (CaCO<sub>3</sub>, calcite).

The TG-curves in Fig. 4 and the values in Table S3† show that CEM\_GGBS\* has the highest content of both pore water and chemically bound water in CSH (mass change until 400 °C). In contrast, the lowest content was determined for CEM\_SiO<sub>2</sub>\* that is attributed to chemically bound water in silanol groups, which are formed due to the presence of the additive SiO<sub>2</sub>\_nano during hydration. The highest content of portlandite (mass change between 400–600 °C) was determined for CEM\_hyd\*, followed by CEM\_SiO<sub>2</sub>\*. The highest carbonate content (mass change between 600–800 °C) were observed for CEM\_GGBS\*, followed by CEM\_HKP-GGBS\*.

The results of STA (Fig. 4) confirm i) the presumptions based on the phenomenological observations (Fig. 1), ii) the observed hydration products (Fig. 2) and iii) the findings concerning porosity and water absorption (Fig. 3). For example, the advanced crack formation and fragment spalling of CEM\_hyd\* due to freeze-thaw weathering is attributed to the increased freeze-expansion of the high content of free water caused by the high content of capillary pores.

**3.1.5 Relative dynamic E-modulus (RDM).** Fig. 5 shows the RDM-change of the cement composites over the freeze-thaw cycles.

Over all samples, the RDMs decrease steadily with increasing weathering cycle, but with different slope. The RDMs of

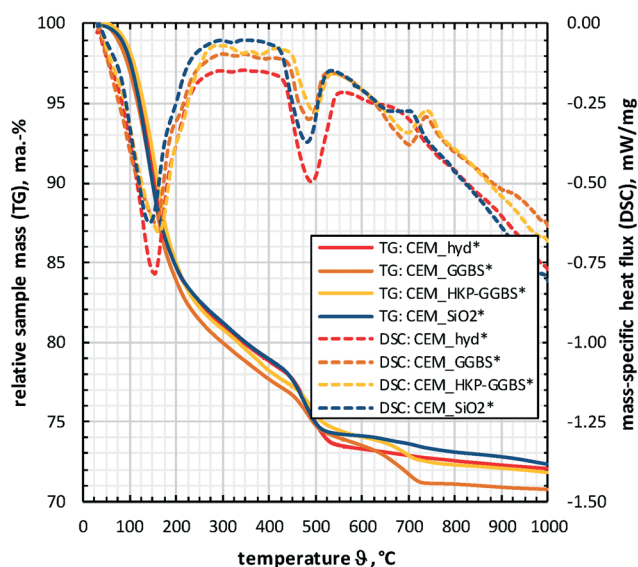


Fig. 4 STA-thermogram (TG-DSC) of the artificially weathered cement composites.





CEM\_SiO<sub>2</sub>\* and CEM\_HKP-GGBS decrease successively from 100% to 35–40% until the end of artificial freeze–thaw weathering. Due to the considerable deteriorations of CEM\_GGBS\* and CEM\_hyd\* no RDM analyses could be performed after the last cycle of artificial freeze–thaw weathering. At FTC 14, the highest RDM-changes were observed for CEM\_hyd\*, followed by CEM\_HKP-GGBS\* and CEM\_SiO<sub>2</sub>\*, while CEM\_hyd\* shows the lowest ones.

### 3.2 Release behaviour

#### 3.2.1 Weathering-induced fragment release into water.

Fig. 6 shows the weathering-induced cumulative area-specific mass of released fragments over the freeze–thaw cycles.

Similar to the RDM analyses, weathering-induced fragment release into water of CEM\_GGBS\* and CEM\_hyd\* could solely characterised until FTC 14, since the structural deteriorations of these samples did not allow the performance of further analyses.

