Metal Additive Manufacturing



Advanced structural analysis of a laser additive manufactured Zr-based bulk metallic glass along the build height

James P. Best^{1,2}, Keita Nomoto^{1,3}, Fan Yang⁴, Bosong Li¹, Moritz Stolpe⁵, Luyang Zeng³, Zach Evenson^{6,7,8}, Christoph Hugenschmidt⁷, Xiaopeng Li¹, Simon P. Ringer³, and Jamie J. Kruzic^{1,*}

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ABSTRACT

Additive manufacturing of bulk metallic glasses (BMGs) has opened this material class to an exciting new range of potential applications, as bulk-scale, net-shaped amorphous components can be fabricated in a single step. However, there exists a critical need to understand the structural details of additive manufactured BMGs and how the glassy structure is linked to the mechanical properties. Here, we present a study of structure and property variations along the build height for a laser powder bed fusion (LPBF) processed Zr-based BMG with composition $Zr_{59.3}Cu_{28.8}Nb_{1.5}Al_{10.4}$ commercially termed AMZ4, using hardness testing, calorimetry, positron annihilation spectroscopy, synchrotron X-ray diffraction, and transmission electron microscopy. A lower hardness, more rejuvenated glassy structure was found at the bottom of the build compared to the middle region of the build, with the structure and properties of the top region between the two. Such differences could not be attributed to variability in chemical composition or crystallisation; rather, the softer bottom region was found to have a larger medium range order cluster size, attributed to

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James P. Best and Keita Nomoto contributed equally to this work

Address correspondence to E-mail: j.kruzic@unsw.edu.au



¹ School of Mechanical and Manufacturing Engineering, University of New South Wales (UNSW Sydney), Sydney, NSW 2052, Australia

² Max-Planck-Institut für Eisenforschung GmbH, Max-Planck-Str. 1, 40237 Düsseldorf, Germany

³ Australian Centre for Microscopy and Microanalysis, and School of Aerospace, Mechanical and Mechatronic Engineering, The University of Sydney, Sydney, NSW 2006, Australia

⁴Institut für Materialphysik Im Weltraum, Deutsches Zentrum für Luft- und Raumfahrt (DLR), 51170 Cologne, Germany

⁵Heraeus Additive Manufacturing GmbH, Heraeusstr. 12-14, 63405 Hanau, Germany

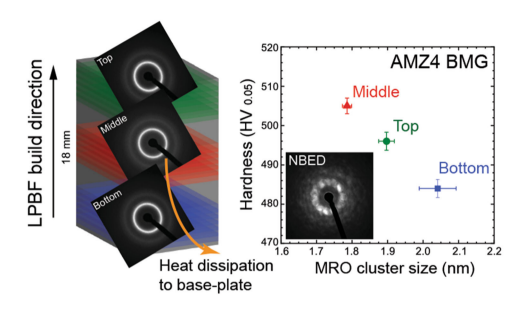
⁶ Physik-Department, Technische Universität München, James-Franck-Str. 1, 85748 Garching, Germany

⁷ Heinz Maier-Leibniz Zentrum (MLZ), Technische Universität München, Lichtenbergstr. 1, 85748 Garching, Germany

⁸ Springer-Verlag GmbH, Tiergartenstr. 17, 69121 Heidelberg, Germany

heat dissipation into the build plate during processing, which gave faster cooling rates and less reheating compared to the steady-state middle of the build. However, at the top of the build less reheating occurs compared to the middle, leading to a somewhat softer and less relaxed state.

GRAPHICAL ABSTRACT



Introduction

Bulk metallic glasses (BMGs) are alloys with no longrange atomic order but with varying degrees of shortand medium-range order (SRO and MRO). This means that there are no classical lattice defect structures (e.g. grain boundaries or dislocations) giving rise to a fundamental change in the plastic deformation mechanisms, with shear transformations and shear bands dominating rather than dislocation slip and twinning [1]. As such, BMG materials have exceptionally high strengths, elastic strain limits, excellent wear resistance, etc. [2, 3]. A significant bottleneck for the widespread utilisation of BMGs, however, is the rapid cooling rates required during processing to 'freeze-in' the amorphous structure [4]. This requirement has traditionally led to difficulties in producing components with large enough dimensions to be industrially relevant for a range of targeted applications (e.g. for aerospace, transportation, biomedical implants, etc.).

To address this issue, laser powder bed fusion (LPBF) has been recently shown to be a promising method for producing large-scale BMG parts with a broad-range of possible geometries [5–10]. However, there is a growing body of evidence that LPBF-processed parts generally have reduced ductility and damage tolerance when compared with cast alloys of the same chemical composition [10–16]. Since those properties are often seen as limiting factors in the application of BMGs [2, 17], such work highlights the importance of understanding the chemical and structural aspects that control the mechanical properties, so that future improvements can be made to LPBF-manufactured BMGs.

