

Article Early Age Performance of OPC-GGBFS-Concretes Containing Belite-CSA Cement Cured at Sub-Zero Temperatures

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Abstract: This study determined how replacing sodium nitrate-based antifreeze admixture (AF) with belite-calcium sulfoaluminate (belite-CSA) cement affects the early age properties of ecological concretes based on ordinary Portland cement (OPC) and ground granulated blast-furnace slag (GGBFS). Concrete specimens were cured at -15 °C and treated in various ways before testing, i.e., no treatment, stored at 20 °C for 12 and 24 h. Generally, the addition of belite-CSA cement shortened the setting time due to the rapid formation of ettringite. The incorporation of 25 wt% of antifreeze admixture (AF) to the OPC-GGBFS concrete cured at -15 °C partially inhibited ice formation and enabled the continuation of hydration processes. This trend was observed for all samples, independent of the applied AF after freezing curing. On the contrary, the addition of 20 wt% of CSA failed to inhibit the ice formation and increased the risk of frost damage for concretes despite the treatment after freezing. These concrete specimens had lower hydration, lower strength, and a more porous binder matrix. The microstructure of the binder matrix was significantly affected by the amount of CSA and extreme negative curing, followed by no notable recovery post-curing at room temperature. Therefore, pre-curing at room temperature for at least 6 h has the potential to avoid frost damage. Concrete containing 25 wt% AF combined with 12 h and 24 h of curing at 20 °C after removal from freezing and prior to testing could enhance the compressive strengths of all concretes. The renewed hydration was indicated as the main influencing factor.

Keywords: ordinary Portland cement (OPC); calcium sulfoaluminate cement (CSA); ground granulated blast-furnace slag (GGBFS); hydration; microstructure—SEM; antifreeze admixture (AF); differential scanning calorimetry (DSC); negative temperature; compressive strength; porosity; UPV

1. Introduction

Winter construction is a very arduous process, particularly in areas where the temperature remains below 5 °C for many months. The three most frequently faced problems are the delay in formwork removal, slower strength development, and freezing of mixing and pore water. At lower temperatures, the hydration rate of the cement is significantly delayed with prolonged setting time and a 20-40% decrease in strength [1–3]. When the temperature drops to -4 °C or lower during freezing, approximately 92% of the fresh water in the concrete's binder matrix is converted to ice, followed by hydrostatic pore pressure [4]. Consequently, a 9% increase in concrete volume, as well as internal and surface microcracks are observed [5–8]. Moreover, pore size and cooling rates are crucial in determining the phase transition of pore water. Larger pore size and slower cooling rates result in faster freezing due to the sufficient time for the ice front to spread and to the low pore vapor pressure [9-13]. A compressive strength of 3.5 MPa developing at an early age considerably limits frost damage. Nevertheless, to overcome such situations, a number of precautionary measures are recommended when the air temperature falls below 5 °C or is below 10 $^{\circ}$ C for at least 12 h throughout the day [14–17]. These include heating the mixing water or aggregates, covering concrete with isolative blankets, or adding heating



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Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). systems to the formworks [18]. Moreover, these prototypes not only increase the total cost of production but also account to add in total CO_2 emissions. The most effective and economical alternative is to use accelerating and/or antifreeze chemical admixtures, which accelerate hydration and increase the early strength, thus preventing the mixing water from freezing and reducing the internal pressure of expanding ice [11,19–23]. Antifreeze agents commonly react with calcium hydroxide to form hydroxy salts, which change the pH and ionic strength. Yet, because of the fusion of the interacting components and chemicals, estimating the eutectic temperature for concrete is relatively complex [24–26].

As commonly known, for every ton of manufactured OPC, between 0.5 and 0.8 tons of CO₂ are emitted, which account for 6–7% of the world's carbon footprint [27,28]. To reduce the negative impact caused by the cement industry and promote sustainable construction, supplementary cementitious materials (SCMs) such as ground granulated blast furnace slag (GGBFS), fly ash (FA), and silica fume (SF) are partially or entirely replacing the OPC. SCMs can delay or accelerate the hydration of OPC, preferably observed at either ambient or accelerated curing temperatures. At freezing temperatures, hydration is further significantly delayed or even inhibited, causing permanent damage to the binder matrix [23,29]. In these circumstances, in OPC-GGBFS systems, the usage of chemical antifreeze agents is advantageous and can considerably delay the formation of pore ice [26]. The ions present in the antifreeze agent often establish an aqueous path within the binder matrix and limit the hydrostatic pore pressure forming during the formation of ice [30,31].

