Effect of sample orientation on the microstructure and microhardness of additively manufactured AlSi10Mg processed by high-pressure torsion

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Abstract

For the first time, high-pressure torsion (HPT) was applied to additively manufactured AlSi10Mg built in two directions (vertical and horizontal) by selective laser melting (SLM), and the influence of extreme torsional strain on the porosity, microstructure and microhardness of the alloy was investigated. ImageJ analysis indicates that significant porosity reduction is achieved by 1/4 HPT revolution (low strain). Optical microscopy (OM) and scanning electron microscopy (SEM) observations reveal the steady distortion and elongation of the melt pools, the continuous elongation of the cellular-dendritic Al matrix and breakage of the eutectic Si phase network with increased HPT revolutions. Microhardness measurements indicate that despite the significant increase in hardness attained from HPT processing, hardness saturation and microstructural homogeneity are not achieved even after 10 HPT revolutions. X-ray diffraction (XRD) line broadening analysis demonstrates increased dislocation densities with increased HPT revolutions, which contributes to the considerably higher hardness values compared to as-received samples.

Keywords Porosity · Microstructure · Microhardness · AlSi10Mg · Additive manufacturing · High-pressure torsion

1 Introduction

Powder bed fusion laser additive manufacturing (L-PBF AM) techniques, such as selective laser melting (SLM) and electron beam melting (EBM) have emerged as attractive methods of fabricating metallic components suitable for a wide range of applications, including automotive and aerospace. These techniques are highly attractive because they are able to manufacture engineering components with intricate structures and tailored microstructures [1]. Compared to traditional manufacturing routes, AM also offers the advantage of short lead times and low material wastage while maintaining accuracy and high resolution for the built structure [2–4]. SLM is one of the most common L-PBF AM technology that builds metallic parts by selectively laser scanning a powder bed in a layer-by-layer fashion (bottom-up approach) through computer-aided design (CAD) until a complete three-dimensional (3D) structure is formed.

To date, a wide range of materials have been processed by SLM, including stainless steels [5–7], Ni-based alloys [8–10], Ti-based alloys [11–13] and Al-based alloys [14–16]. Various reports have suggested improvements in mechanical properties for AM-fabricated parts compared to that of their traditional counterparts, including higher yield and tensile strengths [17, 18], better corrosion resistance [19–21] and enhanced fatigue life [22]. Such improvements are attributed to the unique and very fine microstructure as a result of rapid solidification due to the short laser-material interaction time and high cooling rates of AM processes (10³–10⁸ K s⁻¹) [23–25]. However, the high residual stress, porosity and other defects that often exist in as-manufactured parts cause some concern, which means that some kind of post-processing, e.g. hot isostatic pressing (HIP) and heat treatment before being ready for service [26–28].

On the other hand, severe plastic deformation (SPD) is a unique metal processing technique that introduces large amounts of strain on bulk metallic materials to achieve
extreme grain refinement down to sub-micron (0.1–1 μm) or even nanoscale (<100 nm), collectively termed ultrafine grained microstructure (UFG) [29, 30]. High-pressure torsion (HPT) is one of the most effective SPD techniques in attaining UFG microstructures with large fractions of high-angle grain boundaries via imposing significantly high torsional strains on disk-shaped materials. Various studies have shown improvement in properties of materials processed by SPD processes, in particular by HPT, compared to cast or wrought materials. Examples include higher yield and tensile strengths [31, 32], enhanced wear and corrosion performance [33, 34] and superplasticity at room temperature [35, 36]. The high strengths in SPD-processed metals are largely attributed to the Hall-Petch effect from the UFG microstructure and dislocation strengthening because of large amounts of dislocations introduced during SPD processing. Furthermore, reports have also emerged on attaining pore-free metallic materials via HPT, e.g. in pure Cu [37] and β-Ti alloys [38].

AlSi10Mg is a traditional hypoeutectic Al-Si-based cast alloy that is highly attractive for aerospace, marine and automotive applications due to its light weight, high specific strength, good corrosion resistance and low thermal expansion [39–42]. AlSi10Mg possesses good weldability due to its near Al-Si eutectic composition, in which the size and morphology of eutectic Si significantly influences the overall mechanical properties of this alloy [43, 44]. Coarse and acicular-shaped eutectic Si phase often cause cracks in cast AlSi10Mg (cooling rate < 10² K s⁻¹) when placed under tensile load, resulting in mechanical performance [45]. Therefore, rapid solidification (10³–10⁶ K s⁻¹) is desired to attain refined Si phase that are distributed homogeneously in the Al matrix to achieve better mechanical properties [46, 47]. This can be easily achieved by AM processes, and various studies have indeed shown enhancement in strength and ductility, and other properties of AM-fabricated AlSi10Mg compared to cast or wrought counterparts [48–51].

