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Introduction

Effect of build orientation and heat treatment on the microstructure, mechanical and corrosion performance of super duplex stainless steels fabricated *via* laser powder bed fusion[†]

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In this study, the effect of build orientation (0°, 45° and 90° from a build platform) on microstructural response as well as mechanical and corrosion properties was investigated by comparing laser powder bed fusion-produced samples in the as-built and solution-annealed states. By increasing build orientation, Widmanstätten γ -austenite formation was lowered because of faster cooling and shorter melt tracts, whilst retaining similar δ -ferrite/ γ -austenite phase fractions. This is correlated with improved corrosion performance in the 90° orientation from chemically homogeneous grain boundary γ -austenite. The prevailing δ -ferrite as-built samples exhibit a strong (001) δ -ferrite crystallographic texture in the normal direction across all orientations together with greater hardness and mechanical strength in comparison to solution-annealed samples by virtue of less slip systems in the BCC δ -ferrite structure and fine cellular solidification structure. The 45° build orientation exhibits a greater Widmanstätten γ -austenite content and periodic recrystallisation between scan checkers, contributing to improved mechanical strength and ductility. Solution annealing softened structures, from an increase in the γ -austenite content, via intergranular nucleation or through prior grain boundaries and Widmanstätten needles. The underlying δ -ferrite grain structure and crystallographic texture relationship is retained, although weakened from the recrystallisation process. Tensile strength is reduced compared to the as-built structures and worsened in the 90° orientation due to few Widmanstätten needles, although elongation is significantly increased, and pitting corrosion performance is improved by the removal of stresses and the equilibrium microstructure

Combining both austenitic and ferritic grades, duplex stainless steels exhibit high mechanical strength, corrosion resistance and chloride stress corrosion cracking resistance. These advantageous properties promote their application in oil and gas umbilical manufacturing, pulp and paper mills, desalination

^c School of Mechanical and Aerospace Engineering, Nanyang Technological University, Singapore, 639798, Singapore plants and bridge construction.¹⁻⁴ However, issues pertaining to their sensitivity during solidification to prevent deleterious phase formation, poor machinability induced by high work hardening and limited product availability have restricted their wider adoption in industry.^{5–10} In an attempt to mitigate these challenges, near net-shaped processing routes such as powder metallurgy have been widely researched as avenues to alleviate these processing challenges, although inducing porosities from powder compaction and sintering is detrimental to the mechanical performance of the alloys.^{11,12} Alternatively, additive manufacturing processes applying a localised heat source to the material do not require a sintering stage and produce dense components. Duplex stainless steels have been successfully fabricated via additive manufacturing processes such as directed energy deposition,13-18 wire arc additive manufacturing,¹⁹⁻²⁴ electron beam melting,^{25,26} and laser powder bed fusion (PBF-L).²⁷⁻³⁴ The latter is of particular interest due to the commerciality, versatility, and high design freedom of the system.35,36

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The PBF-L process involves consolidating metallic powder layer by layer via a localised heat source. This heat source and a high-temperature gradient/solidification speed (G/R) ratio yield a refined microstructure with epitaxial to columnar grain growth propagating in the direction of the build,³⁷ with rapid cooling rates promoting fine cellular and dendritic growth within the grains.^{38,39} This columnar grain growth with refined sub cellular growth and high dislocation concentration contribute to the increased mechanical strength of laser-melted samples as compared to the prepared product for a range of alloy systems.⁴⁰⁻⁴⁵ Furthermore, PBF-L processed alloys show improved corrosion performance as compared to their wrought counterparts in a range of solutions contributing from the high solidification velocity, preventing deleterious inclusion formation, refining the grain size and producing a high grain boundary density.46-52

Unlike the wrought material, the anisotropic nature of the additive process can lead to the accentuation of properties at different build orientations. The greater number and reduced cross-sectional area of layers in samples orientated near-toperpendicular with the build platform can result in an accumulative porosity content, hampering mechanical and corrosion performance.53-56 At a microstructural level, the build orientation influences the cooling rate, with faster cooling across smaller cross-sectional areas. The vertical build orientation was reported by Xie et al. to greatly influence the phase ratio and formation of a Ti-6Al-4V alloy, forming harder a"-martensitic needles and an overall less ductile microstructure.57 Furthermore, as columnar grains are orientated along the vertical build direction, parallel with the tensile load, there is limited resistance to dislocation motion imparting lower yield strengths.58 However, recent studies have shown that leveraging the anisotropy of the process to manipulate the crystallographic texture of grains can result in significant mechanical and corrosion property improvements.⁵⁹⁻⁶¹ These textures can be manipulated via tuning the laser parameters including scanning strategy, laser energy and melt pool mode, and processing gas or, *via* build orientation effects.^{59,62–67} The reporting of preferred crystallographic texture control has been seen in many alloys including tantalum, titanium, tungsten, Ni superalloys, steels and stainless steels.⁶⁸⁻⁷⁵ Considering stainless steels, research has been predominantly focused on the influence of the three main crystallographic direction $\langle 001 \rangle$, $\langle 011 \rangle$ and $\langle 111 \rangle$ in the FCC crystal structured γ -austenite-grade 316L. The $\langle 011 \rangle$ and $\langle 111 \rangle$ texture direction was reported by Wang *et al.* to result in better ductility than the $\langle 001 \rangle$ direction, resulting from twinning-induced plasticity (TWIP) effects, while the $\langle 111 \rangle$ texture showed a higher yield strength due to a lower Schmid factor of the grains in the tensile direction.⁵⁹ Considering corrosion resistance, Trisnanto et al. reports it to be highest in the $\langle 111 \rangle$ direction followed by the $\langle 001 \rangle$ direction due to the higher atomic density in these directions and the lower activation energy of corrosion.⁶¹

Unlike the γ -austenitic 316L-grade stainless steel, PBF-Lprocessed duplex stainless steels are predominantly δ -ferritic with a BCC crystal structure, due to both the ferrite-rich precursor powder and the rapid solidification process, restricting the y-austenite growth.^{27,28,76,77} Post-process heat treatments have proven successful in restoring the microstructure equilibrium; however, under the as-built condition, there is less understanding of the anisotropic properties of PBF-L-processed duplex stainless steels.^{30-33,78,79} The influence of build orientation on corrosion properties was studied by Nigon et al., reporting an insignificant difference in corrosion resistance among the PBF-L-processed duplex stainless steel samples orientated, parallel and perpendicular to the building direction, and deciding the chemical composition was the dominant factor determining corrosion resistance.³⁴ For the same orientations, Yadav reported a >60 HV increase in hardness in the samples perpendicular to the build direction, hypothesising the increase related to crystallographic texture, although in both of these studies, the crystallographic texture was not thoroughly evaluated.⁸⁰ More recently, Zhang et al. have correlated the crystallographic texture after solution annealing heat treatments with the mechanical and corrosion response of samples produced perpendicular and parallel to the build direction. Employing a 67° scanning strategy, they noted ferrite to have a predominantly (001) texture and austenite to have a (101)texture in the perpendicular build direction which, in addition to the refinement and equiaxed morphology, contributed to improved yield and ultimate tensile strength. Further, the corrosion resistance was improved due to the finer grain size and higher dislocation density of grains perpendicular to the build direction.81

Evident from the literature is the strong anisotropic effect in PBF-L processing, influencing microstructure phase content and crystallographic texture, and imparting differences in mechanical and corrosion properties. This study aims to expand the knowledge on the mechanical responses of PBF-Lprocessed super duplex stainless steels, ascertaining processstructure relationships considering different building orientations in both the as-built and heat-treated conditions.