However, the cumulative fragment release increases with increasing freeze–thaw cycle over all cements. CEM\_HKP-GGBS\* and CEM\_SiO<sub>2</sub>\* showed a very similar release behaviour to each other with quasi-uniform slope (*i.e.*, quasi-uniform release quantity per cycle) over the whole weathering procedure. The fragment release behaviour of CEM\_hyd\* and CEM\_GGBS\* differ significantly from the ones of CEM\_HKP-GGBS\* and CEM\_SiO<sub>2</sub>\*, *i.e.*, CEM\_GGBS\* shows a higher slope and CEM\_hyd\* a lower one. At FTC 14, CEM\_GGBS\* showed the highest cumulative fragment release (183 g m<sup>-2</sup>), followed by CEM\_HKP-GGBS\* (108 g m<sup>-2</sup>) and CEM\_SiO<sub>2</sub>\* (106 g m<sup>-2</sup>), while CEM\_hyd showed the lowest one (71 g m<sup>-2</sup>).

Interestingly, the behaviour of weathering-induced fragment release into water correlate well with the RDM results given in Fig. 5. This circumstance becomes even more obvious by replacing the cumulative mass of fragments by the cumulative volume of fragments using the dry densities of the cement composites as provided in Table 1.

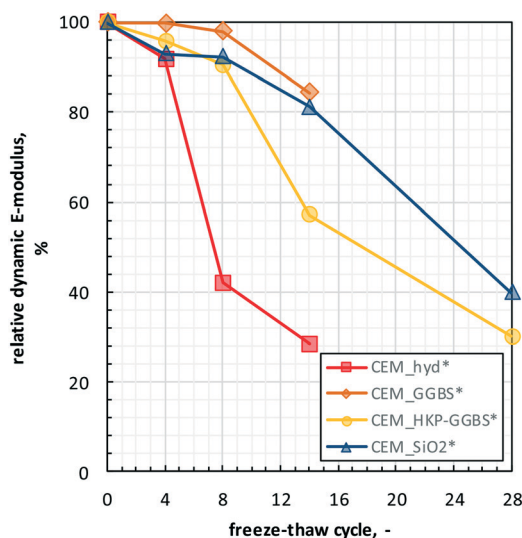


Fig. 5 RDM-change of the cement composites over selected freeze–thaw cycles (*i.e.*, FTC 0, 4, 8, 14, 28) of the artificial freeze–thaw weathering.

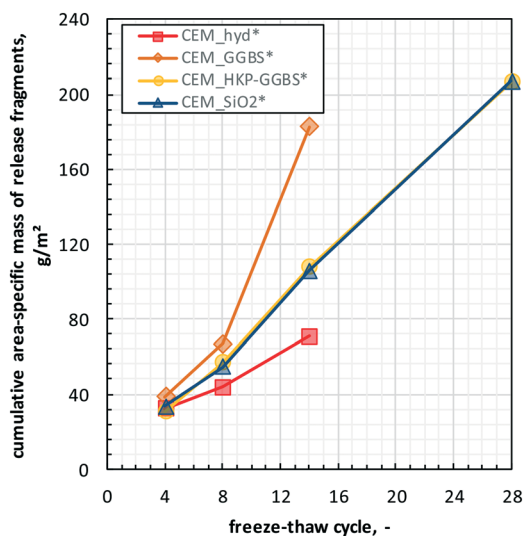


Fig. 6 Cumulative mass of released fragments during artificially weathering of the cement composites over the freeze–thaw cycles.

Finally yet importantly, result for fragment release based on freeze–thaw weathering show in contrast to UV/rain weathering<sup>34</sup> an increased release. This highlights that release testing should be performed with methods that are specific to the material and its susceptibility to the stresses that match the intended use. Roughly generalised, UV irradiation is critical for polymers as well as coatings, whereas freeze–thaw weathering is decisive for cements. Such guidances are implemented in the stepwise decision tree for release testing as developed by ISO TC229, PG29.

**3.2.2 Sanding-induced particle release into air.** In contrast to the weathering induced fragment release into water, sanding-induced particle release into air was characterised not during but before and after artificial freeze–thaw weathering.