For example, two distinct microstructures can be observed in SLM-fabricated AlSi10Mg as a result of the rapid heating/cooling cycle: (a) cellular-dendritic α-Al structures and (b) fine, fibrous eutectic Si phase network around the α-Al phase [48, 49, 52]. Such unique microstructures contribute to the improved mechanical properties of SLM AlSi10Mg, e.g. strength and ductility under quasi-static loading [50, 51, 53]. Kempen et al. [48] established a process parameter window comprising of laser power, P, and scan speed, v to achieve optimum densification levels in SLM AlSi10Mg. Wei et al. [49] found that the porosity in SLM AlSi10Mg can be controlled (but not eliminated) by adjusting the energy density parameter, particularly laser power, P and laser scan spacing, h. Furthermore, Hadadzadeh et al. [54], Chen et al. [55] and Li et al. [56] all agree that the strengthening in SLM AlSi10Mg is caused by Orowan strengthening, Hall-Petch effect, dislocation hardening, or combination of the strengthening mechanisms.

On the other hand, while AlSi10Mg has not been processed by HPT yet, there are vast amount of literature on the HPT processing of other Al–Si-based alloys, e.g. Al-2Si, Al-7Si and Al-12 Si [57–60]. For example, Wang et al. [57] attributed the enhanced hardness and corrosion performance of HPT-processed Al-7Si to breakage of coarse Si particles and intermetallic phases, microstructure homogeneity and increased active sites due to HPT. Mungole et al. [58] found a correlation between microhardness and microstructural homogeneity in HPT-processed Al-7Si. Furthermore, El Aal et al. [59] attributed the improved wear properties of HPT-processed Al-7wt.%Si to increased microhardness due to grain refinement to the nano-scale and the uniform dispersion of Si particles.

In fact, it is only very recently that the combination of SLM and HPT for 316L stainless steel was studied by the current authors [20]. It was found that HPT processing is able to substantially reduce porosity even at low strain levels, achieving nano-scale grain sizes and resulting in enhanced microhardness and corrosion performance. Accordingly, the objective of this study is to investigate the influence of HPT processing on the porosity, microstructure and microhardness of SLM-manufactured AlSi10Mg built on both vertical and horizontal orientations through various number of HPT revolutions (strain levels) at room temperature. Microstructural evolution of the HPT-processed alloy is characterized by optical microscopy (OM), scanning electron microscopy (SEM), Vickers microhardness (HV) measurements and x-ray diffraction (XRD).

2 Materials and methods

Gas-atomised AlSi10Mg powder supplied by Concept Laser was used in this study with the chemical composition shown in Table 1.

A Concept Laser M2 SLM machine was used to additively manufacture the AlSi10Mg samples. The processing parameters chosen were as recommended by concept laser as listed in Table 2. The samples were built in a chamber with N₂ gas environment at room temperature, while an alternating bi-directional scan strategy was used in this study [61].

To investigate the influence of build orientation on the microstructure and microhardness of HPT-processed SLM AlSi10Mg, two 200 mm long cylindrical rods with a diameter of 10 mm each were manufactured in vertical and horizontal orientations, respectively, as shown in Fig. 1. In both cases,

| Chemical composition of AlSi10Mg used in this study (wt%) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Si | Mg | Cu | Mn | Fe | Ni | Zn | Pb | Sn | Ti | Al |
| 10.8 | 0.45 | 0.45 | 0.55 | 0.05 | 0.10 | 0.05 | 0.05 | 0.15 | Bal. |
the laser beam was scanned on the powder bed along the x-y plane and moved vertically upwards along the z-axis.

For HPT processing, the cylindrical rods were sliced into thin disks with thickness of 1 mm and diameter of ~9.8 mm each before being ground down with 800 grit SiC paper to a thickness of ~0.85 mm. Quasi-constrained HPT processing was conducted at room temperature using a HPT facility having two anvils, each containing a circular cavity with depth of 0.25 mm and diameter of 10 mm, respectively [62–64]. The disks were placed inside the cavity of the lower anvil before the anvil is pushed upwards into the cavity of the upper anvil via a compressive force. A small gap existing between the anvils enables some material outflow upon HPT torsional straining. The disks were then deformed under a pressure of 6 GPa by rotating the lower anvil at 1 rpm, from 1/4 to 10 revolutions. The equivalent von Mises strain, \( \varepsilon_{eq} \). Imposed by HPT straining can be calculated by using the following equation [30]:

\[
\varepsilon_{eq} = \frac{2\pi NR}{h\sqrt{3}}
\]

where \( N \) is the number of revolutions, \( R \) the distance from disk centre and \( h \) the initial disk thickness.