The reduced deformation and damage tolerance properties have been attributed in part to various factors such as the presence of crystals, pores, and/or elevated oxygen levels in the bulk amorphous phase [10–16]. However, even for X-ray amorphous samples



studied with synchrotron micro-diffraction, there is evidence demonstrating microscale heterogeneity in the glassy structure of LPBF-processed BMGs related to the melt-pool solidification process [18]. With regard to improving the fracture toughness of BMGs, the creation of heterogeneous glassy microstructures has proven effective [19–22], and indeed, there is evidence that LPBF-produced BMGs are tougher when cracks are forced to propagate across layer-by-layer heterogeneities [11]. Thus, there is a motivation to better understand the glassy structure variations that exist along the build height of LPBF-produced BMG samples, to enable optimised design and rationalisation of mechanical property variations.

Characterisation studies of LPBF-produced BMGs have mainly focused on layer-level structural variations, such as crystallisation and/or chemical variations in the melt-pool and heat-affected zones (HAZs) or structural periodicity that occurs in the amorphous phase across the build layers [13, 18, 23-25]. In contrast, characterisation studies of traditional crystalline alloys produced by LPBF have often revealed variations in the microstructure at different locations along the build height due to the different heat input and thermal histories that evolve as the build progresses [26-28]. However, little is known about variations that possibly occur along the build height of printed BMG pieces, with one report noting similar porosity levels at the top, middle, and bottom sections of a printed sample [7].

Accordingly, in this work the atomic-scale chemical and structural features were studied at distinct locations (top, middle, and bottom) along the 18 mm build height of a LPBF-processed BMG alloy (AMZ4—Zr_{59,3}Cu_{28.8}Nb_{1.5}Al_{10.4}) using microhardness mapping, differential scanning calorimetry (DSC), Doppler-broadening positron annihilation spectroscopy (DB-PAS), synchrotron X-ray diffraction (sXRD), and transmission electron microscopy (TEM) with nanobeam electron diffraction (NEBD). The combination of these analyses demonstrated that no noticeable structural changes in porosity, chemical composition, and nearest neighbour atomic spacing in the glassy structure were observed along the build height on thin samples cut from the bulk free of residual stresses. However, the bottom of the build had a lower average hardness and wider hardness variability compared to the other locations that was associated structurally with a greater relaxation enthalpy and a larger nanoscale MRO cluster size. The results of this study improve our understanding of the processing-structure–property relationships for LPBF additive manufactured BMGs.

Material and methods

Sample fabrication

A LPBF-processed sample was fabricated using commercially available amorphous AMZ4 powder (10–45 µm particle diameter). The LPBF sample was produced by Heraeus Additive Manufacturing GmbH (Hanau, Germany) as described in our previous studies [11, 12] using an Electro Optical Systems (EOS) M290 (with a 400 W Yb-fibre laser) with a classic EOS rotating stripe pattern (67° scan rotation between each layer) and their proprietary optimised LPBF parameters for a layer thickness of 20 µm. A specimen with dimensions of $10 \times 10 \times 18 \text{ mm}^3$ was fabricated, with the longest dimension corresponding to the build direction. The specimen was welded to a \sim 25 mm thick commercially pure Ti base plate using a multiple-pass laser exposure for the first five layers, and no support structures were used. This base plate was screwed into the build-plate holder of the machine. Specimens were cut sufficiently far enough from the plate, at a height of ~ 0.5 mm, so that the initially welded material should have had no apparent influence on the composition of the final tested material. Subsequent mechanical testing and characterisation was conducted at the top, middle, and bottom of the build where the top and bottom refer to within 4 mm of the top surface or base plate, respectively, and middle refers to a central 4 mm along the build direction (Fig. 1).

Mechanical characterisation

Microhardness mapping was conducted on rectangular cross sections at the top, middle, and bottom of the build that were cut perpendicular to the build direction using a low-speed diamond saw (*i.e.* indentation surface corresponds to the x–y plane in Fig. 1). A Vickers microhardness indenter was used in an automated hardness tester (Durascan-80, Struers) to create hardness maps with 625 indents each. An applied load of 0.05 kg and a dwell time of 10 s were used with an indent spacing of 40 μ m as described in our previous studies [20, 22, 29].



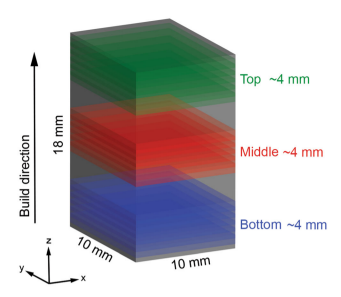


Figure 1 Schematic of the LPBF-processed AMZ4 specimen and the locations of the top, middle, and bottom of the build characterised in this study. With the exception of DB-PAS and SEM/EDS measurements, all specimens were cut perpendicular to the build direction.