Another potential alternative to accelerate hydration processes at low temperatures is the use of calcium sulfoaluminate (CSA) cement. These cements are known for their rapid setting, favorable antifreeze properties, and excellent anti-permeability [11,32–34]. Additionally, CSA required roughly 60% less limestone for its production in comparison with OPC. The clinkering temperatures are lower, ranging from 1250 °C to 1350 °C, with the ability to be lowered to 1100 °C [35,36]. Thus, it exhibits relatively lower density and needs lower grinding energy followed by lower CO_2 emissions [34,37–39]. The CSA is mainly composed of ye'elimite, also called Klein's salt (C_4A_3S), ranging up to 70 wt%, calcium sulfate (anhydrite) (CŜ) 0–25 wt% and belite (C₂S) up to 25 wt% [40]. Calcium monosulfate aluminate (AFm) (C_4ASH_{12}) and gibbsite (AH₃) are the main hydration phases in the reaction of ye'elimite with water. Yet, the presence of anhydrite causes the production of ettringite (AFt) ($C_6AS_3H_{32}$) and gibbsite, resulting in high early strength, rapid drying, and non-expansive large ettringite crystals [32,41,42], while the presence of lime attributes expansive, self-stressing and shrinkage resistance binders [43]. Moreover, the later age strength is often pronounced on optimized anhydrite usage because of slower dissolution kinetics instead of gypsum's faster dissolution [44,45]. Nevertheless, despite the positive impact of the CSAs, it is limited to partial replacements as the main binder from the cost perspective and limited manufactures.

Extensive research on OPC partially replaced by CSA has been studied, resulting in pronounced setting time and strength development as a fast synthesis of AFt and amorphous hydrates [43,46]. The formed hexagonal-shaped thin ettringite plates filled the pore system, consumed excess pore moisture, prevented the large-size ice crystals, minimized the risk of frost damage, and further densified the binder matrix [11,47]. Furthermore, instead of C-S-H and portlandite, the presence of gibbsite in OPC-CSA systems promoted the synthesis of strätlingite (C_2ASH_8) and portlandite via its interaction with alite (C_3S) [48,49]. On the contrary, the belite-rich CSA as the sole binder promoted only the strätlingite (C_2ASH_8) , characterized as slow strength development on reaction with belite (C_2S) and gibbsite (AH_3) [50]. The optimized dosage of CSAs and pre and/or no-standard curing for cold weather pronounced the strength of concrete by 300% placed at -5 °C [11]. Conversely, a 65% reduction in strength was reported for CSA-based specimens cured at -5° for a period of 28 days in comparison to the standard cured concrete, despite the faster hydration and strength development of the CSA [51]. The maximum replacement amount has been up to 20 wt% [11]. The presence of CSA in the OPC system does not change the hydration process of C_3S but rather modulates aluminate dissolution [52,53]. When compared with

a pure OPC system, blended OPC-CSA hydrates faster with a $2.5 \times$ larger first heat-peak flow [11,47,54]. The primary factors were faster wetting, dissolution, and partial hydration. Additionally, sulfates and free lime concentrations were found to be higher in OPC-CSA systems, which resulted in a higher heat release [52]. Gypsum (CSH₂) from OPC accelerated the dissolution of anhydrite (CS) and enhanced the hydration kinetics of ye'elimite [39]. Ettringite crystals were the only reaction products generated on the hydration of ye'elimite in the presence of calcium hydroxide, calcium sulfate, and water. After the complete consumption of calcium sulfates and depletion of calcium hydroxide, the main reactions include the formation of calcium monosulfate aluminate (AFm) (C₄ASH₁₂) and gibbsite (AH₃) as the principal hydration phases on interaction of ye'elimite with water [55].

The tertiary OPC:GGBFS:CSA concrete is yet another approach to minimize the environmental impact. The main hydration phases were the ettringite, monosulfate, gibbsite, and strätlingite found to be comparable to pure CSA-based concrete [36,43,56]. The CSA-based systems showed a lower risk of alkali-silica reaction (ASR) and increased resistance to sulfate attack [43,57]. However, numerous studies found that the dilution effect and limited reactivity of GGBFS in binary GGBFS-CSA mixtures resulted in an overall decline in mechanical properties and durability [36,58,59]. Furthermore, its behavior is unknown when exposed to sub-zero curing conditions. The present study compares the phase transition of the OPC-GGBFS concrete system blended with belite-CSA mineral accelerator cured at -15 °C to that of a similar concrete system blended with chemical antifreeze admixtures. The primary aim was to assess the ice formation and its influence on the microstructure of the binder matrix. Low-temperature differential scanning calorimetry (LT-DSC), ultrasonic pulse velocity (UPV), Scanning Electron Microscopy (SEM), and additional testing on the physical and mechanical properties were performed.

2. Materials and Methods

Rapid hardening Portland cement type CEM I 52.5R identified as Snabbhårdnande cement (SH) from Cementa (Malmoe, Sweden), Merox AB-Sweden (Oxelösund, Sweden) ground granulated blast furnace slag (GGBFS), and commercially available belite-based CSA cement identified as Calumex BeliCem (CSA) from Caltra Nederland B.V. (Mijdrecht, The Netherlands) were used. The used CSA cement contained about 25 wt% of belite (C₂S), Ye'elimite (C₄A₃Ŝ) content over 60 wt% and calcium sulfate (CŜ) at 10 wt%. Tables 1 and 2 show the physical properties and chemical compositions of the used binders and fillers. Jehander Heidelberg Cement Group supplied natural granite aggregates with sizes of 0–4 mm and 4–8 mm. A fine quartz filler Norquartz-45 from Sibelco Nordic (Lille-sand, Norway) and fine sand with a maximum particle size of 150 μ m were provided by Baskarpsand AB (Habo, Sweden). BASF (Rosersberg, Germany) polycarboxylate etherbased superplasticizer "MasterGlenium ACE 30" (SP) was used, and the water to binder (w/b) ratio was 0.42 for all concrete mixes and 0.31 for pastes.

