Microstructural and hardness evolution of additively manufactured Al–Si–Cu alloy processed by high-pressure torsion

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ABSTRACT
Nanostructured Al-9%Si-3%Cu alloy was achieved by direct metal laser sintering (DMLS) and then processed using high-pressure torsion (HPT) processing, which resulted in considerable grain refinement down to 60 nm associated with a substantial dislocation density up to \(6.2 \times 10^{14} \text{ m}^{-2}\) and a significant reduction in the porosity. Hardness measurements across the horizontal and vertical cross sections showed an improvement in the strength homogeneity for processed samples after 10 turns of HPT processing. These results indicate that a controllable ultradisperse-grained microstructure can be achieved by employing additive manufacturing, followed by effective severe plastic deformation processing.

Introduction
Additive manufactured alloys are emerging engineering materials with benefits of controllable microstructures and 3D customized shapes to fit challenges in the industrial needs [1, 2]. Additive manufacturing of aluminium light alloys has received considerable attention because of the capability to control the alloying contents precisely, thus achieving the required properties and performance, as well as their high strength-to-weight ratio. Several techniques have been used in fabrication of additive manufactured aluminum alloys such as selective laser melting for Al–Si–Mg alloy [3, 4], Al–Si–Cu alloy [5], Al–Si alloy [6, 7], and direct metal laser sintering for Al–Si–Mg [8], Al–Cu–Mg/SiC composite [9]. In particular, the Al–Si–Cu alloys are employed in engine parts as engine blocks and pistons [10]. Alloying aluminum with silicon improves the casting fluidity, whereas adding copper and magnesium enhances solid solution and precipitation strengthening [11–13]. Considering the benefits of the additive manufactured alloy, some inevitable drawbacks come within their directly processed microstructures such as high porosity and powder
clusters that can cause serious cracking in the manufactured parts [2, 14]. Recently, new approaches have been adopted to control such drawbacks and alter the properties of the additive manufactured alloys. High-pressure torsion (HPT) was employed as an efficient technique for severe plastic deformation (SPD) to process selective laser melted Al–Si–Mg [3], 316 stainless steel [15] and Co–Cr–Fe–Ni [16]. These studies have focused only on horizontal cross sections of the HPT samples, where the vertical cross sections of these samples have not been investigated. Therefore, the current work focuses on the microstructural homogeneity over both horizontal and vertical cross sections for direct metal laser melted Al-9%Si-3%Cu alloy processed using high-pressure torsion. The study includes the evolution of grain refinement of the aluminium matrix and eutectic silicon phase, and porosity content as measured by different approaches. The distribution of eutectic nano-sized particles and evolution of dislocation density have also been inspected to evaluate the resultant strength in the additive manufactured processed alloy. Evaluating the relationship between the achieved microstructure and subsequent properties in the additive manufactured aluminum alloys processed by HPT will improve the understanding and knowledge of this processing technique in the densification and strengthening of these alloys. As well, this alloy is widely used in the manufacture of engine blocks and parts, therefore, alteration of the microstructure of the additive manufactured alloy will result in improved properties toward better performance under operating conditions at ambient and high temperatures.

