Article

Benchmarking of 316L Stainless Steel Manufactured by a Hybrid Additive/Subtractive Technology

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Abstract: This research study investigated the hybrid processing of 316L stainless steel using laser powder bed fusion (LPBF) processing with high-speed machining in the same build envelope. Benchmarking at four laser powers (160 W, 240 W, 320 W, and 380 W) was undertaken by building additively with machining passes integrated sequentially after every ten deposited layers, followed by the final finishing of select surfaces. The final geometry was inspected against the computer-aided design (CAD) model and showed deviations smaller than 280 µm for the as-built and machined surfaces, which demonstrate the good efficacy of hybrid processing for the net-shape manufacturing of stainless steel products. The arithmetic average roughness values for the printed surfaces, Ra (linear) and Sa (surface), were 11.4 µm and 14.9 µm, respectively. On the other hand, the vertical and horizontal machined surfaces had considerably lower roughness, with Ra and Sa values ranging between 0.33 µm and 0.70 µm. The 160 W coupon contained layered, interconnected lack of fusion defects which affected the density (7.84 g cm⁻³), yield strength (494 MPa), ultimate tensile strength (604 MPa), Young’s modulus (175 GPa), and elongation at break (17.3%). By contrast, at higher laser powers, near-full density was obtained for the 240 W (7.96 g cm⁻³), 320 W (7.94 g cm⁻³), and 380 W (7.92 g cm⁻³) conditions. This, combined with the isolated nature of the small pores, led to the tensile properties surpassing the requirements stipulated in ASTM F3184—16 for 316L stainless steel.

Keywords: hybrid manufacturing; laser powder bed fusion; 316L stainless steel; density; surface quality; X-ray micro-computed tomography; microstructure; hardness; tensile properties; fractography

1. Introduction

In the manufacturing sector, grade 316 L stainless steel (316LSS) is a ubiquitous alloy due to its durability, hygienic efficacy, and resistance to corrosion and high-temperature oxidation. These properties have allowed for the diverse applications of 316LSS, such as in marine engineering, aircraft and land vehicles within the defense sector, as well as in industries processing, medical devices, food, potable water, oil and gas, nuclear waste, chemicals, and petrochemicals, to name a few. The excellent combination of the physical, chemical, and mechanical properties of 316LSS is attributed to its chemistry, consisting of a high percentage of Cr (16–18.5%) and Ni (10–14%), as well as some Mo (2–3%) added to this mix. Considering the recent listing of Cr as a critical raw material (CRM) [1], as well
as the high cost of these alloying elements, the sustainable manufacturing of 316LSS has sparked high research interest in recent additive technologies [2-8]. Luckily, due to the low carbon content (<0.03% maximum), 316LSS exhibits excellent weldability by fusion welding processes and is well-suited for the new paradigm of engineering with additive manufacturing (AM) strategies [9].

On the basis of available AM literature, Fayazfar et al. [10] reviewed powder-based additive technologies for the processing of steels and showed that the vast body of research has concentrated on the laser powder bed (LPB) processing of austenitic 316LSS [11-14]. The review by Baja et al. [11] on the microstructure and properties of steels processed by AM covered the research and challenges over the last decade to develop process windows for 316LSS using LPB technologies. More recently, Haghdadi et al. [15] reviewed the achievements and challenges for the AM of steels and indicated that more research is needed to address distortion (due to residual stresses), anisotropy, and pore formation in additively processed 316LSS. Of further concern for the AM of 316LSS is the low accuracy and high roughness of as-built surfaces that necessitate post-processing to attain high product performance [16] for certification in load-bearing, critical, or extreme environments. Unfortunately, owing to its low carbon and high alloy content, the inclination of this austenitic grade is to work harden at a very rapid rate, which poses considerable difficulties for the finish machining of additive parts to net-shape [17]. Undertaken separately and sequentially after AM, out-of-envelope machining poses additional challenges for aligning, workpiece-holding, and referencing, especially due to the lack of precise geometric datum in the as-built part [18].

Thus, in-envelope hybrid processing—which combines subtractive and additive technologies in a single machine—is an important research area for 316LSS. Hybrid additive/subtractive manufacturing (A/SM) using LPB technology, such as the Matsuura LUMEX Avance-25 system [19], can offer the additive building capability for complex structures in synergy with machined surfaces with high tolerance and low roughness to achieve the minimal waste of high-value alloys and/or CRMs, as well as conformance to high quality and tight geometric requirements. However, the A/SM of 316LSS brings new challenges, including the lack of know-how on the efficacy of dry micro-milling for the finish-machining of surfaces, the allowable engineering design properties, and the material–process–structure–property (MPSP) interrelationships. To date, Avegnon et al. [20] additively built 316LSS using a hybrid machine but undertook out-of-envelope milling to study the effectiveness of energy consumption as a process signature that could be correlated with the microhardness. Afazov et al. [21] used an A/SM machine for building 316LSS, but also investigated machining parameters out-of-envelope, using a stand-alone micro-milling center to assess the impacts on the material removal rates and the surface roughness. By contrast, Ahmad and Enemuoh [22] examined the influence of LPB and in-envelope micro-milling parameters on energy consumption during A/SM and developed an analytical model for the processing of 316LSS, but without considering the efficacy of the parametric set on the part quality and performance. Mutua [23] used an A/SM machine and applied only the LPB process to relate the build parameters to the surface quality, density, microstructure, and microhardness of 316LSS. Thus, a considerable gap exists in the understanding of the hybrid processing capability for 316LSS vis-à-vis surface quality after in-envelope machining, as well as the part quality and mechanical performance possible from the LPB process. The present study was therefore devised to explore hybrid manufacturing of 316LSS to identify a robust process window for A/SM through an evaluation of the powder characteristics, part distortion, surface finish, density, porosity features using X-ray micro-computed tomography (µCT), macro/microstructure, bulk hardness, microhardness, and tensile properties. The mechanical properties and the MPSP interrelationships established in this research undoubtedly expand the current state-of-the-art in hybrid manufacturing know-how on 316LSS, which is of importance for functional improvement of a broad range of products in various industries. In this study, the hybrid A/SM process was conducted on a Matsuura LUMEX Avance-25 system.
2. Experimental Procedure