Binder	Туре	Producer	Fineness cm ² /g	Density g/cm ³		
OPC	CEM I 52.5R	Cementa	5400	3.2		
Slag	GGBFS	Merox AB	4350	2.9		
CSĂ	Belite-CSA	Caltra Nederland B.V	5000	2.9		

The replacement of OPC and/or GGBFS by CSA cement was limited to 20 wt%. Sodium nitrate-based antifreeze admixture "MasterSet AC 220" (AF), accelerator "Master X-Seed 130" (ACC), and air entraining "MasterAir 105" (AE) admixtures from BASF (Rosersberg, Germany) were used [26]. SP, AE, AF, and ACC dosages were held constant at 0.75 wt%, 0.1 wt%, 25 wt%, and 10 wt% of the binder weight, respectively [26]. The amount of water in the chemical admixtures was deducted from the added water.

Chemical (%)	OPC	GGBFS	CSA	Quartz Filler	Fine Sand (B15)
CaO	62.5	30.3	40-44	-	-
SiO ₂	18.8	34	6-10	99.6	90.5
Al_2O_3	5.4	11.6	>32	0.25	4.9
Fe ₂ O ₃	3.2	0.3	<2.2	0.02	0.5
MgO	1.3	12.1	<1.5	-	-
Na ₂ O	0.14	0.5	-	-	1.2
K ₂ O	1.3	0.8	-	-	2
SO ₃	3.7	-	9–10	-	-
Cl	0.03	0.3	-	-	-
LOI	2.6	-0.9	3	0.15	-

Table 2. Chemical compositions of the used binders and fillers.

Tables 3 and 4 show the paste and concrete compositions that were studied. Seven paste mixes and seven concrete mixes were produced. Based on the early age performances of the mixes (fresh and hardened properties), two concrete mixes were subsequently cast and cured at a constant temperature of -15 °C, as indicated in Table 4 (shaded in grey).

Table 3. Mix compositions of the pastes.

Mix	SH CEM I 52.5R (kg/m ³)	GGBFS (kg/m ³)	CSA (kg/m ³)	w/b	Chemical Admi (wt%)		Admixt t%)	ure
CSA (100%)	0	0	400	0.31	SP—0.75			
OPC (100%)	400	0	0	0.31	SP-0.75			
OPC:GGBFS:CSA (80:0:20)	320	0	80	0.31	SP-0.75			
OPC:GGBFS:CSA (0:80:20)	0	320	80	0.31	SP—0.75			
OPC:GGBFS:CSA (40:40:20)	160	160	80	0.31		SP-	-0.75	
OPC:GGBFS:CSA (50:50:0) (0%AF)	200	200	0	0.31	SP—0.75			
OPC:GGBFS:CSA	200	200	0	0.31	SP	AF	ACC	AE
(50:50:0) (25%AF)	200				0.75	25	10	0.1

Table 4. Mix compositions of the concretes.

Mix	SH CEM I 52.5R (kg/m ³)	Slag (kg/m ³) (l	CSA (kg/m ³)	w/b	Chemical Admixture	Filler (kg/m ³)		Aggregate (kg/m ³)	
					(Wt%)	Quartz	B15	0–4	4-8
CSA (100%)	0	0	400	0.42	SP-0.75	90	90	1075	537
OPC (100%)	400	0	0	0.42	SP-0.75	90	90	1075	537
OPC:GGBFS:CSA (80:0:20)	320	0	80	0.42	SP—0.75	90	90	1075	537
OPC:GGBFS:CSA (0:80:20)	0	320	80	0.42	SP0.75	90	90	1075	537
OPC:GGBFS:CSA (40:40:20)	160	160	80	0.42	SP0.75	90	90	1075	537
OPC:GGBFS:CSA (50:50:0) (0%AF)	200	200	0	0.42	SP-0.75	90	90	1075	537
OPC:GGBFS:CSA (50:50:0) (25%AF)	200	200	0	0.42	SP AF ACC AE 0.75 25 10 0.1	- 90	90	1075	537

Initial and final setting times were determined on the paste samples (Table 3) in accordance with the ASTM C191-19 [60]. The heat development due to the hydration reaction (Table 3) was examined using an in-house semi-adiabatic calorimeter. The calorimeter was made of Styrofoam, having a wall thickness of 60 mm, as shown in Figure 1. The PicoLog6 TC-08 instrument with a K-type thermocouple was adopted to record data for 7 days. Temperature was measured by placing the thermocouple into a freshly prepared paste sample with dimensions of 60 mm × 60 mm × 60 mm. The phase transition was studied using a low-temperature differential scanning calorimetry (LT-DSC) type Q100 TA instrument, see Figure 2. Immediately after mixing, paste samples weighing 10 to 12 mg were loaded and sealed in an aluminum crucible, which was placed in the DSC chamber along with an empty reference crucible. For both the freezing and thawing stages, the temperature change rate was 1 °C/min. Applied temperatures ranged from +10 °C to -30 °C. The behaviors of pastes in response to temperature changes were examined using either endothermic or exothermic heat flow. The freezing, melting, and glass transition points have all been determined [26].