Differential scanning calorimetry (DSC)

Samples for the DSC experiments (DSC 8000, PerkinElmer) with mass of $\sim\!20$ mg were cut from sections of the built AMZ4 block using a low-speed diamond saw and ground to a 4000 grit finish using SiC paper to ensure good contact with the bottom of the Al crucible pan. Samples were cleaned in an ultrasonic cleaner and dried prior to measurement. For each DSC experiment, first the instrument baseline was established as stable and reproducible. Then, the BMG sample was stabilised for 1 min at 323 K and then heated from 323 to 863 K at 20 K/min. After cooling, the crystalline baseline was obtained for each sample under the identical heating profile and subtracted from the BMG sample scan.

Doppler-broadening positron annihilation spectroscopy (DB-PAS)

The DB-PAS experiment was performed at the CDB-spectrometer [30] at the high intensity positron source NEPOMUC [31] located at the Heinz Maier-Leibnitz Zentrum (MLZ) in Garching, Germany. The characteristic S-parameter was obtained using coincident Doppler broadening spectroscopy (CDBS), which is defined as the ratio of counts in a fixed area around the maximum and the total counts of the

511 keV annihilation line. S-parameters were measured on polished thin ($\sim 200~\mu m$) cross sections of the samples with a bias voltage set to -30~kV and a positron beam of approximately 100 μm in diameter. Line scans were performed along the build direction, with a step size of 150 μm .

High-energy synchrotron X-ray diffraction (sXRD)

Measurements were performed in the second experimental hutch of the PETRA III P07 beamline at the Deutsches Elektronen-Synchrotron (DESY) in Hamburg, Germany. For measurement, rectangular samples with maximal dimensions of approximately 2 mm (thickness $\sim\!200~\mu\text{m})$ were cut using a low-speed saw from the bottom, middle, and top sections of the built block perpendicular to the build direction. Samples were then ground with 4000 grit SiC paper to remove surface asperities.

Spatially resolved X-ray diffraction line-scans were performed using a micro-focussed monochromatic photon beam with an energy of 98.15 keV and a size of $30 \times 2 \ \mu\text{m}^2 \ (w \times h)$ attained using an Al compound refractive lens system in transmission mode. The resultant X-ray scattering patterns were collected using an exposure time of 4 s on a PILATUS3 X CdTe 2 M flat panel detector placed about 400 mm downstream of the sample. The diffraction patterns showed a fully amorphous structure absent of any intermetallic or oxide crystalline peaks. A 2 µm stepsize along the x-axis was used perpendicular to the sample build direction (x = 0.2 mm), and a 30 μ m step-size was used along the z-axis (z = 0.09 mm), totalling 100×3 diffraction images per sample. The collected 2D diffraction patterns were integrated over the full azimuthal range using the Python pyFAI package [32], with a calibration using CeO₂ standard. Measured intensity data $I_m(Q)$ as a function of scattering vector Q were then processed using PDFgetX3 [33] to allow for the determination of the structure function S(Q). As described in Reference [18], the measurements were corrected for sample absorption, X-ray polarisation, Compton scattering, and any background scattering from the sample container or environment. The peak position of S(Q) was determined by an interpolation using a third-order polynomial in a limited Q range close to the maximum.



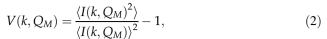
Electron microscopy

Scanning electron microscopy (SEM) with energydispersive spectroscopy (EDS) was done to further confirm the chemical homogeneity in the melt pools and heat-affected zones (HAZs) found in our previous study [18]. The X-ray energy spectra were collected at 20 kV and 0.8 nA with a dwell time of 100 µs using X-Max 80 mm² EDS detector (Oxford Instrument, UK). Transmission electron microscopy (TEM) specimens from the bottom, middle, and top sections of the built specimen were prepared by two different methods: (1) twin-jet electropolishing (TenuPol-5, Struers, USA) with 10% nitric acid in methanol at −20 °C and 30 V and (2) a standard lift-out method using a Xenon plasma focused ion beam (pFIB) scanning electron microscope (PFIB-SEM, Helios G4 PFIB, Thermo Fisher Scientific, USA). The pFIB samples were specifically used for site-specific experiments where TEM lamellae were extracted between four indents of nearly identical microhardness as determined by the microhardness mapping procedures described above. TEM analyses were performed using a JEM-2200FS (JEOL, Japan) at an acceleration voltage of 200 kV. Experiments for calculating TEM-based PDFs were conducted by selected area electron diffraction (SAED). The SAED patterns were acquired at an exposure time of 2 s using an energy slit of 5 eV together with a selected area aperture (10 μm) and entrance aperture (120 µm). The extracted azimuthally integrated diffraction patterns from the SAED patterns were first converted to structure functions, S(Q), and then pair distribution functions (PDFs), G(r), using [34–36]:

$$G(r) = \frac{2}{\pi} \int_{Q_{\min}}^{Q_{\max}} Q[S(Q) - 1] \sin(Qr) dQ,$$
 (1)

and an in-house developed software called EDP2PDF, where the camera length was calibrated at 300 mm in diffraction mode using the (200) plane of gold standard nanoparticles to accurately compare each dataset.