The effect of the performed thermal sample equilibration on the relative surface humidity of the samples as well as the relative content of evaporated substances is shown in the supplement (Fig. S1†). The relative surface humidity of the non-equilibrated cement samples varied between 0.6 wt-% and 1.1 wt-% (0.95 wt-% ± 0.22 wt-%; RSD = 23.1%), whereof the values for the non-weathered cement composites varied between 0.7 wt-% and 1.1 wt-% (1.00 wt-% ± 0.19 wt-%, RSD = 19.4%). This is significantly higher than 0.6–0.8 wt-% (0.69 wt-% ± 0.10 wt-%, RSD = 15.0%) determined for the weathered samples. The relative surface humidity over all cement composites varied after thermal equilibration between 0.3 wt-% and 0.6 wt-% with a significant reduced spreading (0.53 wt-% ± 0.07 wt-%; RSD = 13.9%).

Fig. 7a) provides the sanding-induced area-specific swarf mass determined by gravimetric analyses of the whole sample specimens and comprises therefore the release of both airborne particles and coarse non-airborne fragments, while Fig. 7b) provides solely the area-specific number of released airborne particles. Considering at first the data before freeze–thaw weathering in Fig. 7a), neither the addition of SiO<sub>2</sub>\_nano nor the addition of TiO<sub>2</sub>\_nano showed a significant impact





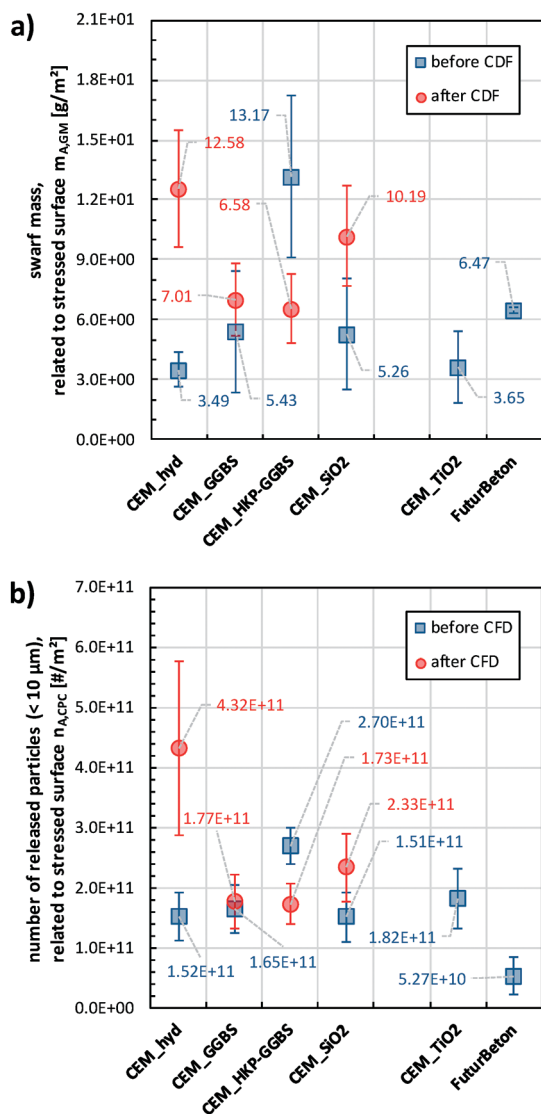


Fig. 7 Sanding-induced area-specific swarf mass based on sample gravimetry (a) and area-specific number of released particles based on condensation nuclei counting (b); error bars = standard deviation of at least eight measurements.

on the area-specific swarf mass as emanating from the comparison between CEM\_hyd, CEM\_TiO<sub>2</sub> and CEM\_SiO<sub>2</sub>. In contrast, the slag containing samples CEM\_GGBS (resp. CEM\_HKP-GGBS) show a slight (resp. considerably) higher area-specific swarf mass than CEM\_hyd. Furthermore, the area-specific swarf mass of FuturBeton is similar to the ones of CEM\_GGBS and CEM\_SiO<sub>2</sub>. The freeze–thaw weathered samples CEM\_hyd\* and CEM\_SiO<sub>2</sub>\* showed in comparison to their non-weathered counterparts an increase in the area-specific swarf mass, whereas quasi no changes were observed for CEM\_GGBS and a decrease in the case of CEM\_HKP-GGBS. After freeze–thaw weathering, CEM\_hyd\* and CEM\_SiO<sub>2</sub>\* showed the highest area-specific swarf mass. The ratios between the mean area-specific swarf mass of the weathered cement composites and their non-weathered counterparts, *i.e.*, the weathering-induced increase or decrease fac-

