Development of a 3D-Printable Ultra-High Performance Fiber-Reinforced Concrete for Digital Construction

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Abstract

This paper presents the systematic development and performance characterization of a non-proprietary 3D-printable ultra-high-performance fiber-reinforced concrete (UHPFRC) for digital construction. Several fresh and hardened properties of the developed 3D-printable UHPFRC matrix (without fiber) and composite (with 2% volume fraction of steel fibers) were evaluated and compared to that of conventionally mold-cast UHPFRC. Additionally, the effects of testing direction on the compressive and flexural strengths of the printed UHPFRC were investigated. The fresh properties of the UHPFRC developed in this study satisfied the criteria for extrudability, buildability, and shape-retention-ability, which are relevant for ensuring printability. The printed UHPFRC exhibited superior flexural performance to the mold-cast UHPFRC due to alignment of the short fibers in the printing direction. The high compressive and flexural strengths, along with the deflection-hardening behavior, of the developed UHPFRC can enable the production of thin 3D-printed components with significant reduction or complete elimination of conventional steel bars.

Keywords

ultra-high-performance fiber-reinforced concrete (UHPFRC); mechanical properties; extrusion; 3D concrete printing (3DCP)
1. Introduction

The automation of construction industry by means of Additive Manufacturing (AM) technologies such as 3D-concrete-printing (3DCP) has been gaining attention over the past decade. In the 3DCP technique, a low-slump cementitious material is extruded from a nozzle mounted on a gantry or a robotic arm to print a concrete structure layer-by-layer, similar to the fused deposition modeling (FDM) technique for printing thermoplastics. The 3DCP can improve the quality and speed of construction and eliminate the need for formwork, and thereby the associated labor, resulting in substantial time and cost savings. Furthermore, the layer-by-layer deposition process of 3DCP stimulates geometrical freedom to build complex multifunctional architectures [1-5]. These practical advantages of the 3DCP technique make it valuable for the construction industry.

One of the main restrictions of the extrusion-based 3DCP is the limited variety of 3D-printable concretes. An ideal printable cementitious material should possess low viscosity to allow uninterrupted flow during extrusion and high yield stress to prevent slumping of the material after deposition. At the same time, the material should set quickly to withstand the weight of the subsequent layers deposited above [6]. Few printable concretes with the above combination of properties have been reported in the literature – for example, normal strength concrete [7], geopolymer concrete [8, 9], earth-based mortar [10], and high-performance fiber-reinforced cementitious composite (HPFRCC) [11, 12]. These materials were designed for a specific printer with particular printing configurations (nozzle type and size, flow rate, pumping specifications, etc.) and will likely not work elsewhere due to change in material constituents or printing configuration. Therefore, application-specific cementitious materials should be systematically developed for extrusion-based 3DCP.

Several researchers have used fibers in concrete to overcome the challenge of using conventional steel reinforcement with the printing process. For instance, polypropylene (PP) micro fibers have been used in the 3D-printable HPFRCC to reduce shrinkage and deformation
in the plastic state [11, 12]. In a previous study, the authors of this paper showed that the 3D-printed geopolymers containing 0.75% and 1% (by volume) of PP fibers exhibited deflection-hardening behavior in bending [13]. Panda et al. [14] investigated the effects of short glass fiber on the mechanical properties of 3D-printed geopolymer concrete. Bos et al. [15] used short straight steel fibers to improve ductility of the normal strength 3D-printed concrete. Although partial reduction of conventional reinforcement (meant for shrinkage, temperature changes, and crack control) has been demonstrated in the previous studies, substantial reduction or complete elimination of conventional steel bars requires printable concretes with significantly greater flexural and tensile strengths.

This study aims to systematically develop a printable ultra-high performance fiber reinforced concrete (UHPFRC) to address the above-mentioned challenges of 3DCP. The ultra-high compressive strength of the UHPFRC enables the printing of complex volumetric profiles with slender cross sections, thereby fully utilizing the geometric freedom offered by the 3DCP technology. In addition, very high flexural strength with a deflection-hardening behavior of the UHPFRC significantly reduces the dependence on the conventional steel bars. Although an ultra-high-performance concrete (UHPC) matrix (without fibers) has been printed before [16] the printing of UHPFRC has not been reported so far. In addition, the reported printable UHPC had a proprietary formulation and its design was not reported. Thus, this research demonstrates the systematic development and 3D-printing of UHPFRC for the first time in the literature.