Figure 1. In-house-built semi-adiabatic calorimeter box cross-section and test setup.



Figure 2. Differential scanning calorimetry (DSC)-Q100 TA.

Concretes were produced using a pan mixer type Zyklos, with 2 min of dry mixing followed by the addition of water blended with superplasticizer and another 2 min of mixing. Workability and compressive strength development were determined using the standards SS-EN 12350-2:2019 and SS-EN 12390-3:2019, respectively [61,62]. The test concrete cubes had dimensions of 100 mm \times 100 mm \times 100 mm. For the preliminary studies, freshly casted cubes covered with plastic foil were placed and cured at an ambient room temperature of approximately 20 °C, while for the main studies, the cubes were placed immediately after casting into the refrigerator and exposed to a constant temperature of -15 °C. The compressive strengths were measured at 1, 3, 7, 14, and 28 days. The loading rate was set to 10 kN/s. Furthermore, to study the impact of the formed pore ice—for each curing period, concrete samples were taken out of the refrigerator at 0 h before testing (instant), 12 h before testing (12 h/bf), 24 h before testing (24 h/bf), and held at room temperature during waiting period [26].

The standard SS-EN 12504-4:2004 following the ultrasonic pulse velocity (UPV) was used to detect the ice formation in the concrete cubes cured at a constant -15 °C [63]. Pundit Lab ultrasonic instrument (Proceq, Zurich, Switzerland) with exponential transducers at 54 kHz was used. The measuring path length was 100 mm. The tests were performed on the 1-day and 28-day-old concrete samples. The test measurements were registered at different time intervals, i.e., immediately after being taken out of the refrigerator (instant), 6 h after (6 h/af), 12 h after (12 h/af), 18 h after (18 h/af) and 24 h after (24 h/af). The concrete samples were stored at room temperature prior to testing.

To examine the microstructure, a scanning electron microscope (SEM)—JSM-IT100 (JEOL Ltd., Tokyo, Japan) was used. The tests were carried out on the 3-day and 28-day-old concrete samples. Cubes were removed from the refrigerator and held at room temperature for 0 h (instant), 12 h (12 h/bf), and 24 h (24 h/bf) before being prepared for the SEM examination. For analysis, concrete samples were extracted and placed in isopropanol for 24 h and another 24 h of drying in a low-vacuum chamber. Samples were impregnated under vacuum with a low-viscosity epoxy resin. After hardening, all samples were polished using grinding plates and three sets of diamond spray with a maximum particle size of 9 μ m, 3 μ m, and 1 μ m. The lamp oil was used as a lubricating and cooling agent [64]. SEM images were obtained using the backscattered electron (BSE) mode detector at 500× magnifications. The images were segmented to separate the porosities based on greyscale histograms [65,66].

3. Test Results and Discussion

3.1. Setting Time

Initial and final setting times are shown in Figure 3. Generally, mixes based on CSA cement showed shorter setting times than those based on OPC cement. Similarly, the mixes blended with CSA cement set faster than the reference OPC. The reference OPC mix had the initial and final setting times of 80 min and 530 min, respectively, see Figure 3. The binary and tertiary mix OPC:GGBFS:CSA—80:0:20 and OPC:GGBFS:CSA—40:40:20 had initial setting times of 9 and 7 min, respectively. This can be related to the fast hydration rate of CSA and the corresponding enhanced ettringite formation and early-age moisture loss [54]. Additionally, gypsum and calcium hydroxide from OPC could accelerate the hydration rate of CSA, as seen by others [41,53]. The initial setting time of the studied CSA cement was 31 min, which could be related to the slow reaction of β -C₂S belite at early ages forming strätlingite. The final setting time of 605 min was in line with the highly reactive α -C₂S as seen by others at later reaction periods [67–70].



Figure 3. Initial and final setting times of the pure and blended binders.

The partial replacement of GGBFS by 20 wt% CSA showed the initial setting time of 70 min. Despite the slow reaction of GGBFS, the CSA stimulated the hydration reaction. Similar amounts of ettringite, monosulfate, and Al(OH)₃ were formed but with a slightly more strätlingite [36]. On the contrary, the final setting time was delayed to 1227 min for a mix containing 80 wt% of GGBFS and 20 wt% of CSA. It was related to the reactivity of slag and a limited amount of formed AFt phases [36,59]. The OPC-GGBFS concrete with no chemical admixtures showed delayed initial and final setting times compared with similar mixes with chemical admixtures. This could be inferred from the reduced hydration heat release of GGBFS, correspondingly decelerated the hydration heat evolution intensity and delayed initial setting. However, the co-existence of calcium hydroxide from OPC hydration considerably reacted with slag to produce more C-S-H at later ages. Additionally, chemical admixtures incorporated a mix containing C-S-H seeds, boosted the hydration rate at early ages, and shortened the induction period and more C-S-H nuclei sites [71,72].