Experimental procedures

PBF-L setup

A Renishaw AM250 PBF-L machine, with a Nd:YAG fibre laser heat source with a maximum output of 400 W, was used to manufacture samples atop a mild steel substrate heated to 170 °C. The samples were processed in an argon environment with an oxygen content <0.5%. Super duplex stainless steel powder of grade UNS S32750/SAF 2507 and spherical morphology was processed with the following composition: 24.6Cr, 6.9Ni, 3.88Mo, 0.9Mn, 0.5Si, 0.31N, 0.04Co, 0.02Al and 0.02C (wt%). Cubic samples produced with laser powers from 300 to 400 W in 25 W increments and three laser scan speeds of 350, 400 and 450 mm s⁻¹ were studied by Archimedes density measurements to determine the favourable processing condition for part density. From this study, a laser powder of 325 W, with a scan speed of 350 mm s⁻¹ hatch spacing of 120 µm and a powder layer thickness of 50 µm was selected, with a resulting



Fig. 1 (a) Checkerboard scan strategy and (b) tensile sample build orientation. Arrows denote build direction (BD), loading direction (LD) and normal direction (ND).



Fig. 2 (a) Volumetric powder particle size distribution; inset: SEM image of powders. (b) XRD pattern of the powder.

laser energy density of 154.8 J mm⁻³. A checkerboard scan strategy was employed, consisting of four 5×5 mm checkers each containing either vertical or horizontal scan lines, alternating adjacent to each other and followed by a single border scan, spaced at 100 µm from the consolidated tracts following steps 1–10, as depicted in Fig. 1(a). Scan lines in each checker starts from the centre and terminates at the border. Scan lines for each checker reverse after each pass with a spacing of 120 µm. The (n + 1) layer scan lines are rotated 90° to the (n + 1)th layer, as shown in Fig. 1(a).

Tensile test samples are placed atop the build platform along their long edge in three orientations $(0^{\circ}, 45^{\circ} \text{ and } 90^{\circ})$, as presented in Fig. 1(b). One set of each parameter and orientation was produced atop the substrate, with four identical builds produced for validation purposes and two subjected to post-process solution annealing.

Post-process heat treatment conditions

A post-process solution annealing heat treatment was applied to restore the microstructural equilibrium of the laser melted duplex stainless steel. An Abar Ipsen HR-50 vacuum furnace was employed on samples attached to the substrate to minimise warping due to the internal stress of the samples. Conditions were selected to exceed the critical cooling rate of 0.8–0.9 °C s⁻¹ to prevent the formation of the deleterious precipitates.⁸² The solution annealing temperature of 1150 °C was selected as it resulted in the greatest γ -austenite volume fraction from a range of temperatures on UNS S32750-grade super duplex stainless steel weldments.⁸³ Thus, the samples were heat treated at 1150 °C for 30 min, prior to a high-pressure nitrogen quench with an average cooling rate of 1.72 °C s⁻¹ and, a 5 °C s⁻¹ cooling rate over the first 500 °C.

Powder analysis

Powder size and distribution were measured using Malvern Morphologi 4 static automated imaging equipment with $> 36\,000$ particles measured. The volumetric particle size distribution is presented in (Fig. 2(a)), with a scanning electron microscopic (SEM) image in the inset acquired using a JEOL 7800F Prime field emission scanning electron microscope, showing the presence of some satellites after the atomisation process. The crystallographic phases of the atomised powder were determined by X-ray diffraction (XRD) using a Bruker D8 Advance diffractometer with a CuKa radiation source ($\lambda =$ 0.154 nm), which we operated at 40 kV at an emission current of 40 mA. The XRD scanning speed was set to 0.05° per min across the 2θ range between 20° and 90° and the XRD patterns were indexed using the Match! 3 software. The base powder is predominantly δ -ferritic after atomisation (Fig. 2(b)), with some small γ -austenite peaks on account of the high cooling rate during atomisation restricting the γ -austenite formation.²⁸

Mechanical and corrosion testing setup

Tensile tests were conducted using a Tinius Olsen H50KS, following the ASTM standard E8M for rectangular samples at a crosshead speed of 0.2 mm min⁻¹, without the use of an extensometer. A sub-size sample defined with a gage length, width and thickness of 25, 6 and 3 mm respectively was used as the base model. Tensile testing conditions and sample sizes are similarly reported in other PBF-L works for comparable alloys.^{84–86} Fracture surface characterisation was established from 6 mm tall sections cut off one half of each broken sample. A Hitachi SU-70 SEM was used for higher magnification imagery of the fracture surfaces up to $20\,000\times$ magnification.

Instrumented indentation technique was used to investigate the variation of micromechanical hardness in samples. An Anton Paar MCT3 system equipped with a Berkovich indenter tip was used at room temperature with 100 indentations taken on each sample in a 10×10 matrix at a spacing of 40 µm between indents. A maximum load (P_{max}) of 100 mN was applied for each indent with a hold time of 10 s at P_{max} . The Oliver–Pharr method was employed to determine the maximum load, contact depth and projected area based off the loadingdisplacement curves obtained. The following equations were used to calculate HVIT as follows:

$$S = \left(\frac{\mathrm{d}P}{\mathrm{d}h}\right)_{\mathrm{max}} \tag{1}$$

$$h_{\rm c} = h_{\rm max} - 0.75 \frac{P_{\rm max}}{S} \tag{2}$$

$$A = 24.5h_{\rm c}^{2}$$
 (3)

$$HVIT = \frac{P_{max}}{A}$$
(4)

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where *S* is the contact stiffness, P_{max} is the maximum load applied during indentation, h_{max} is the maximum indentation depth at maximum load, h_c is the contact depth, *A* is the projected contact area and HVIT is the hardness. The results are represented by hardness distribution maps in the OriginPro graphing software.

Cyclic potentiodynamic polarisation (CPP) tests were conducted in a 3.5 wt% NaCl solution to evaluate the corrosion response of the alloys before and after solution annealing in a chloride-containing aqueous environment. A BioLogic potentiostat controlled by the EC lab software was employed. A flat cell with a three-electrode configuration was used. An exposed surface of 1 cm² of the alloy was used as the working electrode, while a silver-silver chloride reference electrode and a platinum mesh counter electrode were also included in the setup. The samples were mounted in an epoxy resin and the opposite side connected electrically with copper wires. The exposed surface of the sample was prepared by grounding with silicon carbide paper diminishing grit to 2000, followed by ultrasonic cleaning with ethanol. Prior to testing, the samples were conditioned in a 3.5 wt% NaCl electrolyte for 1 h to obtain a relatively stable open circuit potential. Polarisation scans were conducted linearly from 0.25 V below the open circuit potential to 1 $V_{Ag/AgCl}$ at a scanning rate of 1 mV s⁻¹ with a limiting current density of 1 mA cm⁻². The tests were conducted under the naturally aerated condition at room temperature (25 $^{\circ}$ C), with each test repeated three times for reproducibility.

Microstructural characterization

After the testing samples were sectioned using a diamond saw from the grip area of each dog-bone sample. These samples were hot mounted in a conductive resin and ground with SiC papers of diminishing grit from 180 to 2400 and, polished on felt pads using diamond pastes from 6 to 1 µm. A final oxide polishing solution (0.04 µm) stage was incorporated for electron back scatter diffraction (EBSD) followed by immersion in ethanol in an ultrasonic bath. The EBSD images were taken using a JEOL 7800F Prime field emission SEM with an Oxford Instruments Symmetry S2 detector. Measurements were taken at accelerating voltage of a 20 kV and a probe current of 35 nA at a step size of 0.2 µm. The AztecCrystal software was used for EBSD image analysis with the MTEX Matlab toolbox software used to determine the relationships between phases and Schmid factor analysis. Microstructural imaging was taken by electro-etching the sample with a 60% HNO3 - 40% H2O solution at an applied voltage of 10 V for 6 seconds, followed by rinsing with ethanol. Imaging was carried out using an Olympus BX51M optical microscope at 200× magnification.

We confirmed the presence of crystallographic phases in the bulk by means of XRD using the same Bruker D8 Advance diffractometer and process presented in the Powder analysis section above. The indexing of XRD patterns was carried out using Match! 3 software, and the Rietveld refinement was calculated using the Topas V5 software.