The fresh and hardened properties of the developed 3D-printable UHPFRC were reported in this article. Fresh properties of extrudability and buildability were assessed qualitatively, whereas workability and shape-retention-ability were quantified experimentally. Hardened properties, including apparent porosity, bulk density, compressive strength, flexural strength and interlayer bond strength, of the printed UHPFRC matrix (without fiber) and composite (with fiber) were determined through standard tests. The hardened properties of the printed UHPFRC were also compared with those of the conventionally mold-cast UHPFRC. In addition, the effects of testing direction on the mechanical properties of the printed UHPFRC were investigated. The
practical applicability of the developed UHPFRC was validated by 3D-printing a relatively large component with a complex geometry.

2. Materials and mix proportions

In this research, the binder for the UHPFRC consisted of general purpose (Type GP) Portland cement, conforming to AS 3972 [17] and silica fume (micro silica), conforming to AS 3582 [18] with specific surface area of 18 m²/g. The chemical compositions of cement and silica fume are presented in Table 1.

Table 1 Chemical compositions of cement and silica fume

| Oxides (wt%) | Cement ¹ | Silica fume ² |
|-------------|----------|---------------|
| Al₂O₃       | 4.47     | 0.33          |
| SiO₂        | 20.34    | 98.4          |
| CaO         | 62.91    | 0.01          |
| Fe₂O₃       | 4.58     | 0.31          |
| K₂O         | 0.29     | 0.01          |
| MgO         | 1.24     | --            |
| Na₂O        | 0.31     | 0.01          |
| P₂O₅        | --       | 0.2           |
| TiO₂        | --       | 0.01          |
| SO₃         | 2.58     | --            |
| LOI*        | 3.27     | 1             |

*LOI = Loss on ignition

¹ Adopted from [7]
² Provided by the supplier

Three types of sieve graded silica sands – each with different particle sizes – were used in this study. The fine, medium, and coarse silica sands had median particle sizes (D₅₀) of 176 µm, 498
µm, and 840 µm respectively. A polycarboxylate ether-based superplasticizer (SP) was used to adjust the workability. Sodium carboxymethyl cellulose water-soluble polymer powder was used as a viscosity modifying agent (VMA) to adjust the rheological properties of the mixture with respect to 3D printing process. Brass-coated short straight steel fibers with the properties given in Table 2 were used in the UHPFRC at a volume fraction of 2%.

Table 2 Properties of steel fibers

| Diameter (µm) | Length (mm) | Young's modulus (GPa) | Density (g/cm³) | Strength (MPa) |
|---------------|-------------|-----------------------|-----------------|----------------|
| 200           | 13          | 200                   | 7.85            | 2500           |

Two types of methods were used to quantify the particle size distributions of the dry granular raw materials. Sieve analysis as per ASTM C136 [19] was carried out for all three types of sands. Particle size distributions of cement and silica fume were measured using the laser ultrasonication technique with a particle size analyzer. The particle size distributions of the dry granular materials are shown in Figure 1.
The mix proportions of the UHPFRC matrix were calculated using the modified Andreasen and Anderson (modified A&A) \textsuperscript{[20]} model for maximizing the particle packing density of the dry materials. The modified A&A packing model has been successfully applied for the mixture optimization in the design of conventional concrete \textsuperscript{[21]}, lightweight concrete \textsuperscript{[22]} and self-compacting concrete \textsuperscript{[23]}. Ragalwar et al. \textsuperscript{[24]} investigated the influence of the distribution modulus ($q$) on the properties of the UHPFRC matrix. Yu et al. \textsuperscript{[25]} recommended the value of $q = 0.23$ for the production of UHPFRC mixes. The mathematical equation for the target particle size distribution curve in the modified A&A model is presented in Equation 1.

$$ P_{\text{tar}}(D) = \frac{D^q - D_{\text{min}}^q}{D_{\text{max}}^q - D_{\text{min}}^q} $$

where $D$ is the particle size, $P_{\text{tar}}(D)$ is the ratio of total dry particles with diameter smaller than $D$, $D_{\text{max}}$ and $D_{\text{min}}$ are the maximum and minimum particle sizes of the mix, respectively. The
parameter $q$ is called the distribution modulus, which controls the curvature of the $P_{tar}(D)$. Physically, $q$ represents the fraction between the coarse and fine particles of the mixture. In general, greater $q$ leads to a coarser mixture. The $P_{tar}(D)$ used in this study utilized $D_{max}$, $D_{min}$, and $q$ of 1.2 mm, 0.1 µm, and 0.23, respectively. The proportions of ingredients were iteratively calculated by minimizing the sum of squared-error between $P_{tar}(D)$ and the composite particle size distribution at discrete particle sizes. The target curve and the composite particle size distribution of the UHPFRC matrix are presented in Figure 2.