Materials and methods

Table 1 shows the compositions of the Al-9%Si-3%Cu alloy manufactured by direct metal laser sintering (DMLS). The alloy was manufactured using a Concept Laser M2 machine under nitrogen atmosphere at room temperature using the following parameters: 200 W of laser power, 1000 mm/s of scan speed, 200 μm of hatch spacing, and 40 μm of layer thickness. The alloy was additively manufactured as a cylindrical rod of 150 mm in length and 11 mm in diameter. This alloy was built along the vertical direction relative to the manufacturing area as illustrated in Fig. 1. A wire discharge machine was employed to reduce the diameter of the rod alloy to 9.8 mm and cut to disks of thickness 1 mm. The disks were thinned to thicknesses of 0.85 mm by 800 grit abrasive paper in order to be placed between the HPT anvils. Under a quasi-constrained condition, HPT processing was achieved [17, 18], using the following parameters: 6.0 GPa applied pressure, 1 rpm rotational speed and number of turns (N) of 1/2, 1, 3, 5 and 10 turns at room temperature. The final average thickness of the HPT processed sample after 10 turns was ~ 0.7 mm. The following investigations were carried out on as-received and processed samples: microstructural observations, X-ray diffraction analysis, porosity and microhardness measurements. The samples were mechanically ground and polished using abrasive papers of (800–4000) grit and 1 μm diamond paste, respectively. These samples were etched using Keller’s reagent for 10 s and then cleaned with ethanol and compressed air, respectively. Then, the microstructures were observed through the horizontal and vertical cross sections that are represented in Fig. 1 using optical microscopy (OM, OLYMPUS-BX51) [19]. Scanning electron microscopy (SEM, FEI Quanta 200) was conducted on the horizontal cross sections of the etched samples. Elemental mapping of the as-received alloy was obtained using the energy-dispersive spectroscopy (EDS) to determine the chemical composition with weight fractions. Transmission electron microscopy (TEM, JEOL JEM-3010) was conducted to observe the microstructure at the edge region of each disk. Samples of 3 mm in diameter and 100 μm in thickness were taken from these areas and then subjected to electropolishing in 30% vol. of nitric acid in methanol at 20 V and 243 K using twin-jet electro-polishing facility (Struers). Patterns of the X-ray diffraction (XRD), size of crystallite domains and density of dislocations were attained from X-ray analysis of the as-received and processed samples using an XRD analyzer (Bruker, D2 Phaser) using a Kα-Cu with wavelength of 0.154 nm and a step size of 0.02° from 2θ range of 30–90°. The recorded data were refined.
using MAUD software using Rietveld refinement of the full peak X-ray profile based on aluminum peaks [19, 20]. The dislocation density (\(\rho\)) was computed via XRD data through the following relationship [18]:

\[
\rho = 2\sqrt{3}\varepsilon/d_b, \quad \text{where} \quad \varepsilon \text{ is the average microstrain, } d_b \text{ is the average crystallite size; both values have been gained by MAUD software, and } b \text{ is the vector of Burgers for the aluminum (0.286 nm [21]).}
\]

The area fraction of the eutectic silicon phase particles was calculated by employing the SEM micrographs and ImageJ analysis software. Area fraction of particles refers here to the particles total number times their average area divided by the field area on SEM micrograph [22]. Random locations on each SEM micrograph for each etched sample were selected as field areas, which transformed into black and white field areas using ImageJ software to calculate the area fraction of particles.

The porosity was measured through both Archimedes principle approaches and ImageJ analysis of OM micrographs [23, 24]. In the Archimedes method, an average of five readings for each mass measurement was recorded using a digital balance to four-digits accuracy [24, 25]. In ImageJ software analysis of OM micrographs, areas in OM micrographs of (3000 \times 3000) mm\(^2\) across the horizontal cross sections, and (1000 \times 1000) mm\(^2\) across the vertical cross sections were taken for the as-received and processed samples. This procedure was repeated over several OM micrographs for different cross sections to ensure more reliable measurement of porosity. The porosity was calculated according to the ASTM E562-01 [26] and average pore diameter was obtained using ImageJ software analysis in the pixelated OM micrographs. Vickers microhardness was recorded for the as-received and processed samples across both polished horizontal and vertical cross sections using a microhardness tester with 100 gf load for 15 s, and the recorded results were constructed as colour-coded maps using similar procedures as detailed in Ref. [19].