Archimedes density analysis was performed using the samples cut for metallographic evaluation, with measurements



Fig. 3 (a) Density and porosity with respect to build orientation in both the as-built and solution-annealed states. (b) Porosity images based on optical micrographs of sample cross-sections. The inset denotes build direction (BD) and loading direction (LD).

taken using a Mettler Toledo XS204 mass balance in air and in ethanol repeating each measurement three times, with results represented in Fig. 3(a). Using a reference density of 7.7 g cm⁻³, densities for all samples exceeded 98.5%, with improvements in density recorded after solution annealing due to the greater austenite fraction.⁸⁷ Micrographs taken at the tensile cross-section are binarized and presented in Fig. 3(b), with the corresponding porosity fraction measured using ImageJ, image processing software, which is reported in Fig. 3(a). Negligible porosity is evident in the cross-section with a volume fraction <0.06% for all samples and orientations.

Results and discussion

Microstructure

Crystallographic phases present in the as-built and solutionannealed samples were determined *via* XRD patterns, presented in Fig. 4. The dual phases of δ -ferrite and γ -austenite were observed in both samples, however, with different relative intensities. The as-built samples (Fig. 4(a)) show high intensities for the δ -ferrite phase in comparison to the γ -austenite phase, expected due to the δ -ferritic nature of the powder and fast cooling during PBF-L, restricting the γ -austenite growth.^{28,76} Unlike other PBF-L works processing the same alloy composition and powder supplier, γ -austenite peaks are observed at a 2 θ of 50.2° for all build orientations and, at a 2 θ of 73.9° for the 0 and 45° build orientations.^{28,32,88} These peaks are also observed in the powder XRD (Fig. 2(b)), indicative of retention after the PBF-L process. After solution annealing, the γ -austenite peak intensity greatly increases, evident in Fig. 4(b),



Fig. 4 XRD patterns of (a) as-built and (b) solution-annealed samples at each build orientation.

with comparable peak profiles across all build orientations. The relative phase fractions of all samples were evaluated by the Rietveld refinement, which are summarised in Table 1, and comparatively evaluated against the EBSD phase maps in this section.

The photomicrographs of the as-built and the solutionannealed samples used for the tensile tests are presented in Fig. 5 in order to establish the structure and property relationships, with the arrows inserted denoting the build direction (BD) and micrographs taken out of plane of the tensile direction (TD). The as-built microstructures consisting of long columnar δ -ferrite grains with sporadic γ -austenite precipitation along the grain boundaries (only evident under SEM) and the intersecting black Widmanstätten γ -austenite needles are observed in Fig. 5. These needles are observed in similar works and reported to grow from allotriomorphic γ -austenite, originally forming along grain boundaries,⁷⁶.⁸⁹ This growth is reported to follow a diffusional mechanism with the dominant Kurdjumov–Sachs phase relationship, with growth occurring between 700 and 1100 °C subjected to rapid cooling.^{90,91}

The underlying δ -ferritic grain morphology is retained in the solution-annealed samples similarly noted by Nigon et al. though, compositionally, γ -austenite precipitation is increased with finer Widmanstätten y-austenite needles and inter-granular precipitates evident in white.92 The retention of this underlying δ -ferritic grain morphology after solution annealing is reported by Balluffi et al. to be due to the preferential nucleation and growth of γ -austenite at δ -ferrite grain boundaries.⁹³ Widmanstätten γ -austenite formation is attributed to the holding temperatures between 700 and 1100 °C, corresponding to the solution annealing cycle employed in this study.90 Phase maps and corresponding phase content fractions in Fig. 6 and Table 1 show comparable austenite fractions for each orientation with only a slight increase in the 45° orientated sample on account of its greater austenite content in the underlying as-built microstructure.

Phase fraction

The primarily δ -ferritic microstructure in laser-melted samples produced in the as-build state is clearly evident in the phase maps presented in Fig. 6(a), (c) and (e). Austenite growth can be seen in limited regions along grain boundaries, intragranular and as Widmanstätten needles in all build orientations. The 45° build orientation shows a slightly greater γ -austenite content than that of the other orientations, as noted in Table 1.



150µm

Fig. 5 Cross-sectional micrographs of as-built and solution annealed samples in the; (a)–(b) 0° , (c)–(d) 45° and (e)–(f) 90° build orientations. Inset denotes build direction (BD) and loading direction (LD).

However, comparative works on PBF-L processing of the same alloy exhibit lower γ -austenite contents (<1%) with growth exclusively allotriomorphic along grain boundaries.^{32,88,94–97} The Rietveld Refinement of XRD patterns (Fig. 4(a)) validate the greater γ -austenite content in the as-built samples, with the γ -austenite content as great as 9.47% in the 90° orientated sample (Table 1). The greater γ -austenite content observed in this work is due to the pulsed laser and keyhole melt mode in the equipment used herein, which results in a deep melt pool with greater cooling rates, up to 10⁷ K s⁻¹, compared to conductive mode melting, promoting Widmanstätten austenite growth and increasing relative austenite content in the sample.⁹⁸⁻¹⁰⁰

After solution annealing, the γ -austenite content increases significantly, up to 59.46%, with nucleation intergranular, along grain boundaries and as Widmanstätten needles, as shown in Fig. 6(b), (d) and (f). The phase ratios were validated by Rietveld Refinement of XRD patterns (Table 1), with

Table 1 Phase fraction of samples built at 0°, 45° and 90° under the as-built and solution-annealed conditions

		As-built (%)			Solution-annealed (%)			
Phase		0 °	45°	90 °	0 °	45°	90°	
δ-Ferrite	EBSD Rietveld refinement	$\begin{array}{c} 94.47 \pm 1.19 \\ 90.59 \pm 0.65 \end{array}$	$\begin{array}{c} 91.83 \pm 1.34 \\ 92.54 \pm 0.37 \end{array}$	$\begin{array}{c} 92.20 \pm 0.86 \\ 90.53 \pm 0.84 \end{array}$	$\begin{array}{c} 43.06 \pm 0.54 \\ 46.02 \pm 0.63 \end{array}$	$\begin{array}{c} 40.89 \pm 0.35 \\ 46.72 \pm 1.02 \end{array}$	$\begin{array}{c} 45.14 \pm 0.28 \\ 42.36 \pm 0.73 \end{array}$	
γ-Austenite	EBSD Rietveld refinement	$\begin{array}{c} 5.53 \pm 1.19 \\ 9.41 \pm 0.65 \end{array}$	$\begin{array}{c} 8.17 \pm 1.34 \\ 7.46 \pm 0.37 \end{array}$	$\begin{array}{c} 7.80 \pm 0.86 \\ 9.47 \pm 0.84 \end{array}$	$\begin{array}{c} 56.94 \pm 0.54 \\ 53.98 \pm 0.63 \end{array}$	$\begin{array}{c} 59.11 \pm 0.35 \\ 53.28 \pm 1.02 \end{array}$	$\begin{array}{c} 54.85 \pm 0.28 \\ 57.64 \pm 0.73 \end{array}$	



Fig. 6 Cross-sectional microstructural phase maps of as-built and solution annealed samples in the; (a)–(b) 0° , (c)–(d) 45° and (e)–(f) 90° build orientations. The inset denotes build direction (BD) and loading direction (LD).

analogous results to EBSD phase maps. As mentioned in the previous section, the underlying δ -ferritic grain morphology after solution annealing is retained after solution annealing, with the preferential nucleation of grain boundary γ -austenite along prior δ -ferrite grain boundaries. Yet overall γ -austenite is homogeneously dispersed in all samples, with a greater preference for Widmanstätten formation in the 0 and 45° orientation, given the slower cooling rate observed in these samples by virtue of their longer melt tract lengths in the build plane, 5 and 4.2 mm respectively and the greater average scanned area for each layer.¹⁰¹ The 90° sample has a short 2.5 mm-long tract in the build plane and a smaller scanning area for each layer, resulting in greater cooling rates and less Widmanstätten yaustenite formation from prior δ -ferrite grain boundaries.^{90,102} Further evidence of this is exhibited in the Grain boundary interface orientation relationship section (Fig. 12).