2.1. Powder Characteristics

The starting material used in this study was a commercial nitrogen gas-atomized SUS 316LSS powder (Matsuura stainless steel 316 L, St. Paul, MN, USA) with a nominal particle size of \(-45/10 \mu m\) and an elemental composition (in wt.%) of 0.01% C, 12.52% Ni, 0.70% Mn, 17.18% Cr, 2.04% Mo, 0.93% Si, 0.004% S, 0.008% P, 519 ppm N, 405 ppm O, and balance Fe. Powder characterization was undertaken with a scanning electron microscope (SEM) (Hitachi SU3500, Fukuoka, Japan) to investigate the surface morphology. As revealed in Figure 1, the as-received powder showed a high volume of spherical, regularly-shaped particles, with a smooth surface and a few satellites. Moreover, a limited number of irregular and elongated particles was observed. The particle size distribution of the starting powder was assessed using a laser diffraction analyzer (Horiba LA-920, Kyoto, Japan); the three characteristic mean volume-weighted diameters of the fine (D_{10}), median (D_{50}), and coarse (D_{90}) particle fractions were determined to be 15 \mu m, 22 \mu m, and 36 \mu m, respectively.

The flowability of the 316LSS powder was assessed, respectively, with Hall and Carney funnels (Qualtech Products Industry-HFM1800 SS, Denver, CO, USA) by following the specifications in the ASTM B213 [24] and ASTM B964 standards [25]. The measured flow times through these funnels were 28.6 s (Hall) and 4.5 s (Carney) for 50 g of 316LSS powder. The apparent density of the powder was assessed to be 4.32 g·cm\(^{-3}\) (Hall) and 4.36 g·cm\(^{-3}\) (Carney) in accordance with ASTM standards [26,27]. Also, the static flow behavior of the powder was assessed from the conical heap of powders using image analysis. The static angle of repose for the 316LSS powder was comparable statistically to 34° and sufficient to minimize defects during LPB AM by encouraging good powder spreading, according to Carr’s classification of powder flowability [28].

A GranuDrum\(^{\circledR}\) rotating drum instrument was employed to quantify the cohesion occurring in the 316LSS powder during drum rotation by measuring the cohesive index of the particles from the fluctuations of the avalanche interface [29,30]. The analysis consisted of rotating a transparent drum filled with 50 cm\(^3\) of powder at angular velocities that ranged from 2 to 30 revolutions per minute (RPM) to induce powder flow. The measurements were obtained from backlighting the rotating drum and capturing images of the avalanche at different times using a CCD camera. In total, 40 images of the rotating drum were taken at an interval of 1 frame per second for each RPM. The built-in GranuDrum\(^{\circledR}\) software detected the air–powder interface location automatically and also computed the average interface position, as well as the fluctuations from this average position for each angular velocity. The fluctuations of the air–powder interface position were then directly related to the cohesion inside the drum and designated as the cohesion index. The evolution in
the cohesive index for the 316LSS powder as a function of the rotating speed (Figure 2) indicated a relatively low cohesive index, which is inherently characteristic of good powder flowability, according to the findings [31] for powders with a cohesive index lower than 24 that exhibited free-flowing behavior and resulted in a homogenous and uniform layer during the spreading of the powder in the LPB process.

Figure 2. Cohesive index as a function of the rotation speed of the drum.

2.2. Hybrid Additive/Subtractive Processing

A/SM with the 316LSS powder was undertaken in a Matsuura LUMEX Avance-25 system that combines the LPB technology with high-speed milling. The powder bed platform or build plate was fabricated from 4140 wrought steel and had dimensions of $X = 125$ mm, $Y = 175$ mm, and $Z = 30$ mm. This build plate was demagnetized to a magnetic field less than 0.2 Gauss using a surface demagnetizer (Electro-Matic model A13-1, R. B. Annis, Elmatco, Chicago, IL, USA) and then placed on the A/SM worktable. During the A/SM process, the temperature of the build platform was set to 50°C, and the build chamber was purged with nitrogen gas (of purity not lower than 97%) to prevent the oxidization of the molten pool during laser melting. The deposition of layers on the build platform involved single-direction laser scanning with a 90° rotation after every layer to build prismatic rectangular blocks (dimensions of $X = 75$ mm $\times Y = 25$ mm $\times Z = 25$ mm) using different laser power (P) conditions—namely, 160 W, 240 W, 320 W, and 380 W—for both the rastering/infill and contouring passes (one of each per layer). The laser scan speed ($v$) was 700 mm·s$^{-1}$ and 1400 mm·$s^{-1}$, respectively, for the infill and contour passes, while the values for the other parameters were fixed at a spot diameter ($d$) of 200 µm, a layer height ($h_L$) of 50 µm, and hatch spacing ($h_S$) of 120 µm. Thus, with these fixed and varied parameters, for the laser powers selected in this study of 160 W, 240 W, 320 W, and 380 W, the corresponding infill energy density ($E_{\text{density}}$) values were $38.1$ J·mm$^{-3}$, $57.1$ J·mm$^{-3}$, $76.2$ J·mm$^{-3}$, and $90.5$ J·mm$^{-3}$, and the contour $E_{\text{density}}$ values were $19.0$ J·mm$^{-3}$, $28.6$ J·mm$^{-3}$, $38.1$ J·mm$^{-3}$, and $45.2$ J·mm$^{-3}$, respectively, using Equation (1):

