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1	Workability, strength and shrinkage of ultra-high-performance seawater, sea sand
2	concrete with different OPC replacement ratios
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9	Abstract
10	An experimental investigation on the chemical, physical, mechanical, and shrinkage of seawater
11	and sea sand-based ultra-high-performance concrete (UHP-SWSSC) with supplementary
12	cementitious materials (SCM) (i.e., slag and silica fume) is reported. Several mixes were designed
13	with varying proportions of SCMs (25%, 37.5%, 50%, and 62.5% of binder), aggregate source,
14	and water-to-binder ratio. Heat evolution, density, workability, compressive strength development,
15	and long-term autogenous and drying shrinkage of UHP-SWSSC were monitored. Seawater
16	accelerates cement hydration as reflected in the heat evolution, and consequently, dictates the
17	early-age strength, and autogenous shrinkage. SCM addition although limits the early-age strength
18	development offers a comparable 90 days strength. The chloride content increases from marine
19	resources and may limit the application to non-structural components. Nonetheless, a UHP-
20	SWSSC mix with 50% OPC replaced by 37.5% slag and 12.5% silica fume is recommended in
21	this study, which can achieve satisfactory workability, long-term strength, and shrinkage
22	properties.

- 23 Keywords: Ultra-high-performance seawater sea sand concrete, OPC replacement ratio, water-to-
- 24 binder ratio, strength, workability, shrinkage.

## 25 **1. Introduction**

Being the most widely utilized construction material, the demand for ordinary Portland cement 26 27 (OPC) concrete in infrastructure industries has significantly increased over the past few decades. The total amount of cement produced worldwide in the year 2020 was 4.1 billion tons, which is 28 almost three-fold compared to the cement production in 1995 [1]. This huge demand and 29 30 consequently, the production, has not only exacerbated the scarcity of natural resources (i.e., fresh 31 water and river sand) by overexploiting them as raw materials but also worsened the ecological 32 balance through substantial  $CO_2$  emission in the manufacturing of OPC [2]. An alternative to the 33 consumption of limited natural resources is to utilize seawater and sea-sand concrete (SWSSC) which can be a sustainable substitute to depleting fresh water and river sand. In addition, proper 34 management of hazardous industrial by-products, including ground granulated blast furnace slag, 35 silica fume, fly ash, etc. is a potential challenge [3] with a noteworthy possibility of being reused 36 37 and recycled as supplementary cementitious materials (SCMs) in concrete manufacture. Blast 38 furnace slag and silica fume are by-products of steel and ferrosilicon manufacturing industries, respectively [4], which possess pozzolanic reactivity and thus, may be used as SCMs by replacing 39 OPC in SWSSC. However, concerns remain with the use of chloride enriched marine resources in 40 41 concrete as this can potentially stimulate corrosion in the embedded reinforcing steel. Alternative hybrid construction approaches utilizing seawater and sea-sand concrete and corrosion resistant 42 43 materials such as fiber reinforced polymers (FRP) and stainless steel, nonetheless, possess an 44 enormous potential in future sustainable construction practice e.g., [5-9], particularly in the coastal 45 regions.

Relevant studies have demonstrated that the utilization of sea-sand [2, 10] and seawater [11-13] in
concrete increases early-age compressive strength, lowers long-term compressive strength [14,

15], makes the concrete less workable [16], and increases drying shrinkage [17, 18] compared to 48 fresh water and river sand counterparts due to the salt and sea-shell contents present in them. 49 Application of seawater to cure SWSSC reduces the compressive strength and modulus of 50 elasticity as opposed to conventional freshwater curing [15]. The expansion of SWSSC because of 51 52 alkali-silica reaction (ASR) which ultimately induces cracking in concrete can be substantially 53 mitigated by utilizing SCMs (i.e., blast furnace slag, fly ash, silica fume etc.) [19]. Hence, 54 utilization of such by-products from manufacturing industries can mitigate the concrete reactivity 55 concerns and thus advocate sustainability in current construction practice.

56 Ultra-high-performance concrete (UHPC) has recently emerged as an efficient alternative to normal strength concrete due to its superior mechanical and excellent durability characteristics, 57 with high demand in infrastructures designed for long service life. According to some available 58 59 literature, UHPC is a new generation concrete that possesses an ultra-high strength of at least 60 120 MPa [20, 21]. UHPC is defined by its relatively high binder content and significantly lower 61 water-to-binder ratio compared to conventional concrete. Enhancement of microstructure, improvement of homogeneity, acceleration of hydration reactions, toughness improvement, and a 62 63 significant reduction in porosity are the key manufacturing principles of UHPC [22, 23]. Uniform 64 and dense microstructure and improved interfacial transition zone (ITZ) lead to the superior performance of UHPC compared to conventional concrete. Utilization of SCMs [22, 24] and use 65 of water reducing superplasticizers particularly containing polycarboxylates [25] helps to achieve 66 67 superior strength and maintain workability even though the water-to-binder ratio is kept relatively low. High binder content and use of superplasticizers in UHPC produces high heat during 68 69 hydration and causes higher shrinkage strains, which is the prime reason behind its susceptibility 70 to cracking [24, 26]. The portion of autogenous shrinkage is significantly larger in UHPC than normal concrete due to the presence of high binder content and a very low water-to-binder ratio.
Partial replacement of mixing water by crushed ice [27], applying heat curing [23], limiting the
dosage of superplasticizer [26], and incorporation of super absorbent polymers [28], etc. are some
innovative approaches to mitigate shrinkage in UHPC.

The above-mentioned papers on UHPC mainly dealt with concrete using fresh water and river 75 sand. Natural and artificial seawater, as well as sea sand, have been successfully used in the 76 77 preparation of ultra-high-performance concrete which achieved satisfactory mechanical 78 characteristics [29]. In consideration of the review above, there is a lack of research on the strength, 79 workability, and shrinkage of ultra-high-performance seawater and sea sand concrete (UHP-SWSSC), OPC-slag based UHP-SWSSC, let alone the influence of OPC replacement ratio on the 80 81 strength, workability, and dimensional stability (e.g., volumetric change of concrete due to 82 shrinkage). This paper explores the possibility to develop a UHP-SWSSC from local marine 83 resources in Australia which utilize industrial by-products such as blast furnace slag and silica 84 fume to replace OPC. Several mixes were prepared by varying the cement replacement ratio (0% 85 as the control mix, 25%, 37.5%, 50%, and 62.5%), aggregate type, and water-to-binder ratio. The 86 impact of these parameters on the compressive strength and workability was assessed. UHPC with 87 a similar mixing ratio but utilizing tap water and river sand as well as washed Sydney beach sand 88 were prepared and analyzed to compare with the seawater and sea sand UHPC. The influence of 89 different types of aggregates, cement substitution ratio, and water-to-binder ratio on early age and 90 long-term shrinkage properties of UHPC was determined. Finally, an appropriate UHP-SWSSC 91 mix was selected based on its physical, mechanical, and dimensional stability characteristics.

## 92 2. Raw Materials, Mix Proportions, and Methods

### 93 2.1. Raw Materials

A general-purpose ordinary Portland cement conforming to AS 3972-2010 [30] was used in this 94 study with two supplementary cementitious materials (SCMs) - ground granulated blast furnace 95 slag (GGBFS) and undensified silica fume conforming to AS 3582.2-2016 [31] and 96 97 AS/NZS 3582.3-2016 [32], respectively. The slag was sourced from Australian Steel Mill Services (ASMS), Port Kembla, NSW, and the silica fume was sourced from Simcoa, plant Kemerton, 98 Western Australia. Chemical oxide compositions of the binders (Table 1) were determined by X-99 100 ray fluorescence (XRF) spectroscopy in an AXIOS WDXRF instrument and SUPERQ software 101 was used for data analysis.

## 102 [Table 1 near here]

Natural seawater and sea sand were collected from Malabar beach of Sydney, NSW, Australia for
fabricating UHPC. The cations and anions in seawater were detected through inductively coupled
plasma – optical emission spectrometry (ICP-OES) and ion chromatography (IC), respectively. **Table 2** compares the ion concentrations of the natural seawater used in this study with that of the
world average values [29]. The seawater collected from Malabar beach was found to be reasonably
close to the corresponding world average values. Tap water was used in mixing and curing of tap
water river sand and tap water washed Sydney beach sand UHPC.