Crystallographic texture

The crystallographic texture has been widely reported to significantly influence the mechanical and corrosion responses of the alloy. The cross-sectional crystallographic texture maps of tensile samples in the as-built state for each orientation under the as-built and solution-annealed conditions, taken along the normal direction (ND), rolling direction (RD) and transverse direction (TD) are presented in Fig. 7. A strong $\langle 001 \rangle$ crystallographic texture of δ -ferrite-BCC is present in the normal direction for all build orientations as reflected in the high multiples of uniform density (MUD) values of 8.42, 7.12, and



Fig. 7 Cross-sectional crystallographic texture maps of the as-built samples in the; (a)–(c) 0°, (d)–(f) 45° and (g)–(i) 90° build orientations along the normal direction (ND), rolling direction (RD) and transverse direction (TD). The inset denotes the build direction (BD) and loading direction (LD).

6.76 for the 0°, 45° and 90° samples respectively in Fig. 10(a), (d) and (g). The $\langle 001 \rangle$ texture preference is characteristic to cubic ferrous alloys, as it is the fast growth direction following the direction of heat flow.¹⁰³ The vertical heat flow within the melt pool along its centreline facilitates this $\langle 001 \rangle$ orientation, with alignment of the melt pools contributing to epitaxial growth along the BD.^{63,70} Although a checkerboard scan strategy was employed in this work, whereby the scan direction rotates by 90° after each layer, Carter *et al.* reported the presence of long elongated grains with a strong $\langle 001 \rangle$ texture along the build direction.¹⁰⁴

Evident in the rolling (RD) and the transverse direction (TD) (Fig. 7(b, c), (e, f) and (h, i)) is the change in preferential crystallographic texture with build orientation. For the 0 and 90° orientated samples, the δ -ferrite-BCC grains exhibit a strong (001) texture; however, the 45° orientated sample reveals a (203) texture in the {011} plane family as RD and TD maps are taken 45° to the loading direction similarly reported for comparable cubic alloys with the same build direction.^{105,106}

Given the lower grain count and wider statistical variance, γ -austenite-FCC exhibits a lower texture strength under the asbuilt condition (Fig. 10) with maximum MUD values <4.31, probably due to the lower γ -austenite and wider statistical variance. However, it is interesting to note the relationship between δ -ferrite-BCC and γ -austenite-FCC. Both phases exhibit the same $\langle 001 \rangle$ texture in the ND, a result of the fast growth direction of the phase. However, in the RD, the texture is 45° out of plane with the corresponding δ -ferrite-BCC phase.

As reported in the prior section, the δ -ferrite grain morphology in the as-built samples is retained after the solution annealing process, evident in the crystallographic texture maps of Fig. 9, showing the underlying grain build directions and in the inverse pole figures of Fig. 10, showing similar



Fig. 8 Inverse pole figures of the as-built sample texture maps presented in Fig. 7 in the; (a)–(c) 0° , (d)–(f) 45° and (g)–(i) 90° build orientations. Inverse pole figures are out-of-plane, in the tensile direction (TD).

crystallographic textures.^{92,93} The result of this is the same δ -ferrite and γ -austenite texture direction after annealing, however, with a different texture strength. On average, the texture strength decreases after solution annealing for all build orientations and texture directions, as observed in other PBF-L works.^{31,33} Xiang *et al.* reports this due to both the recrystallisation and the formation of grains without a preferred orientation and high-solution annealing temperatures driving γ -austenite nucleation and randomness of preferential crystallographic orientation.³³ The crystallographic orientation relationship



Fig. 9 Cross-sectional crystallographic texture maps of the solutionannealed samples in the; (a)–(c) 0° , (d)–(f) 45° and (g)–(i) 90° build orientations along the normal direction (ND), rolling direction (RD) and transverse direction (TD). The inset denotes the build direction (BD) and loading direction (LD).



Fig. 10 Inverse pole figures of solution-annealed sample texture maps presented in Fig. 9 in the; (a)–(c) 0° , (d)–(f) 45° and (g)–(i) 90° build orientations. Inverse pole figures are out-of-plane, in the tensile direction (TD).

between the δ -ferrite and γ -austenite is important to the texture strength and is studied in detail in the following section.

Grain interface orientation relationship

On the whole, δ -ferrite: γ -austenite interfaces under both the asbuilt and solution-annealed conditions exhibit a misorientation peak of $45-45.6^{\circ}$ (Fig. 11(a)-(f) and 12(a)-(f)) fitting the orientation relationships of Kurdjumov-Sachs (K-S), Nishiyama-Wassermann (N-W), Pitsch (P), Greninger-Troiano (G-T) and Bain (B), as presented in Table 2.33,107 Given a tolerance in the orientation relationship of 2° from ideal, the N-W orientation relationship is more prominent in the as-built and solution-annealed samples (Table 3) especially in the 0° and 45°-orientated samples with average length fractions >21% with the remaining orientation relationships categorised as others.^{107,108} However, when comparing the build orientations, it can be observed that the 0° sample contains comparable length fractions of δ -ferrite: γ -austenite interfaces with the N-W and K-S relationship. This difference in length fractions can be explained by the relative increase in Widmanstätten γ -austenite in the 0° and 45° build direction samples (Fig. 11(a), (b) and 12(a) and (b)), which follow the N-W orientation relationship as compared to the K-S orientation relationship of grain boundary austenite. Although the difference between the N–W and K–S orientation relationship is $\leq 3.13^{\circ}$, the preferred N–W orientation relationship of Widmanstätten γ -austenite:ô-ferrite is not widely reported, with classic works highlighting the preferential K-S orientation relationship possibly due to difference in unit cell spacings.^{91,109,110} Considering the interfaces between similar phases before and after solution annealing, δ-ferrite:δ-ferrite interfaces exhibit a misorientation peak at 45-46° (Fig. 11(g)-(i) and 12(g)-(i)) with a misorientation peak at 60° for γ -austenite: γ -austenite interfaces (Fig. 11(j)-(l) and 12(j)-(l))

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Fig. 11 As-built samples: (a)–(c) orientation relationships of δ -ferrite: γ -austenite interfaces; (b) misorientation angle histograms for the (d)–(f) δ -ferrite: γ -austenite, (g)–(i) δ -ferrite: δ -ferrite and (j)–(l) γ -austenite: γ -austenite grain boundaries at each build orientation.



Fig. 12 Solution-annealed samples: (a)–(c) orientation relationships of δ -ferrite: γ -austenite interfaces, (b) misorientation angle histograms for the (d)–(f) δ -ferrite: γ -austenite, (g)–(i) δ -ferrite: δ -ferrite and (j)–(l) γ -austenite: γ -austenite grain boundaries at each build orientation.

representing $\Sigma 3$ ($\langle 111 \rangle$) grain boundaries, notably reported in the hot rolled duplex stainless steel alloy with formation in the $\langle 111 \rangle$

direction, as this constitutes the configuration with the minimum energy.^{107,111,112}

Instrumented indentation mapping

Instrumented indentation mapping of samples (Fig. 13 and 14) is used to connect microstructures and local property response. In general, the as-built microstructure is harder than the solution-annealed microstructure, exhibiting a peak hardness of 268.3 HVIT in comparison to 230.2 HVIT after solution annealing. The higher hardness in the as-built samples is reported due to the fine cellular structure, columnar grain growth and residual stress.^{114,115} The decrease in hardness is widely reported to be the result of recrystallisation after solution annealing with the greater volume of the softer γ austenite phase.^{33,76,80,116,117} Thus, troughs in hardness maps for both the as-built and solution-annealed samples occur with the hardness indent within a γ -austenite grain. In contrast, peak hardness measurements in the as-built samples (Fig. 13) are observed at δ -ferrite: δ -ferrite grain boundaries and at Widmanstätten γ -austenite sites, the latter reporting greater hardness values due to greater nitrogen content solid solutions strengthening the phase.^{118–120}

In the case of solution-annealed samples (Fig. 14), hardness peaks occur within δ -ferrite grains with a shorter range between the peak and trough values, by virtue of the recrystallisation and stress relieving of the microstructure.³³ It may also be noted that there is no distinct correlation between the hardness and the crystallographic texture for the as-built and solution-annealed samples as comparable textures in the normal direction are observed for each build orientation.