In this paper, disk samples sliced from the vertically built rod are denoted with \( N-V \), while disk samples sliced from the horizontally built rod are denoted as \( N-H \), where \( N \) corresponds to the number of HPT revolutions; e.g. 1/4-V (1/4 HPT revolution, vertical), 1-H (1 HPT revolution, horizontal). On the other hand, as-received disks are denoted as AR-V (vertical) and AR-H (horizontal) respectively.

The porosity content in the as-received and HPT-processed disks was analysed using ImageJ analysis software based on images taken from OM and SEM observations following the procedures described in Ref. [20]. The microstructures of the as-received and the HPT-processed disks were characterized via OM and SEM after being ground using different grits of SiC paper, polished using 3 and 1 \( \mu \)m Al\(_2\)O\(_3\) suspensions to a mirror-like surface finish and finally etched with Keller’s reagent. OM and SEM observations were conducted on the disk surface, i.e. the circular cross-section of the rods. Thus, for vertically built samples, the disk surface is parallel to the x-y plane, while the disk surface for horizontally built samples is parallel to the x-z plane.

Two different sets of microhardness measurements were recorded using Future Tech FM-300 Vickers hardness testing machine under an applied load of 500 gf with a dwell time of 15 s. The results are expressed in two ways: as plots of hardness against distance from the centre of the disk and as 2D colour-coded contour mapping plots throughout the disk surface. Firstly, a series of HV measurements were taken across the diameter of the disks with a distance of 0.3 mm between one indentation point to another. To improve the accuracy of the measurements, four further HV measurements were made with distances of 0.15 mm around each initial indentation point. The five readings were averaged and the error bars were recorded. Secondly, microhardness measurements were taken throughout the whole surface of the disks in a rectilinear grid mapping pattern determined by x- and y-coordinates, with coordinates (0, 0) defined as the centre of the disks. The indentation points are placed 0.3 mm between each other.

The phase composition and dislocation density, \( \rho \), of the as-received and HPT-processed disks were determined by x-ray diffraction (XRD) analysis using Rigaku SmartLab X-ray Diffractometer, with 10 steps per degree and a count time of 1 s per step on the instrument using a slit length of 5 mm, which is equipped with a graphite monochromator using CuK\(\alpha\) radiation. The phase composition was determined by the XRD peaks and peak broadening data, while the dislocation density was estimated from the microstrain, \( \varepsilon \) and crystallite size, \( D_c \) using materials analysis using diffraction (MAUD) software based on the Rietveld refinement method [65–67].
3 Results and discussion

3.1 Porosity evolution

Spherical and non-spherical pores were observed on the polished samples shown in Fig. 2a. The spherical pores are also known as gas-induced porosity, which could be caused by the entrapment of inert gas in the melt pool during the melting of powder or may already exist inside the initial raw powder and then remain in the finished structure [68, 69]. The irregular-shaped pores are known as process-induced porosity, which could arise from incomplete melting of the powder as the result of insufficient energy, or due to spatter ejection from the powder bed upon contact with the laser beam [25].

The SEM image in Fig. 2b shows an unmelted powder region causing lack-of-fusion porosity (a type of process-induced porosity) as the result of inadequate energy contact at that particular powder bed region to melt the powder. Etched OM image in Fig. 2c shows a mix of gas-induced pores (dashed yellow circle) and process-induced pores within the melt pool microstructure of the vertically built samples (AR-V), while etched OM image in Fig. 2d indicate that process-induced pores (solid red circle) are more apparent in the horizontally built samples (AR-H). The lack of gas-induced porosity in the AR-H samples may be the result of the continuous melting and re-melting of successive layers during the SLM process, which can cause the entrapped gas to eject away from the melt pool upon contact with the laser beam [70]. On the other hand, upon scanning a single layer of powder bed for the AR-V samples, it could be more difficult for the entrapped gas to move out of the melt pool since their movement are restricted by the compact powder distribution within the layer.