3.2. Hydration Heat Development

Figure 4 shows the hydration heat development of pastes in the first 7 days. All the mixes showed exothermic heat releases immediately after mixing with water and a shoulder formation before the main peak. Mixes containing CSA cement had shorter induction periods, see Figure 4 which equaled 140 min, 9 min, 8 min, and 25 min, respectively, for the pastes CSA (100%), OPC:GGBFS:CSA (80:0:20), OPC:GGBFS:CSA (40:40:20), and OPC:GGBFS:CSA (50:50:0) (25%AF). These trends corresponded to the measured setting times shown in Figure 3. This could be related to the faster dissolution and partial hydration of gypsum and ye'elimite forming ettringite and amorphous AH_3 [52,73]. The sulfates and calcium hydroxide originating from OPC are yet other factors enhancing the hydration of the OPC-CSA system [52]. As a result of the binding of gypsum sulfates to C_3A , a shallow peak equivalent to AFm was also found in the OPC-CSA mix pastes at a later age [74]. The longer induction period for the mix containing 100 wt% of the CSA could be related to the cement's chemical composition, especially the composition including 25 wt% of belite. This phase is known for its slow hydration, which presumably delayed the early-age reactions. The following ultimate peak temperature corresponded to the formation of more ettringite and strätlingite [70].



Figure 4. Semi-adiabatic temperature developments of the pure OPC, CSA, and blended OPC:Slag:CSA pastes.

As shown in Figure 4, the paste OPC:GGBFS:CSA (0:80:20) had a very low ultimate hydration peak despite the presence of the CSA cement. This could be related to the ye'elimite being replaced by slag [36]. Earlier research showed that decreased amounts of AFt, AFm, and AH₃ phases in systems containing higher amounts of GGBFS (>30 wt%) were linked to the dilution effect and low reactivity [36,58,59]. The paste OPC:GGBFS:CSA—50:50:0 mixes with no chemical admixtures showed a delayed hydration peak compared with other tested paste samples. A similar mix containing 25 wt% AF [OPC:GGBFS:CSA—50:50:0] showed a main but shallow peak much earlier after 60–65 min. The delay corresponded to the slow reactivity of slag.

3.3. Fresh and Hardened Concrete Properties (Early Age)

Mixes containing OPC:GGBFS (50:50 wt%) and 25 wt% of the AF showed a maximum slump of 20 cm, while the reference OPC concrete showed a minimum of 3 cm, see Figures 5 and 6. The workability was affected by the binder's fineness and incorporated chemical admixtures [49]. Fine binder, in conjunction with chemical admixtures, adheres more to the binder particles to pronounce the admixture effect and consequently shows higher workability. The formation of ettringite crystals at an early age could reduce it. The OPC:GGBFS:CSA (40:40:20) mix showed a higher slump of 13 cm compared with the mix containing OPC:CSA (80:20), which reached only 5 cm. Despite similar CSA dosages, the presence of slag was favorable due to its coarser particles and lower reactivity, while the mutual hydration reaction of OPC:CSA, the calcium sulfate, and calcium hydroxide originating from OPC accelerated the hydration of CSA at a very early age, resulting in the formation of ettringite crystals which caused a moisture loss [52,57].



Figure 5. Measured slumps immediately after mixing the concretes.

Pure OPC (100%), CSA (100%), and binary mix with OPC:CSA (80:20) concretes achieved a maximum 1-day compressive strength of 42 MPa, see Figure 6. The used belite-CSA cement contained 25 wt% of belite (C₂S), Ye'elimite (C₄A₃Ŝ) content over 60 wt%, and calcium sulfate (CŜ) at 10 wt%. Consequently, the strength gain could be attributed to the formation of more C-S-H/C₂ASH₈ and Portlandite but a limited formation of AFt phases [40,49]. Furthermore, relative to the CSA (100%) concrete, the binary blend OPC:CSA (80:20) showed lower strength at all ages except at 1 day. The differences in the strengths were 12.84% at 3 days and 10% at 7 days. The high early age strength was related to the rapid dissolution and accelerated hydration rate due to the co-existence of two types of cement and, consequently, the rapid formation of ettringite crystals [54]. However, these



overlapping ettringite deposits on the C-S-H particles over time caused an adverse effect on the microstructure by the occurrence of microcracks [49,54].

Figure 6. Compressive strength developments of the concrete cubes cured at ambient temperature.

The binary mix GGBFS:CSA (80:20) achieved the lowest compressive strengths between 6–7 MPa at all ages, which complies with earlier results [36,75,76]. Some connected the trend to the degradation of physiochemical properties of the CSA cement due to the presence of GGBFS [77]. Furthermore, the decreased strength development was also linked with the high GGBFS contents over 30 wt%, which could limit the formation of ettringite, strätlingite phase, and also an increased porosity [29,59,78]. Other researchers have shown that mixes containing less than 30 wt% of GGBFS could reach comparable strengths as pure CSA-based concrete [36]. Ettringite, monosulfate, strätlingite, and AH₃ were observed as the principal hydration products when the amount of slag added to CSA based binder system was optimal, consequently inhibiting the porosity growth and promoting densification of the concrete microstructure [36].