110 [Table 2 near here]

111 [Figure 1 near here]

The sea sand was sieved through a 1.18 mm sieve to separate the stones, seashells, organic 112 materials, and other impurities such as portions of aquatic plants, branches, man-made pollutants, 113 114 etc., as per Willie and Boisvert-Cotulio [33] recommendations. The processed sand is presented in Figure 1. The total mass of impurities was 2.85% of the natural sea sand. Natural river sand was 115 obtained from Taren Point, New South Wales. The washed beach sand was collected from beaches 116 117 outside Sydney and washed thoroughly with fresh water to make it salt-free. Particle size 118 distribution of different sands determined according to AS 1141.11.1-2009 [34] are compared in 119 Figure 2. Sea sand with and without impurities is termed as 'sea sand before processing' and 'sea 120 sand after processing', respectively. Natural river sand was found to be well graded, while the sea sand was gap graded with a significant portion of its particles retaining on 0.3 mm and 0.6 mm 121 sieves. To avoid any effect of aggregate gradation in this study, natural river sand with the same 122 123 particle size distribution as the sea sand was used. This modified river sand is termed as 'processed river sand' in this study. Modification of the sea sand by eliminating larger-sized particles does 124 125 not alter its particle size distribution. The fineness of sea sand was found to be in between washed Sydney beach sand and natural river sand. 126

# 127 [Figure 2 near here]

The chemical composition of the sands was determined using the X-ray fluorescence (XRF) spectroscopy technique and presented in **Table 3**. It is evident that the sands are rich in silica. **Table 4** lists various physical properties of the sands used in this study including particle densities (i.e., oven-dry density, saturated surface dry or SSD density, apparent density), bulk densities (uncompacted and compacted), percentage of water absorption, and percentage void determined following relevant standards [35-37]. Natural river sand was found to possess slightly higher oven dry as well as SSD particle density compared to marine sands, due to the presence of a bigger proportion of larger-sized aggregates (especially over 1.18 mm) in natural river sand and lightweight sea-shell fragments in sea sand. Water absorption of sea sand is the highest (3.46%), whereas natural river sand has the lowest water absorption (1.68%). Difference between the uncompacted and compacted bulk density was found to be higher in natural river sand, because of the presence of aggregates with various size ranges. Percent voids were measured to be very similar among different sand types (ranging between 39 - 41%).

141 The surface morphology of different sand particles and SCMs were assessed through SEM 142 micrographs as shown in **Figure 3**. The procedure for microstructural analysis is discussed in 143 **Section 2.3.3**.

144 [Figure 3 near here]

# 145 [Table 3 near here]

MasterGlenium SKY 8700, a polycarboxylate-based high range water reducer (HRWR) conforming to the requirements of AS 1478.1 [38], was used for batching UHP-SWSSC for its excellent cement dispersion and slump retention ability along with its suitability to be used with self-compacting concrete.

## 150 [Table 4 near here]

## 151 2.2. Mix Proportions, Batching, and Specimen Preparation

A total of 12 UHPC and one normal strength concrete mixes summarized in **Table 5** were investigated in this study. No steel fiber was added to the UHPC in order to produce an economically viable mix and eliminate the chances of steel corrosion. Seven UHPC mixes were prepared with seawater and sea sand (UHP-SWSSC), two from washed Sydney beach sand and 156 tap water (UHP-TWWBSC), and the remaining three utilized river sand and tap water (UHP-TWRSC). Five mix ratios with varying proportions of OPC replacements were chosen for UHP-157 SWSSC. UHP-SWSSCC with 100% OPC (termed as 'control mix') along with 25%, 37.5%, 50%, 158 and 62.5% of OPC replacement by ground slag and silica fume were produced. The control mix 159 160 was developed based on the recommendations provided by Willie and Boisvert-Cotulio [33] and 161 through a trial-and-error process until a suitable mix with satisfactory flowability and strength was 162 achieved. Although the ratio of OPC to supplemental material (ground slag in this study) is 163 suggested to be 1:0.25 by weight in [33], this study explores the possibility to utilize slag at a much 164 higher proportion (up to a ratio of OPC: slag = 1:1.33). In the OPC replaced mixes, the portion of silica fume was kept constant at 12.5% of the total binder (except OPC-slag binary mix with 25% 165 substitution of OPC). Total binder content, aggregate content, water-to-binder ratio, and 166 167 superplasticizer-to-binder ratio were kept the same for all the mixes with different OPC replacements (1200 kg/m<sup>3</sup>, 1000 kg/m<sup>3</sup>, 0.2, and 0.02, respectively). Two more seawater and sea 168 169 sand UHPC mixes were prepared by altering the water-to-binder ratio (0.15 and 0.25) of the 50% 170 OPC replaced mix. Table 5 also presents the label for each of the concrete mixes (termed as 'mix ID'); each label consists of the type of aggregate used, percentage of OPC replacement, and water-171 172 to-binder ratio. For example, a mix ID of SS-50-0.2 refers to the UHPC mix where sea sand has been used as aggregate, with an OPC substitution of 50% and a water-to-binder ratio of 0.2. 173

# 174 [Table 5 near here]

For batching, the cement, SCMs and SSD fine aggregate were dry mixed for ten minutes at low speed (50 rpm) in a laboratory pan mixer capable of controlling mixing speed. Water and HRWR were mixed separately as per admixture supplier recommendations and later added to the dry materials over 30 seconds. Afterward, mixing was continued for an additional 8 minutes at low 179 speed and another 2 minutes at medium speed (100 rpm) until a workable mix was attained. UHPC prepared in this study was found to be highly flowable and self-compacting, except for the mixes 180 181 with significant OPC replacement and a very low water-to-binder ratio (particularly, SS-50-0.15). Immediately after mixing, the flow spread of the fresh UHPC was measured on a flow table as per 182 ASTM C1437-15 [39] guidelines and the fresh density was measured following procedures 183 184 outlined in ASTM C 138 [40]. According to the relevant standard [21], the flow table test was 185 modified to adjust to UHPC's fluid nature. Concrete was poured into a mold in a single layer 186 without tamping, the table was not dropped, and the fresh concrete was allowed to flow. The 187 diameter of the flow spread was measured and reported. Simultaneously, standard 50 mm  $\times$  50 mm cube specimens were prepared and compacted on a vibrating table running at low 188 189 frequency. The molds were then stored in a temperature and moisture-controlled cabinet at 23°C 190 and relative humidity of 50% for  $24 \pm 2$  hours. After the specified time, the concrete specimens 191 were demoulded, and water cured. Two sets of specimens from each mix were prepared; one set 192 was cured in a 23°C lime-saturated water bath while the other was cured in natural seawater at the 193 same temperature. A set of 25 mm  $\times$  25 mm  $\times$  285 mm prisms were also prepared from each batch to monitor UHPC's shrinkage behavior. 194

# 195 2.3. Testing Procedures

# 196 2.3.1. Heat of hydration

The heat evolved in the first 72 hours of hydration of some representative UHPC pastes was captured in a TAM Air isothermal calorimeter operating at 23°C. From **Table 5**, the paste portion of SS-0-0.2, RS(P)-0-0.2, and SS-50-0.2 were chosen for the test to evaluate the effects of seawater and SCM on the reaction kinetics of UHPC. For this purpose, about 12 g paste sample was handmixed in a 20 ml glass ampoule and sealed before quickly transferring the specimen along with an inert reference sample (oven-dried quartz sand) in the calorimeter chambers. Each reference sample mass was determined to have the same heat capacity as the UHPC paste sample. The time elapsed between the addition of the mix water and the start of the test was adjusted during data analysis. Each test was replicated to confirm the repeatability of the method.

### 206 2.3.2. Mechanical properties and shrinkage

For each mix proportion and curing condition studied, the 1, 3, 7, 28, and 90 days compressive 207 208 strength of 50 mm  $\times$  50 mm cube specimens were determined following the requirements of 209 ASTM C109/C109M-2020 [41]. Meanwhile, length change of prism specimens was measured up 210 to an extended period of 120 days to determine the long-term drying and autogenous shrinkage of 211 UHPC. The total shrinkage samples were air-dried after demoulding in a moist chamber at a 212 constant conditioning temperature of 23°C and 50% relative humidity while the autogenous 213 shrinkage samples were properly sealed with aluminum tape to prevent any loss of moisture and 214 thus obtain their autogenous shrinkage at the same temperature and relative humidity. The total 215 and autogenous shrinkages were determined by measuring the length change of the specimens 216 according to AS 1012.13-2015 [42]. Drying shrinkage was determined from the difference 217 between the respective total shrinkage and autogenous shrinkage.