Porosity measurement was conducted on polished but unetched AR-V and AR-H samples using ImageJ software to determine the initial porosity level before HPT processing. The average porosity content for AR-V samples is 0.766 ± 0.023% and 0.695 ± 0.019% for AR-H samples. Despite the slight discrepancy between porosity level in AR-V and AR-H samples, they indicate high solidification levels (> 99%) were attained for samples built in both orientations. Figures 3 and b show the distribution of pore diameter in the as-received disks before HPT processing, which indicates a wide, but roughly similar distribution of pore diameter within the 0–90 μm range for both sample orientations. The slightly larger pore diameter distribution in AR-H samples may be caused by the larger process-induced pores in the horizontally built samples compared to those in the vertically built samples. The average pore diameter is 19.38 ± 15.64 μm for AR-V and 19.50 ± 17.7 μm for AR-H. Most of the pores lie between 5 and 10 μm and 5–15 μm for AR-V and AR-H, respectively.

After 1/4 revolution of HPT processing, the porosity content is reduced to 0.045 ± 0.011% and 0.037 ± 0.008% for 1/4-V and 1/4-H, respectively, yielding ~ 94% decrease compared to the as-received samples. Figures 3c and d show the distribution of pore diameter for the disks processed through 1/4 HPT revolution. The results indicate a much narrower distribution of pore diameter ranging from 0 to 40 μm and

**Fig. 2** a OM image showing types of porosity in the as-received AlSi10Mg. b SEM image showing unmelted powder region. c Dashed circle showing gas-induced porosity in AR-V sample. d Process-induced porosity at the melt pool boundaries (MPB) in AR-H sample.
significantly reduced pore diameters, with an average of 5.52 ± 4.81 µm for 1/4-V and 4.33 ± 3.62 µm for 1/4-H. Majority of the pores are in the sizes of 0–5 µm, and to a much lesser extent, between 5 and 15 µm. Only small amounts of pores in the sizes of 15–40 µm (< 10 counts) were observed for both 1/4-V and 1/4-H disks. Such dramatic reduction in porosity level was also observed by Yusuf et al. [20] in their study for SLM-fabricated and then HPT-processed 316L SS. These observations indicate that the HPT-imposed shear strain is able to effectively ‘close’ the pores, significantly reducing pore sizes for larger pores (40–90 µm) and possibly eliminate smaller pores (0–20 µm).

It is widely accepted that the generation and growth of pores in metals mainly depend on large levels of stress triaxiality under hydrostatic tension experienced, either under loading or during processing [71, 72]. On the other hand, the application of hydrostatic pressure has been found to be crucial in eliminating porosity. For example, Nakasaki et al. [73] observed that rolling a steel billet in multiple passes via a constant hydrostatic pressure was able to eliminate the centre pores that initially existed in the steel billet. Wang et al. [74] found that the rate of pore closure during hot rolling of steels are influenced by rolling process parameters and the pore location relative to the rolling contact surfaces.

In addition to hydrostatic pressure, the application of shear strain also promotes the collapse and closure of pores, enabling a sound bonding to be achieved throughout the pores after they come into close contact atomically as the result of both hydrostatic pressure and shear strain [75, 76]. Hydrostatic pressure and shear strain are two main features in severe plastic deformation (SPD) techniques to yield UFG microstructures and promote strong metallic bonding within the processed materials at both atomic and macroscopic scales [77, 78].

In addition, Qi et al. [37] systematically studied the generation and healing of porosity in HPT processing of pure Cu and explained the pore elimination mechanism in terms of grain refinement. The compressive pressure (from pushing the lower anvil to the upper anvil) and even small amount of torsional strain could cause the pores to collapse and stretch...
their internal surfaces together with the generation of UFG microstructures. Upon higher torsional strain, true sub-micron grains with clear grain boundaries start to form and the pores continuously elongate parallel to the shearing direction, fragmenting the pores and creating intimate atomic contact within the internal surface of the pores. Finally, at even higher torsional strain, internal friction within the internal pore surfaces creates strain gradients and high strain localisation, thus closing the pores. In this study, HPT has been proven as a very effective approach to reduce porosity via the strong metallic bonding that brings together and closes the pores as the result of the combination of hydrostatic pressure and torsional shear strain imposed.

3.2 Phase composition and dislocation density

XRD analysis Fig. 4 shows both Al and Si peaks are present in the AR-V and AR-H samples. A weak peak of Mg2Si intermetallic is also detected at ~ 40°, which might precipitate from the diffusion of Mg and Si as the result of repeated heating cycles experienced during SLM processing [79]. Such heating cycles resemble the heat treatment process during the initial stages of age hardening of Al–Mg–Si alloys [80]. Upon HPT processing, broadening and shifting of peaks could be observed in both vertically and horizontally built samples (except for (1 1 1) orientation), similar to other studies on HPT-processed metals [81, 82]. These are mainly caused by lattice defects, e.g. internal microstrain, dislocations, small crystallite sizes and additional grain boundaries introduced via the HPT-imposed torsional strains [83] (Fig. 4).