![Figure 2 UHPFRC mix design curve vs. modified A&A target curve](image)

Several trials of the UHPFRC matrix and composite were conducted for determining the appropriate SP and VMA contents to satisfy the extrudability, buildability, and workability requirements. The mix proportions of the printable UHPFRC are presented in Table 3.
Table 3 Mix proportions of the printable UHPFRC

| Binder | Silica sand | Water | SP   | VMA | Fiber |
|--------|-------------|-------|------|-----|-------|
| Cement | Silica fume | Fine  | Medium | Coarse | 0.16  | 0.015  | 0.003  | 2%   |
| 0.7    | 0.3         | 0.4   | 0.3   | 0.3  | 0.16  | 0.015  | 0.003  | 2%   |

Note: All numbers are mass ratios of the binder weight, except the fiber content (volume fraction).

3. Experimental methods

3.1. Mixing

A 20L Hobart mixer was used for mixing. First, dry materials were mixed for 5 minutes. Approximately 75% of the water was added and mixed for 5 additional minutes. At this point, the remaining water was mixed with the SP and gradually added to the mixture. The mixing was continued for another 9 minutes until a good rheology was achieved for facilitating homogeneous fibers dispersion. The fibers were then gradually added and mixed for 10 minutes. Finally, the VMA powder was added and mixed for an additional minute. The total mixing time was 30 minutes. Visual assessment by the authors confirmed the absence of fiber segregation in the fresh mixture.

3.2. Printing and casting

As shown in Figure 3 (a), a small-scale custom-made 3D printer was used in this study to simulate the extrusion-based 3DCP process. A piston type extruder having a 45º nozzle opening of 30 mm (W) × 15 mm (H) was used in the printer. The extruder was a metallic cylinder with a diameter of 50 mm and a length of 600 mm. Figure 3 (b) shows the extrusion of the printable UHPFRC matrix through the nozzle. Two-layer specimens measuring 30 mm (W) × 30 mm (H) × 280 mm (L) were printed at a constant speed with a print-time interval (delay time) of 2 minutes. In addition, 25 mm cubes and prisms with the dimensions of 25 mm × 25 mm × 280 mm were conventionally mold-cast for comparison purposes. It should be noted that the mixture
proportions and mixing procedure of the conventionally mold-cast specimens were identical to those of the printed specimens.

![Extruder and Steel frame](image)

45° nozzle

UHPFRC matrix

Figure 3 (a) Small-scale custom-made 3D printer used in the study, (b) Extrusion of 3D-printable UHPFRC matrix

### 3.3. Curing

After completion of the printing process, the printed specimens were immediately placed inside a sealed container for 24 hours to reduce water loss. The conventionally mold-cast specimens were covered with plastic sheets and de-molded after 24 hours. An elevated temperature curing procedure was adopted in this study to accelerate the hydration reactions. At the end of 24 hours, all printed and mold-cast specimens were transferred to a water tank and cured at the room temperature (20 ± 3°C) for seven days. The specimens were then put inside an oven and heat cured for four days in water at 90°C, followed by two days in air at 90°C. According to Ranade et al. [26], the 14-day compressive strength of the samples cured using the above procedure is equivalent to the 90-day compressive strength of the samples cured in the ambient conditions. At the end of the accelerated curing process, the surfaces of the printed specimens were ground to have flat surfaces for testing.

### 3.4. Test methods

#### 3.4.1. Fresh properties
Flow tests in accordance with ASTM C1437 [27] were conducted to measure the workability of the fresh UHPFRC matrix and composite. Two orthogonal diameters of the mixture flow were recorded before and after the drops of the flow table.

A qualitative method was adopted to evaluate the extrudability of the UHPFRC matrix and composite through a piston type extruder. The acceptance criterion for the extrudability test was the successful extrusion of five single layers (not stacked) of the fresh mixtures through the nozzle without any obstruction such as blockage, tearing, segregation or bleeding. Each of the five layers was at least 250 mm long with a cross-section of 30 mm (W) × 15 mm (H) extruded using the small-scale printer with a piston type extruder (described in section 3.2). Similar approaches were used elsewhere [8, 11] to evaluate extrudability.