tor, are determined to be 0.50 for CEM\_HKP-GGBS, 1.29 for CEM\_GGBS, 1.94 for CEM\_SiO<sub>2</sub> and 3.61 for CEM\_hyd.

Fig. 7b) provides the results of airborne particle release characterisation, *i.e.*, the determined release quantities based on condensation nuclei counting (CNC). The area-specific number of released particles correlate in most cases well with the above-discussed results concerning the area-specific swarf mass (see Fig. S3†). The area-specific number of released particles varied before CFD weathering between  $2 \times 10^{10}$  #/m<sup>2</sup>– $4 \times 10^{11}$  #/m<sup>2</sup> and after CFD weathering between  $1 \times 10^{11}$  #/m<sup>2</sup>– $7 \times 10^{11}$  #/m<sup>2</sup>. The ratios between the mean area-specific numbers of released particles of the weathered cement composites and their non-weathered counterparts are determined to be 0.64 for CEM\_HKP-GGBS, 1.07 for CEM\_GGBS, 1.54 for CEM\_SiO<sub>2</sub> and 2.84 for CEM\_hyd. This is exactly the same order with similar weathering-induced release ratios as

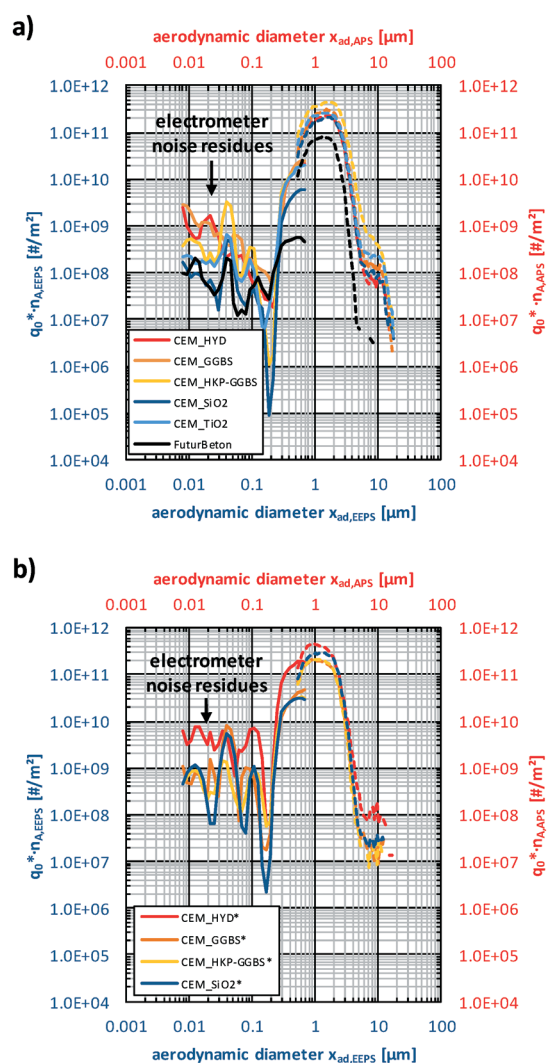


Fig. 8 Non-normalised, transformed distribution densities of released airborne particles over aerodynamic equivalent diameter before (a) and after (b) CFD weathering. EEPS data with electrometer-noise-correction, APS data with non-Stokesian-correction and EEPS equivalent-diameter-conversion using a mean effective particle density of  $1.76 \text{ g cm}^{-3}$ , solid lines = EEPS data; dotted lines = APS data.