The dislocation density $\rho$ can be calculated from the lattice microstrain based on the following equation [84, 85]:

$$\rho_{XRD} = \frac{2\sqrt{3}(\varepsilon^2)^{1/2}}{Dc b}$$

where $D_c$ is the average crystallite size, referred to as coherently scattered domains (CSD), and $b$ is the burgers vector ($b = 0.286$ nm for Al [55]). The values of $<\varepsilon^2>$ and $D_c$ are obtained through the Rietveld refinement method applied in MAUD software, and the corresponding values of $\rho$ calculated based on Eq. 2 are shown in Table 3. It is revealed that the values of $\rho$ for as-received samples built in both orientations are already in the order of $1 \times 10^{14}$ m$^{-2}$, much higher than $10^8$ or $10^{10}$ m$^{-2}$ in conventional cast or wrought metals [86]. Such high $\rho$ values for as-built SLM samples are consistent with other studies on SLM AlSi10Mg [54] and pure iron [87], which are attributed to the numerous fine cellular structure colonies that store a large amount of dislocations, providing more sites to inhibit dislocation motions, and leading to relatively higher hardness compared to cast or wrought metals [86]. Upon HPT processing through 1 and 10 revolutions, the dislocation densities further increase to the region of $\sim 3.5-5 \times 10^{14}$ m$^{-2}$. On the other hand, the crystallite sizes dramatically decrease from $\sim 280$ to $\sim 55-57$ nm after 1 HPT revolution, before only slightly decreasing to $\sim 51-52$ nm after 10 HPT revolutions. The decrease in crystallite size is common in SPD-processed materials because the X-ray coherency is broken due to the small misorientations in the actual grains as the result of large shear strains applied during the process [83]. Therefore, the XRD analysis actually measures the crystallite sizes, rather than actual grain sizes.

For HPT-processed samples, the total dislocation density was calculated as the sum of geometrically necessary dislocations (GNDs) and statistically stored dislocations (SSDs). This is due to the presence of a strain gradient throughout the disk, with higher strains imposed at the edge compared to the centre region of the HPT-processed disk as the result of radial dependence, in accordance with Eq. 1 [81, 88].

3.3 Microhardness measurements for as-received and HPT-processed samples

Figures 5a and b show that the as-received samples exhibit reasonable homogeneity in the distribution of HV values across the diameter of the disks in both build orientations. It is worth noting that the HV values for SLM AlSi10Mg from this study (130–150 HV) are higher than that of their traditionally manufactured counterparts, e.g. 95–105 HV in as-high pressure die cast (HPDC) parts [89] and 64–70 HV in as-cast parts [90]. Higher hardness in SLM is also observed in various literatures and is commonly attributed to the finer microstructures attained as the result of significantly higher cooling rates/rapid solidification achieved in SLM due to the short laser-material contact time, thereby preventing recrystallisation or coarsening of the final microstructure [1, 41, 44, 91].

However, the HV values for AR-V (average: ~145 HV) is slightly higher than that of AR-H (average: ~136 HV), indicating anisotropy in SLM AlSi10Mg between both build orientations. Similar anisotropies are also observed in other AM materials, in which their properties differ with respect to build direction [2, 68, 91, 92]. Several studies have identified possible causes for anisotropy in AM materials, including level of defects, residual stress, local heat transfer conditions, scan strategy, grain orientation, and to a lesser extent, crystallographic texture [22, 42]. For example, Yadroitsev et al. [93] attributed the lower elastic modulus in IN 625 specimens built vertically (parallel to z-axis) to the higher defect content due to larger concentration of residual stresses compared to the horizontal ones (parallel to x-y plane). Frazier et al. [2] explained that lack-of-fusion pores is the reason for lower ductility of Ti alloys in the z-direction. Carroll et al. [92] observed lower ductility in Ti alloys built horizontally as a result of the orientation of prior-/β grains parallel to the build (z-direction). Chlebus et al. [94] ascribed the anisotropy in fracture modes...
and tensile properties of SLM IN 718 to the different orientations of the columnar grains with respect to loading directions. Accordingly, the lower hardness for AR-H samples than AR-V could be attributed to the larger content of lack-of-fusion pores (process-induced porosity) at the melt pool boundaries (MPB) of AR-H compared to AR-V (Fig. 2c, d). This may lead to grain structure heterogeneity and structural discontinuities, which could weaken the bonding of the surrounding microstructure, thereby affecting the macroscopic mechanical behaviour [42].