Results

Microstructural observations

The optical microstructures of the as-received material and that obtained after 10 HPT turns in the central region of the disks are shown, respectively, in Fig. 2a–c. These micrographs were taken over the horizontal cross sections of the disks. The as-received sample in the horizontal disk plane clearly shows melt pools with average width of 150 \(\mu\)m and different lengths, showing the island scanning strategy perpendicular to the build direction of the rod. Both process-induced and gas-induced porosities were observed in the polished and etched as-received samples, as indicated, respectively, by solid and dashed circles (Fig. 2a, b). However, a different pool morphology was observed in the samples after HPT processing, where these pools have been sheared along with the direction of HPT strain as the number of turns increased up to 10 turns as indicated by curved dashes arrows in Fig. 2c. The microstructures over the vertical cross sections of the as-received (parallel to the build direction) and HPT-processed samples are shown in Fig. 2d–f. A fish-scale like morphology is shown in the as-received sample, whereas an elongated microstructure is exhibited in the HPT-processed sample, as indicated by the straight arrows. SEM observations of etched samples revealed two main phases, dark \(\alpha\)-Al matrix and white eutectic Si continuous network, as seen in Fig. 3a. Process-induced porosity embedded with a few nonmelted powder particles can also be seen in Fig. 3b. The morphology of the alloy was altered following HPT processing as seen in Fig. 4a–f, where the eutectic phase has undergone a gradual fragmentation at the peripheries of processed disks as the number of HPT turns increased as exhibited in Fig. 4a, b. In addition, this phase has been sheared clearly toward the torsional HPT strain as indicated by yellow solid arrows in Fig. 4c–f. Finally, this phase showed relatively homogeneous distribution of eutectic nano-sized particles at the peripheries of processed samples as indicated by the area fraction measurements and illustrated in Fig. 5.

TEM observations of the HPT-processed microstructures at the edges of disks showed extensive grain refinement at early stages of HPT processing as revealed in Fig. 6a–c, with grain size of 150 nm after 1/2 turn, down to 60 nm after processing up to 10 turns. XRD patterns for the as-received sample exhibited several peaks, identified as belonging to aluminum and silicon, confirming the EDS data establishing the compositional analysis of the current alloy, as indicated in Fig. 7a. Post-HPT processing, aluminum (111) and (200) peaks continued to appear, silicon (220) and (311) peaks
disappeared and a shift in peak locations was observed for the processed sample in comparison with the as-received sample as illustrated in Fig. 7b. The fitted XRD pattern of sample processed for 10 turns using the Rietveld refinement procedure, and the values of microstrain and lattice parameter in the as-received and processed samples are shown in Fig. 7c, d.

**Porosity measurements**

A comparison of porosity between the as-received samples and after 10 HPT turns shows a significant reduction in the porosity fraction after HPT processing to about 73, 90 and 96%, respectively, based on measurements conducted by the Archimedes method, ImageJ analysis on (1) the horizontal and (2) vertical cross sections at the peripheries of the disks, as shown in Fig. 8. Considerable reduction in porosity fraction and pore diameter was noticed with increasing number of HPT turns in the horizontal cross sections at the peripheries of the disks, as illustrated in Fig. 9. Porosity fraction and average pore diameter are 0.3% and 7 µm for the as-received sample and 0.03% and 0.5 µm in the edge region of the horizontal cross section of the sample processed for 10 turns.