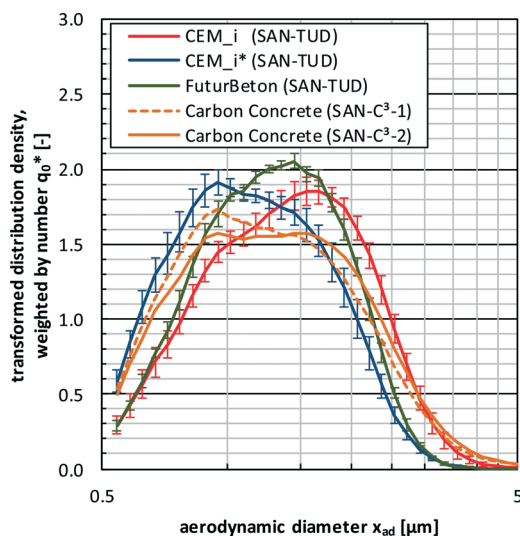


Fig. 9 Averaged normalised size distributions with standard deviation based on TOF over all non-weathered cement composites (CEM\_i;  $n = 8$  analyses  $\times$  5 composites = 40), weathered cement composites (CEM\_i\*;  $n = 8$  analyses  $\times$  4 composites = 32) and FuturBeton ( $n = 8$  analyses) in comparison to workplace measurements data of high-strength, double-layered carbon concrete determined by Bienkowski *et al.*<sup>31</sup>

determined for the area-specific swarf mass rate. In contrast to the cement/concrete swarf mass ratios (0.5–2.0), the cement/concrete particle release ratios varied from 2.9 up to 5.1, *i.e.*, the concrete FuturBeton released significant less particles than the cement composites. All these findings are in line with the release data determined by TOF, which cover a nominal size range of 0.5–20  $\mu\text{m}$ , as shown in the supplement (Fig. S2†).

Fig. 8 shows the mean non-normalised particle size distributions (PSD) of the released airborne particles over the aerodynamic equivalent diameter based on DEMA and TOF for all

analysed samples on logarithmic-graduated ordinate, while corresponding PSDs with linear-graduated ordinate are provided in the supplement (Fig. S5†). The supplement contains also additional notes on the performed correction and conversion procedures for the given PSDs.

However, Fig. 8 shows at first that the number-weighted PSDs determined by DEMA supplement well the ones of TOF, whereby the highest amounts (*i.e.*, >50%) of released particles were detected within the measurement range of TOF (see also parity plot in Fig. S4†). Furthermore, the PSDs show nearly no release quantities below 200 nm. Excepting the PSD of FuturBeton, all PSDs contain a characteristic peak around 10  $\mu\text{m}$ . The comparison of the central peak positions before and after freeze–thaw weathering of the cement composites show a slight weathering-induced shift towards finer particles. Since there are nearly no differences in the normalised PSDs between the different non-weathered cement composites (CEM\_i) and between the different weathered ones (CEM\_i\*), Fig. 9 provides the mean number-weighted PSDs based on TOF for CEM\_i, CEM\_i\* in comparison to FuturBeton. Here, the shift towards lower sizes of the weathered cement composites CEM\_i\* in comparison to their non-weathered counterparts became more visible. Additionally, Fig. 9 provides TOF-PSDs determined during workplace measurements based on real professional sanding (diamond based sanding machine DG150, Hilti) of carbon concrete composites as reported by Bienkowski *et al.*,<sup>31</sup> which are very similar to the ones determined in this study. Bienkowski *et al.* showed furthermore that the given size distributions correlate also well with the ones determined for professional stemming, cutting and drilling (core drilling, solid drilling), which are not shown here.