In comparison to as-received samples, it is clear that the HV values in HPT-processed samples are significantly higher, increase with distance from the centre to the edge of the disk and with increased HPT revolutions. A relatively strong hardening was obtained after 1/4 revolution, by a factor of ~2 for both sample orientations. It is also apparent that the majority hardness increment was achieved from 1/4 revolution, a common trend observed in many HPT-processed metals [95]. However, the increase in hardness is not homogeneous, with the central region exhibiting low HV values compared to the peripheral regions. This is expected because of the radial dependency of HPT-induced torsional strain in accordance with Eq. 1. Additional hardening was achieved with increasing number of revolutions and the highest HV value attained after 10 revolutions at the peripheral region was 218 HV and 210 HV for samples built vertically and horizontally, respectively. However, the hardness distribution remains inhomogeneous across the disks even after 10 revolutions for samples built in both orientations, indicating that hardness saturation has not been achieved, even at extreme torsional strain imposed at the highest number of revolutions.

Nevertheless, the HV values are slightly higher in vertically built samples than that of horizontally built samples for all conditions, indicating that some level of anisotropy might still exist even after 10 revolutions of HPT processing. However, it is possible that this discrepancy is due to the grain orientation rather than porosity, since it is already established that the pores that were initially present in the as-received samples were significantly reduced after HPT.

The variation of HV values with equivalent strain $\varepsilon_{eq}$ calculated from Eq. 1 is shown in Fig. 6a, b. It is apparent that the hardness varies monotonically with the imposed strain. However, microhardness saturation is still not achieved for samples built in both directions even after 10 HPT revolutions. Other studies have also confirmed the difficulty of achieving complete homogeneity in some high-strength materials even after 20 revolutions [96–98]. They attributed this heterogeneity to the difficulty to attain true sub-micron or nano-scale grains at the central region or near the centre of the disks, even via extreme torsional strains.

Colour-coded contour maps are generated based on the microhardness measurements from the rectilinear grid mapping pattern to provide visualization of the distribution of individual HV values for the as-received and HPT-processed disks. Figure 7a shows the HV mapping for as-received samples, and Fig. 7b–e for HPT-processed samples through 1 and 10 revolutions. The colour-coded legend on the right of Fig. 7a defines the range of HV values represented by the different colours in the contour maps (from 140 to 220 HV).

A significant evolution of HV values could be observed after 1 HPT revolution for both 1-V (Fig. 7b) and 1-H (Fig. 7c) disks compared to that of the as-received samples (Fig. 7a). This is indicated by the transition from the blue region of low HV values

| Table 3 Dislocation density parameters obtained through MAUD analysis of the XRD data |
|---------------------------------|---------------------------------|
| Sample | Crystallite size, $D_c$ (nm) | Dislocation density, $\rho$ ($\times 10^{14}$ m$^{-2}$) |
|-------|-------------------------------|----------------------------------|
| AR-V  | 288 ± 23                      | 1.23                             |
| AR-H  | 285 ± 20                      | 1.18                             |
| 1-V   | 57 ± 4                        | 3.82                             |
| 1-H   | 55 ± 6                        | 3.75                             |
| 10-V  | 52 ± 2                        | 5.11                             |
| 10-H  | 51 ± 3                        | 4.92                             |

Fig. 4 XRD patterns for as-received and HPT-processed disks for samples built in the a vertical and b horizontal directions.
to green–orange region of higher HV values. However, within the disks processed through 1 HPT revolution, Fig. 7b, c demonstrates that the sample built in vertical orientation shows higher HV values (~190–205 HV) compared to that built in horizontal orientation (~175–195 HV). These are indicated by the cooler colours in 1-H compared to 1-V, especially at the peripheral regions, with larger area in the centre of the disk extending over a width of ~4 mm exhibiting lower HV values. The HV values further increase after 10 revolutions, as indicated by the transition from green–orange to green–red region in Fig. 7d, e. It is apparent that 10-V disks possess higher HV values (~190–220 HV) than 10-H disks (~190–210 HV), as illustrated by the hotter colours at the peripheral regions and smaller area of cooler colours at the central region. Nevertheless, these contour maps further demonstrate that hardness saturation and microstructural homogeneity could still not be fully achieved, particularly at the central region even though the area with low HV values shrink with increasing torsional strain values.