Similar strength values were measured at all ages for the concretes based on OPC:GGBFS (50:50) with 25 wt% AF and OPC:GGBFS:CSA (40:40:20). In both cases, the used AF admixture and CSA cement were the key elements for synergistic hydration reaction and increased strength. GGBFS, as a partial replacement of OPC, underwent a pozzolanic reaction with Ca(OH)₂ to produce more C-S-H gel corresponding to the later age strength. Additionally, the synthetic C-S-H seeds contained in the added chemical admixtures accelerated the hydration and early age strength. On the other hand, the presence of the CSA cement supported the formation of ettringite crystals and other phases (C-S-H/strätlingite, monosulfate, and AH₃) during the hydration reaction of ye'elimite, alite, and belite. This considerably amplified the early age strength. Other researchers observed that increasing dosages of CSA in combination with OPC and GGBFS enhanced the strength and durability properties [76]. Additionally, the usage of GGBFS promoted the cohesiveness of the matrix and densified the microstructure [43].

3.4. Differential Scanning Calorimetry (DSC)

The phase transitioning was determined using differential scanning calorimetry (DSC). Two events were observed, i.e., exothermic peak during freezing and endothermic peak during thawing, Figures 7 and 8, respectively. In general, based on the eutectic point of the used chemicals and/or mineral accelerators, the freezing point of an aqueous solution



can be determined [24,26]. However, due to the fusion of interacting components and chemicals, estimating the eutectic temperature for concrete is complicated.

Figure 7. Exothermic heat flow of the pure OPC, CSA, and blended OPC:GGBFS:CSA pastes at a freezing rate of 1°/min using DSC.



Figure 8. Endothermic heat flow of the pure OPC, CSA, and blended OPC:GGBFS:CSA pastes at a thawing rate of 1°/min using DSC.

For both freezing and thawing procedures, a single dominant peak was observed [13]. The OPC:GGBFS (50:50) mix with no chemical admixture showed the earliest freezing point of all mixes at around -12 °C. While the similar mix but with 25 wt% of AF admixture, the freezing point was lowered to around -29 °C. Reference mixes, OPC (100%) and CSA

(100%) showed the freezing point at -14 °C. While the mixes with the presence of 20 wt% of CSA cement further decreased this point to around -16 °C. The depressed freezing temperature, in all cases, could be associated with the dissolved ionic concentration in the pore solution. The latter "concentration effect" results in a decreased pore size and enhanced network structure, according to Raoult's law and Clausius–Clapeyron relation [11]. Pore solution present in larger pores freezes faster, whereas in smaller pores, the vapor pressure is increased, which depresses the freezing point [9,10]. The presence of CSA cement resulted in the formation of a higher amount of ettringite crystals in the first few minutes. Consequently, more water was absorbed, leading to an increase in the ionic concentration, followed by a decrease in the pore size, due to the formation of ettringite, and limited the ice formation [11,79]. Several studies showed an adverse effect for lower replacements of OPC with CSA cement, ≤ 10 wt%, on the hydration process and exposure to freezing temperature [11].

Similarly, upon thawing endothermic peak related to the energy absorption in pores where ice has melted can be seen in Figure 8. The OPC:GGBFS (50:50) paste without admixtures showed a well-defined peak at around 0 °C [30,80]. While all pastes containing CSA cement showed a dominant endothermic peak immediately after 0 °C and a shoulder before it. The OPC:GGBFS (50:50) mix containing 25 wt% of AF showed two peaks forming at different temperatures. This could be related to the geometry of the pore system and the concentration of the dissolved ions [7,13]. The early peak generated during melting could be related to the melting of ice located in finer pores, while the later peak to the ice formed in larger pores [13,81,82]. Melting of ice is a continuous process, but at extreme freezing temperatures, it is limited [5,83].

3.5. Compressive Strength Development—Concrete Cured at -15 °C

Further study focused only on two mixes, i.e., mix OPC:GGBFS:CSA—(40:40:20) and OPC:GGBFS:CSA—(50:50:0) (25%AF). The choice was based on various factors, including fresh and hardened properties, carbon footprint, and cost-effectiveness.

Figures 9–11 show the strength development of concrete cubes cured at constant -15 °C, measured at different intervals. In general, for all variants, the concrete mix OPC:GGBFS:CSA—(50:50:0) containing 25 wt% of AF had higher strength values in comparison with the concrete containing 20 wt% of CSA and no AF. The maximum ultimate 28 days compressive strength of 35 MPa was achieved for samples containing AF and exposed to 20 °C for 24 h before testing (24 h/bf), see Figure 11. Irrespective of the age and test intervals, the concrete samples with AF showed a continued strength increase for the entire measured 28-day period. This could be related to the use of high concentric AF admixture, which limited the pore ice formation and correspondingly supported the hydration process. The difference in the measured strength values between samples that were tested immediately after their removal from the freezer and samples that were held for 12 h at 20 °C tended to decrease with age in the case of concretes containing 25% AF, see Figures 9 and 10. This may be related to advancing hydration and, consequently weaker effect of the false strength due to the ice formation. The observed strength results for both mixes followed the trends observed in the differential scanning calorimetry test.