218 2.3.3. Microscopic imaging

A Hitachi S3400 scanning electron microscope (SEM) was utilized for investigating the microstructure of 28 days mature UHPC. For this purpose, about 5 mm thick slices of samples were cut using a slow-rotating diamond saw. Specimens were immersed in 100% ethyl alcohol to stop the hydration and impurities were eliminated by ultrasonic cleansing. The imaging surface of the specimens was gold coated in an Emitech K550x gold sputter coater. For imaging raw binders and sands, the particles were attached on an adhesive carbon tape and gold coated. The accelerating current for imaging was set at 30 amps with a beam strength of 20 kV and a working distance of
approximately 5 mm under a high vacuum mode. Images of the microstructure were captured using
the backscattered electron (BSE) mode of the electron microscope.

#### 228 **3. Results and Discussions**

### 229 **3.1.** Heat of Hydration

The heat evolution in the first 72 hours of cement hydration (normalized per unit mass of cement) 230 231 in the presence of tap water, seawater, and SCMs (slag and silica fume) is illustrated in Figure 4. 232 The typical stages of cement hydration such as the dissolution of cement particles as identified in 233 the initial peak right after the addition of water, induction peak, and the acceleration peak marking the hydration of cement [43] are observed in Figure 4(a). The temporal change in the occurrence 234 235 of the acceleration peak suggests that seawater accelerates cement hydration. The acceleration 236 peak shifts by at least 6-8 hours from seawater hydration of cement along with reaching about 15% 237 higher heat flow. A similar phenomenon is also observed when 50% cement is replaced by 37.5% 238 slag and 12.5% silica fume. However, the mechanism behind them is different.

# 239 [Figure 4 near here]

When seawater is utilized in the mix, the abundance of chloride ions chemically binds with the calcium, aluminate, and ferrous phases of cement [44], forming insoluble calcium oxychloride phase and Friedel's salt [45]. When SCMs replace 50% of cement, it causes a similar effect, but from the well-explored filler effect [46, 47]. The slag and silica fume particles act as fillers between cement particles and offer extra nucleation sites for hydration products to occupy [46, 47]. As such, the heat evolved per unit mass of cement is higher than cement-only paste. This is also reflected in **Figure 4(b)** which shows a significantly higher cumulative heat generated per unit mass of cement confirming the filler effect of SCMs. The heat of hydration of cement from seawater and tap water mixing after 72 hours of hydration is somewhat similar, but seawater mixing generates more heat in between 12-30 hours. Thus, it is expected that the 1-day strength of seawater composite could exceed that of tap water, however, may follow a similar strength development in later ages of hydration.

### 252 **3.2.** Properties of Fresh Concrete

253 The fresh and 28 days SSD density of UHPC are summarized in Table 6. The SSD density of 254 concrete was found to be greater than its fresh density for all the mixes due to the continuing water 255 absorption during the curing period. UHPC was found to possess around 6.6% greater unit weight than normal strength concrete because of its significantly compact microstructure from the 256 257 selection of lower water-to-binder ratio [48]. The unit weight of seawater sea sand UHPC was 258 found to be marginally higher than tap water river sand and washed beach sand UHPC, for both 259 100% OPC and 50% OPC replaced mix. However, when sea sand is replaced with washed beach 260 sand and river sand, the fresh densities drop by 0.35% and 0.9%, respectively. The use of seawater 261 may have resulted in the slight increase in density from the formation of more hydration products 262 as captured in the heat evolution curves in Figure 4.

# 263 [Table 6 near here]

The incorporation of SCMs reduced the unit weight of UHPC, possibly due to the lower densities of SCM compared to OPC. The percentage of OPC replacement has been found to have a linear relationship with the decrease in a unit weight of both fresh and 28-day SSD unit weight. From **Table 6**, an OPC replacement of 25%, 37.5%, 50% and 62.5% yielded in a reduction of 1.15%, 3.49%, 3.99% and 4.01%, respectively in the unit weight of fresh UHP-SWSSC. The incorporation of silica fume had a greater effect on unit weight reduction compared to ground slag, due to the significantly lower relative density of silica fume than cement and slag used in this study. These findings are consistent with previous studies of Zain et al. [48] and Turk [49].

272 The UHPC mixes reported in **Table 5** were trialed to possess excellent workability in general; their flow spread is summarized in Figure 5. The high flowability was achieved due to the 273 utilization of polycarboxylate-based superplasticizer at a relatively high dosage than recommended 274 275 for normal grade concrete. Regardless, the effects of slag, silica fume, and the use of different 276 types of sand on the flowability can still be comparatively assessed. In general, the presence of 277 ground slag decreased the flowability of all UHPC mixes. For instance, a 12.5% reduction in the flow spread resulted from the 25% replacement of cement with slag (Figure 5(a)). The angular 278 279 shape of slag particles (shown in Figure 3(d)) together with the accelerated calcium reaction is 280 possibly the primary reason behind the reduced flowability. However, the workability of UHPC 281 improved when silica fume was added in addition to slag as partial replacements of OPC. The 282 ternary blends produced better workability compared to the OPC-slag binary blend from the 'ball 283 bearing' and lubricating effect of the spherical silica fume particles, as demonstrated in Figure 3(e) 284 [50-52]. The addition of 12.5% silica fume in SS-37.5-0.2 increased the flow spread by 12.7% 285 compared to the OPC-slag binary mix. However, with the increase of slag content in ternary blends, the flow degraded. The SS-62.5-0.2 mix with 50% ground slag and 12.5% silica fume 286 yielded a 19.4% lower flow spread compared to the control mix. This particular mix was viscous 287 288 and required high-frequency vibration for proper compaction.

289 [Figure 5 near here]

290 From Figure 5(b), the flowability of UHPC slightly declined from the use of sea sand and seawater. The uniformity and size fractions of sea sand particles together with the presence of 291 irregularly shaped seashells and other micro-impurities may have led to a reduced flow [2]. 292 Figure 3 confirms that washed beach sand particles are round in shape with a relatively smooth 293 surface, while the river sand particles are mostly angular with rough edges and varying size 294 295 fractions. The angularity and roughness of sea sand somewhat lie in between the washed beach sand and river sand, which is also reflected in the flowability of UHPC ranging in the same order. 296 297 It is evident from **Figure 5(b)** that the control UHP-SWSSC mix had a flow spread of 360 mm, in 298 contrast to 390 mm and 345 mm for washed Sydney beach sand and processed river sand UHPC with identical mix proportions. The 50% OPC replaced mixes also showed a similar trend. 299

300 In addition, the flow spread increased with the increase in the water-to-binder ratio, as expected 301 (Figure 5(c)). 50% OPC replaced UHP-SWSSC with a water-to-binder ratio of 0.15, 0.2, and 0.25 302 produced slump spreads of 185 mm, 340 mm, and 360 mm, respectively. The SS-50-0.15 mix was 303 found to be highly viscous, and the workability was challenged in specimen making; this also 304 yielded a 45.6% reduced spread compared to the SS-50-0.2. Chen et al. [53] reported that UHPC's 305 extremely low water content is insufficient to fill the inter-particular void spaces and is readily 306 absorbed around the particle surfaces while leaving the voids empty. This study found a water-to-307 binder ratio of 0.2 to be optimum for lubrication of the granular materials at a superplasticizer dose 308 of 24 kg/m<sup>3</sup>. Overall, the workability of all the UHPC mixes in this study was found to be similar 309 in comparison with the targeted 280 mm – 340 mm flow spread of Willie and Boisvert-Cotulio 310 [33]. The results are also consistent with the flowability of seawater and sea sand UHPC studied 311 by Teng et al. [29].

### 312 3.3. Compressive Strength

**Table 7** summarizes the compressive strength of all mixes investigated in this study under both
 313 314 tap water and seawater curing regimes; the results are the mean of the strength of three identical specimens. The strength of conventional concrete cylinders under tap water curing was also 315 316 assessed for comparison purposes. The 1-day strength was determined just after demoulding of 317 specimens and before storing them in the curing chamber. The standard deviation is well within 318 5% of the compressive strength value, which indicates lesser variability and better homogeneity 319 of respective UHPC mixes. The following section highlights the influence of several mix 320 proportion parameters on the compressive strength of UHPC.