3.4 Microstructure observations for as-received samples

Figures 8a and b show low-magnification OM images of the as-received samples built vertically (AR-V) and horizontally (AR-H), respectively, revealing distinct melt pool morphologies for samples built in different orientations. The AR-V sample (Fig. 8a) exhibits alternating bi-directional melt pool patterns corresponding to the scan strategy used in this study, while the AR-H sample (Fig. 8b) illustrates a fish scale (semi-circle) morphology corresponding to the layer-by-layer laser interaction with the powder bed. Good inter- and intra-layer overlapping is obtained for both samples, indicating a good bonding within a single layer and between successive layers [99].
Figures 8c and d exhibit SEM images of AR-V and AR-H, respectively, while Fig. 8e shows a higher magnification SEM image of Fig. 8c, displaying much clearer SLM AlSi10Mg microstructures. They consist of very fine cellular-dendritic $\alpha$-Al matrix (size from $\sim$0.5 to 1 $\mu$m) surrounded by thin fibrous eutectic Si network (thickness $<$ 0.2 $\mu$m). Such fine microstructures are attributed to the rapid solidification of SLM process through repeated fast melting/cooling cycles, kinetically favouring the cellular solidification morphology and the extended solubility of Si into Al [22, 100]. Sub-micron Si particles that did not fuse together with the eutectic Si network could also be observed in the as-received samples.

The MPBs (dashed red lines in Fig. 8c, d) and the areas close to the MPBs have relatively coarser and more elongated microstructures, compared to the central region of the melt pools. This could be ascribed to the relatively different solidification rates throughout the melt pool, even though the solidification rates in SLM process is generally high ($10^3$–$10^8$ Ks$^{-1}$). In particular, the region around the MPBs and/or at the overlapping region between MPBs experiences longer laser-material interaction time due to the melting and re-melting of nearby powder bed in a single layer (AR-V) and of successive powder layers (AR-H). Hence, these regions could maintain at higher temperatures for longer periods, causing slower cooling rates and thus allowing the microstructures to coarsen. This phenomenon is common in SLM AlSi10Mg, which has been observed in other AM alloys [22, 41, 44].
In addition, partially broken and coarsened eutectic Si network could also be observed at the MPBs (dashed red lines in Fig. 8c, d) due to the lower solidification rate at that region. Mg$_2$Si precipitates have not been found from the SEM observations in this study, despite the presence of weak XRD peaks for this intermetallic.

### 3.5 Microstructural evolution after HPT

Figure 9 displays representative OM images of HPT-processed disks through 1 and 10 revolutions for samples built in both orientations at the centre and edge (3 mm from centre) of the disks. These images show the evolution of melt pools as a result of HPT processing. The melt pools are observed to be slightly distorted and oriented at the centre after 1 revolution for 1-V (Fig. 9a) and 1-H (Fig. 9c) disks, compared to the shape and morphology of melt pools observed in AR-V (Fig. 8a) and AR-H (Fig. 8b). However, the melt pools at the edge of the disks become significantly sheared and extremely elongated for 1-V (Fig. 9b) and 1-H (Fig. 9d) disks, resulting in the loss of shape and morphology of the as-received melt pools.

These observations provide clear indications of the inhomogeneous strain distribution across the diameter of the disks due to the radial dependency of shear strain imposed during HPT in accordance with Eq. 2. Based on this equation, the edges experience higher amounts of torsional strain compared to the centre of the disks, and this is confirmed via the heavily deformed melt pools at the edges after one turn corresponding to $\varepsilon_{eq}$ of $\sim$ 13.6 (1-V, Fig. 9b and 1-H, Fig. 9d) compared to
that at the centre (1-V, Fig. 9a, 1-H Fig. 9c). This is also in agreement with the higher HV values at the peripheral regions compared to the central region (Fig. 5a, b), thus affirming the application of microhardness measurements as an effective method to evaluate the extent of microstructural evolution in HPT-processed materials [101–103].

After 10 revolutions ($\varepsilon_{eq} \approx 136$), the increased torsional strain at this stage further deformed the melt pools at the centre, such that they are extremely oriented, and the initial shape and morphology are no longer apparent although not completely eliminated as shown in Fig. 9e for 10-V and Fig. 9g for 10-H. In addition, the shape and morphology of the
melt pools at the edges are completely lost for samples built in both directions (10-V, Fig. 10f and 10-H, Fig. 10h).