Tensile properties

Engineering stress-strain curves in Fig. 15 show the difference in mechanical response of the super duplex alloy under the asbuilt and solution-annealed conditions. Comparing the two graphs and result summary in Table 4, it is clear that the δ-ferrite-dominated as-built sample has a higher ultimate tensile strength (UTS) (792.2-904.4 MPa) than the solutionannealed samples (677.5-804.5 MPa). The trade-off in ductility is clear with strain at facture values of 30.6-50.1% in the solution-annealed samples reducing to 15.4-29.8% in the asbuilt samples. Similar observations were made in other duplex stainless steel works processed by PBF-L.^{30,31,88} The greater UTS in the as-built samples is expected with the BCC δ -ferritic structure given less slip systems compared to γ-austenite, additionally the fine cellular solidification structure, high dislocation concentration, and the grain pinning further attribute to the improved performance.¹²¹ Essentially mechanical strength is comparable to the work by He et al. for the same alloy, with an improvement in elongation at fracture under both the as-built and solution-annealed conditions.⁹⁷

The as-built samples orientated at 45° and 90° (Fig. 15(a)) exhibit favourable properties with the high UTS (871.2 and 875.9 MPa) values and the greatest strain to fracture (26.6 and 22.2%). Given the comparable phase ratios across all build orientations (Fig. 6), this increase in strength is probably the

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Table 2 Orientation relationships of the δ -ferrite: γ -austenite interfaces with parallelism conditions and angle/axis pairs of the two phases^{107,113}

Parallelism	Minimum angle/axis
$\{111\}\gamma//\{110\}\delta$	$42.85^{\circ}~\langle 0.968 0.178 0.178 angle$
$(110)\gamma/(111)\delta$ $\{111\}\gamma/(110)\delta$	$45.98^\circ\;\langle \textbf{0.976} \hspace{0.1cm}\textbf{0.083} \hspace{0.1cm}\textbf{0.201}\rangle$
$(112)\gamma//(110)\delta$ $\{100\}\gamma//(011)\delta$	$45.98^{\circ}\;\langle 0.083\; 0.201\; 0.976 angle$
$\langle 011 \rangle \gamma / / \langle 111 \rangle \delta$ $\{111\} \gamma / / \{110\} \delta$	$44.23^{\circ}~\langle 0.973~0.189~0.133 angle$
$\langle 123 \rangle \gamma / / \langle 133 \rangle \delta$ $\langle 100 \rangle \gamma / / \langle 100 \rangle \delta$	45° $\langle 1 0 0 angle$
	$\begin{array}{c} \mbox{Parallelism} \\ & \{111\}\gamma // \{110\}\delta \\ \langle 110\rangle\gamma // \langle 111\rangle\delta \\ \langle 111\rangle\gamma // \{110\}\delta \\ \langle 112\rangle\gamma // \langle 110\rangle\delta \\ \langle 100\rangle\gamma // \langle 111\rangle\delta \\ \langle 011\rangle\gamma // \langle 111\rangle\delta \\ \langle 111\rangle\gamma // \langle 110\rangle\delta \\ \langle 123\rangle\gamma // \langle 133\rangle\delta \\ \langle 100\rangle\gamma // \langle 100\rangle\delta \\ \langle 100\rangle\gamma // \langle 110\rangle\delta \\ \rangle \end{array}$

Table 3 Grain boundary length fractions of the δ -ferrite: γ -austenite interfaces for the dominant orientation relationships

	As-built			Solution-annealed		
Orientation relationship	0°	45°	90°	0 °	45°	90 °
Kurdjumov–Sachs (K–S) Nishiyama–Wassermann (N–S)	$\begin{array}{c} 12.4 \pm 1.2 \\ 21.2 \pm 12.4 \end{array}$	$\begin{array}{c} 9.6 \pm 2.1 \\ 24.6 \pm 12.9 \end{array}$	$\begin{array}{c} 13.4 \pm 0.9 \\ 10.9 \pm 2.2 \end{array}$	$\begin{array}{c} 27.6 \pm 1.2 \\ 28.2 \pm 1.1 \end{array}$	$\begin{array}{c} 24.3 \pm 0.6 \\ 28.9 \pm 4.2 \end{array}$	$31.7 \pm 3.5 \\ 11.0 \pm 4.6$





Fig. 13 Instrumented indentation mapping of the as-built samples with corresponding phase, and crystallographic texture maps in the ND and RD, taken in the; (a)–(d) 0°, (e)–(h) 45° and (i)–(l) 90° build orientations.

Fig. 14 Instrumented indentation mapping of solution-annealed samples with corresponding phase, and crystallographic texture maps in the ND and RD, taken in the; (a)–(d) 0°, (e)–(h) 45° and (i)–(l) 90° build orientations.

result of lower residual stress and the number of built layers for each orientation. The chessboard scan strategy is reported to have higher residual stresses than a 90° rotating scan vector, and these stresses are tensile on the upper surface and can be close to the yield stress of the material.^{122–124} The residual stresses are at their highest with the 0°-orientated sample as it has the longest laser irradiation time per layer and greater exposed surface area. Elongation at fracture is greatest in the samples orientated 45° to the build platform, and similar observations were reported in cubic alloys processed by PBF-L. As tensile testing is orientated at an angle to the melt pool tract-tract and layer-layer boundaries, the failure projection path is reported to improve ductility in the 45° sample.^{105,125} As opposed to PBF-L reported in the literature whereby the 90° orientated samples are built vertically along the build direction, resulting in preferential fracture pathways as there is limited resistance to dislocation motion of the grains and alignment of melt pool boundaries perpendicular to the testing direction, which act as locations of weakness due to poor interfacial bonding of the consolidated layers, the 90° samples considered



Fig. 15 Tensile results for (a) as-built and (b) solution-annealed samples at each build orientation.

herein are rotated along their length, contributing to improved strength and ductility.^{58,125}

The ratio of δ -ferrite to γ -austenite is a greater influence on tensile properties in solution-annealed samples, with 72.4 MPa spread in UTS values across the build orientations. However, the 90° sample presents poorer elongation properties, which are probably the result of austenite formation type, given that the 90° sample contains less of the stronger Widmanstätten austenite.¹¹⁹ This is a result of the faster cooling rate during laser melting and less Widmanstätten austenite in the as-built structure.

The influence of the crystallographic texture on mechanical properties can be evaluated by Schmid's law as follows:

$$\tau_{\rm RSS} = \sigma m \tag{5}$$

where $\tau_{\rm RSS}$ is the resolved shear stress, σ is the applied stress and *m* is the Schmid factor.⁵⁹ For a constant τ , the smaller the Schmid factor (*m*), the larger the applied stress in the phase.¹²⁶ The onset of plastic deformation occurs when a critical value of the resolved shear stress ($\tau_{\rm CRSS}$) is reached, which can be defined by the following equation:

$$\tau_{\rm CRSS} = \sigma_{\rm v} m_{\rm max} \tag{6}$$

where by τ_{CRSS} is the critical resolved shear stress, σ_y is the single crystal yield strength and m_{max} is the maximum Schmid factor along a specified crystal orientation for each slip system.⁵⁹ The calculated m_{max} Schmid factors are presented in Table 5 for each phase and slip system.¹²⁶ For an identical critical resolved shear stress (τ_{CRSS}) observing the maximum Schmid factor (Table 5), it can be determined from that the yield strength (σ_y) for the $\langle 001 \rangle$ and $\langle 011 \rangle$ crystallographic orientations is similar and, that the $\langle 111 \rangle$ orientation differs in yield strength by $\sim 1.5 \times .^{59}$

The Schmid factors for each sample in the direction of loading (ND) were calculated from the EBSD data, and are

Table 5 Maximum Schmid factors along each crystallographic orientation for slip systems in δ -ferrite and γ -austenite