**Hardness measurements**

The microhardness distribution recorded on the horizontal and vertical cross sections of the as-received and processed samples are illustrated in Figs. 10 and 11. The as-received sample showed a Vickers microhardness about 120 Hv. After HPT processing, a gradual increasing evolution in the hardness was recorded throughout these cross sections. The hardness evolution started at a faster rate in the peripheral areas of the deformed disks than in the central regions. A fairly saturated homogeneity in hardness was obtained for samples processed up to 10 turns across the horizontal and vertical cross sections, where the hardness value was about 240 Hv, twice that for the as-received sample. In general, the maximum values of achievable hardness after HPT processing on the horizontal cross sections of HPT disks were 180, 200, 206, 220 and 240 Hv in the samples processed for 1/2, 1, 3, 5 and 10 turns, respectively. A similar increase was obtained in the maximum achievable hardness across the vertical cross sections of HPT disks of 170, 190, 200, 210 and 230 Hv in the samples processed for 1/2, 1, 3, 5 and 10 turns, respectively. The trend of hardness evolution was correlated with the development in crystallite size obtained from XRD analysis and TEM of the grain size as shown in Fig. 12.
Vickers microhardness has shown a significant improvement with increasing number of HPT turns that is accompanied with a considerable decrease in the size of crystallites from 320 nm in the as-received sample to 30 nm in the sample deformed for 10 HPT turns. The same trend has been observed in TEM analysis where the minimum grain size appeared to be 60 nm. The hardness was also correlated with the dislocation density as illustrated in Fig. 13. The as-received sample gave a density of dislocation of $1.1 \times 10^{14}$ m$^{-2}$, whereas this value has increased to $6.2 \times 10^{14}$ m$^{-2}$ in the sample processed for 10 HPT turns. Empirical Hall–Petch and Taylor relationships [27, 28]: $H_v = H_0 + kd^{-1/2}$ and $H_v = H_0 + \frac{zMGb\rho^{1/2}}{C}$, respectively, were calculated for the current processed alloy. $H_v$ represents the Vickers microhardness, $H_0$ is the stress of friction in terms of hardness, $k$ is a constant, $d$ is the TEM grain size, $z$ is a constant $(0.3)$, $M$ is factor of Taylor (3), $G$ is the modulus of shear $(26 \text{ GPa})$, $b$ is the vector of Burgers $(0.286 \text{ nm})$ and $\rho$ is the density of dislocation that computed by XRD. The results calculated from both Hall–Petch and Taylor were normalized by the experimental recorded hardness in a similar manner to a recent study [21], then these results are plotted in Fig. 14 in order to show the distinct (separate) contributions of grain size and dislocation density on the resultant strengthening.

Discussion

Microstructure morphology and grain refinement

The microstructure of the additive manufactured Al-9%Si-3%Cu aluminium alloy was significantly refined into nanoscale grain sizes by HPT at room temperature as showed in the OM, SEM and TEM observations (Figs. 2, 3, 4, 6). High-pressure torsion has introduced significant alteration in the microstructure morphology, grain size and density of dislocations in the alloy after processing. These changes were indicated via the refinement of $\alpha$-Al matrix, fragmentation and uniform distribution of Si eutectic phase (Fig. 5), and accumulation of high density of dislocations within the refined microstructure (Fig. 13).

The HPT processing has changed the microstructure morphology from the large melt-pool morphology to ultrafine grains (Fig. 2). This is ascribed to the influence of torsional shear strain imposed by the HPT that has resulted in refinement of the grains inside the melt pools, rotation of the grains in the direction of torsional shear strain, disappearance of the melt pools and finally generation of ultrafine grains as the torsional deformation proceeded. This is consistent with many previous investigations [3, 15, 29]. It was shown that the two-phases materials processed by HPT to exhibit shear vortices phenomenon [30–33]. These local vortices structures appear at a high level of strain HPT deformation, where the plastic flow of the processed material is variable from one region to another on the processed HPT disk due to heterogenous distribution of HPT strain between the centre and edge regions. The visualization of strain heterogeneity is affected by the amount of phases in processed material, where a two-phases material with high percentage of both phases and different plasticity would show more visual vortices during HPT such as in 50% asutentic-50% ferrite duplex steel [30, 31], Cu-28%Ag alloy [32] bronze/niobium composite [33]. In the current processed alloy, the amount of the second phase (silicon) was about 9% within the alloy, thus less visualization of these vortices was observed compared to the aforementioned investigations especially at the preliminary stage of HPT deformation. These vortices were observed at the high number of turns (10 turns) at the central regions of the processed disk as shown in Fig. 2c. These vortices were assumed to be driven primarily by the local plastic instability that arises from the localized blocking of shear deformation at the central regions of HPT-processed samples. Agglomeration of fine precipitates and second-phase particles could cause such localized deformation blockage and heterogeneity distribution of the shear strain, which can be estimated through minor difference in the microhardness distribution at the central regions compared to the peripheral regions of HPT-processed samples [33–35]. Therefore, these vortices were seen clearly on the centre regions over horizontal cross sections (Fig. 2c) and along the whole vertical cross sections (Fig. 2f) in the processed sample for 10 turns.