SEM-images of contemporaneous electrostatically precipitated airborne particles, as exemplarily shown in Fig. 10 for

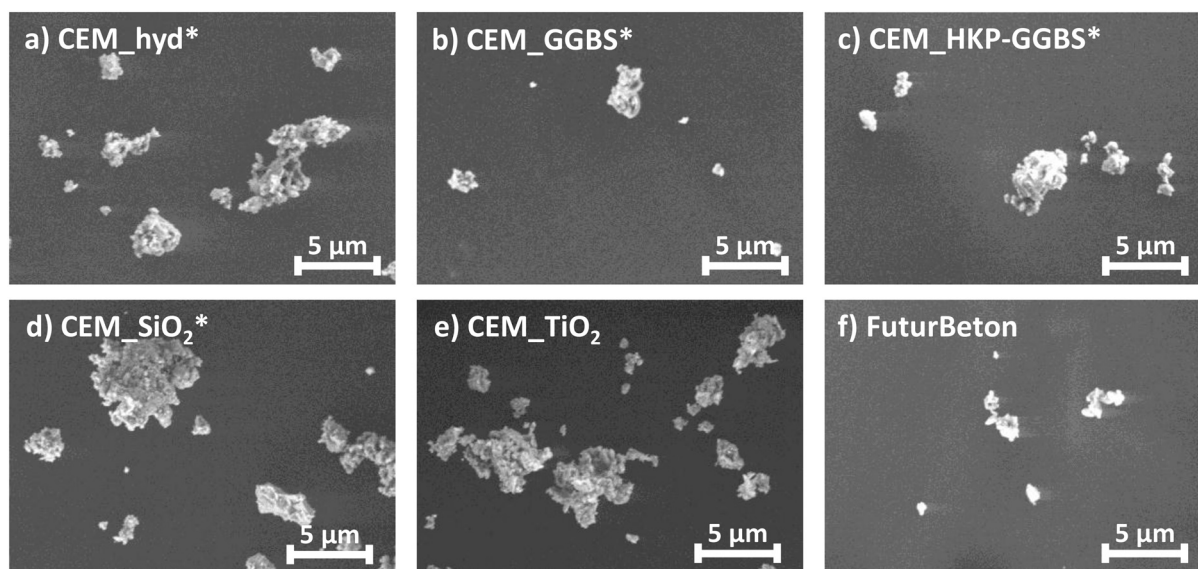


Fig. 10 SEM-images of electrostatic precipitated airborne particles released during sanding from a) CEM\_hyd\*, b) CEM\_GGBS\*, c) CEM\_HKP-GGBS\*, d) CEM\_SiO<sub>2</sub>\*, e) CEM\_TiO<sub>2</sub> and f) FuturBeton.



the different cement/concrete composites, confirm the findings from DEMA and TOF, *i.e.*, mean geometric size of released particle between 200 nm and 5  $\mu\text{m}$ . For the purpose of comparison, it should be noted here that the ratio between geometric particle diameter and aerodynamic diameter amounts to appr. 0.7.

## 4. Conclusion

To characterise the impact of weathering on integrity, internal structure and particle release from micro- and nanostructured cement composites (not to be confused with mortar or concrete), a comprehensive testing program was performed before, during and after standardised freeze–thaw weathering (CDF). Cement paste composite samples with different additives have been manufactured with non-compositional cement paste samples as reference.

Freeze–thaw cycling caused considerable deteriorations, which were significantly modulated by the different additives *via* changes of the cement micromorphology. The susceptibility to freeze–thaw deterioration decreased in the order: CEM\_GGBS > CEM\_hyd (no additive control)  $\gg$  CEM\_SiO<sub>2</sub> > CEM\_HKP-GGBS. The deteriorations are attributed to residual moisture and capillary sucked water in the pores that expands by freezing, and thus lead to crack formation and spalling. For CEM\_GGBS, the action by residual water dominated, whereas for CEM\_hyd the suction into capillary pores dominated. CEM\_SiO<sub>2</sub> and CEM\_HKP-GGBS were more resilient due to their moderate content of either gel or capillary pores and because their residual water content, which was reduced by the characteristic hydration products formed with the SiO<sub>2</sub> and HKP-GGBS additives. The loss of bulk mechanical integrity ranked in a different order: CEM\_hyd > CEM\_HKP-GGBS > CEM\_GGBS > CEM\_SiO<sub>2</sub>. Since the relative dynamic *E*-modulus describes a relative change with regard to the initial conditions of a material, it cannot predict micro- and nano-scale fragment release quantities.