#### 321 [Table 7 near here]

## 322 3.3.1. Effect of seawater and sea sand

Figure 6 shows the comparative compressive strength development of UHPC composed of 323 324 different types of sand and mixing water under tap water curing conditions. The incorporation of sea sand and seawater in UHPC can produce relatively higher compressive strength at an early 325 age, particularly, up to 28 days. The control mix SS-0-0.2 in this testing regime yielded 326 327 substantially high early-age strength, with no cement replacement. However, comparative 328 discussion can be made for the other mixes in which, the sand type is the only variable. Among 329 the 50% OPC substituted mixes, the seawater and sea sand mix (SS-50-0.2) produces 29.1%, 330 16.8%, and 13.9% larger compressive strength values in comparison with tap water and washed 331 beach sand mix after 1, 3, and 7 days, respectively. Although in the early ages, the difference is 332 more pronounced, the long-term (90 days) strength of all mixes is similar. The seawater and sea 333 sand mix reached 62.5% of its 90-day strength at an age of only 3 days, whereas the river sand and washed beach sand mixes utilizing tap water in mixing achieved slightly over half of their long-334

term strength at three days. This indicates that seawater accelerates the hydration of the binder, 335 336 providing early-age strength gain to UHPC. This is also reflected in the heat of hydration curves 337 in **Figure 4** which shows greater heat release from the use of seawater. For tap water mixing, the hydration continues beyond 28 days, although at a slower pace, but reaches a similar strength as 338 that of seawater mixed UHPC. The higher early-age compressive strength is due to the abundance 339 340 of chloride content in sea sand and seawater which accelerates the hydration reaction initially. Faster hydration generates a greater amount of hydration products, such as calcium silicate hydrate 341 342 (C-S-H), which blocks the pores in the resulting concrete and thus enhances the early strength 343 development [2, 16].

### 344 [Figure 6 near here]

345 However, the development of strength gain in UHP-SWSSC slows down with time, especially after 28 days. The UHP-SWSSC mix with 50% cement substitution (SS-50-0.2) yields the highest 346 strength of 136.8 MPa after 28 days of tap water curing, which is a mere increase (6% and 2.2%, 347 respectively) compared to UHP-TWWBSC and UHP-TWRSC counterparts. Similar trends were 348 349 also observed when mixes were cured in a seawater environment. This deterioration in strength 350 gain occurs as a result of leaching out of soft hydration products with time [11, 14]. Long-term strengths, especially after 90 days of curing were found to be almost identical, which aligns with 351 352 the findings of the studies of Mohammed et al. [12]. Overall, a slight increase in the early age 353 strength and a slight reduction in the long-term strength of SCM-based UHP-SWSSC were 354 consistent with the results of Teng et al. [29].

## 355 *3.3.2.* Effect of OPC replacement by ground slag and silica fume

It is apparent from **Figure 7** that in general, UHPCs produce significantly higher strength (about 4 times) than normal strength concrete with a water-to-binder ratio of 0.35. The ratio of strength between OPC-based UHPC and NC was found to be higher at an early age (a strength ratio of 4.7 after 1-day), whereas a slightly reduced strength ratio of 3.4 has been found for long-term 90-day compressive strength. This indicates the faster development of strength in UHPC with high binder content at a significantly lower water-to-binder ratio.

# 362 [Figure 7 near here]

It has been found that early-age strength decreases with the increase in OPC replacement. UHPC 363 364 with no cement replacement yielded a compressive strength of 85.2 MPa at day 1, whereas 25%, 365 37.5%, 50%, and 62.5% OPC replaced mixes produced respectively 57 MPa, 38.9 MPa, 37.3 MPa, 366 and 22.9 MPa strength at the same age. OPC substitution by SCMs, such as silica fume and ground slag, was found to yield lower strength initially, up to as early as 7 days of casting. The slower rate 367 of pozzolanic reaction of ground granulated blast furnace slag at room temperature curing hinders 368 369 the early age strength development [54]. However, OPC substituted mixes tend to generate more 370 strength compared to cement-based mixes as time goes on, especially after 28 days of curing and beyond. In our study, 28-days, as well as 90-days strengths of all UHP-SWSSC mixes, were found 371 to be quite similar, with a slight increase of strength for higher OPC replacements. For example, 372 373 the 50% OPC replaced mix reaches a compressive strength of 136.8 MPa after 28 days of tap water 374 curing, while the 100% OPC variant has a slightly lower strength of 127.4 MPa. The strength ratios 375 of 62.5% cement replaced mix to control mix were 0.27, 0.71, 0.95, 1.03, and 1.05 after tap water 376 curing periods of 1, 3, 7, 28, and 90 days correspondingly. The short-term strengths were 377 considerably lower in comparison with the control mix, while slightly higher long-term strengths

were observed. Silica fume with significantly smaller particle size has a filler effect which improves the particle packing density. The highly pozzolanic slag and silica fume convert the Ca(OH)<sub>2</sub> into secondary C-S-H and thus enhance the interfacial bond, which eventually leads to more homogeneous ITZ and therefore, an increased strength [54, 55]. Overall improvement of microstructure in SCM incorporated UHPC blends enables them to achieve higher strength compared to OPC based UHPC.

#### 384 [Figure 8 near here]

UHP-SWSSC mixes with different proportions of cement replacement show identical trends when cured under a seawater environment (**Figure 8**). The mix with 50% OPC replacement yields 56.3% and 24.9% lower strength after 1 and 3 days of curing compared to the control mix, however it generates 10.8% and 7.5% higher strength after a curing period of 28 and 90 days, respectively.

# 389 3.3.3. Effect of water-to-binder ratio

390 Three mixes of 50% cement replaced UHP-SWSSC were compared with varying water-to-binder 391 ratios, i.e., 0.15, 0.2, and 0.25. In general, an increase in the water-to-binder ratio causes a 392 reduction in compressive strength (Figure 9). This result is consistent with the findings of Zhang 393 and Zhang [56], where it was found that the reduction of water content in the UHPC matrix 394 densifies the hydration film produced around the surface of cementitious particles, which prohibits further hydration reaction. Additionally, lowering the water-to-binder ratio reduces the porosity in 395 396 microstructure, which eventually leads to greater strength. However, early age strength was seen 397 to be impacted more compared to long-term strength. Increasing the water-to-binder ratio from 398 0.15 to 0.25 reduces the 1-day and 3-days strength by 59.8% and 20.2%, whereas the strength 399 reductions were found to be 6.4% and 3.5% after 28 and 90 days, respectively.

Interestingly, the SS-50-0.2 mix with a water-to-binder ratio of 0.2 yielded marginally higher 28-401 402 day strength compared to the counterpart SS-50-0.15 with a lower water-to-binder ratio. In the 403 case of extremely low water content, the available water is completely absorbed within the particle surfaces, thereby leaving the voids empty. If the water content in the cementitious matrix is lower 404 than a minimum margin, the strength does not increase due to the void spaces present in the 405 406 microstructure and due to the unavailability of water for hydration of cement particles [53, 56]. 407 The UHPC mix with a very low water-to-binder ratio of 0.15 was found to be sticky and viscous, 408 with a flow spread of only 185 mm (**Figure 5**( $\mathbf{c}$ )). The cross-section exhibited visible air voids as shown in Figure 10(a), which could not be eliminated through vibration. The porous 409 410 microstructure with an extremely low water-to-binder ratio inhibits its ability to gain long-term 411 ultra-high strength. In contrast, cross-sections of SS-50-0.2 and SS-50-0.25 mixes were more 412 homogeneous with almost no visible air voids, as shown in **Figure 10(b)** and **Figure 10(c)**. The 413 optimum water-to-binder ratio in order to achieve maximum long-term compressive strength was 414 found to be 0.2.

# 415 [Figure 10 near here]

### 416 *3.3.4.* Effect of curing condition

The influence of seawater curing on the compressive strength of UHPC comprised of different aggregates is depicted in **Figure 11** and **Figure 12**. Overall, seawater incorporated as curing water has a negative impact on the strength of UHPC. The 100% OPC-based UHP-SWSSC samples cured in tap water yielded higher strength compared to curing in seawater, 3.3%, 8.8%, 13.5%, and 13.8% strength enhancement after curing periods of 3, 7, 28, and 90 days, respectively

(Figure 11). River sand and washed Sydney beach sand-based UHPC experienced similar 422 deterioration in strength while cured in seawater, with 8.67% and 9.31% of reduction, respectively 423 after a prolonged seawater curing period of 90 days. The loss of strength in seawater-cured cement-424 based UHPC has been observed to escalate with the curing age. The degradation of strength gain 425 may have occurred because of the formation of expansive (such as ettringite and Friedel's salt) as 426 427 well as soft leachable compounds in the cementitious microstructure due to seawater penetration during prolonged exposure [14]. Development of micro-cracks due to chloride and sulfate attacks 428 429 from harsh seawater environments as well as salt crystallization is known to occur in concrete with 430 high cement contents [57]. The combined phenomena explain the deterioration of long-term strength development of UHPC under seawater curing regime. 431

## 432 [Figure 11 near here]

Similar behavior has been observed in SCM-based UHP-SWSSC. In comparison with seawater 433 434 curing, 50% OPC replaced UHP-SWSSC mixes attain a compressive strength increase of 18.5%, 9.6%, 10%, and 9.7% respectively after the age of 3, 7, 28, and 90 days in tap water curing 435 (Figure 12). Contrary to OPC-based UHPC, early age strength is more impacted than long-term 436 strength in slag and silica fume-based UHPC, regardless of the sand type utilized. The addition of 437 SCMs furthers the hydration reaction due to their enhanced pozzolanic activity, which continues 438 to densify the microstructure as time progresses. Therefore, SCM-based UHPC experiences lower 439 440 compressive strength loss under a harsh seawater environment [58].