Grain refinement has happened extensively in the additively manufactured alloy in the preliminary stages of the HPT deformation where the average value of grain size has reached 60 nm (Fig. 6c) and the average value of eutectic particle size decreased...
Figure 2 OM observations over the horizontal cross sections of the disks at central regions showing: (a and b) polished and etched as-received samples and c processed sample for 10 turns. The OM observations over the vertical cross sections of the disks showing: d as-received sample, e and f processed sample for 10 turns at the central and peripheral regions, respectively.
to 175 nm after 10 turns (Fig. 5). Formation of subgrains observed in the processed sample up to 10 turns (Fig. 6c) could be ascribed to the dislocation accumulation and tangling in these dark contrast areas [36]. The deformation-induced twinning on Si eutectic plane (220) [37, 38] has provided deformation continuity from the relatively ductile α-Al matrix to the brittle Si eutectic phase under such heavy HPT
imposed strains [39]. The existence of deformation-induced twinning was assumed to occur across the α-Al matrix/Si eutectic phase interfaces of different crystallographic orientation [40]. This interface plays a role in strengthening the microstructure, where the accumulation of moving dislocations via the Al/Si interface during HPT deformation contributes to additional strengthening in the processed alloy [41, 42].

Figure 4 SEM observations of the processed samples at various HPT turns over the horizontal cross sections of the processed disks.
A finer microstructure has been obtained in the additively manufactured Al-9%Si-3%Cu alloy compared to the as-cast Al-7%Si alloy processed using HPT where grain size of 300–700 nm is reported [43], or in as-cast Al-11%Si-1%Cu alloy processed using ECAP with a reported grain size of 200–400 nm [44], or as-cast Al-7%Si alloy processed using HPT with a reported grain size of 87 nm [45]. This can be attributed to the initial microstructure of the current alloy compared to these counterparts. The as-cast Al-7%Si alloy was prepared from casting of pure aluminum with master Al–Si alloy with relatively large initial grain size of 50 µm, and then a relatively lower strain was imposed via only 5 turns in HPT [43]. However, the current alloy has been processed using 10 turns at the same applied pressure for both alloys that indicates the direct impact of imposed strain on the extent of the grain refinement [18].

It is familiar that the imposed strain by HPT is larger compared to ECAP, therefore, a finer microstructure with larger dislocation density is expected compared to that using ECAP as seen in this investigation when compared to the as-cast Al-11%Si-1%Cu alloy [18, 46, 47]. The ratios of silicon and copper in the Al–Si–Cu alloy have influence on the level of grain refinement. These elemental additions are expected to decrease the stacking fault energy for the alloy and activate deformation-induced twinning and dislocation structures resulting in an increase in grain refinement [48, 49]. Therefore, the current Al-9%Si-3%Cu has achieved a finer microstructure than an Al-7%Si alloy despite the HPT applied pressure being relatively lower (6 GPa) in the current work compared to the counterpart work (8 GPa) that was used for the Al-7%Si alloy, which indicates the important role of alloying elements in grain refinement processes over simply considering the HPT applied pressure [38, 50]. The grain refinement observed by TEM is relatively consistent with the crystallite size data obtained by XRD measurements (Fig. 12), which confirms the effectiveness of HPT processing at room temperature in the additively manufactured Al-9%Si-3%Cu alloy.