Instead, the form, identity and rate of release were assessed directly. The highest fragment release into water was observed for CEM\_GGBS, as expected from the macroscopic deterioration. Released fragments were sharp-edged from CEM\_GGBS, but rounded-edged from the other materials. The sanding-induced airborne particle release was assessed prior and after freeze–thaw weathering, and were benchmarked against a cement containing TiO<sub>2</sub>-nano and against a high performance concrete (FuturBeton). Before freeze–thaw weathering, no modulation of airborne particle release quantities by the nanomaterial additives was observed, but the micron-scale GGBS additive induced a significant increase. In general, all the cements showed a significantly higher release than the tested concrete and then other materials such as coatings or plastics tested previously in the same setup.<sup>21</sup> Comparing the weathered samples to their non-weathered controls, the release by sanding shifted towards finer particles, and scaled in quantity by a factor of 0.6 (decrease: CEM\_HKP-GGBS); 1.1 (quasi no change: CEM\_GGBS); 1.5 (increase: CEM\_SiO<sub>2</sub>); 2.8 (increase: CEM\_hyd).

Beyond the above structure–property–activity relationships, that are specific to cement, we contributed here a case study on degradation processes of minerals that induce the release of fragments at the micro- and nanoscale, contributing to the environmental background of geogenic, biogenic and anthropogenic fragments at the micro- and nanoscale.<sup>53</sup> Nanomaterial additives had the potential to modulate the release rates, but the order of magnitude was primarily determined by the combination of degradation process and matrix.

## Authors' contributions

All authors discussed the initial outline of the manuscript. First literature collection was based on the field of research separately for each author.

BF (ZÖZ) performed sample preparation, laboratory analyses, data interpretation, drafting and critical revision for important intellectual content concerning cement pastes and concretes. DG (TUD) performed airborne particle release analyses, data evaluation, data interpretation, drafting and critical revision of the manuscript. MS (TUD) performed data interpretation and critical revision for important intellectual content of the manuscript. BS (IUTA) performed data interpretation and critical revision for important intellectual content of the manuscript. BS (BASF) performed CDF testing and associated analyses. WW (BASF) conducted study design, supervision of the study, data interpretation, drafting and critical revision of the manuscript.

## Endnotes

Wording and definitions of this article is based on the ISO/TS 80004 series of standards<sup>54</sup> and the recommendation of the European Commission<sup>55</sup> concerning the definition of nanomaterials. The graphical representation, the used symbols as well as the vocabulary to describe particle size distributions are in line with ISO 9276 series<sup>56</sup> and ISO 26824:2013.<sup>57</sup> In the case of transformed size distributions, the logarithm to the base of 10 was operated to calculate logarithmic class widths.

## Abbreviations

APS	Aerodynamic particle sizer
BET	Mass-specific surface area based on Brunauer–Emmett–Teller theory
CEM_i	Collective designation for all non-weathered cement composites
CEM_i*	Collective designation for all weathered cement composites
CDF	Capillary suction of de-icing solution and freeze thaw
CNC	Condensation nuclei counting
CPC	Condensation particle counter
CSH	Calcium silicate hydrates
DEMA	Differential electrical mobility analysis
DSC	Differential scanning calorimetry





EEPS	Engine exhaust particle sizer
FTC	Freeze–thaw cycle
GBS	Granulated blastfurnace slag
GGBS	Ground granulated blastfurnace slag
GM	Gravimetry
HKP	High kinetic processing
MIP	Mercury intrusion porosimetry
PSD	Particle/pore size distribution
RDM	Relative dynamic <i>E</i> -modulus
RSD	Relative standard deviation
SEM	Scanning electron microscopy
STA	Simultaneous thermal analysis
TG	Thermogravimetry
TOF	Time of flight spectrometry
XRF	X-ray fluorescence

## Conflicts of interest

The authors declare that they have no competing interest, neither in financial nor in non-financial regard.

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