441 [Figure 12 near here]

## 442 **3.4.** Shrinkage

### 443 *3.4.1. Effect of sand type*

444 Figure 13 compares the total, autogenous, and drying shrinkage of UHPC comprising of three different types of sand and that of conventional concrete. UHPCs experienced significantly higher 445 autogenous shrinkage, and therefore higher total shrinkage compared to normal concrete 446 447 (Figure 13 (a, b)). The 7 day early-age and 120 day long-term autogenous shrinkage of UHP-SWSSC was found to be 15 and 6 times larger than that of normal concrete, respectively. As the 448 449 negligible water content in UHPC is not enough for complete hydration of binder contents, greater 450 surface tension is created in the fine capillary pores of the cementitious matrix, which leads to larger autogenous shrinkage [23, 59]. UHPC experiences a very quick self-desiccation during the 451 first few days after casting and thus obtains a very high early autogenous strain rate. The early 452 453 growth (first 7 days) of autogenous shrinkage is slower in plain concrete. While normal concrete 454 reaches only 28% of its maximum autogenous shrinkage gained over 120 days testing period, 455 UHP-SWSSC reaches 72% of its maximum autogenous shrinkage in the same duration (Figure 14). It can be observed from Figure 15 that the proportion of autogenous shrinkage is 456 insignificant in NC (< 20% of total shrinkage) but it is significant in UHPC (over 40% in UHP-457 458 SWSSC and over 60% in UHP-TWRSC as well as UHP-TWWBSC).

## 459 [Figure 13 near here]

UHPC manufactured from seawater and sea sand in this study yielded considerably higher drying
shrinkage strain than tap water and river sand or washed beach sand counterparts (Figure 13(c)),
whereas the autogenous shrinkages were quite similar (Figure 13(b)). After 120 days of testing
period, the drying shrinkage of UHP-SWSSC was 973.7 με, compared to 288 με and 347.3 με for
UHP-TWWBSC and UHP-TWRSC respectively. Due to the presence of excessive sodium and

465 calcium chloride in seawater sea sand concrete, the morphology of the pore structure densifies, 466 which increased the number of micropores. This eventually increases the capillary tension and 467 therefore generates substantially high drying shrinkage [60, 61]. Additionally, the presence of sea-468 shell contents may have accounted for larger drying shrinkage in sea sand-based concrete, 469 particularly due to the lower rigidity of sea-shell particles [62].

470 Although autogenous shrinkage dominates the total shrinkage in washed beach sand and river sand 471 UHPC mixes (with a ratio of 120 days autogenous to drying shrinkage to be 2.23 and 1.89, 472 respectively), the drying shrinkage strain component was observed to be larger in the seawater sea 473 sand variant (ratio of autogenous to drying shrinkage was 0.7). Over the whole testing timeframe, autogenous shrinkage was less prominent in UHP-SWSSC (around 40-45% of the total shrinkage), 474 475 whereas it formulated the major portion of the total shrinkage in river sand and washed beach sand 476 UHPC (around 65-70%). It is evident from Figure 14 that sea sand-based UHPC achieves 93% of 477 its maximum autogenous shrinkage after 28 days from casting, whereas washed beach sand and 478 river sand UHPCs reach respectively 75% and 73% at the same point of time. The increased 479 amount of calcium chloride as a result of reactions between sodium chloride and calcium 480 hydroxide in the pore solution of seawater-based UHPC is the prime reason behind its accelerated 481 hydration [63]. Autogenous shrinkage strain in UHP-SWSSC reaches a plateau after 21 days 482 indicating the completion of accelerated hydration, whereas it takes around 100 days to stabilize in UHP-TWRSC and UHP-TWWBSC. Therefore, UHP-SWSSC may be susceptible to early-age 483 484 cracking due to its high early-age autogenous shrinkage.

485 [Figure 14 near here]

486 Overall, the growth of drying shrinkage in UHP-SWSSC is slower than river sand and washed 487 beach sand UHPCs. Drying shrinkage strain in UHP-SWSSC stabilizes after 84 days of casting, 488 while UHP-TWRSC and UHP-TWWBSC take only around 21 days to attain plateau. Very small 489 pores in the compact microstructure of UHP-SWSSC continue their capillary actions for an 490 extended time, which leads to the gradual increase of the drying shrinkage.

## 491 [Figure 15 near here]

## 492 *3.4.2. Effect of SCMs*

It was observed from Figure 16(a) that replacement of OPC by SCMs tends to decrease the overall shrinkage of seawater and sea sand-based UHPC. Although the autogenous shrinkage was found to moderately increase (Figure 16(b)), the drying shrinkage considerably reduced when OPC was partially replaced by slag and silica fume (Figure 16(c)).

# 497 [Figure 16 near here]

498 A more detailed comparison on the growth of autogenous shrinkage suggests that the autogenous 499 shrinkage in OPC-slag-silica fume trinary mix was lower than OPC-slag binary mix. However, 500 both types of mixes yielded greater autogenous shrinkage compared to cement-only UHPC. 501 Incorporation of slag and silica fume is known to enhance the autogenous shrinkage of concrete 502 due to their ability to refine the microstructure into a finer pore capillary system, which increases 503 the degree of self-desiccation, leading to a higher autogenous shrinkage [64, 65]. Generally, the 504 effect of silica fume in hydration is much more pronounced in fresh water-based UHPC compared 505 to that of slag, especially at early ages. However, due to the alkaline environment and presence of 506 a high quantity of sulfates in the pore solution of seawater mixed UHPC matrix, the hydration 507 reaction of slag accelerates and the reactivity of silica fume with cement reduces [63]. The rapid

508 dissolution of slag and decelerated influence of silica fume in the hydration kinetics of UHP-509 SWSSC explains the larger autogenous shrinkage of OPC-slag UHPC compared to OPC-slag-510 silica fume ternary blends. However, as the slag content in ternary mixes increased, the autogenous shrinkage values also increased. After 120 days, the binary (25% OPC substitution by only slag) 511 512 and ternary (37.5% OPC replacement by 25% slag and 12.5% silica fume) mixes yielded 50.9% 513 and 6.7% higher autogenous shrinkage, respectively compared to the control mix. Blast furnace 514 slag in UHP-SWSSC significantly elevates autogenous shrinkage, while further addition of silica 515 fume balances this surge. Figure 17 demonstrates the SEM images of shrinkage cracks in the 516 microstructure of control UHP-SWSSC (Figure 17(a)) and the blend with 25% OPC replaced by slag (Figure 17(b)). As the samples right after casting were sealed in closed plastic tubes and 517 518 remained sealed until the day of SEM analysis, the shrinkage that occurred in the specimens was 519 primarily autogenous shrinkage. A prominent network of autogenous shrinkage cracks was seen 520 in the microstructure of 25% OPC replaced UHPC compared to UHPC with 100% OPC, where 521 autogenous shrinkage cracks were almost nonexistent.

# 522 [Figure 17 near here]

The inclusion of slag and silica fume accelerates the growth of autogenous shrinkage development as well. This is from the higher degree of reaction of cement from the addition of SCMs as captured in the heat evolved per gram of cement (shown in **Figure 4**). In the control mix, drying shrinkage grows more rapidly than autogenous shrinkage (88% of drying and 50% of autogenous shrinkage develops in the first 7 days, as demonstrated in **Figure 18**). However, with more portion of OPC being replaced by slag and silica fume, autogenous shrinkage starts to grow quicker at early ages due to elevated interaction of slag in the cement-slag-silica fume ternary blends. For a 50%, OPC replaced mix, 72% autogenous shrinkage develops after 7 days as opposed to 68% dryingshrinkage.