The MRO cluster size was measured by nanobeam electron diffraction (NBED) and fluctuation electron microscopy (FEM) [37–41]. For each measurement, 225 NBED images were converted to mean images and a normalised variance image using [37]:



where *I* is the diffracted intensity from NBED, *V* the normalized variance, k the scattering vector, Q_M the radius of a virtual objective aperture, and < > represents the average over all 225 diffracted images. A nanobeam diameter, R, range of 0.7–1.6 nm was used in this study and Q_M is given by $Q_M = 0.61/R$ [37]. The annular integrated variance and mean were calculated on a pixel by pixel basis [42] as a function of the k-value using PASAD tools [43] and a custom script in DigitalMicrograph [44]. The first two peak positions of $V(k, Q_M)$ were read at approximately 3.6 nm⁻¹ and 4.5 nm⁻¹ using a custom MATLAB script, and the error analysis for the MRO cluster size is explained in our previous study [29]. The pair persistent model was then used to extract MRO cluster/paracrystal diameter in our BMG samples [40]. In this model, the plot of the calculated values of Q_M^2/V against Q_M^2 shows a linear trend [38]:

$$\frac{Q_M^2}{V(k, Q_M)} = \frac{1}{P(k)\Lambda^3} + \frac{4\pi^2}{P(k)\Lambda} Q_M^2,$$
 (3)

where Λ is a characteristic length scale:

$$\Lambda = \frac{1}{2\pi} \sqrt{\frac{m}{c}},\tag{4}$$

that can be extracted from the slope m and intercept c of the line fit to Q_M^2/V versus Q_M^2 . Gibbson et al. [40] suggested that the Λ corresponds to the radius of ordered regions. If we assume it as the radius of gyration for spheroidal regions, the width W of the cluster/paracrystal (MRO in this case) can be estimated from the relation.

$$W = \sqrt{10}\Lambda. \tag{5}$$

Further experimental details can be found in our previous study [29].

Results

Our previous investigations of a comparable LPBF-processed AMZ4 sample indicated uniform porosity and chemistry along the build height, though fluctuations in structure and elastic modulus exist layer-by-layer corresponding to the melt pool thickness [18]. Analysis of other samples produced by the same LPBF-processing parameters have further confirmed



the uniformity in chemistry and porosity [11], and no chemical variations were found after doing further EDS in this study (e.g. see Supplementary Materials Fig. S1). Figure 2a–c shows the microhardness map for the top, middle, and bottom of the build on the sample cross-sections, respectively. Some indent points were measured as very soft spots (e.g. below 400 HV_{0.05}) due to pores produced by the LPBF process and were not included in subsequent averaging. Comparing the overall hardness for the three build positions, the bottom region had the lowest average hardness (491.6 \pm 9.7 HV_{0.05}), which can be seen in the light blue and green colour distributed on the map, while the top and middle regions have an average hardness $HV_{0.05}$ of 494.7 ± 8.3 and 497.4 ± 8.4 , respectively. Figure 2d–f shows Vickers microhardness distribution for each build location. While all three distributions overlap, the bottom of the build contains an enhanced relative frequency of lower hardness points (\sim 470–490 HV_{0.05}) and there is a perceptible shift of the hardness distribution to lower values compared to the top and middle of the build.

Representative DSC curves are displayed in Fig. 3. Although two repeat measurements were performed per sample position, the collected data were found to be repeatable and thus single curves are compared for clarity. Differences between all curves were found to be small; however, some variation was observed for the relaxation enthalpy prior to the glass transition temperature, $T_{\rm g}$, and also in the crystallisation temperature, T_x . The relaxation enthalpy $(\Delta \phi)$ was determined for each curve by integrating the area between the DSC curve and a horizontal line drawn starting at the heat flow at 230 °C. The DSC data from the bottom region of the build had the highest relaxation enthalpy at approximately -8.9 J/g, followed by the top region at approximately -7.1 J/g. The middle region showed the lowest relaxation enthalpy at approximately -6.6 J/g, along with a slight T_x shift to higher temperatures ($\Delta 0.7$ °C). The increased relaxation enthalpy magnitude in the lower ~4 mm of the build suggests a significantly