$$E_{\text{density}} = \frac{P}{(v \cdot h_L \cdot h_S)} \quad (1)$$

After depositing ten successive layers, remnant metal powder around the build was suctioned/removed and surface milling was then iterated [19]. A feed rate of 450 mm·min$^{-1}$ with a 0.1 mm radial depth of cut for the vertical side-walls and 4 mm for the horizontal top surface, based on the manufacturer’s recommended settings, was applied. The four prismatic blocks produced by A/SM consisted of two as-built (B) faces and three built-and-machined (B&M) faces, as displayed in Figure 3.
2.3. Sample Preparation and Qualification Testing

An optical three-dimensional (3D) scanning system (ATOS Core, GOM, Germany) was used to inspect the geometrical dimensions of the four prismatic rectangular blocks (still attached to the build platform) after A/SM to assess the conformity/accuracy against the 3D CAD model. Then, the blocks were separated from the build platform using electro-discharge machining (EDM) (FANUC Robocut C400iB, Oshino-mura, Yamanashi, Japan) with a brass wire of 0.2 mm in diameter. A portable Surftest SJ-210 (4 mN type profilometer, Mitutoyo Aurora, IL, USA) with a tip radius of 2 µm was utilized to assess the surface quality of the blocks on both the as-built, as well as the built-and-machined faces by measuring the primary profile to calculate the roughness parameters, which included the arithmetic mean height (Ra), root mean square height (Rq), and maximum height (Rz). It is noteworthy that the total measured length of the primary profile was 4 mm and the roughness profile was derived from the primary profile by suppressing the long wave component using the high-pass filter with a cut-off of λc = 0.8 mm. Moreover, a 3D laser scanning confocal microscope (Keyence VK-X250, Osaka, Japan) was used to measure the equivalent areal roughness parameters—namely Sa, Sq, and Sz—on the different surfaces in accordance with ISO 25178-2 [32].

Each block was then divided into two sections (S1 and S2) alongside the XZ plane, as illustrated in Figure 4a. Samples were extracted from section S1 to evaluate the relative density, porosity by X-ray µCT, and tensile properties, while section S2 was utilized to extract samples for metallography and hardness testing. The subsequent EDM of section S1 was used to extract three samples through the thickness (in the BD) of each block (160 W, 240 W, 320 W, and 380 W), as illustrated in Figure 4a. These were then machined to dog-bone-shaped tensile samples with a 25 mm gage length (Figure 4b), according to the guidelines for a standard sub-size geometry in ASTM E8M-21 [33]. Prior to static tensile testing, these samples were inspected non-destructively for porosity and density using three methods: Archimedes [34], helium gas pycnometry [35], and X-ray µCT. Density measurements, using the Archimedes and He gas pycnometry methods, were undertaken with an AND BM-500 density measurement kit and an Anton Paar Ultrapyc 5000 system, respectively. For the calculation of relative density, a theoretical value of 8.0 g/cm³ was used for 316LSS [36]. The pore size and porosity distribution were evaluated on the gage length of the tensile samples using an HMXST 225 X-ray µCT system (Nikon Metrology Inc., USA) equipped with a Perkin-Elmer 1621AN CsI (2000 × 2000 pixels, 40 cm × 40 cm, 200 µm/pixel) detector panel. The X-ray source was operated with a voltage of 120 kV, a current of 58 μA, and an exposure time of 1000 ms with a 0.25 mm Ag filter. The scans were undertaken at two magnification levels: (1) a low magnification level of 10.9X to permit the visualization of the full gage length, which gave a voxel size of 18.4 µm; and (2) a higher magnification scan at 46X, which gave a voxel size of 4.4 µm. For image analysis,
Dragonfly software was utilized for 3D reconstruction to analyze the volume and size distribution of the pores using manual segmentation.

Figure 4. Schematic diagrams showing (a) extraction methodology, and (b) the geometry of the tensile samples.

The twelve tensile samples—three for each laser power (160 W, 240 W, 320 W, and 380 W) that were extracted from the top, middle, and bottom of section S1 in the four as-built blocks—were then tested using a 250 kN testing frame integrated with a laser extensometer (MTS Systems Corporation, Eden Prairie, MN, USA). Tensile testing was performed at room temperature using displacement control at a rate of 0.4 mm min\(^{-1}\) until rupture. The tensile properties of 316LSS that were evaluated from the engineering stress–strain curves included the yield strength or 0.2% proof stress (YS), ultimate tensile strength (UTS), percent elongation (EL), and Young’s modulus (YM). The fractured surfaces of the tensile samples were examined after testing using an SEM at 15 keV to understand the role of the different pore structures (isolated, interconnected).