### 532 [Figure 18 near here]

533 In general, the percentage of autogenous shrinkage with respect to total shrinkage increases with 534 an increase in OPC replacement percentage. This applies especially to the OPC-slag binary mix, 535 where autogenous shrinkage was 53% of the total shrinkage at 120 days, compared to 33% in the 536 control mix (Figure 19). OPC replaced by only slag (or high content of slag) produces a high value 537 of autogenous to drying shrinkage ratio. For instance, a 25% substitution of OPC by slag yielded 538 an autogenous to drying shrinkage ratio of 1.14, compared to 0.49 for the control mix. Autogenous 539 shrinkage was most significant in SS-62.5-0.2 (50% slag and 12.5% silica fume) as a result of its 540 high slag inclusion (autogenous to drying shrinkage ratio of 1.28).

Drying shrinkage decrease with the increase of cement substitution by slag and silica fume unlike 541 542 autogenous shrinkage, which is consistent with relevant literature [66]. This is attributed to the relatively denser microstructure in slag-based UHPC. A compact matrix contains lesser 543 544 microstructural pores, which keeps water loss due to evaporation to a minimum and therefore leads 545 to a decreased drying shrinkage. It was also noticed that drying shrinkage in silica fume added mixes grew slower than control and OPC-slag binary mixes. After 7 days of casting, the SS-62.5-546 0.2 ternary mix develops 72% of its ultimate drying shrinkage, compared to 88% and 94% growth 547 in control and 25% slag replaced mixes, respectively (Figure 18). This delayed growth of drying 548 549 shrinkage is probably because silica fume further densifies OPC-slag binary system and yields a 550 more compact microstructure. In general, drying shrinkage becomes less prominent with more 551 percentage of OPC supplanted by slag and silica fume.

### 552 [Figure 19 near here]

### 553 3.4.3. Effect of water-to-binder ratio

**Figure 20** illustrates the effect of the water-to-binder ratio on the shrinkage of UHP-SWSSC. From **Figure 20(c)**, it is observed that the drying shrinkage in the UHPC with higher water content is larger than the lower water-to-binder ratio counterparts (701.6  $\mu\epsilon$  in SS-0.50-0.15 compared to 1090  $\mu\epsilon$  in SS-50-0.25 after 120 days of casting). More pores are formed in a concrete mix with larger water content, which produces higher surface tension in the pore structure during evaporation [26].

### 560 [Figure 20 near here]

561 Several literatures have reported an increase in autogenous shrinkage with the reduction of water-562 to-cement ratio in high-performance concrete [59], ultra-high performance concrete [64], and 563 reactive powder concrete [26] as a result of enhanced internal drying, self-desiccation, and 564 capillary tension. However, a quite opposite finding has been obtained in this research regarding 565 seawater and sea sand-based UHPCs. From Figure 20 (b), the autogenous shrinkage was observed 566 to increase with the increase in the water-to-binder ratio of UHP-SWSSC. Autogenous shrinkage 567 grew faster when the water-to-binder ratio was higher, especially at early ages, which may have occurred due to the effect of seawater on the ternary blend of OPC, slag, and silica fume. For 568 example, 69% and 80% of maximum autogenous shrinkage were reached in the first 7 days for 569 570 mixes with the water-to-binder ratio of 0.15 and 0.25, respectively (Figure 21). As discussed in 571 Section 3.4.2, the incorporation of seawater in pastes with a low water-to-binder ratio can increase 572 the reactivity of slag, especially at early ages [63]. A higher amount of seawater-induced ions and higher alkalinity in mixes with larger water content can accelerate the hydration rate of slag to a 573

greater extent, which means enhanced self-desiccation and greater autogenous shrinkage. This mechanism also explains the rapid initiation of autogenous shrinkage in higher water-to-binder ratio mixes. It is evident from **Figure 21** that the development of autogenous shrinkage is lower than drying shrinkage during the first 7 days in the SS-50-0.15 mix. However, for SS-50-0.25 mix with higher water content, growth of autogenous shrinkage after 7 and 28 days was significantly higher than drying shrinkage. No clear trend regarding the proportions of autogenous and drying shrinkage was obtained in mixes with different water-to-binder ratios (**Figure 22**).

# 581 [Figure 21 near here]

Although autogenous shrinkage usually governs the overall shrinkage in reactive powder concrete, overall drying shrinkage was found to be dominant in seawater and sea sand-based UHPC in this study, with a couple of exceptions of OPC-slag binary mix (SS-25-0.2) and a large portion of slag incorporated ternary mix (SS-62.5-0.2). The relatively lower proportion of autogenous shrinkage in UHP-SWSSC is advantageous particularly in terms of early-age autogenous shrinkage-induced cracking.

588 [Figure 22 near here]

## 589 4. Selection of a Proper Mix based on Physical, Mechanical and Dimensional Stability

590 **Properties** 

591 Physical, mechanical, and dimensional stability properties of the industrial by-product based 592 seawater and sea sand UHPC mixes are compared in **Table 8** to evaluate their performance. 593 Properties of the individual mixes were compared against the control mix (SS-0-0.2). An increase 594 or a decrease in the properties with respect to the control mix has been quantified by a positive or 595 negative percentage, respectively. It is evident from the comparison that the OPC replaced UHP-

SWSSC mixes possessed slightly lower fresh as well as 28-day unit weight. Workability of UHP-596 SWSSC can be an issue if a high proportion of slag is added (>37.5% of the total binder) or the 597 598 water-to-binder ratio is kept relatively low (<0.2). Although early age compressive strengths were smaller than the control mix, strength increased with time and exceeded the control mix after 90 599 days. Overall, autogenous shrinkage was higher (except SS-50-0.15) and drying shrinkage was 600 601 lower with respect to the control mix. When added together, total shrinkage was also lower compared to the control mix (except SS-50-0.25). Comparing the mechanical and dimensional 602 603 stability properties, the SS-50-0.2 mix is recommended in this study. Compared to the control mix, 604 this mix achieved slightly higher long-term strength (7.47%) and lower total shrinkage strain (10.95%) without compromising the workability. Although a marginal reduction in workability 605 (5.56%) and a moderate increase in the autogenous shrinkage (15.92%) was observed, this blend 606 607 yielded satisfactory performance among other UHP-SWSSC mixes with different OPC substitution percentages and water-to-binder ratios. 608

609 XRF analysis of ground concrete powder reveals the chloride content of mature UHPC samples. 610 Most importantly, the chloride content of the seawater and sea sand UHPC mix was determined to be 8.0 kg/m<sup>3</sup>, as opposed to 0.72 kg/m<sup>3</sup> for tap water and river sand-based UHPC. Chloride content 611 612 found in UHP-SWSSC was found to be 10 times higher than the AS 1379 [67] limits of 0.8 kg/m<sup>3</sup> 613 for normal class concrete. However, the experimental results of this study provide evidence that 614 marine resources can still be utilized in UHPC fabrication for unreinforced applications or may be 615 embedded with corrosion-resistant reinforcement, such as FRPs. The durability study of seawater and sea sand UHPC mixes which is currently underway by this research group will provide critical 616 617 data for a well-informed cost-benefit-sustainability analysis.

618 [Table 8 near here]

619 **5.** Conclusions

The following conclusions can be drawn within the limited scope of this preliminary study on thepotential of using seawater and sea sand in UHPC fabrication:

- Seawater accelerates cement hydration and releases more heat from the formation of Friedel's salt in the early ages of hydration. This is also reflected in the early-age compressive strength.
- Incorporation of seawater and sea sand in UHPC yields marginally higher fresh and hardened
   density compared to river sand and washed beach sand counterparts, whereas industrial by products such as ground slag and silica fume reduce the unit weight of UHP-SWSSC.
- The use of sea sand and seawater in UHPC slightly decreases its workability. However, silica fume in an OPC-slag-silica fume ternary blend offers better workability compared to OPCslag binary blends. A sufficiently workable mix at a water-to-binder ratio of 0.2 can be achieved under an optimized superplasticizer dose with a maximum 50% OPC replacement with SCMs.
- Sea sand and seawater UHPCs provide a higher rate of strength development especially up to
   28 days, beyond which, the strength development slows down to reach similar 90 days
   strength. Slag and silica fume offer a similar 90 days strength from the delayed secondary
   hydration products from their pozzolanic reactivity. The optimum water-to-binder ratio to
   achieve maximum long-term compressive strength was found to be 0.2.
- Seawater curing lowers the long-term strength of OPC-based UHPC and the early age strength
   of SCM incorporated UHPC. However, UHPC with SCMs experiences marginal deterioration
   under seawater environment compared to cement-only counterpart.
- UHP-SWSSC produces high early-age autogenous shrinkage and considerably higher drying shrinkage than tap water and river sand or washed beach sand variants because of the

accelerated hydration of cement. SCM incorporation in the mix also increases autogenous
 shrinkage from the accelerated hydration of cement particles, but significantly reduces drying
 shrinkage from densifying the UHPC matrix, and hence reduces the overall shrinkage.