In general, the exposure to higher temperatures caused the pore ice to melt and consequently accelerated the hydration process, leading to higher strength values. On the contrary, no strength increase was observed for samples containing 20 wt% of the CSA cement. The measured values were between 10 MPa to 15 MPa, irrespective of age, Figures 9–11. The observed strength could be partially related to the initial hydration of CSA cement and larger amounts of formed ice [11]. Furthermore, concrete samples with 20 wt% of the CSA cement, subjected to 20 °C for 24 h prior to testing, showed a significant decline in compressive strength development with age, see Figure 11. This trend could be related to the formation of ettringite crystals that eventually resulted in more porous and microcracked microstructure, as seen by others [54]. Additionally, at a constant freezing temperature, the concentration and strength of the ice lenses have been observed to increase

exponentially [84]. Consequently, the internal pressure created by the developed pore ice could result in microcracking of the already-formed binder matrix [85–87]. Alzaza et al. reported that concretes with 40 wt% of OPC with CSA cement where no pre-curing was applied showed a limited strength development for concrete cured at -5 °C [79]. Similar investigations demonstrated that pre-curing for at least 6 h at room temperature could mitigate the harmful effects of the negative temperature [11,54].



Figure 9. Strength development of the "Instant" removed concrete samples cured at -15 °C.



Figure 10. Strength development of the "12 h/bf" removed concrete samples cured at -15 °C.

In comparison, the early age strength of concrete tested immediately after their removal from the freezer (1-day instant) and containing 20 wt% of the CSA cement was over 10 MPa versus 0 MPa for the concrete OPC:GGBFS concrete with 25 wt% of the AF. This clearly indicates that a portion of the strength was derived from formed ice as well as ettringite formation [11]. The presence of formed ice was later confirmed by the UPV measurements shown in Figures 12 and 13.



Figure 11. Strength development of the "24 h/bf" removed concrete samples cured at -15 °C.



Figure 12. Average Ultrasonic Pulse velocities (UPVs) were measured on the 1-day-old concrete samples.



Figure 13. Average Ultrasonic Pulse Velocities (UPVs) were measured on the 28-day-old concrete samples.

3.6. Ultrasonic Pulse Velocity (UPV)

To identify and validate the ice formation within the binder matrix, UPV measurements were performed on the 1-day and 28-day-old concrete samples kept at ambient temperature for varying periods after being taken out from the freezer, as shown in Figures 12 and 13. In the present study, a denser concrete microstructure and a higher pulse velocity (m/s) were related to either pore ice + binder matrix or binder matrix alone. In contrast, lower observed velocity (m/s) may indicate a less compact microstructure due to either the presence of empty pores and/or due to the presence of microcracks. Other researchers have measured the pulse velocities of 3828 m/s in solid ice and 1485 m/s in water [30,88]. Immediately after being removed from the freezer, the concrete samples containing 20 wt% of the CSA cement showed UPV values of 4768 m/s and 4510 m/s for the ages of 1 day and 28 days, respectively. These values decreased significantly over time on exposure to 20 °C, see Figures 12 and 13. These trends are related to the melting of formed ice [88]. Moreover, during longer freezing periods, ice nucleation can cause irreversible damage to the binder matrix [19,89]. Despite thawing for 12 h or longer, the concrete containing 20 wt% of the CSA cement showed the lowest increase in the UPV, a relatively negligible percentage growth compared with the concrete containing 25 wt% AF. After a thawing period of 24 h, the 1-day and 28-day-old samples incorporating 20 wt% of the CSA cement had an UPV of 4000 m/s. This could indicate that the water released from the melted ice promoted hydration of Portland cement, which products partially fill the pores. These results, in combination with the measured compressive strength values, further confirm that the incorporation of 20 wt% of the CSA cement did not prevent the formation of ice in pores.

The UPVs for the 1-day and 28-day-old concrete samples containing 25 wt% of the AF were considerably lower at 3367 m/s and 4464 m/s when measured immediately after being taken out of the freezer, see Figures 12 and 13. Exposure to ambient conditions increased the UPV, unlike observed for the concrete samples containing 20 wt% of the CSA cement. This could be directly related to the depression of the freezing point of water and, thus, to the limited ice growth [71,90]. The application of a thawing period of 24 h resulted in the maximum UPVs of 4587 m/s and 4695 m/s for the 1-day and 28-day-old concrete samples, respectively. The thawing accelerated the hydration process, which resulted in the formation of a dense binder matrix.

3.7. Scanning Electron Microscope (SEM)

Figures 14 and 15 show the SEM images of the concrete samples OPC:GGBFS:CSA— 40:40:20 and OPC:GGBFS:CSA—50:50:0 (25 wt.% of AF), which were cured at -15 °C for 3 and 28 days, respectively. The samples containing 20 wt% of the CSA cement contained more unhydrated particles regardless of the thawing periods, see Figures 14a–c and 15a–c. Due to the rapid dissolution of the CSA cement, the areas filled with ettringite were often surrounded by empty voids in all studied samples. In Figures 14 and 15, red color boxes indicate the areas with detected ettringite crystals, and yellow boxes indicate the unhydrated CSA cement. As described earlier, this could be linked to the significant amount of water consumed by forming ettringite, leading to moisture loss at the early stages of hydration [54]. The low reactivity of GGBFS could cause a disruption in the CSA cement physicochemical properties, leading to higher porosity and reducing strength, see Figures 16–18 [59]. Earlier results concluded similar trends. SEM images of concrete samples containing 25% of the AF showed significant recovery in the binder matrix on exposure to room temperature are shown in Figures 14d-f and 15d-f. Ettringite was observed in the 3-day-old concrete samples, but it disappeared later. In comparison, the 25% AF incorporated samples showed a pronounced hydration rate, followed by dense microstructure and compact morphology.