By contrast, samples for metallography and hardness testing (Figure 4a) were extracted from the S2 sections of the four blocks produced at the different laser powers. For conceptualizing the 3D macro/microstructures of the 316LSS, metallographic samples from the three orthogonal planes (X-Y, X-Z, Y-Z) were extracted for each laser power (160 W, 240 W, 320 W, and 380 W) using a precision low-speed diamond cut-off saw. The metallographic preparation of the samples involved mounting in cold resin, followed by grinding and polishing automatically to a finish of 0.04 µm, as detailed in [37]. While the characteristics of the pores were examined on polished surfaces, the general microstructure was revealed by immersing the samples in an electrolytic etch at room temperature for 30 s in a solution of 10% oxalic acid with a voltage of 15 V at a current of 1.6 A using an austenitic stainless steel cathode.

The phase analysis of the as-built 316LSS samples was investigated by X-ray diffraction (XRD) and magnetic induction methods. A Bruker D8 Discover diffractometer (Karlsruhe, Germany) with Co-K\(\alpha\) radiation (wavelength of 0.178897 nm) at a voltage of 35 kV and 45 mA was used to obtain XRD patterns of the as-built 316LSS samples at room temperature. The diffraction angle (2\(\theta\)), at which the X-rays impinged on the polished surface of the cross-sections prepared for metallography, was varied from 20° to 110° at a scan step of 0.005 s. The DiffracPlus software of the diffractometer allowed the initial processing of the diffraction pattern to differentiate the peaks corresponding to the austenite (\(\gamma\)), delta (\(\delta\))-ferrite, and martensite (\(\alpha’\)) phases. Moreover, the \(\delta\)-ferrite content in the samples was examined using a Feritscope\(^{\circledR}\) (Model MP3C, Helmut Fisher GmbH). This non-destructive inspection device is based on a magnetic induction method that was originally designed to measure the \(\delta\)-ferrite content in austenitic stainless steel welds. In this magneto-inductive test method, an electromagnetic field generated by a coil interacts with the magnetic constituents in the sample, including \(\delta\)-ferrite and/or any retained/strain-induced \(\alpha’\)-martensite in 316LSS. The magnetization of these phases induces an electrical potential difference in a second
coil and this output voltage is linearly related to the magnetic permeability of the sample. It has been reported that this method is a direct and reliable way to quantify low amounts (roughly 0.1%) of \( \delta \)-ferrite \[38\]. To obtain a reliable quantitative measurement, a calibration was performed using certified standard sets (Helmut Fischer) with traceability to The Welding Institute (TWI, UK) secondary standards, according to the method described in DIN EN ISO 8249 \[39\] and ANSI/AWS A4.2 \[40\]. The Feritscope measurements were performed on the as-built and/or machined surfaces. Roughly ten different points were measured for each laser power and reported as the percentage of ferrite.

The hardness of the 316LSS samples produced using different laser powers was assessed using Rockwell and Vickers testing according to specifications given respectively in ASTM standards E18-20 \[41\] and E384-17 \[42\]. The bulk hardness of the samples was evaluated by Rockwell testing on the C-scale, which is typically used for steels. For each laser power (160 W, 240 W, 320 W, and 380 W), at least 5 measurements were performed—using a diamond rounded-tip cone indenter (ground at 120°) with a diameter of 0.2 mm and a minor load of 10 kg, followed by a major load of 150 kg—to calculate the average Rockwell C hardness (HRC). Vickers microhardness testing was carried out on polished (mirror-finished) metallographic samples using an automated hardness tester (Struers DuraScan 80, Ballerup, Denmark) \[43\] with a load of 500 g, an indent spacing of 0.8 mm, and a dwell period of 15 s. For each sample condition, the reported Vickers hardness profile across the cross-sections was averaged from three measurements for each data point.

3. Results and Data Analysis

3.1. Printed Geometry Versus CAD Model Qualification

During LPB processing, the repetitive cycles of rapid, layer-wise heating and cooling induce large temperature gradients and thermal stresses that can distort the as-built geometry relative to the CAD model. To evaluate the geometric stability and accuracy of the four blocks built by A/SM, inspection with an optical 3D scanner (ATOS Core) was undertaken (before EDM from the build platform), so as to obtain 3D coordinate measurement data of the build in the form of a 3D mesh for analysis and comparison with the reference CAD data using a best-fit algorithm. Figure 5 describes the inspection results for the geometry of the four 316LSS blocks, charted via a contour map with color levels representing deviations ranging from 0.3 mm to −0.3 mm. The as-built surfaces (i.e., B faces labeled with deviation values in Figure 5a) exhibited geometric deviations ranging from 80 \( \mu \)m to 280 \( \mu \)m, which could be significantly decreased after machining (i.e., B&M faces labeled with deviation values in Figure 5b) to values under 10 \( \mu \)m. This demonstrates the high efficacy of the in-envelope milling process for dimensional and shape accuracy. The lower accuracy of the as-built surfaces concurs with findings from thermal simulations that have pointed to the key LPB process parameters \( P, \nu, h_L, h_S \) affecting the part geometry \[44\].