A UHP-SWSSC mix with 50% OPC replaced by 37.5% slag and 12.5% silica fume is
 recommended in this study, which can achieve satisfactory workability, long-term strength,
 and dimensional stability properties.

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(a) (b) Figure 1. Preparation and modification of sea sand (a) Impurities in sea sand, (b) sea sand after processing.



Figure 2. Particle size distribution curves of different sand aggregates (cumulative passing).



(c) Figure 3. SEM images of different sand particles and SCMs (a) Sea sand, (b) washed beach sand, (c) river sand, (d) ground slag, (e) silica fume.



Figure 4. (a) Heat flow and (b) cumulative heat captured in an isothermal condition in the first 72 hours of hydration.



Figure 5. Comparison of workability of fresh UHP-SWSSC (by flow table test) based on (a) percentage of OPC replacement, (b) water to binder ratio, (c) different sand types.



Figure 6. Effect of seawater and sea sand as mixing water and aggregate (tap water curing).



Figure 7. Effect of OPC replacement on cube compressive strength of UHP-SWSSC (tap water curing).



Figure 8. Effect of OPC replacement on cube compressive strength of UHP-SWSSC (seawater curing).



Figure 9. Effect of water to binder ratio on cube compressive strength (seawater curing).



Figure 10. Cross-sections of UHP-SWSSC with different water to binder ratio (a) SS-50-0.15, (b) SS-50-0.2, (c) SS-50-0.25.



Figure 11. Effect of seawater curing on UHPC (100% OPC) with different aggregate types.



Figure 12. Effect of seawater curing on UHPC (50% OPC replacement) with different aggregate types.



Figure 13. Effect of sand types on shrinkage of UHPC (a) Total shrinkage, (b) Autogenous shrinkage, (c) Drying shrinkage.



AS – autogenous shrinkage, DS – drying shrinkage.

Figure 14. Percentage of autogenous and drying shrinkage development among mixes with different types of sand.



Figure 15. Proportions of autogenous and drying shrinkage at 120 days (different types of sand).



Figure 16. Influence of OPC replacement on shrinkage of UHPC (a) Total shrinkage, (b) Autogenous shrinkage, (c) Drying shrinkage.



Figure 17. Autogenous shrinkage cracks in the microstructure of UHPC (a) 100% OPC, (b) 25% OPC replaced by slag.



AS – autogenous shrinkage, DS – drying shrinkage.

Figure 18. Percentage of autogenous and drying shrinkage development among mixes with different OPC replacement.



Figure 19. Proportions of autogenous and drying shrinkage at 120 days (percentage of OPC replacement).



Figure 20. Influence of water-to-binder ratio on shrinkage of UHPC (a) Total shrinkage, (b) Autogenous shrinkage, (c) Drying shrinkage.



AS – autogenous shrinkage, DS – drying shrinkage.

Figure 21. Percentage of autogenous and drying shrinkage development among mixes with different water to binder ratio.



Figure 22. Proportions of autogenous and drying shrinkage at 120 days (water to binder ratio).

# **Figure captions**

Figure 1. Preparation and modification of sea sand (a) Impurities in sea sand, (b) sea sand after processing.

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Figure 14. Percentage of autogenous and drying shrinkage development among mixes with different types of sand.

Figure 15. Proportions of autogenous and drying shrinkage at 120 days (different types of sand).

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Figure 21. Percentage of autogenous and drying shrinkage development among mixes with different water to binder ratio.

Figure 22. Proportions of autogenous and drying shrinkage at 120 days (water to binder ratio).

Chemical properties (wt. %)	Portland cement	GGBFS	Silica fume
SiO <sub>2</sub>	19.37	32.13	92.18
Al <sub>2</sub> O <sub>3</sub>	4.88	13.66	0.09
Fe <sub>2</sub> O <sub>3</sub>	3.4	0.36	0.01
CaO	63.45	42.56	0.1
MgO	0.88	5.51	0.28
Na <sub>2</sub> O	0.48	0.31	0.22
K <sub>2</sub> O	0.37	0.31	0.26
TiO <sub>2</sub>	0.31	0.54	< 0.01
SO <sub>3</sub>	2.36	3.24	0.01
P <sub>2</sub> O <sub>5</sub>	0.06	0.01	0.1
Mn <sub>3</sub> O <sub>4</sub>	0.05	0.15	< 0.01
Loss on ignition (LOI)	4.25	0.95	6.25

Table 1. Chemical compositions of cementitious materials.

Ion (in mg/L)	Natural seawater from Malabar,	World average of seawater
	NSW	Teng et al. [29]
F -	10.6	1.3
Cl -	19131	19352
Br -	71.7	67.3
SO4 <sup>2-</sup>	2658	2712
PO4 <sup>3-</sup>	nil	nil
$NO_2^-$	nil	nil
NO <sub>3</sub> -	nil	nil
Li <sup>+</sup>	0.296	nil
Na <sup>+</sup>	10691	10784
$Mg^{2+}$	891.6	1283.7
$Al^{3+}$	0.112	nil
Si <sup>4+</sup>	0.140	nil
K <sup>+</sup>	427.4	399.1
Ca <sup>2+</sup>	419.5	412.1
Fe <sup>2+</sup>	0.002	nil

Table 2. Chemica	compositions of natural	seawater.
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Chemical properties (wt. %)	Sea sand	Washed beach	River sand
		sand	
SiO <sub>2</sub>	88.80	86.00	95.90
Al <sub>2</sub> O <sub>3</sub>	0.70	1.00	1.08
Fe <sub>2</sub> O <sub>3</sub>	0.44	0.28	0.23
CaO	4.58	5.32	0.03
MgO	0.17	0.16	< 0.01
Na <sub>2</sub> O	0.20	0.20	0.01
K <sub>2</sub> O	0.12	0.35	0.18
TiO <sub>2</sub>	0.06	0.03	0.08
SO <sub>3</sub>	0.04	0.02	< 0.01
P <sub>2</sub> O <sub>5</sub>	0.02	0.02	< 0.01
Mn <sub>3</sub> O <sub>4</sub>	< 0.01	< 0.01	< 0.01
SrO	0.03	0.02	0.04
Loss on ignition (LOI)	4.26	5.05	0.85

Table 3. Chemical compositions of different types of sand
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Physical properties	Sea sand	Washed beach	River sand	River sand
		sand	(natural)	(processed)
Oven dry density $(t/m^3)$	2.41	2.44	2.51	2.41
SSD density $(t/m^3)$	2.50	2.51	2.55	2.46
Apparent density (t/m <sup>3</sup> )	2.63	2.63	2.62	2.55
Uncompacted bulk density (t/m <sup>3</sup> )	1.47	1.42	1.50	1.47
Compacted bulk density (t/m <sup>3</sup> )	1.68	1.62	1.76	1.67
Water absorption (%)	3.46	2.95	1.68	2.35
Void (%)	38.94	41.73	40.42	39.04

Table 4. Physical properties of different types of sand.