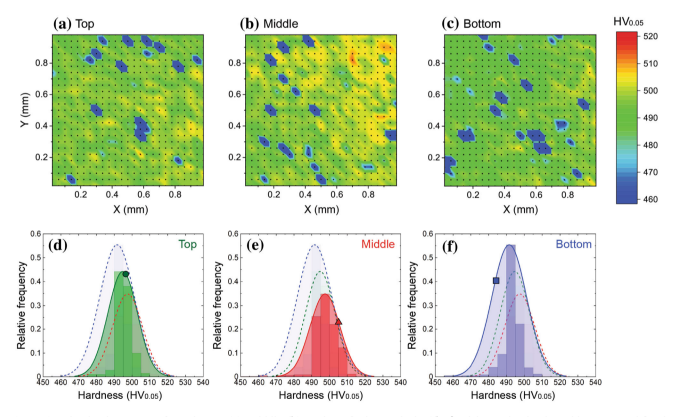
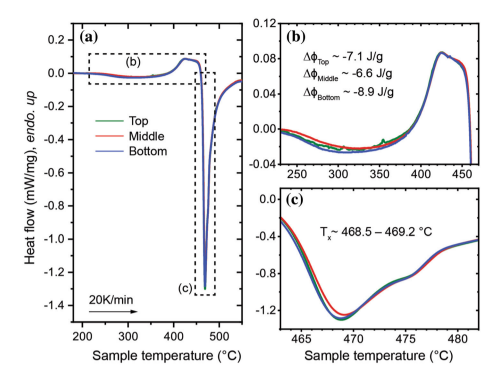


Figure 2 Microhardness maps from the top (a), middle (b), and bottom (c) of the LPBF-processed AMZ4. Approximately 625 indent points were measured for each map, and indents affected by pores (dark blue spots below 400 HV_{0.05}) were removed from

further analysis. (d-f) Vickers microhardness histogram and fitted normal distribution from the top, middle, and bottom of the sample. Symbols in (d-f) indicate the hardness values where FIB samples were taken for TEM analysis of the MRO.



Figure 3 DSC traces for samples taken from the top, middle, and bottom of the LPBF AMZ4 build (a). Magnified view of the DSC curves near the glass transition temperature (b) and crystallisation transition (c). Also shown are the relaxation enthalpy $(\Delta \phi)$ values and the crystallisation temperature range (T_x) .



less relaxed structural state with higher levels of free volume that should reduce the flow stress [45]. Indeed, comparisons of the hardness and relaxation enthalpy results in this study confirm the less relaxed state at the bottom of the build is softer.

To investigate this structural difference observed from DSC in increased detail, DB-PAS was used to monitor free volume difference from the bottom of the build in comparison to the middle portion, i.e., where the largest differences are expected. Figure 4 shows the measured S-parameter obtained from DB-PAS along the build direction over a distance of 10 mm, for two parallel line-scans separated by 100 μm. The S-parameter scatters around a mean value of ~ 0.578 , where the scattering can be attributed to the layered structure present in the printed sample. However, unlike in our previous sXRD measurements [18], no clear periodicity of the variation related to the layer thickness or HAZ can be recognised, owing to the relatively large size of the positron beam which is on the order of the layer thickness. Nevertheless, for the scanning position in the range corresponding to the bottom part of the build, the mean S value is perceptibly higher than that measured in the middle of the build and is emphasised with a smoothed adjacent-averaged curve as the mean values are within a single deviation. It should be noted that in our previous DB-PAS

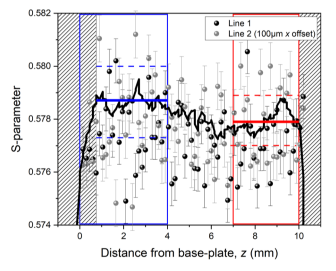


Figure 4 S-parameter obtained from DB-PAS on the LPBF-processed AMZ4 specimen. Two line-scans are shown over a scan distance of ~ 10 mm from the base plate covering the transition from bottom (blue box) to middle (red box) of the specimen. Hashed areas indicate where the measurement was affected by scattering from the Al frame holding the samples. Mean and standard deviations are indicated for both regions, while an 8-pt. adjacent average for both data sets (black line) highlights the subtle transition of the S-parameter from the bottom to middle of the build.

investigation between LPBF and cast AMZ4, the rapidly cooled LPBF material has an S-parameter 0.5% greater than the cast [12]; differences here along



build height are expected to be small. However, the slightly higher S-parameter at the bottom of the build indicates a state of more trapped excess free volume, which is in line with that observed in the DSC measurement as well as the lower average hardness value found for that region.