Similar to the extrudability test, a qualitative method was adopted to evaluate the buildability of the UHPFRC matrix and composite. Seven layers of the UHPFRC matrix and composite were printed using the small-scale printer (described in section 3.2). Each layer had the dimensions of 15 mm (H) × 30 mm (W) × 280 mm (L) and the delay time between layer depositions was approximately 2 minutes. The acceptance criterion for buildability was the successful layer upon layer build-up of the fresh mixtures without any vertical distortion, layer failure or excessive deformation of the lower layers.

A quantitative test was set up to determine the shape-retention-ability (SRA, i.e. the ability of the fresh mixtures to hold their shapes after extrusion). In each test, the fresh mixture was filled into a plastic cylinder of 60 mm diameter and 60 mm height and tamped lightly. The plastic cylinder was lifted after two minutes (to simulate the delay time of 2 minutes used in the printing process) and the SRA of the freestanding mixtures was monitored using a camera. The fresh UHPFRC composite sample immediately after lifting of the plastic cylinder is shown in Figure 4. Circular steel plates of 60 mm diameter, each weighing 170 grams were loaded gradually one after the other on top of the freestanding fresh mixture until its failure. Before the application of circular steel plates, a plastic plate measuring 100 mm in diameter and 10 grams in weight was gently placed on top of the fresh mixture to distribute the pressure uniformly on
the fresh mixture. The images of the fresh samples were recorded at each stage of loading using the camera. These images were then processed digitally to quantify the change in diameter and height of the fresh sample.

Figure 4 Fresh UHPFRC composite sample immediately after lifting of the plastic cylinder for determining the shape-retention-ability
The SRA of the fresh UHPFRC matrix and composite is defined in Equation 2. SRA_d and SRA_h determine the dimensional variations with respect to the diameter and the height of the UHPFRC sample, respectively.

\[
SRA_d = \frac{d_L}{d}; \quad SRA_h = \frac{h_L}{h}
\]

where \(h_L, d_L\) are the height and diameter of the bottom of the freestanding UHPFRC sample at a given load \(L\) and \(h, d\) are the height and diameter of the bottom of the freestanding UHPFRC sample before load application.

### 3.4.2. Hardened properties

To measure the compressive strength of the printed UHPFRC matrix and composite, 25 mm cube specimens were extracted from the printed specimens by sawing and were loaded in three directions as shown in Figure 5. At least six printed cubes were prepared for testing in each direction. To compare the compressive strength of the printed UHPFRC with that of the mold-cast UHPFRC, at least six 25 mm mold-cast cube specimens were also tested. A constant load rate of 0.33 MPa/s was applied for all the compression tests.

![Figure 5 Loading directions for compression test of printed specimen (not to scale)](image-url)
To measure the modulus of rupture (MOR or flexural strength) of the printed UHPFRC matrix and composite, 280 mm × 25 mm × 25 mm printed specimens were tested in the perpendicular and lateral directions. At least six specimens were prepared for testing in each direction. Similar to the compression tests, at least six mold-cast specimens with the dimensions of 280 mm × 25 mm × 25 mm were prepared to compare the MOR of the printed UHPFRC with that of the mold-cast UHPFRC. A three-point flexure test with a span of 160 mm and a displacement control rate of 1 mm/min was used. Figure 6 shows the loading directions of the flexural test specimens.

![Figure 6 Loading directions for flexural test of printed specimen (not to scale)](image)

To measure the inter-layer bond strength of the UHPFRC matrix and composite, at least six 50 mm × 25 mm × 25 mm specimens were sawn from the printed layers. Two T-shape metallic brackets were glued to the top and the bottom of each specimen using an epoxy so that the specimens could be hung under uniaxial pull-out load without shear support. In addition, small notches of 5 mm depth (see Figure 7) were cut from the edges at the layer interface to increase the probability of the failure at the interface. The specimen was pulled in the direction...
perpendicular to the layer interface under displacement control at the rate of 1 mm/min. Care was taken to eliminate eccentricity while aligning the specimens vertically.