The processed alloy has undergone considerable deformation that has been stored within the microstructure in the form of lattice distortions, which was detected as broadening and shifting in the XRD peaks in the processed samples for different numbers of HPT turns (Fig. 7b) [51, 52]. The HPT deformation has caused a shift to the lower diffraction angles that led to an increase in the lattice spacings and microstrain values as shown in Fig. 7d. This expansion in the distorted lattices can be attributed to the presence of tensile strains within these lattices due to potential interstitials of ultraﬁne precipitates of intermetallic phases and deformation-
induced twinning within the crystallographic planes, as well as the lattice curvature due to the heavy dislocation density [53–55]. However, the expansion in the distorted lattices has been found to reduce the stacking fault energy, thus leading to occurrence of deformation-induced twinning that contribute to strengthening in the processed alloy [56–58].

**Porosity elimination**

The as-received sample contained (1) gas-induced and (2) process-induced porosities. (1) has a spherical shape, as observed in the OM micrographs (Fig. 2a, b) and arises from gas molecule dissolution into the melt powders during the additive manufacturing process, and then their re-nucleation on cooling. (2) tends to have irregular shapes as seen in OM micrographs and occurs due to lack-of-fusion in some areas of the powders due to inadequate fusion in these spots and/or spattering of powders from these spots by the high vapor pressure of the melt [1]. A significant porosity reduction has been achieved in the HPT processed additive manufactured alloy as indicated by Archimedes and ImageJ analysis procedures (Figs. 8, 9).

This reduction in porosity has been observed in the processed samples across the horizontal and vertical cross sections. Several factors in HPT processing have controlled the porosity reduction: hydrostatic...
Figure 7  
(a) Patterns of XRD in the as-received and processed samples at various HPT turns,  
(b) Magnified part of (a) showing peak shifting,  
(c) the scattered data are the XRD profile for the alloy and the solid colorful line profile is the Rietveld fitted data, and  
(d) the variation in the microstrain and lattice parameter in the as-received and processed samples.
pressure and torsional shear strain. The hydrostatic pressure is due to the state of stress when the alloy comes into contact with anvils. This pressure contributes to prevent cracking and then confines the material under processing, thus enhancing the solidity of the processed microstructure [59, 60]. It has been shown that increasing HPT pressure has resulted in reduction in the interconnected porosity in pure copper up to 6 GPa [61].

The current alloy was processed at 6 GPa producing several microstructural features that also contributed to coalescence of pre-existing voids and pores during HPT torsional deformation. The morphology of both matrix and eutectic phases was altered significantly as the torsional shear strain increased, as observed in OM and SEM micrographs (Figs. 2, 3, 4). The torsional HPT strain has contributed to microstructural reorientation and refinement including α-Al and eutectic Si phase, which has resulted in filling the pores as the torsional strain increases with number of turns [3, 18, 62, 63]. The microstructural evolution leads to pore closure, the reduction in porosity fraction and pore diameter occur at a relatively faster rate at the peripheral areas than the central region of the deformed disks as a result of the HPT imposed strain being higher at the peripheries than the central regions of the disk [3, 18, 19]. It was found that HPT has an effective impact on vacancy annihilation and then porosity elimination by means of rearrangement of dislocation structures [60, 64].

The difference in porosity data between Archimedes and ImageJ analysis procedures (Fig. 8) can be attributed to the nature of each test regarding the porosity counting. In Archimedes method, the closed pores inside the alloy were not filled with the immersion liquid leading to inaccurate volume measurement. This exclusion of closed pores has resulted in the difference of porosity data between Archimedes and ImageJ analysis [65, 66]. However, a relatively higher consistency between the aforementioned procedures was obtained with the higher HPT deformation, where more pores closure is expected to occur under the conditions of severe HPT pressure and torsion [3]. Both procedures taken together have provided a reliable estimation of porosity and density of the additive manufactured alloy combining the impact of the total volume of sample considered and different cross sections under imaging [23, 67].