Phase	Slip system	Crystallographic orientation	Maximum Schmid factor
		$\langle 001 \rangle$	0.408
	$\{110\}$ $\langle 111 \rangle$	$\langle 011 \rangle$	0.408
		$\langle 111 \rangle$	0.272
δ-Ferrite	$\{112\}$ $\langle 111 \rangle$	$\langle 001 \rangle$	0.471
_		$\langle 011 \rangle$	0.471
		$\langle 111 \rangle$	0.314
	$\{123\}$ $\langle111\rangle$	$\langle 001 \rangle$	0.463
		$\langle 011 \rangle$	0.463
		$\langle 111 \rangle$	0.307
γ-Austenite		$\langle 001 \rangle$	0.408
_	$\{111\}$ (110)	$\langle 011 \rangle$	0.408
		$\langle 111 \rangle$	0.272

presented in Fig. 16 and Table 6. A Schmid factor between 0.40 and 0.49 in δ -ferrite is evident in the as-built samples, indicative of an ease in grain slip of this phase.¹²⁷ The 0° build orientation exhibits the lowest Schmid factor across each of the slip planes (Fig. 16(a)). This can be explained by the stronger (001) crystallographic texture (Fig. 8(a)) in this orientation. For the primary BCC slip system, $\{110\}$ $\langle 111 \rangle$, the maximum Schmid factor (m_{max}) for the BCC (δ -ferrite) unit cell is 0.408 (Table 5) in the $\langle 001 \rangle$ direction, identical to that of the 0° build orientation and as such exhibits a single crystalline-like texture. The larger Schmid Factor in the 45° and 90° build orientations can be attributed to the considerable volume of aberrant grains in the samples.⁵⁹ The difference in Yield strength between build orientations is 0.2-4.6%, similar to the difference between equivalent Schmid factors (0.9-5.3%) across all slip systems. This suggests that the difference in yield strength is influenced by crystallographic texture. However, the 0° build orientation sample with its lower Schmid factor exhibits the lowest yield strength of all build orientations, probably induced by the high residual tensile stress of the chessboard scan strategy, close to the yield stress of the material.¹²²⁻¹²⁴ Additionally, both the BCC δ -ferrite and FCC structured γ -austenite exhibit high Schmid factors (Table 6) in comparison to maximum values reported (Table 5), indicative of a less ordered crystallographic texture. Similar observations were made in FCC γ -austenitic alloys processed by PBF-L, containing equivalent Schmid factors with analogous texture strengths.^{126,128,129} These factors suggest a low likelihood of crystallographic texture strengthening by both the BCC δ -ferrite and FCC γ austenite phases.

After solution annealing, the BCC δ -ferrite Schmid factor increased along the {110} (111) slip system for all build

Table 4 Tensile results of samples built at 0°, 45° and 90° under the as-built and solution-annealed conditions								
	As-built			Solution-annealed				
Sample	0 °	45°	90°	0 °	45°	90°		
Yield strength (MPa) Ultimate tensile strength (MPa) Elongation at fracture (%)	$\begin{array}{c} 691.7 \pm 27.4 \\ 801.5 \pm 12.5 \\ 18.0 \pm 2.6 \end{array}$	$725.1 \pm 29.4 \\ 871.2 \pm 17.9 \\ 26.6 \pm 3.2$	$\begin{array}{c} 723.5 \pm 20.7 \\ 875.9 \pm 28.5 \\ 22.2 \pm 0.8 \end{array}$	$545.0 \pm 22.1 \\789.4 \pm 15.2 \\45.1 \pm 2.4$	$510.6 \pm 8.2 \\ 764.1 \pm 24.6 \\ 44.9 \pm 5.2$	$523.3 \pm 11.4 \ 753.8 \pm 21.6 \ 34.4 \pm 3.8$		



Fig. 16 Schmid factor maps for both phases under the as-built and solution-annealed conditions in the; (a)–(d) 0° , (e)–(h) 45° and (i)–(l) 90° build orientations, taken in the normal direction (ND). The inset denotes the build direction (BD) and loading direction (LD).

orientations indicative of recrystallisation weakening the $\langle 001 \rangle$ preferred orientation (Fig. 16(b), (f) and (j)).³³ Recrystallisation also lead to narrowing of the difference in Schmid factors between build orientations (0.9–1.9%) for the BCC and FCC slip systems. The equivalent Yield strength difference is between 2.4 and 6.3%, suggesting a lower probability of crystallographic texture strengthening.

Differences in ductility can be attributed to the formation of deformation twinning, delaying necking and improving

ductility.^{41,128} The formation of twins is greatly influenced by the crystallographic texture and can provide insights into the tensile response of different build orientations.^{59,128}

In duplex stainless steels, it is reported that FCC structured γ -austenite strengthens duplex stainless steels from dislocation formation and twin hardening.¹³⁰ Meanwhile, the BCC δ -ferrite deforms easier by the slip of dislocations due to the considerably higher stacking fault energy.^{131,132} As such, ductility is directly related to the γ -austenite volume content and its crystallographic texture. The stress level required to initiate twinning in an alloy is known as the critical stress for deformation twinning ($\sigma_{\rm TW}$) and can be calculated for the γ -austenite phase from Byun's formula as follows:¹³³

$$\sigma_{\rm TW} = \frac{2\gamma_{\rm e}}{m_{\rm TW}b_{\rm p}} \tag{7}$$

where $\gamma_{\rm e}$ is the effective stacking fault energy, $m_{\rm TW}$ is the Schmid factor for deformation twinning which is equivalent to the Schmid factor for the leading Shockley partial dislocation $(m_{\rm L})$ and $b_{\rm p}$ is the Burgers vector of a Shockley partial dislocation.^{128,134,135} The Burgers vector $(b_{\rm p})$ of a partial dislocation was calculated by measuring the lattice constant (*a*) obtained from XRD pattern analysis (Fig. 4). Applying eqn (8), the Burgers vector was calculated to be 0.148 nm for both the as-built and solution-annealed γ -austenite:¹³⁶

$$b_{\rm p} = \frac{a}{\sqrt{6}} \tag{8}$$

The effective stacking fault energy (γ_e) in tension can be determined using the following equation:^{137,138}

$$\gamma_{\rm e} = \gamma_0 + \frac{(m_{\rm T} - m_{\rm L})}{2} \sigma b_{\rm p} \tag{9}$$

The stacking fault energy at the zero-stress state is represented as γ_0 , $m_{\rm T}$ and $m_{\rm L}$ are the trailing and leading partial dislocation Schmid factors and σ is the applied stress level. The Schmid factor maps of the leading $(m_{\rm L})$ and trailing $(m_{\rm T})$ partial dislocations are presented in Fig. 17. The stacking fault energy at the zero-stress state (γ_0) can be empirically calculated from the chemical composition of the alloy and was estimated to be ~16.59 mJ m⁻², comparable to other duplex stainless steel works.^{139–141} The critical stress for deformation twinning (σ_{TW}) is presented in Table 7 in addition to the true stress at yielding. Evident for all samples is the true stress at yield 1.6–2.2 \times greater than the critical stress for deformation twinning. This difference is greatest in the as-built samples due to the dominant stronger BCC δ-ferritic structure. Given that the critical stress for deformation twinning (σ_{TW}) is considerably lower than the true stress at yield, it can be inferred that deformation twinning occurs before dislocation slip, similarly noted in other additive-manufactured duplex stainless steels.^{126,142} Assessing the critical stress for deformation twinning results against ductility (Fig. 15), the greatest critical stress for deformation twinning for either of the build condition correlates with the lowest ductility. This is reported in other PBF-L works to be induced by crystallographic texture; however, a comparable

Table 6 Schmid Factors of samples built at 0°, 45° and 90° under the as-built and solution-annealed conditions, along the loading/normal direction (ND)

	Slip system	Schmid factor						
		As-built (%)			Solution-annealed (%)			
Phase		0°	45°	90°	0 °	45°	90°	
δ-Ferrite	$\{110\}$ $\langle111 angle$	0.40 ± 0.03	0.46 ± 0.01	0.47 ± 0.01	0.46 ± 0.02	0.47 ± 0.02	0.47 ± 0.02	
_	112 111	0.47 ± 0.03	0.49 ± 0.01	0.49 ± 0.02	0.49 ± 0.01	0.49 ± 0.01	0.47 ± 0.03	
	$\{123\}$ $\langle111\rangle$	0.48 ± 0.02	0.49 ± 0.01	0.49 ± 0.01	0.49 ± 0.01	0.49 ± 0.01	0.48 ± 0.02	
γ-Austenite								
	$\{111\}$ $\langle110 angle$	0.45 ± 0.04	0.46 ± 0.03	0.46 ± 0.04	0.45 ± 0.03	0.46 ± 0.04	0.46 ± 0.03	

texture strength and crystallographic orientation ($\langle 001 \rangle$) is observed in γ -austenite (Fig. 8 and 10) irrespective of build orientation for each build condition,¹²⁸ indicating the low probability of ductility enhancement by crystallographic texture.