![Inter-layer bond strength test setup](image)

Figure 7 Inter-layer bond strength test setup

The apparent porosity and bulk density of the hardened UHPFRC matrix and composite were measured in accordance with ASTM C20 [28]. For each mixture, at least six specimens were collected from the residual pieces of the UHPFRC specimens tested under flexure. Details of the test can be found in Nematollahi et al. [13]. The apparent porosity and bulk density of each
UHPFRC mixture was calculated according to [28]. These properties of the mold-cast UHPFRC matrix and composite were also measured for comparison.

4. Results and Discussion

4.1. Fresh properties

The flow (spread diameter) of UHPFRC matrix and composite are presented in Figure 8. The spread diameters of the UHPFRC matrix and composite before drop of the flow table were close to the bottom diameter of the mini-slump cone used in this study (100 mm), indicating the fresh mixtures had almost zero slump before drop of the flow table, which is desirable for extrusion-based 3DCP. The spread diameter after the drop of the flow table satisfies the printability region recommended in [29]. As expected, the workability was slightly reduced with the addition of the fibers due to weak adhesive forces between the fibers and the matrix and increase in the surface area per unit volume of the composite that slightly increases the adsorption of the mix water.
Figure 8 Flow test results of the UHPFRC matrix and composite

Figure 9 shows five singly printed layers of the UHPFRC matrix and composite, each having a length of at least 250 mm. No blockage, tearing, segregation or bleeding was observed during the extrusion process, which indicates the developed UHPFRC matrix and composite passed the extrudability criterion. Figure 10 shows seven stacked printed layers of the UHPFRC matrix and composite. No vertical distortion, layer failure or excessive deformation of the lower layers was observed, which indicates the developed UHPFRC matrix and composite passed the buildability criterion.
Figure 9 Extrudability test results of the printable UHPFRC: (a) matrix; and (b) composite

Figure 10 Buildability test results of the printable UHPFRC: (a) matrix; and (b) composite

Figure 11 shows the photos of the UHPFRC matrix and composite samples under increasing vertical load for testing their SRA. Figure 12 plots the SRA indices (SRA₁ and SRA₂) of the UHPFRC matrix and composite as a function of the vertical load. Both the horizontal and vertical distortions (SRA₁ and SRA₂) of the UHPFRC composite were smaller than those of the UHPFRC matrix at all vertical loads. It can also be observed that the UHPFRC matrix sample failed at a vertical load greater than about 0.7 kg, which is lower than the load carried by the UHPFRC composite sample (1.2 kg). Therefore, the use of fibers improves SRA and increases the vertical load capacity of the fresh mixture.
Figure 11 Shape-retention-ability test results of the printable UHPFRC: (a) matrix; and (b) composite

Figure 12 Shape-retention-ability indices of the printable UHPFRC matrix and composite
4.2. Hardened properties

Figure 13 presents the results of the compression tests. The compressive strength of the mold-cast UHPFRC was higher than that of the 3D-printed UHPFRC in all loading directions. This is explained by the greater porosity of the printed UHPFRC due to lack of confinement and vibration (during printing) compared to the mold-cast UHPFRC (see Figure 19). The apparent porosity and bulk density results of both mold-cast and 3D-printed UHPFRC are presented in the subsequent section. The compressive strength of the printed UHPFRC is dependent on the loading direction. The pressure exerted during the extrusion process contributes to improved compaction of the UHPFRC in the longitudinal direction. As a result, the highest compressive strength is obtained in the longitudinal direction. In contrast, due to negligible confinement in the lateral direction (in-plane of the layer but perpendicular to the printing direction), the lowest compressive strength is observed in this direction. Due to the weight of the top layer, the bottom layer of the printed UHPFRC is slightly compacted. This results in larger compressive strength in the direction perpendicular to the layers, but it is still smaller than the compressive strength in the longitudinal direction. It should be pointed out that the anisotropic behavior of the printed UHPFRC is an inherent feature of the layer-by-layer manufacturing process. The quantitative difference in the strength values in different directions generally depend on many parameters including printing configurations and printing materials [30]. Similar anisotropic behavior has been reported in the literature for normal strength concrete [7], high performance concrete [12] and geopolymers [8, 13].
The MOR results are presented in Figure 14. The MORs of the UHPFRC matrix and composite in the perpendicular direction are slightly higher than the respective MORs in the lateral direction. This is consistent with the previous studies where the primary load resisting component was either normal strength concrete [17], high performance concrete [10], or geopolymer concrete [7]. The bottom layer carrying the tensile stress during a flexural test was compacted more in the perpendicular direction compared to the lateral direction due to the self-weight of the top layer, which explains the greater MOR in the perpendicular direction compared to that in the lateral direction.
Figure 14 MOR of the UHPFRC matrix and composite