Figure 10 illustrates representative SEM images of HPT-processed disks through 1 and 10 revolutions for samples built in both orientations at the centre and edge (3 mm from centre) of the disks. After HPT processing through 1 revolution, at the centre of the disks (1-V, Fig. 10a and 1-H, Fig. 10c), the cellular-dendritic $\alpha$-Al matrix was compressed, while the eutectic Si network was refined and elongated. Some of the eutectic Si networks were also broken into fine Si particles. On the other hand, most of the eutectic Si networks are broken into fine Si particles that are dispersed more uniformly in the Al matrix at the edge of the disks (1-V, Fig. 10b and 1-H, Fig. 10d). After 10 revolutions, Fig. 10e–h exhibits the complete...
breakage of the eutectic Si network into fine Si particles. At this stage, the size of the fine Si particles significantly decreases, while the volume fraction obviously increases and more homogeneously distributed throughout the Al matrix compared to that after 1 revolution. This is because of the continuous breakage of the eutectic Si network after 10 HPT revolutions, resulting in the formation of new fine Si particles.

The observation of eutectic Si network breakage into fine Si particles upon increased HPT-imposed torsional strain is consistent with other studies on the HPT processing of traditional cast Al-7Si alloy [58, 104], which can be attributed to two factors [104, 105]: (i) increase in temperature due to friction generated during HPT straining may lead to nucleation and growth of fine Si particles or (ii) the severe plastic deformation that could cause fragmentation and redistribution of larger Si particles into smaller segments in the Al matrix.

To evaluate the extent of the microstructural evolution due to HPT, ImageJ software was used to characterize the cellular-dendritic network observed via SEM. Values for average diameter of the cells in the as-received and HPT-processed disks are listed in Table 4 for samples built in both directions. It could be seen that the average cell diameter for as-received samples is ~0.75 μm, which then decreases to ~0.50 μm (centre) and ~0.37 μm (edge) after 1 HPT revolution for samples built in both orientations.

On the other hand, Fig. 11 shows the evolution of the circularity of the cellular-dendritic network in as-received and HPT-processed disks for samples built in the horizontal orientation. Values closer to 1 indicate increasingly circular shape, while values closer to 0 represents increasingly elongated morphology. The circularity of the cells for the as-received sample ranges from 0.5 to 0.95 (average: 0.756 ± 0.07) and

| Sample | Centre (μm) | Edge (μm) |
|--------|-------------|-----------|
| AR-V   | 0.75 ± 0.15 | –         |
| AR-H   | 0.74 ± 0.21 | –         |
| 1-V    | 0.50 ± 0.05 | 0.38 ± 0.11 |
| 1-H    | 0.51 ± 0.08 | 0.36 ± 0.13 |

Fig. 11 | Circularity of the cellular-dendritic structure for as-received and HPT-processed disks in horizontally built samples
after 1 HPT revolution, decreases to 0.25–0.75 (average 0.473 ± 0.10) at the centre and to 0.15–0.60 (average 0.351 ± 0.08) at the edge of the disks. The lower average circularity at the edge compared to at the centre of the disks after 1 HPT revolution (by ~ 25%) further confirms the radial influence of the torsional strain imposed via HPT. The diameter and circularity of the cellular-dendritic structure could not be measured after 10 HPT revolutions because most of the eutectic Si networks are broken down into finer Si particles. Nevertheless, the reduced average diameter and circularity of the cells with increased HPT revolutions are attributed to the large shear strains due to the HPT-induced torsional strain, which have also been observed by Yusuf et al. in their study of 316L SS fabricated by SLM and then processed by HPT [20].

4 Conclusion

The evolution of microstructure and microhardness of AlSi10Mg fabricated by SLM in both vertical and horizontal orientations and then processed by HPT through 1/4 to 10 revolutions were investigated through OM, SEM, microhardness measurements, and XRD. The main conclusions can be drawn as follows:

1. Although both spherical (gas-induced) and non-spherical (process-induced) pores are present in vertically built samples, process-induced porosity is more apparent in horizontally built samples.
2. The microstructure of as-received SLM AlSi10Mg consists of cellular-dendritic Al matrix surrounded by fine, fibrous eutectic Si phase network for samples built in both build orientation.
3. The lower HV values in horizontally built samples are attributed to the larger fraction of process-induced pores at the fish-scale MPB, compared to vertically built samples.
4. HPT processing for 1/4 revolution significantly reduced the porosity by ~ 94% compared to the as-received samples.
5. HPT processing significantly increases the hardness of the alloy, but microstructural homogeneity was not achieved even after 10 HPT revolutions.
6. As the number of HPT revolutions increase, the cellular-dendritic Al matrix becomes increasingly refined and elongated, while the eutectic Si network is continuously broken into fine Si particles and dispersed uniformly throughout the Al matrix.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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