Evident after tensile testing as-built samples in the 45° build orientation is periodic necking (Fig. 18), with a 5 mm peak-topeak distance, consistent with the checker length in the checkerboard scan strategy. The cause of this "wavy" necking is probably due to the greater remelting present at the boundary of each checker, seen in prior work to greater recrystallisation and enlarged grains.⁷⁶ A cross-sectional crystallographic texture map in the normal direction is presented in Fig. 18(b) with a visible difference in grain size from the bottom to the top, and the small grains at the bottom result from the melting of the unsupported powder layer, enlarging with increasing solid layers. Altogether, a weak $\langle 101 \rangle$ crystallographic texture is observed for the δ -ferrite grains (MUD < 2.61) and complementary (001) γ -austenite grain orientation (MUD < 3.87), resulting from EBSD imaging at 45° to the build orientation. Similar crystallographic textures were observed in highly deformed wrought duplex stainless steels, with strain-induced $\langle 101 \rangle$ texture development.¹⁴³ The contribution of this texture on mechanical properties can be evaluated from Schmid factor analysis. The Schmid factor map in Fig. 18(c) provides insights into the contribution of crystallographic texture on mechanical properties. Schmid factors of 0.44, 0.47 and 0.48 are calculated for the $\{110\}$ $\langle 111 \rangle$, $\{112\}$ $\langle 111 \rangle$ and $\{123\}$ $\langle 111 \rangle$ (δ -ferrite) slip systems, indicative of an ease in slip. It may be noted that Schmid factor (0.47) in the $\langle 011 \rangle$ direction of the $\{112\} \langle 111 \rangle$ slip system is identical to the maximum Schmid factor (m_{max}) presented in Table 5, suggesting single crystalline-like texture. This texture is probably induced by the tensile deformation, where twinning of the γ -austenite leads to hardening of the $\{112\}$ $\langle 111 \rangle$ slip system in δ -ferrite.¹⁴⁴ However, a less ordered crystallographic texture of the γ -austenite can be deduced from the high Schmid factor of 0.44. The Kernel average misorientation (KAM) map presented in Fig. 18(d) represents the distribution of dislocations in the necked region, with a decrease in dislocations from the centre of the necked region to the remelted-recrystallised contour region of the checker. Recrystallization by laser remelting is reported in other PBF-L works to result in increased elongation and strength, by increasing

the bonding between melt pools and through heterodeformation-induced strengthening between the recrystallised and non-recrystallised structures.^{145,146} This periodic necking is not as predominant in the 0°-orientated sample probably due to the thinner gage section in the build direction, though it was lightly evident in the 90°-orientated samples, corroborating the mechanical property response reported above.

The fractured surfaces of both the as-built and solutionannealed samples are presented in Fig. 19. The as-built samples show lesser necking and ductility than those of the solutionannealed samples and show evidence of cleavage fractures consistent with the brittle fracture modes induced by the dominant BCC δ -ferrite microstructure. The areas of ductility are evident in the as-built samples at 0° and 90° along the upper face, as shown in Fig. 19(a) and (e) as regions of necking and at the centre of the 0° sample as a cone-shaped fracture. Necking is less visible in the 45° sample, however, as fracture occurred at the inter-checker boundary. Comparably, solution-annealed samples show a ductile trans-granular fracture with sufficient necking along the gage width, as evident in Fig. 19(b), (d) and (f). The solution annealing treatment results in a change in the microstructure and possible recrystallisation of the austenite phase growing within the ferrite matrix. This is expected to enhance the ductile nature of the specimens, which is clearly evident looking at the necking and other features of the photographs presented in Fig. 19(b), (d), and (f). It may also be observed that cracks are initiated and evidently dispersed in the central regions of the fractured surfaces, but there was insufficient energy to allow them to grow and propagate the fracture across the surface.

Higher magnification images of the fracture surfaces highlight the micro-sized dimple fracture features, reported by Sufiiarov *et al.* to constitute ductile transgranular fractures and suggested as comparable in size to solidification cells, suggesting cell size of the order of 0.3–1.2 µm comparable to other PBF-L-processed duplex alloys.^{77,96,147,148} Enlarging of these dimples is evident after solution annealing with sizes between 0.8 and 4.5 µm, notable in laser-melted alloys after heat treatment processing formed as a result of the recrystallisation process and equilibrium microstructure formation.¹⁴⁹ Dimple morphologies also vary between build orientations, with the as-built sample at 45° showing evidence of smaller and more equiaxed dimples than the larger and more elongated





Fig. 18 Periodic necking in the as-built 45° sample: (a) optical micrograph of sample surface showing spatial necking along sample thickness, (b) crystallographic texture map in the normal direction (ND) and corresponding inverse pole figures, (c) Schmid factor map and (d) KAM map.

5mm

whereby tensile tearing produces more elongated dimples.¹⁵⁰ It may be noted that the fracture directions are different in different samples, as already noted. Further, the direction of the growth of the cellular structures is also different from one build orientation to the other. It is understood that the variations in the dimple sizes from one build orientation to the other, as noted in Fig. 19, are the results of the combined effects of these two aspects: the direction of growth of the cells and the direction of fracture and the consequent direction at which the fractured cells are observed.

the build direction (BD) and loading direction (LD). dimples in 0° and the 90° samples. As Janssen *et al.* reports,

uniaxial loading results in equiaxed dimple morphology,

dislocations for γ -austenite in the; (a)–(d) 0°, (e)–(h) 45° and (i)–(l) 90° build orientations, taken in the normal direction (ND). The inset denotes

Table 7 True stress at yield and critical stress for deformation twinning of γ -austenite in the samples built at 0°, 45° and 90° under the as-built and solution-annealed conditions

	As-built			Solution-annealed		
Sample	0°	45°	90°	0°	45°	90°
True stress at yield (MPa) Critical stress for deformation twinning (MPa)	$\begin{array}{c} 878.2 \pm 28.8 \\ 450.8 \pm 3.3 \end{array}$	$\begin{array}{c} 918.9 \pm 27.2 \\ 421.8 \pm 3.3 \end{array}$	$\begin{array}{c} 932.2 \pm 32.2 \\ 422.1 \pm 3.7 \end{array}$	$\begin{array}{c} 726.2 \pm 14.0 \\ 433.4 \pm 1.9 \end{array}$	$\begin{array}{c} 684.8 \pm 16.8 \\ 437.2 \pm 2.3 \end{array}$	$\begin{array}{c} 703.6 \pm 16.5 \\ 444.3 \pm 2.1 \end{array}$



Fig. 19 Fracture surfaces of the as-built and solution-annealed samples in the; (a)–(b) 0° , (c)–(d) 45° and (e)–(f) 90° build orientations at (I) low magnification (35×) and (II) high magnification (5000×).