To inspect the quality of the surface finish on the four 316LSS blocks produced by A/SM, the average values of the different linear profile and areal roughness parameters were evaluated on the as-built and machined surfaces, as tabulated in Table 1. In this study, both areal and profile parameters were measured. The former has the competitive advantage of extending the evaluated dataset laterally over a surface area. However, the latter method directly traces the surface with a mechanical probe and provides comparable data to the longstanding norms/practice used in manufacturing industries for specifying surface quality required for geometric dimensioning and tolerancing (GD&T) \[45,46\].
106.4 μm, respectively, at the lowest and highest conditions of 160 W (19.0 J · mm⁻³) and 380 W (45 J · mm⁻³). By contrast, intermediary conditions at laser powers of 240 W (57.1 J · mm⁻³) and 320 W (76.2 J · mm⁻³) resulted in more even surfaces with Sz values of 85–90 μm. The typical Ra (11.12 µm to 11.73 µm) and Sa (13.23 µm to 16.88 µm) values measured in the present study for the as-built surfaces of 316LSS corroborated well with the reported Ra of 9.95 µm by Mutua [23] and 14.05 µm by Avegnon et al. [20] for samples built using the Matsuura LUMEX Avance system.

Figure 5. Contour plots of the geometric deviations on the (a) as-built (B) and (b) built-and-machined (B&M) surfaces of the four blocks fabricated at the different laser powers.

Table 1. Profile and areal roughness parameters measured on the different vertical side-walls and horizontal top-faces of the four blocks.

| Condition/Face       | Power (W) | E_density (J · mm⁻³) | Profile (µm) | Areal (µm) |
|----------------------|-----------|----------------------|--------------|------------|
|                      |           | Infill               | Contour      | Ra         | Rq         | Rz         | Sa         | Sq         | Sz         |
| As-built vertical    | 160       | 38.1                 | 19.0         | 11.67      | 14.34      | 62.07      | 16.88      | 21.75      | 144.77     |
| side-walls           | 240       | 57.1                 | 28.6         | 11.22      | 13.59      | 57.52      | 13.23      | 16.63      | 84.94      |
|                      | 320       | 76.2                 | 38.1         | 11.12      | 13.14      | 56.38      | 13.38      | 16.15      | 89.80      |
|                      | 380       | 90.5                 | 45.2         | 11.73      | 14.39      | 62.45      | 16.15      | 19.45      | 106.40     |
| Machined vertical    | 160       | 38.1                 | 19.0         | 0.33       | 0.43       | 2.32       | 0.42       | 0.54       | 5.04       |
| side-walls           | 240       | 57.1                 | 28.6         | 0.38       | 0.48       | 2.37       | 0.38       | 0.48       | 6.09       |
|                      | 320       | 76.2                 | 38.1         | 0.62       | 0.78       | 3.98       | 0.38       | 0.50       | 5.50       |
|                      | 380       | 90.5                 | 45.2         | 0.65       | 0.81       | 3.88       | 0.70       | 0.87       | 6.86       |
| Machined horizontal  | 160       | 38.1                 | 19.0         | 0.36       | 0.48       | 2.64       | 0.35       | 0.45       | 5.35       |
| top-face             | 240       | 57.1                 | 28.6         | 0.47       | 0.55       | 2.57       | 0.36       | 0.47       | 5.02       |
|                      | 320       | 76.2                 | 38.1         | 0.59       | 0.76       | 4.00       | 0.36       | 0.46       | 4.11       |
|                      | 380       | 90.5                 | 45.2         | 0.55       | 0.72       | 3.58       | 0.45       | 0.54       | 4.45       |
For the as-built vertical side-wall surfaces, the roughness parameters (Table 1) showed a minor dependence on the laser power over the process window studied in this work, namely with the contour $E_{\text{density}}$ ranging from 19.0 J mm$^{-3}$ to 45 J mm$^{-3}$. The surface roughness on the as-printed vertical side-walls exhibited $S_z$ values of 144.77 µm and 106.4 µm, respectively, at the lowest and highest conditions of 160 W (19.0 J mm$^{-3}$) and 380 W (45 J mm$^{-3}$). By contrast, intermediary conditions at laser powers of 240 W (57.1 J mm$^{-3}$) and 320 W (76.2 J mm$^{-3}$) resulted in more even surfaces with $S_z$ values of 85–90 µm. The typical Ra (11.12 µm to 11.73 µm) and Sa (13.23 µm to 16.88 µm) values measured in the present study for the as-built surfaces of 316LSS corroborated well with the reported Ra of 9.95 µm by Mutua [23] and 14.05 µm by Avegnon et al. [20] for samples built using the Matsuura LUMEX Avance-25 system.

After in-envelope micro-milling of the four blocks, the roughness (Table 1) on the vertical side-walls had Ra and Sa values that ranged from 0.33 µm to 0.65 µm and 0.35 µm to 0.7 µm, respectively. The in-envelope machined horizontal top-faces were comparable with Ra and Sa values ranging from 0.36 µm to 0.59 µm and 0.35 µm to 0.45 µm, respectively. Relative to the as-built surfaces (having average values for Ra and Sa of 11.4 µm and 14.9 µm, respectively), these Ra and Sa values after the in-envelope micro-milling of the 316LSS surfaces were more than an order of magnitude lower and similar to finish qualities achievable on polished surfaces [47,48]. It is noteworthy that Afazov et al. [21] reported minimized Ra values between 0.55 µm and 0.65 µm when using optimized milling parameters on an out-of-envelope system (i.e., Kern Evo Ultra-Precision CNC Machine) to machine 316LSS samples as-built using a Matsuura LUMEX Avance-25 system. The comparable roughness attained in the present study demonstrates the good capability, processing flexibility, and high efficacy possible with in-envelope hybrid processing.