**Strength evolution**

The evolution in strength of the additive manufactured Al–Si–Cu alloy after HPT processing was expressed in terms of hardness development. The microstructural heterogeneity in the processed samples at a comparatively low number of HPT turns has caused a diversity in the microhardness distribution (Figs. 10, 11). The gradual increasing homogeneity in the strength was achieved using additional HPT straining at higher turns, because of a progressive development of microstructural homogeneity in the
HPT processed samples over the horizontal and vertical cross sections [18, 29, 45].

A higher HPT imposed strain has contributed to microstructural and strength homogeneity in the through-thickness direction of the HPT samples [18, 68]. The Al-9%Si-3%Cu additively manufactured alloy has showed a comparatively higher level of strengthening than the Al-7%Si, where all

Figure 10 Color-coded hardness maps for the as-received and processed samples over the horizontal cross sections.
Figure 11 Color-coded hardness maps for the as-received and processed samples over the vertical cross sections.
The aforementioned alloys were processed by HPT [45, 57]. This can be ascribed mainly to the level of grain refinement, dislocation density and existence of second-phase fine particles. A high fraction of high-angle grain boundaries will be generated with the progress of the torsional straining, which will restrict the dislocation motion during plastic deformation, resulting in strengthening of the microstructure [18, 28, 69].

The Al-9%Si-3%Cu additively manufactured alloy has exhibited smaller ultrafine-grain microstructure and higher strength rather than the Al-7%Si alloy, which reflects the dependence of strengthening effect on grain size reduction (Figs. 12, 13). The improvement in strength of the processed alloy was associated with the progress of equivalent imposed strain [18, 69]. The dependency of strength on grain size for the alloy was expressed using Hall–Petch equation and the dislocation strengthening as expressed by the calculated hardness from Taylor equation.

Figure 12 Variation in the crystallite size, grain size and hardness with the HPT turns.

Figure 13 Variation in the dislocation density and hardness with the HPT turns.

Figure 14 A comparison between the grain boundary strengthening as expressed by the calculated hardness from Hall–Petch equation and the dislocation strengthening as expressed by the calculated hardness from Taylor equation.

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the grain refinement and dislocation activity as calculated by the Hall–Petch and Taylor relationships, respectively. Hall–Petch relationship refers to the strengthening from grain boundaries that act as obstacles for dislocation motion [74–76]. Taylor relationship describes strengthening by dislocation–dislocation interactions during accumulation at the grain boundaries [28, 77]. By comparison of the calculated strengthening from the grain size reduction and dislocation activity as presented in Fig. 14, it is clear that the grain boundaries have a relatively larger impact on the strengthening rather than the dislocation forest. The remaining contribution to the strengthening is inferred to have come from the pinning and accumulation of moving dislocations at the nano-sized particles of the Si eutectic phase [28, 63, 77].

At low HPT strains, the dislocation–dislocation interactions are generally considered to be more effective in strengthening which is associated with lower misorientation between the grain boundaries. At higher HPT strains, the obstacle resistance of grain boundaries is considerable and increases with higher misorientation between the grain boundaries [28, 47, 78]. However, the current processed alloy showed a significant grain refinement at the preliminary stages of HPT deformation that caused a dominating effect by strengthening from the grain boundaries over the dislocation–dislocation interactions [20, 28]. The current data showed a clear dependency of the strength on both TEM estimated grain size and density of dislocation observed in the deformed alloy. However, the calculations of crystallite size were obtained with some potential errors compared to the TEM grain size data, that could be the reason for the minor difference between the aforementioned mechanisms [27, 28, 79].

A substantial dislocation density has been achieved in the processed samples on increasing the HPT torsional straining up to 10 turns. Dislocation density has accumulated effectively in the processed alloy under room temperature-HPT rather than elevated temperature-HPT [18, 19, 80, 81]. There are many factors that could affect the evolution of dislocations during HPT processing: applied HPT pressure, volume fraction of the nano-sized particles and stacking fault energy [18, 19].