Figure 14. SEM images of the 3-day-old concrete samples at distinct time intervals: (**a**–**c**) OPC: GGBFS:CSA—40:40:20; (**d**–**f**) OPC:GGBFS:CSA—50:50:0 (25 wt.% of AF). (**Note:** Red color boxes indicate the areas with the detected ettringite crystals, and yellow boxes indicate the unhydrated CSA cement).



Figure 15. SEM images of the 28-day-old concrete samples at distinct time intervals: (**a**–**c**) OPC: GGBFS:CSA—40:40:20; (**d**–**f**) OPC:GGBFS:CSA—50:50:0 (25 wt.% of AF). (**Note:** Red color boxes indicate the areas with the detected ettringite crystals, and yellow boxes indicate the unhydrated CSA cement).



Figure 16. Pore segmentations on the 3-day-old concrete samples at distinct time intervals: (**a**–**c**) OPC: GGBFS:CSA—40:40:20; (**d**–**f**) OPC:GGBFS:CSA—50:50:0 (25 wt.% of AF). (**Note:** Black color indicates the pores).



Figure 17. Pore segmentations on the 28-day-old concrete samples at distinct time intervals: (**a**–**c**) OPC:GGBFS:CSA—40:40:20; (**d**–**f**) OPC:GGBFS:CSA—50:50:0 (25 wt.% of AF). (**Note:** Black color indicates the pores).



Figure 18. Developments of pore % on the concrete samples at distinct age and time intervals: (a) 3-day-old samples; (b) 28-day-old samples.

The porosity of the studied concretes was separated from SEM images using greyscale histograms and ImageJ win-64 software, see Figures 16 and 17. High porosity was found in the samples containing 20 wt% of the CSA cement and significantly lower porosity in the 25% AF using concrete. For 3-day and 28-day-old concrete samples containing 20 wt% of the CSA cement, the porosity varied from 9% to 11% and from 6% to 9%, respectively, while the porosity for the 25 wt% AF concrete ranged from 3% to 4% regardless of the used curing methods, see Figure 18a,b. The high porosity of the 20 wt% CSA concrete could be related to the formation of ettringite, as described earlier [54]. Over time, the porosities of these concretes dropped slightly, which could be related to the continuous hydration of all present binders, including CSA, OPC, and GGBFS. However, it could be assumed that the formation of C-S-H from OPC was limited to some extent due to the presence of CSA cement and higher content of Al ions in the pore solutions [91]. Replacement of binder by 20 wt% of CSA cement in the samples impacted the nucleation of ice in pores, see Figure 7. Other researchers showed that an excessive amount of CSA (>20 wt%) and extreme curing $(>-5 \circ C)$ decelerated the hydration rate and strength development if no pre-curing was adopted [79]. On the contrary, lower porosity in the 25% AF was a result of well-controlled hydration of OPC and pozzolanic activity of GGBFS, which was not hindered by low temperatures due to the application of AF admixture. The SEM observations and the earlier test results from the present study are in line and confirm that pore ice formation in the samples containing 20 wt% CSA and partially inhibited with the samples using 25 wt% AF chemical admixture.

4. Conclusions

This study aimed to determine if belite-CSA cement could be used as a partial replacement of OPC and/or GGBFS in concretes cast at subfreezing temperatures. The following conclusions can be drawn.

- The partial replacement of either OPC or GGBFS with 20 wt% of the belite-CSA cement shortened the initial setting time. This was attributed to the faster hydration of the CSA cement and the related ettringite formation.
- OPC:GGBFS mixes containing either 20 wt% CSA or 25 wt% AF showed similar strength developments when cured at 20 °C.

- An optimum dosage of sodium nitrate-based antifreeze (AF) considerably reduced the freezable water and promoted hydration.
- CSA cement appeared to induce some microcracks within the binder matrix, presumably due to the ettringite formation, which supported the formation of ice and the development of false strength.
- Pre-curing concrete mixes containing 20 wt% CSA for at least 6 h at room temperature before their exposure to -15 °C could mitigate some of the negative effects.
- OPC-GGBFS concrete containing 25 wt% of the AF admixture cured at -15 °C showed densification of the binder matrix after the application of extended post-curing at room temperature. On the contrary, no densification was observed for the concrete with 20 wt% of the CSA cement.
- Hydration tended to resume with the formation of C-S-H and ettringite crystals for mixes containing 25 wt% AF when the extended post-curing at room temperature was applied.
- Prolonged curing at -15 °C increased the porosity of concrete containing 20 wt% of CSA cement, which resulted in a lower strength gain due to the increased formation of ice.

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