Mix ID	Replacement	Aggregate	Mixing water	Water to	OPC	Ground	Silica	Sand	Coarse	Water	HRWR
	of OPC (%)			binder		Slag	fume		aggregate		
				ratio	kg/m <sup>3</sup>						
SS-0-0.2	0	Sea sand	Seawater	0.2	1200	0	0	1000	N/A	240	24
SS-25-0.2	25	Sea sand	Seawater	0.2	900	300	0	1000	N/A	240	24
SS-37.5-0.2	37.5	Sea sand	Seawater	0.2	750	300	150	1000	N/A	240	24
SS-50-0.2	50	Sea sand	Seawater	0.2	600	450	150	1000	N/A	240	24
SS-62.5-0.2	62.5	Sea sand	Seawater	0.2	450	600	150	1000	N/A	240	24
SS-50-0.15	50	Sea sand	Seawater	0.15	615	461.3	153.7	1025	N/A	184.5	24.6
SS-50-0.25	50	Sea sand	Seawater	0.25	585.7	439.3	146.4	976.2	N/A	292.9	23.43
WBS-0-0.2	0	Washed beach sand	Tap water	0.2	1200	0	0	1000	N/A	240	24
WBS-50-0.2	50	Washed beach sand	Tap water	0.2	600	450	150	1000	N/A	240	24
RS(P)-0-0.2	0	River sand	Tap water	0.2					N/A		
		(processed)			1200	0	0	1000		240	24
RS(N)-50-0.2	50	River sand (natural)	Tap water	0.2	600	450	150	1000	N/A	240	24
RS(P)-50-0.2	50	River sand	Tap water	0.2					N/A		
		(processed)			600	450	150	1000		240	24
NC-0-0.35	0	Washed beach sand	Tap water	0.35	380	0	0	1102	840	133	1.9

Table 5. Mixture proportions of materials for UHPC and normal strength concrete.

SS – Sea sand, WBS – Washed beach sand, RS(P) – Processed river sand, RS(N) – Natural river sand, NC – Normal strength concrete.

Mix ID	Fresh density	SSD density					
	kg/m <sup>3</sup>	kg/m <sup>3</sup>					
SS-0-0.2	2400.8	2421.0					
SS-25-0.2	2373.3	2392.0					
SS-37.5-0.2	2316.8	2335.4					
SS-50-0.2	2305.0	2323.0					
SS-62.5-0.2	2304.8	2315.8					
SS-50-0.15	2308.6	2315.4					
SS-50-0.25	2247.5	2264.8					
WBS-0-0.2	2392.3	2424.4					
WBS-50-0.2	2285.1	2291.2					
RS(P)-0-0.2	2379.4	2394.2					
RS(N)-50-0.2	2322.2	2338.1					
RS(P)-50-0.2	2264.3	2299.1					
NC-0-0.35	2252.2	2270.8					
SS – Sea sand, WBS – Washed beach sand, RS(P) – Processed river sand, RS(N) – Natural river sand, NC – Normal strength concrete.							

Table 6. Fresh and SSD unit weight of UHPC and normal strength concrete.

Mix ID SS-0-0.2 SS-25-0.2 SS-37.5-0.2	Curing	1 day	(MPa)	3 days (MPa)		7 days (MPa)		28 days (MPa)		90 days (MPa)		
	water	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	
Mix ID SS-0-0.2 SS-25-0.2 SS-37.5-0.2 SS-50-0.2 SS-62.5-0.2 SS-50-0.15 SS-50-0.25 WBS-0-0.2 WBS-0-0.2 WBS-50-0.2 RS(P)-0-0.2 RS(P)-50-0.2 RS(P)-50-0.2 NC-0-0.35	TW curing	05 72	1.62	106.33	1.74	113.27	6.00	127.43	0.58	141.13	6.10	
	SW curing	85.23	1.05	102.93	2.96	104.10	2.36	112.20	4.69	124.27	2.70	
88.25.0.2	TW curing	57.01	2 1 6	99.86	3.35	118.40	3.31	131.35	4.17	144.39	0.59	
33-23-0.2	SW curing	57.01	5.10	99.67	2.48	111.66	5.17	117.27	9.48	126.47	2.66	
55 27 5 0 2	TW curing	38.94	1 5 1	93.40	1.31	117.17	5.15	138.63	7.02	143.23	7.27	
SS-37.5-0.2	SW curing		1.51	91.50	0.78	112.03	0.15	129.00	3.20	130.80	8.52	
SS-50-0.2	TW curing	27.78	0.22	91.63	2.50	116.53	1.48	136.80	2.00	146.50	9.90	
	SW curing	57.20		77.30	1.41	106.30	2.94	124.33	5.92	133.55	3.04	
SS-62.5-0.2	TW curing	22.85	1 10	75.60	6.47	107.07	3.00	131.10	5.79	148.15	0.21	
	SW curing		1.10	80.77	1.76	101.60	3.39	113.07	4.87	139.10	3.67	
SS 50 0 15	TW curing	52.43	4.95	91.67	3.88	112.50	0.56	128.05	0.07	140.25	6.86	
55-50-0.15	SW curing			88.83	9.67	110.83	6.43	122.50	3.80	133.77	1.05	
SS-50-0 25	TW curing	21.09	21.00	0.67	73.03	1.90	94.43	7.25	124.20	3.01	128.97	2.32
55-50-0.25	SW curing		0.07	70.89	2.80	90.67	8.14	114.73	7.58	129.05	0.35	
WDS 0.0.2	TW curing	48.93	18 03	1 5 1	101.07	1.85	106.97	4.01	116.70	9.35	138.47	1.29
WDS-0-0.2	SW curing		1.51	101.59	1.59	101.60	5.31	116.60	4.00	126.47	3.04	
WBS 50.0.2	TW curing	28.98	274	78.40	4.26	102.33	6.38	129.10	6.21	147.57	2.50	
WDS-50-0.2	SW curing		5.74	65.83	1.71	93.43	4.40	111.70	9.81	131.80	4.00	
$PS(D) \cap O 2$	TW curing	60.07	0.81	101.68	5.91	121.90	0.14	133.50	0.53	148.30	5.62	
KS(I)-0-0.2	SW curing	00.07	0.81	99.47	4.90	108.43	8.33	134.97	3.08	134.50	3.31	
<b>PS(N) 50 0 2</b>	TW curing	28.28	0.80	76.80	0.80	103.73	8.17	131.53	5.98	142.27	6.82	
KS(IN)-30-0.2	SW curing	20.70	0.80	74.53	1.15	95.97	2.25	113.00	10.65	130.07	1.50	
<b>DS(D)</b> 50.0.2	TW curing	22 55	0.56	78.58	3.96	114.30	2.51	133.97	3.75	146.70	4.95	
NS(1)-50-0.2	SW curing	55.55	0.50	70.45	3.59	96.90	1.13	120.20	5.71	139.23	2.50	
NC-0-0.35	TW curing	19.10	0.43	24.40	1.44	30.13	1.41	36.54	0.89	41.20	1.61	

Table 7. Compressive strength of UHPC and normal strength concrete (cube specimens).

SS – Sea sand, WBS – Washed beach sand, RS(P) – Processed river sand, RS(N) – Natural river sand, NC – Normal strength concrete, TW – Tap water, SW – Seawater, SD – Standard deviation.

	Variation in Unit		Variation	Variati	on in comp	ressive	Variation in maximum				
	W	eight	in	strengtl	strength (seawater curing)			shrinkage			
Mix ID	Fresh unit weight	28-day unit weight	workability	1-day	28-day	90-day	AS	DS	TS		
SS-0-0.2	0%	0%	0%	0%	0%	0%	0%	0%	0%		
SS-25-0.2	-1.14%	-1.20%	-12.50%	-33.11%	+4.52%	+1.77%	+50.90%	-35.07%	-7.15%		
SS-37.5-0.2	-3.50%	-3.53%	-1.39%	-54.32%	+14.97%	+5.26%	+6.72%	-18.37%	-10.14%		
SS-50-0.2	-3.99%	-4.04%	-5.56%	-56.26%	+10.81%	+7.47%	+15.92%	-22.01%	-10.95%		
SS-62.5-0.2	-4.00%	-4.34%	-19.44%	-73.19%	+0.77%	+11.94%	+52.64%	-41.60%	-10.67%		
SS-50-0.15	-3.84%	-4.36%	-48.61%	-38.48%	+9.18%	+7.64%	-4.05%	-43.80%	-30.76%		
SS-50-0.25	-6.38%	-6.45%	0%	-75.25%	+2.26%	+3.85%	+39.18%	-12.70%	+4.33%		
WBS-50-0.2	-4.82%	-5.36%	-2.78%	-66.00%	-0.45%	+6.06%	+5.48%	-76.82%	-49.88%		
RS(P)-50-0.2	-5.68%	-5.03%	-6.94%	-60.63%	+7.13%	+12.04%	+7.38%	-71.99%	-46.07%		
NC-0-0.35	-6.19%	-6.20%	N/A	-77.59%	-67.43%	-66.85%	-80.60%	-54.78%	-63.26%		

Table 8. Percentage variation of physical, mechanical and stability properties of different mixes compared to reference UHP-SWSSC.

AS – autogenous shrinkage, DS – drying shrinkage, TS – total shrinkage.

NC-0-0.35 was cured in tap water.

A positive (+) value indicates an increase, and a negative (-) value indicates a decrease.