### 3.2. Density Characteristics

The density of the as-built 316LSS was measured by the Archimedes and helium gas pycnometry methods, and the data are tabulated in Table 2 for the laser powers used in this study. The lowest densities of 7.84 g cm$^{-3}$ (Archimedes) and 7.97 g cm$^{-3}$ (gas pycnometry) were measured for the lowest laser power of 160 W (38.1 J mm$^{-3}$). On the basis of these density measurements, the relative density ranged between 98.0% and 99.6%, indicating a porosity percentage of 0.4%–2%. An examination of the X-ray µCT scans of the 160 W sample (Figure 6a) revealed that this porosity was related to balling and/or lack of fusion defects (i.e., black regions in images) that were large, irregular, and interconnected. It is worth mentioning that the relative density of 98%, measured for the 160 W condition on the basis of the average Archimedes density, resembles more closely the porosity level in the µCT scans. This points to an overestimation of the density values by the helium gas pycnometry method, which may be attributed to the substantially lower dynamic viscosity of helium (by 40 times) relative to water, which leads to a higher penetration of the open porosity, as explained in [49].

| Laser Power (W) | Infill $E_{\text{density}}$ (J mm$^{-3}$) | Archimedes Density (g cm$^{-3}$) | STD | Pycnometer Density (g cm$^{-3}$) | STD |
|----------------|-------------------------------------|---------------------------------|-----|-------------------------------|-----|
| 160            | 38.1                                | 7.84                            | 0.01| 7.97                          | 0.05|
| 240            | 57.1                                | 7.96                            | 0.03| 8.02                          | 0.04|
| 320            | 76.2                                | 7.94                            | 0.01| 8.02                          | 0.05|
| 380            | 90.5                                | 7.92                            | 0.02| 8.04                          | 0.04|
By contrast, for the 240 W (57.1 J·mm⁻³), 320 W (76.2 J·mm⁻³), and 380 W (90.5 J·mm⁻³) conditions, the presence of small spherical and isolated pores were observed in the µCT scans (Figure 6b–d). On the basis of the Archimedes density values, the relative densities for the 240 W (99.5%), 320 W (99.3%), and 380 W (99.0%) conditions were higher than the 160 W condition (98.0%). Moreover, the pores were observed to be finest for the 240 W condition, as shown in Figure 7, and some minor coarsening was apparent with increasing laser power (Figures 6b–d and 7). Specifically, increasing the laser power from 160 W to 240 W led to a higher infill $E_{\text{density}}$ that improved the densification of the powder during LPB processing, as revealed in Figure 6 by the absence of the lack of fusion pores and in Figure 7 by the small size of the pores, i.e., less than 30 µm. Further increases in the laser power to 320 W resulted in relatively similar porosity characteristics, as those observed for the 240 W condition, but the size of the pores was larger, ranging up to 60 µm. At the maximum power of 380 W (and infill $E_{\text{density}}$ of 90.5 J·mm⁻³) used in this study, the coarsening of the pores was obvious, with typical sizes of up to 100 µm. Nonetheless, on the basis of the µCT scans (Figure 6) and the standard deviations measured on the density datasets (Archimedes and gas pycnometry in Table 2), the relative density and porosity percentage can be considered statistically comparable (from a t-test with an alpha level of 0.05) for the 240 W, 320 W, and 380 W laser powers when factoring in the error margins of the test methods. This process window to achieve near-full density in 316LSS by A/SM concurs reasonably well with the highest relative density of 99.2% reported by Mutua [23] at 320 W and 77.92 J·mm⁻³.
3.3. Macro/Microstructures

Figure 8 illustrates the results of reconstructing representative 3D macrostructures from two-dimensional (2D) optical images of the etched metallographic cross-sections in the three orthogonal planes—the X–Y plane in the transverse direction (TD) which is normal to X–Z and Y–Z planes in the BD—of the as-built 316LSS produced at the different laser powers. The presence of these irregularly shaped pores at the melt pool boundaries can be seen in Figure 8a, along with some indication of balling behavior due to the insufficient melting of the 316LSS powder at the low applied laser power (160 W) and infill $E_{\text{density}}$ (38.1 J mm$^{-3}$). The presence of such defects can be mainly attributed to insufficient bonding between the layers and/or scan tracks due to the low penetration of the laser energy [7], which results in a smaller melt pool size that leaves some powder particles un-melted or semi-melted in the 160 W sample. Similarly, insufficient energy during the powder bed processing of other metals, including maraging steels [50], as well as titanium and aluminum alloys [51–53], has also been noted to lead to melt pool discontinuities, which result in a lack of fusion defects, consisting of pores with irregular morphologies and un-/semi-melted powders.

By contrast, the 3D macrostructures of the 316LSS samples produced at 240 W (Figure 8b), 320 W (Figure 8c), and 380 W (Figure 8d) revealed no cracking and a very limited number of small, isolated pores with spherical morphologies. Typically, such non-uniformly distributed spherical pores in the microstructure of 316LSS have been reported as unavoidable gas-induced porosity originating from gas entrapment in the powder feedstock [54], as well as solidification shrinkage [55].