According to Figure 14, the MORs of the printed specimens are significantly higher than those of the mold-cast specimens. This observation is in contrast to the compressive strength results presented in Figure 13, which is attributed to the printing process. As pressure is exerted in the longitudinal direction during printing, the steel fibers tend to align in the direction of concrete flow (the longitudinal direction). The printing nozzle size of 15 mm – which is only slightly larger than the length of steel fiber (13 mm) – aligns the fibers in the plane of the layer. The alignment of short fibers in the printing direction due to the extrusion process has been reported in the literature for 3D printed fiber-reinforced geopolymer composites [13, 31] and strain-hardening cementitious composites [32, 33]. On the other hand, the mold-cast specimens show a more random orientation of fibers in all the three dimensions [34].
Figure 15 Flexural stress versus mid-span deflection graphs of the UHPFRC matrix and composite

Figure 15 presents the flexural stress versus mid-span deflection graphs for the UHPFRC matrix and composite specimens. The deflection-hardening effect of steel fiber reinforcement is clearly visible in this figure, as all the composite specimens exhibited significantly greater load carrying capacity (after first-crack) than the respective matrix specimens. Figure 16 presents the flexural toughness values calculated from the area under the flexural stress versus mid-span deflection curves presented in Figure 15 up to mid-span deflection of 2.5 mm or complete failure, whichever occurs first. The superior flexural performance of the 3D printed UHPFRC to conventional mold-cast UHPFRC can be clearly seen in Figures 18 and 19. Furthermore, Figure 16 shows that the flexural toughness of the UHPFRC composites are about 1-2 orders of magnitude greater than the respective matrices.
Figure 16 Flexural toughness of UHPFRC matrix and composites

The inter-layer bond strength results are presented in Figure 17. The UHPFRC composite shows stronger bonding between the layers than the matrix. This shows that the inclusion of fibers improves the inter-layer bond strength, despite the fiber orientation being predominantly parallel to the inter-layer surface. Figure 18 shows the inter-layer bond stress versus displacement curves of the UHPFRC matrix and composite samples. Unlike the UHPFRC matrix sample, a displacement-hardening behavior was noticed in the UHPFRC composite specimen, which may be attributed to the development of multiple fine cracks in the printed layers adjacent to the inter-layer surface. When the inter-layer surface cracked, it still failed in a brittle manner as observed by the sudden drop of load in the composite specimen at the end of the stress-displacement curve in Figure 18, which supports the fact that there were no fibers bridging across the inter-layer surface. It should be noted that the observed significantly higher inter-layer bond strength in the printed UHPFRC composite as compared to that of the printed UHPFRC matrix shall be investigated comprehensively in our future research. The inter-layer bond strength of the UHPFRC observed in this study is significantly higher than the values reported in the literature for normal strength concrete [35], geopolymer concrete [8, 36], fiber-
reinforced concrete [12], fiber-reinforced geopolymer concrete [13] due to the superior mechanical properties of the UHPFRC.

Figure 17 Inter-layer bond strength of 3D-printed UHPFRC matrix and composite

Figure 18 Inter-layer bond stress versus displacement curves of 3D-printed UHPFRC matrix and composite
Figure 19 shows the apparent porosities of the UHPFRC matrix and composite. For both the mold-cast and 3D-printed UHPFRCs, the apparent porosity of the composite samples was significantly higher than that of the matrix. This is due to a fiber-induced damage effect causing greater air entrapment in the composite sample, and thereby higher porosity of the composite than the matrix [37]. Similar results were reported for strain-hardening geopolymer composites reinforced by different polymeric fibers [38, 39]. For both the matrix and composite samples, the apparent porosity of the 3D-printed UHPFRC was slightly higher than that of the mold-cast UHPFRC. This is due to the unconfined lateral expansion and the lack of vibration and compaction in the 3D-printed UHPFRC samples.