An increase in the HPT pressure obstructs the migration of defects that suppresses the annihilation of these defects [64, 82]. It was found that fragmentation and distribution of second phase particles of nano-sizes within the processed microstructure have resulted in additional obstruction of the dislocations’ motion [41, 83]. In this research, considerable refinement and distribution in the Si eutectic phase during HPT were associated with relatively homogeneous distribution of these eutectic nano-sized particles over the whole of the processed samples. This behavior of the second phase provides an additional strengthening of the processed alloy when tested at room temperature and enhances the superplastic performance at higher temperatures [17, 84].

Stacking fault energy of pure aluminium is normally high and relatively lower for its alloys, thus addition of alloying elements has shown a reduction in this energy resulting in easier formation of partial dislocations and deformation-induced twinning [85, 86]. The current alloy has silicon, copper and magnesium as alloying elements which not only reduce the stacking fault energy of aluminum, but also provide additional sources for dislocations and thus strengthening [87, 88]. The high contents of Si and of Cu in the Al-9%Si-3%Cu compared with the Al-7%Si have another impact on strengthening in the current processed alloy. These elements normally form several hard intermetallic phases (such as Al2Cu and Al6Fe2Si) in addition to the eutectic Si [89, 90].

As a result, the volume fractions of these phases in the Al-9%Si-3%Cu alloy are higher than in Al-7%Si alloy. The alloying elements were added in relatively precise manner during the direct laser melting process, which is consider as a controllable-microstructure manufacturing process. Therefore, all these elemental structures were subjected to extensive refinement down to nanoscale during HPT and acted as barriers to dislocations motion. Accordingly, the strengthening in the Al-9%Si-3%Cu alloy has increased with the higher volume fractions of these hard nano-sized secondary phases after HPT [91–93].

The increment in volume fraction of the eutectic nano-sized particles with progress in torsional HPT deformation will result in a reduction in the mean free path of the accumulated dislocations. Therefore, more barriers for dislocation mobility will appear that would improve the strength [73, 94, 95]. A similar trend of increasing the dislocation density has been found with increasing the fraction of alloying element in the additively manufactured alloys Al-3.5%Si-1.5%Mg and Al-3.5%Si-2.5%Mg [81]. A comparison of the current HPT processed Al-Si–Cu alloy with a Al–Si–Cu/SiC alloy composite subjected to
HPT [96], revealed that the current processed additively manufactured alloy has a comparable strength without addition of SiC hard particles. This also indicates the impact of combination of severe plastic deformation with controllable-microstructure additively manufactured alloy.

Conclusions

1. Effective grain refinement (60 nm) and dislocation density \((6.2 \times 10^{14} \text{ m}^{-2})\) have been obtained by room temperature-HPT processing at higher HPT turns (10 turns) for the additively manufactured Al-9%Si-3%Cu alloy.

2. A considerable porosity reduction has been measured via Archimedes method (75%) and image analysis (96%) at higher HPT turns (10 turns) and linked to the hydrostatic pressure and torsional deformation.

3. Strength homogeneity was obtained over both horizontal and vertical cross sections at higher HPT turns (10 turns), which is attributed to the evolution in microstructural homogeneity with further HPT deformation.

4. The ultrafine-grained additively manufactured alloy showed a good consistency between the grain size and dislocation strengthening mechanisms. The content of alloying elements, nano-sized and distribution of second-phase particles have additional effects on the resultant strengthening.

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Author’s contribution

Ahmed S.J. Al-Zubaydi contributed to formal analysis, methodology, Writing—original draft, investigation. Philippa A.S. Reed & Nong Gao contributed to supervision, conceptualization, writing—review & editing.

Declaration

Conflict of interest  The authors state that they have no competing financial interests or known personal relationships that could appear to affect the work presented in this article.

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