Material addition layer-upon-layer during the LPB process resulted in a macrostructure composed of overlapping scanning tracks of the laser beam, as illustrated by the optical microscopy images in Figure 8 of the etched 316LSS samples. In particular, the rastering scan pattern and the layer-by-layer rotation (at an angle of 90°) applied during LPB processing were especially evident in the X–Y planes of the near-full-density 316LSS samples, as revealed in Figure 8b–d. For the sample produced at 240 W (infill $E_{\text{density}}$ of 57.1 J mm$^{-3}$), the average width of the scan tracks was ~94 μm. With increasing laser power and $E_{\text{density}}$, the average width of the scan tracks increased roughly to 100 μm at 320 W (76.2 J mm$^{-3}$) and 195 μm at 380 W (90.5 J mm$^{-3}$). By contrast, the cross-sections of these melted scan tracks appeared as curved “troughs”, and both the X–Z and Y–Z planes in the near-full density 316LSS samples showed overlapping patterns of half-ellipse-shaped
molten pool boundaries, as displayed in Figure 8b–d. For the 240 W sample, the melt pool beads were ~135 µm in width and ~128 µm in depth. With increasing laser power, the average dimensions of the melt pool beads increased in width to 148 µm and 162 µm for the 320 W and 380 W samples, respectively, but the aspect ratio remained similar. This agrees with reported studies on the LPB processing of steels that showed increasing bead width with increasing laser power [56,57] and was attributed to the dependence of the melt pool dimensions on the LPB process parameters, including the $E_{\text{density}}$, $d$, $h_L$, $h_S$, etc. [58].

At higher magnification, the visible molten pool boundaries showed good metallurgical bonding without the presence of any microvoids and/or microcracking. A fine cellular solidification structure was observed within the individual molten pools, as shown in Figure 9a,b for the BD and TD of the as-built 316LSS. These solidification cells were
inhomogeneous in size and orientation and their growth was confined within the molten pool boundaries. In the 316LSS samples, the different remnant cells within the epitaxially grown grains had an average spacing of ~0.58 μm. Similar fine cellular structures have been reported for 316LSS produced additively, and attributed to the rapid solidification of the molten pool due to the high cooling rates (on the order of \(10^3\)–\(10^6\) K·s\(^{-1}\)) in the LPB process [59,60].

**Figure 9.** Representative optical microscopy images in as-built 316LSS (a) along the BD and (b) normal to the BD (TD).

The phase analysis of the 316LSS samples indicated that the primary solidification phase was face-centered cubic γ-austenite without any detectable presence of δ-ferrite, as indicated by the representative XRD patterns given in Figure 10a. A similar result was obtained through the measurement of the δ-ferrite content using a Feritscope, with average readings between 0.13% and 0.17% that were at the detection limits of this method and indicate a predominant austenitic structure. Considering the equivalent Cr/Ni ratio (1.38) and a solidification rate of ~20 mm·s\(^{-1}\) obtained from the Kurz–Giovanola–Trivedi (KGT) model of 316LSS, using the measured average cell size of ~0.58 μm, the dominant phases can be predicted using the solidification map reported in the literature [61]. Figure 10b shows that the samples used in this study are well into the fully austenitic region, further supporting the XRD and Feritscope results. Many studies on the LPB processing of 316LSS have observed a predominant austenitic structure [5,7,11,62–64], but some have also reported small amounts of δ-ferrite or α′-martensite [65–67]. The rapid solidification of the molten pool due to the high cooling rates has been reported to favor the formation of γ-austenite [68], while slow cooling rates (as a result of thermal cycling, for example) tend to promote δ-ferrite stabilization, especially in micro-segregated regions [69,70].
Figure 10. Representative XRD patterns of the 316LSS samples were produced using LPB A/SM (a). (b) Solidification map for austenitic stainless-steel welds [61].

3.4. Mechanical Properties

Using the Rockwell C-scale and Vickers methods, the bulk hardness and microhardness of the 316LSS were measured for the different laser power (and infill $E_{\text{density}}$) conditions, as given in Table 3. The average hardness of the 316LSS was lowest (11.0 ± 0.5 HRC and 206.9 ± 17.0 HV$_{0.5}$) for the lowest applied laser power of 160 W (infill $E_{\text{density}} = 38.1$ J·mm$^{-3}$), which exhibited the lowest densities of 7.84 g·cm$^{-3}$ (Archimedes) and 7.97 g·cm$^{-3}$ (gas pycnometry) or highest porosity percentage values of 0.4% (gas pycnometry) and 2% (Archimedes). In addition, the measured Vickers hardness of 207.1 ± 17.6 HV$_{0.5}$ in the TD and 206.6 ± 16.3 HV$_{0.5}$ in the BD indicated insignificant anisotropy in the hardness of 316LSS (with the lowest $p$-value of 0.9958). However, the microhardness plotted across the sample cross-section (Figure 11) indicated considerable scatter in the data for the 160 W laser power that could be attributed to the distributed defects in the 316LSS structure (Figures 6a and 8a). Specifically, the local hardness near any large inter-layer and/or interconnected pores (186.2 HV$_{0.5}$) was lower by 16% relative to the remaining locations (221.3 HV$_{0.5}$) in the 316LSS. Thus, improved densification (to near-full density) at the higher laser powers of 240 W (infill $E_{\text{density}} = 57.1$ J·mm$^{-3}$), 320 W (infill $E_{\text{density}} = 76.2$ J·mm$^{-3}$), and 380 W (infill $E_{\text{density}} = 90.5$ J·mm$^{-3}$) resulted in an increase of more than 30% in the average Rockwell C hardness and 9% in the average Vickers microhardness relative to the 160 W condition. In the range of 240 W to 380 W, the influence of the laser power on the hardness was not obvious, as the bulk hardness increased from 14.2 to 17.9 HRC, while the Vickers microhardness was statistically comparable (from a t-test with an alpha level of 0.05) within the narrow range of 226.1 HV$_{0.5}$ to 235.6 HV$_{0.5}$.