Figure 5 shows the position of the first structure function S(Q) maximum as a function of scan position from three representative sXRD scans for the top, middle and bottom parts of the build. Despite using a scanning direction perpendicular to our previous work, similar periodic variations caused by heterogeneities from the HAZs can be recognised [18]. This was attributed to the 3-dimensional shape of the melt pools and data collection is expected to cross many HAZs for almost any scan direction. The samples exhibited structure factors characteristic of fully amorphous materials. There was negligible average variation in S(Q) peak positions when comparing the various line scans, indicating that significant differences in oxygen content between build locations can be ruled out based on our previously reported observations of S(Q) for AMZ4 samples with varying oxygen content [46]. Furthermore, no significant statistical differences in S(Q) peak positions were observed for the different regions of the build. Averaging all scans for the bottom, middle, and top part of the build (300 data points per region) gives mean positions of the first structure function maximum of $2.597 \pm 0.002 \text{ Å}^{-1}$, $2.596 \pm 0.004 \text{ Å}^{-1}$, and $2.596 \pm 0.002 \text{ Å}^{-1}$, respectively, which does not

directly correspond to the trends observed in microhardness, DSC, or DB-PAS experiments.

Figure 6 shows representative bright-field (BF) high-resolution TEM (HRTEM) images and selected area electron diffraction (SAED) patterns of the LPBF AMZ4 samples from the top, middle, and bottom of the build. A typical amorphous structure was observed for all of the samples. By searching a large field of view around the edge of the electropolished TEM lamella, no evidence of nanocrystalline phase formation was found at the melt-pools, HAZs, or the matrix. Similarly, searching even larger areas using SEM and EDS further confirmed the melt pools and HAZs were compositionally homogeneous (see Supplementary Materials Fig. S1). While sub-micrometre sized crystals could be found in some HAZs by searching in the SEM, they were isolated and few enough to not appear in the XRD patterns nor in the TEM samples and they did not significantly affect the DSC crystallisation peak magnitudes. Thus, the difference in the hardness response with the build position observed in Fig. 2 was not attributed differences in chemical composition or crystallinity at the micro- or nanoscale.

Figure 7 compares the TEM-based structure functions, S(Q), and PDFs for the top, middle, and bottom of the build. The observed structural changes in the DSC measurement did not result in measurable shifts in the TEM-based S(Q) and PDF nearest neighbour peak positions. While TEM-based PDF analysis has the advantage of collecting local structural information via SAED patterns, the ability of the subsequent

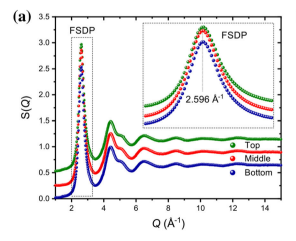
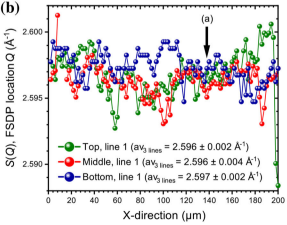


Figure 5 High-energy X-ray micro-diffraction results for the S(Q) scanned on thin cross-sections of material with data collection steps perpendicular to the build direction. Shown are



representative S(Q) curves extracted from the diffraction data (a), along with peak positions for the first significant diffraction peak (FSDP) from a line scan over the sample (b).



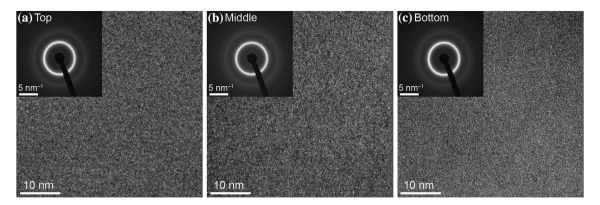
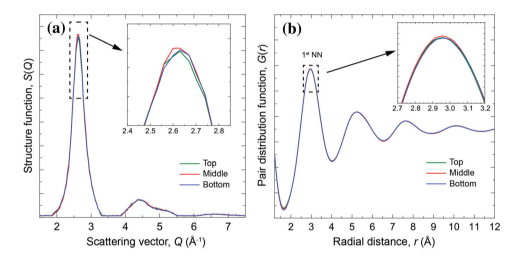


Figure 6 BF-HRTEM images and SAED patterns at different regions of the build: a top, b middle, and c bottom.

Figure 7 TEM-based **a** structure function and **b** pair distribution function for the top, middle, and bottom of the build. The 1st NN represents 1st nearest neighbour interatomic distance.



PDF analysis from SAED patterns to detect the fine structure is relatively insensitive compared with that from synchrotron X-ray diffraction analysis [47–49]. Similar to the synchrotron X-ray diffraction results seen in Fig. 5, there were no clear trends of changing TEM-based structure function or nearest neighbour position with the build height that correspond to the observed trends in hardness, relaxation enthalpy, and S-parameter.