Corrosion response

The corrosion responses of the samples produced under the asbuilt and solution-annealed conditions were studied by cyclic potentiodynamic polarisation (CPP), as presented in Fig. 20. Comparing the microstructures of samples under the as-built and solution-annealed conditions, the results show an overarching improvement in corrosion resistance after solution annealing, with a pitting potential (E_{Pit}) of solution-annealed samples of, on average, 29 mV higher than that in the as-built state, signifying increased pitting resistance. The passive region of the cyclic potentiodynamic polarization (CPP) curves for solution-annealed samples illustrates fewer fluctuations and spike-like features. This suggests a reduced formation of metastable pitting in the alloy. Furthermore, the lower passive current density indicates a slower dissolution rate of annealed samples than that of the as-built samples. This can be



Fig. 20 Cyclic potentiodynamic polarisation curves for (a) as-built and (b) solution-annealed samples at each build orientation in 3.5 wt% NaCl.

predominantly attributed to the increased γ -austenite content in the solution-annealed samples. The increased austenite content, as both Widmanstätten needles and inter-granular precipitates, decreases the corrosion rate of the alloy as the greater Ni and N contents in the austenitic phase increase passivation ability and reduce the oxidation rate.^{78,151} The removal of residual stress is also reported to improve the corrosion resistance of PBF-L-processed alloys.^{89,152,153}

A similar CPP response is observed for all the as-built samples, as shown in Fig. 20(a). Only a slight variation in CPP response is observed for the three orientations, with the 45° and 90° orientations showing relatively high pitting potentials (E_{Pit}) , 11 mV and 18 mV respectively, compared to that built with a 0° build orientation. The lower volume of Widmanstätten γ -austenite in the 90° sample (Table 3) results in improved corrosion properties, given the structure depletion in corrosion-resistant elements and the formation of secondary γ -austenite.^{154–156} Thus, concluding that the ratio of γ -austenite to δ -ferrite is more influential to corrosion response than the crystallographic texture, given comparable phase ratios across the build orientations. Similar observations were made by Nigon et al. for the standard duplex stainless steel processed by PBF-L.³⁴ In general, the pitting potential values of the asbuilt samples are comparable to the work by Akilan et al. for the same alloy with similar part densities.94

In contrast, the solution-annealed samples reveal a different trend, as illustrated in Fig. 20(b). Specifically, the sample orientated at 0° exhibits a slightly compromised passive corrosion behaviour, evident in its one order of magnitude higher passive current density and less steep slope of the CPP curve compared to other heat-treated alloys. This is probably the product of the $\langle 011 \rangle$ γ -austenite texture (Fig. 10(a)) retained from the as-built microstructure, which is reported by Trisnanto et al. to result in lower corrosion resistance due to its lower atom packing density and greater surface energy than the other orientations,61,157 leading to higher corrosion susceptibility among the solution-annealed samples. The 45° and 90° built direction samples demonstrate the highest pitting potential, 27 mV higher than that of the 0° sample. This can be attributed to the strong texture of δ -ferrite in this sample, given the comparable phase ratios for all orientations.

The comparable $E_{\rm corr}$ and $I_{\rm corr}$ values between the solutionannealed and as-built samples suggest a similar corrosion behaviour during the active dissolution stage prior to the passivation, which is likely influenced by chemical phase variations within the alloys.^{34,158} However, in the context of corrosion of DSS, evaluating the pitting potential is paramount, as it indicates the material's resistance to localised corrosion. Yet despite their δ -ferrite-dominated microstructure, the asbuilt samples exhibited promising pitting corrosion resistance because of their solute-rich ferritic structure containing Cr, Ni and Mo alloying elements beneficial to corrosion resistance.¹⁵⁹

In addition to the phase content and crystallographic texture, other microstructural features may influence the corrosion resistance of PBF-L-processed alloys, and one such feature is porosity. It is widely reported that increasing the porosity

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leads to increased pitting characteristics including pit initiation and pit re-passivation behaviour.^{46,160–164} It is reported that porosity is particularly detrimental above 1%, likely increasing the susceptibility to environmentally assisted cracking.^{46,160–164} While difficult to completely assess each microstructural feature's influence on corrosion in isolation, porosity is minimal ranging from 0.01 to 0.05%. Furthermore, annealed samples with a slightly higher porosity (45° and 90° build orientation) exhibit lower pitting susceptibility, indicated by lower pitting potential (E_{Pit}) values. Thus, porosity plays a less dominant role in the corrosion resistance of PBF-Lprocessed duplex stainless steel.

Conclusions

In this work, the mechanical and corrosion responses of PBF-Lprocessed super duplex stainless steels built in three build orientations have been examined and related to their underlying microstructures under both the as-built and heattreatment conditions. From these findings, the structure-property relationships can be ascertained and the following conclusions are drawn.

The as-built microstructures are predominantly δ -ferritic (>90.53%), on account of the δ -ferritic powder material and rapid cooling of the PBF-L process, limiting the γ -austenite growth. Analogously, the γ -austenite ratios (~7.97%) are discernible for each build orientation. However, a greater volume of Widmanstätten γ -austenite needles is visible in 0 and 45° orientations, given the slower cooling rates from longer melt tract lengths. The shorter scanning area of the 90° orientation exhibits a greater tendency for grain boundary γ -austenite growth. A strong $\langle 001 \rangle$ crystallographic texture is observed in the normal direction of δ -ferrite for all orientations, representing the fast growth direction following the direction of heat flow. Along the rolling direction, the 45° orientation exhibits a $\{011\}$ plane texture for δ -ferrite.

Solution annealing yields a considerable increase in the γ austenite ratio (>53.28%) with nucleation along prior grain boundaries and as Widmanstätten needles, in addition to intergranular growth. Thus, the underlying δ -ferritic grain morphology is retained after the annealing process, retaining the as-built crystallographic texture however, at a lower strength resulting from recrystallisation and new grain growth without preferential orientation.

Interfaces between δ -ferrite and γ -austenite grains exhibit a misorientation peak at 45–45.6° and follow the Kurdjumov–Sachs and Nishiyama–Wassermann orientation relationship for grain boundary and Widmanstätten γ -austenite respectively. An equivalent misorientation peak (45–46°) was observed for δ -ferrite: δ -ferrite interfaces in addition to a misorientation peak of 60° for γ -austenite: γ -austenite interfaces, indicative of Σ 3 grain boundaries.

The as-built $\delta\mbox{-ferritic}$ microstructure is harder due to the fine cellular structure, columnar grain growth and

high residual stress, with hardness peaks occurring at δ -ferrite: δ -ferrite grain boundaries and solid solutionstrengthened Widmanstätten γ -austenite. Local hardness peaks are observed in the δ -ferrite phase of solution-annealed samples; however, the average hardness decreases after recrystallisation.

The as-built samples exhibit a higher ultimate tensile strength up to 904.4 MPa owing to the BCC δ -ferrite structure with less slip systems and fine cellular solidification structures, though with a trade-off in elongation (<29.8%). Solution annealing restored elongation up to 50.1% by stress relieving and increased ductile FCC γ -austenite content but reduced the tensile strength to <804.5 MPa. Promising tensile strength and elongation are observed in the as-built 45° orientation by virtue of lower residual stress and periodic recrystallisation between scanning checkers. Microstructure equilibrium after solution annealing generated comparable tensile strengths for each build orientation, but with the 90° orientation, exhibiting decreased elongation owing to a lower Widmanstätten γ -austenite content.

The mixed-mode fracture is apparent in the as-built samples with brittle and ductile trans-granular fracture features, with ductile fracture dominant after solution annealing. Microdimple-fractured surfaces are observed in both the as-built and the solution-annealed states, enlarging from 0.3 to 1.2 μ m in the as-built state to 0.8–4.5 μ m after heat treatment from stress relieving and increased γ -austenite content.

Solution annealing fractionally improved pitting corrosion resistance, a result of the increased γ -austenite content as both Widmanstätten needles and inter-granular precipitates and, through the removal of residual stress. The 90° build orientation exhibited slightly higher pitting corrosion resistance in the as-built condition, probably due to the greater chemically homogeneous grain boundary γ -austenite content.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Karl Peter Davidson: conceptualisation, data curation, investigation, formal analysis, writing original draft, review and editing. Ruiliang Liu: data curation, writing review and editing. Chenyang Zhu: data curation, formal analysis. Mehmet Cagiciri: data curation, formal analysis. Li Ping Tan: data curation, formal analysis. Alpravinosh Alagesan: data curation. Sarat Singamneni: supervision, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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