![Graph showing apparent porosity of UHPFRC matrix and composite](image)

**Figure 19** Apparent porosity of the UHPFRC matrix and composite
Figure 20 Bulk density of the hardened UHPFRC matrix and composite

The bulk densities of the UHPFRC matrix and composite are presented in Figure 20. For both the mold-cast and 3D-printed UHPFRCs, the density of the composite samples was slightly higher than that of the corresponding matrix due to the high density of the steel fibers. For both the matrix and composite samples, the density of the 3D-printed UHPFRC was slightly lower than that of the mold-cast UHPFRC. This result is in agreement with the slightly higher porosity of the printed UHPFRC specimens (see Figure 19).

5. 3D-printing a complex curvilinear component

A relatively large-scale component with a complex geometry was 3D-printed to ascertain the applicability of the developed 3D-printable UHPFRC in practice. In this regard, a gantry printer with an effective printing space of 1.8 m × 1.6 m × 1.8 m was used. An augur type extruder with a circular nozzle of 30 mm diameter was attached to the gantry system. The printing speed and extrusion rate of the printer were 30 mm/s and 1.3 L/min. The fresh UHPFRC composite was
successfully extruded through the nozzle in a single attempt without any obstruction caused by blockage, tearing, segregation or bleeding. Figure 21 shows the printed structure consisted of 25 layers. The length of each layer was approximately 2 m and the layer height was about 10 mm. From the elevation view of the layers, no signs of vertical distortion, layer failure or excessive deformation were observed in the built-up layers. Thus, the suitability of the developed 3D-printable UHPFRC for production of relatively large-scale component with complex geometries is experimentally established. The detailed test results of the printed curvilinear structure will be published in a future article.

![Figure 21 3D printing of a complex curvilinear component using the developed 3D-printable UHPFRC mixture: (a) during 3D-printing process; (b) top-view; and (c) side-view](image)

6. Conclusions

This study demonstrates the mix design development and performance characterization of a non-proprietary 3D-printable ultra-high-performance fiber-reinforced concrete (UHPFRC) mixture for 3D construction printing applications. Conventionally mold-cast UHPFRC matrix and composite were prepared for comparison purposes. The fresh properties of workability, extrudability, buildability, and shape-retention-ability of the developed 3D-printable UHPFRC matrix (without fibers) and composite (with steel fibers) were investigated. The hardened properties of compressive, flexural and inter-layer bond strengths, as well as apparent porosity
and bulk density of the developed 3D-printable UHPFRC matrix and composite were measured. Further, effects of the loading direction on the compressive and flexural strengths of the printed UHPFRC matrix and composite were also studied. The main conclusions of the study are:

1. The presence of short straight steel fibers in the 3D-printable UHPFRC influenced its fresh properties. While the workability of the UHPFRC composite was slightly lower than that of the UHPFRC matrix, the shape-retention-ability of the composite was significantly better than that of the matrix. Both the UHPFRC matrix and composite had good extrudability and buildability based on the qualitative criteria selected in this study.

2. (a) The compressive strength, modulus of rupture (MOR) and flexural toughness of the developed UHPFRC composite were significantly enhanced due to the addition of short steel fibers as compared to those of the UHPFRC matrix. All UHPFRC composite samples exhibited clear deflection-hardening behavior in bending. (b) However, the apparent porosity also increased with the addition of the steel fibers. Both (a) and (b) are true for both the mold-cast and 3D-printed UHPFRCs.

3. The compressive strength of the printed UHPFRC was dependent on the loading direction. It was the highest in the longitudinal direction, followed by that in the perpendicular direction, and the strength was the lowest in the lateral direction of loading. This pattern is true for both the matrix and composite specimens. The compressive strength of the mold-cast UHPFRC was higher than the printed UHPFRC, which is also true for both the matrix and composite samples. This is due to better compaction and confinement of the mold-cast specimens.

4. For both the matrix and composite samples, the MOR of the printed UHPFRC in the perpendicular direction was slightly higher than that in the lateral direction. In contrast to the compressive strength results, the printed UHPFRC samples exhibited superior flexural performance to the mold-cast UHPFRC samples. The MORs of the printed UHPFRC specimens in both perpendicular and lateral directions were significantly higher than those of the mold-cast specimens. This is attributed to the alignment of short fibers in the printing
direction during the extrusion process, resulting in the 2D orientation of steel fibers in the plane of the printed layers, as opposed to a more random 3D fiber distribution in the mold-cast specimens.

5. The inter-layer bond strength of the printed UHPFRC composite was significantly higher than that of the matrix, despite the fact that there was no apparent bridging of the inter-layer surfaces by the steel fibers. Further research is needed to fully understand the underlying reasons for this behavior.

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