The microhardness maps in Fig. 2 and the DSC results in Fig. 3 showed variations in the hardness and structure between the top, middle, and bottom of the build, while the short-range order nearest neighbour distances and local chemistry were identical. To better understand the structure and property relationship, the medium-range order (MRO) cluster size results were examined using NBED experiments and FEM as presented in Fig. 8. Figure 8a shows an example NBED pattern from the top of the build. With more than 200 NBED patterns, the normalised variance patterns were generated and converted

into a 1D profile, V, as a function of scattering vector, k, as shown in Fig. 8b. The different nanobeam sizes, R, from 0.7 nm to 1.6 nm allow us to generate Q_M^2 V versus Q_M^2 plots (Q = 0.61/R) to extract the MRO cluster size. Figure 8c displays the MRO cluster size using electropolished TEM samples from the top, middle, and bottom of the build. The MRO cluster size from the bottom position was larger than that from the top and middle. Note that the hardness values for the electropolished TEM samples were unknown and the NBED patterns were collected from different locations near the edge of the thin area of the samples. In Fig. 8b, the TEM lamellae were taken between four indents of nearly identical hardness to correlate the MRO cluster size and the local hardness. The specific local hardness values for each position were plotted on the curves with circle (top), triangle (middle), and square (bottom) marks in Fig. 2d–f. A clear trend was found of increasing MRO cluster size with decreasing hardness. The linear



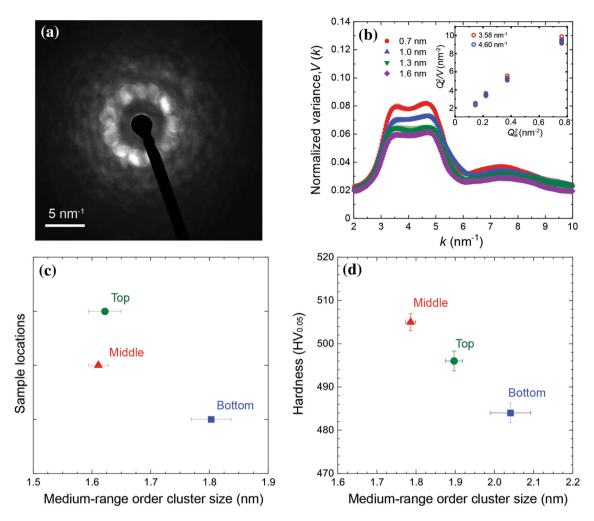


Figure 8 Nanobeam electron diffraction (NBED) and fluctuation electron microscopy (FEM). (a) Example of NBED pattern using a 1.0 nm electron nanobeam. (b) Normalised variance profiles and Q_M^2/V versus Q_M^2 plots (inset). The electron beam diameter was varied from 0.7 nm to 1.6 nm to obtain the plots and the normalised variance profiles were azimuthally integrated into 1D profiles. The peak positions of the normalized variance profiles were read at 3.58 nm⁻¹ and 4.60 nm⁻¹. The sample used in (a) and

(b) was from the top of the build. Medium-range order (MRO) cluster size from the top, middle, and bottom of the LPBF AMZ4 sample using (c) electropolished TEM samples of unknown hardness and (d) site-specific TEM samples of known hardness produced via FIB lift-out. The error bars represent standard deviation for the hardness and standard error for the MRO cluster size.

correlations between the local hardness and MRO cluster size were the same found in Ref. [29].

Discussion

The present study identifies variability in the BMG hardness and atomic-scale structure when comparing the top, middle, and bottom locations of a LPBF build with dimensions $10 \times 10 \times 18 \text{ mm}^3$. It is important to note that these differences could not be attributed to differences in chemical composition, crystallinity,

or porosity at the micro- or nanoscale. Additionally, the thin specimens cut for the presented analyses are also free of residual stresses, evidenced though azimuthal integration of *I*(*Q*) over 30° increments on comparably thin sXRD foils published in Ref [18]. Uniform density/porosity along the build [50] and a lack of apparent crystallisation [14] has also been reported by other studies of AMZ4 produced at the same vendor using the same LPBF processing conditions.

In this study it was found that the bottom of the build was softer with a wider microhardness



distribution that was shifted to lower values. Furthermore, the DSC curves and DB-PAS results in Figs. 3 and 4 indicated that the bottom of the build had a larger relaxation enthalpy, lower crystallisation temperature, and increased free volume. These collective results suggest that a softer, higher energy, and more rejuvenated glassy structure is present at the bottom of the build while the hardest, lowest energy, and most relaxed glassy structure occurs in the middle of the build.

To further understand the structural differences between the build locations, both sXRD and TEM were used. While our previous work found large differences in the sXRD-based structure function and PDF peak positions when comparing as-cast to LPBF-processed AMZ4 [11] and subtle differences could be observed in the structure function across melt pools [18], differences between build locations were not detectable in the present work in both the sXRD and TEM diffraction data. However, structural differences could be observed in the MRO as revealed by NBED experiments in the TEM.

Our initial NBED experiments using electropolished TEM samples revealed the bottom sample to have larger MRO cluster size than the top and middle samples (Fig. 8a); however, those experiments examined random locations in a microstructure of variable hardness, as observed in Fig. 2. Our recent results have shown of the MRO cluster size increases linearly with decreasing local hardness, and that correlation should be independent of the processing history of the specimen [29]. This motivated a second set of NBED experiments where a FIB was used to extract TEM lamella of known hardness from the hardness maps (Fig. 2) of each region. To match the rank order of the average hardness, the softest sample was taken from the bottom and the hardest taken from the middle region. The exact hardness values chosen relative to the overall hardness distribution for each case can be seen in Fig. 2d-f. Figure 8b shows a linear relationship between the local hardness and MRO cluster size that is in good agreement with our previous study [29]. Moreover, a larger nanoscale MRO cluster size can be considered the structural signature of the softer, more rejuvenated state found at the bottom of the build, and to a lesser extent at the top of the build.

In the LPBF process, the base-plate at the bottom of the build acts as a heat sink, helping to dissipate the applied heat by the laser energy. Thus, the bottom of the build is expected to have a higher cooling rate relative to middle locations in the build where a steady state was reached, as has been revealed by simulations of the LPBF process [27, 51, 52]. Higher cooling rates will naturally lead to a more rejuvenated glassy state which is associated with lower hardness; indeed, the relaxation enthalpy for LPBF produced AMZ4 is much higher than for its cast counterpart [12]. Both the hardness maps and the DSC results confirm this with the latter showing larger relaxation enthalpy in the bottom region. As the build progresses into the middle region, more heat must be dissipated in the part leading the slower cooling rates and cyclic reheating [27, 51]. For BMGs, this will lead to a more relaxed structural state forming in the middle of the build, and if the energy density is high enough, large amounts of crystallisation will occur in the HAZs [6]. As a result, a harder and more relaxed structure is found in the middle region and that structure should persist if a steadystate flow of heat input and dissipation is achieved. Finally, simulations have shown a slight uptick in cooling rate at the top section of LPBF builds along with a reduced amount of heat input and fewer thermal cycles relative to the steady state [52], which should result in a slightly less relaxed structure, as is observed in Figs. 2 and 3.

It is important to note that while some structural relaxation of lower layers may occur due to cyclic heat input, there does not appear to be a large amount of relaxation occurring since the relaxation enthalpy everywhere in the LPBF build remains much higher than for cast material [12]. Thus, it is thought that the cooling rate is the dominant factor controlling the structural state, and that the temperature of the solidified material must quickly fall to a low enough fraction of T_g such that continuous relaxation does not occur during the build. This is identical to reports of microstructure heterogeneity along the build height in crystalline metals, where finer scale grains, second phase cell structures, and/ or martensitic structures are observed with higher hardness at the bottom of the build due to the higher cooling rate, yet the subsequent heat input during the build is insufficient to coarsen the bottom region microstructure relative to the higher locations [53-57].

Overall, the variability in the structure and hardness seen with build height for the Zr-based BMG in this study can be explained by the variability of heat



dissipated in the sample. To achieve more uniform microstructure and mechanical properties throughout the build, further investigations will be required, but one strategy might be to adjust the energy density at the beginning and end of the build within the available processing window to ensure a more uniform heat dissipation, while avoiding crystallisation and printing defects. In that regard, simulation studies will be an invaluable tool to guide energy density selection.

Conclusions

In this work, variations in structure and properties of a LPBF-processed BMG along its build height were elucidated using a range of advanced material characterisation techniques. These measurements revealed the structure and property differences that occur at the top, middle, and bottom regions of an LPBF-processed $Zr_{59.3}Cu_{28.8}Nb_{1.5}Al_{10.4}$ bulk metallic glass sample, where the following conclusions may be drawn.

- 1. The bottom region of the build was found to have a softer, higher energy, more rejuvenated glassy state than the middle region of the build, with the top region being intermediate in structure and properties.
- Differences in hardness at different locations could not be attributed to variability in chemical composition, porosity, or crystallinity between the regions.
- 3. The structural signature of the softer bottom region was found to be a larger medium range order cluster size, as revealed by nanobeam electron diffraction experiments in the TEM, together with increased free volume evidenced by DSC and DB-PAS measurements.
- 4. Variability in the structure and properties with build height were attributed to differences in the heat dissipation at various locations in the sample. The bottom of the build experienced heat dissipation into the build plate, which gave faster cooling rates and less reheating compared to the steady-state middle of the build. Finally, at the top of the build less reheating occurs, leading to a less relaxed state compared to the middle region.

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Declarations

Conflict of interest The authors declare that there is no conflict of interest associated with this research.

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