Table 3. Average hardness values of as-built 316LSS.

| Laser Power (W) | Infill $E_{\text{density}}$ (J/mm$^3$) | Rockwell Hardness (HRC) | Vickers Microhardness in TD (HV$_{0.5}$) | Vickers Microhardness in BD (HV$_{0.5}$) |\(\text{Infill E}_{\text{density}} \times 10^3\) |
|-----------------|-----------------------------|-------------------------|----------------------------------|----------------------------------|-----------------------------|
| 160             | 38.1                        | 11.0                    | 207.1                            | 206.6                            | 14.9                        |
| 240             | 57.1                        | 14.2                    | 229.6                            | 231.2                            | 17.1                        |
| 320             | 76.2                        | 16.5                    | 226.1                            | 227.8                            | 19.0                        |
| 380             | 90.5                        | 17.9                    | 235.6                            | 234.7                            | 20.5                        |
To analyze these measured data against benchmark values for conventionally manufactured 316LSS, the stipulations in relevant ASTM standards were examined, as given in Table 4. For instance, the maximum hardness of 95 HRB (or ~18 HRC), as specified in both ASTM A240-20a [71] and ASTM A666-15 [72] for 316LSS, has been specified to ensure the adequate efficacy of the solution annealing heat treatment to reduce excess hardness from cold working, to dissolve carbide precipitates, and/or fully transform any retained (high hardness) δ-ferrite to austenite within the through-thickness of wrought products. Representative values for hot-rolled (75.8 HRB) and annealed (73.2 HRB) 316LSS, as reported by Song et al. [73], are consistent with these ASTM standards and point to the highest hardness possible when processing 316LSS additively. However, for CF3M, the cast equivalent of wrought 316LSS, ASTM A743-21 [74], does not stipulate any hardness maximum, but a value of 179 HB (~89 HRB or ~10 HRC) after solution annealing was reported in [75]. Similarly, for additively manufactured 316LSS by powder bed processing (laser or electron beam), no requirements for hardness are specified in ASTM F3184–16 [76], but values between 76.5 HRB [77] and 25.5 HRC [78,79] have been reported and support

Figure 11. Variation in the microhardness of as-built 316LSS produced at different laser powers: (a) TD and (b) BD.
reasonably the bulk hardness values measured in the present study. These higher hardness values measured for 316LSS manufactured additively (relative to castings, for example) have been linked to the fineness of the cellular microstructure (Figure 9) that results from the rapid cooling conditions during LPB processing [68].

Table 4. Specification of the mechanical properties of 316LSS in ASTM standards.

|                | Rockwell (HRB) | Rockwell (HRC) | YS (MPa) | UTS (MPa) | EL (%) |
|----------------|----------------|----------------|----------|-----------|--------|
| ASTM A240-20   | ≤95            | 18             | ≥170     | ≥485      | ≥40    |
| Annealed [71]  |                |                |          |           |        |
| ASTM A666-15   | ≤95            | 18             | ≥170     | ≥485      | ≥40    |
| Annealed [72]  |                |                |          |           |        |
| ASTM A743-21   | N/A            | 10             | ≥205     | ≥485      | ≥30    |
| Cast CF3M [74,75] |              |                |          |           |        |
| ASTM F3184-16  | N/A            | N/A            | ≥205     | ≥515      | ≥30    |
| LPB processing [76] |            |                |          |           |        |

*a* Converted from chart and/or calculator in [79]. N/A = Not available.

Also, several studies have examined the microhardness of as-built 316LSS to understand the effect of anisotropy, process parameters, and density. For instance, Kamarlah et al. [80] investigated different orientations (0°, 45°, and 90°) in 316LSS produced by selective laser melting (SLM) using a layer thickness of 30 µm on an SLM 125 HL system and reported comparably similar microhardness values, ranging from 209 HV0.5 to 212 HV0.5, which indicated low anisotropy in the as-built condition. Tolosa et al. [81] also explored the effect of the material anisotropy and oriented the built samples (SLM 250 Realizer) in every possible direction to examine the impact on mechanical properties; though the individual values ranged from 215 HV to 255 HV, no trend with orientation was identified and a mean value of 235 HV was reported. Cherry et al. [8] analyzed the effect of E density on the porosity and hardness of 316LSS produced on a Renishaw AM250 system and reported increasing hardness with decreasing porosity until a maximum of 225 HV10 at 125 J·mm−3. Sun et al. [82] investigated the influence of ν and hS on the SLM (250 HL) of 316LSS at a fixed E density (~104.5 J·mm−3) with an hL of 50 µm. The microhardness of their as-built near-full-density parts was between 213 HV1 and 220 HV1. The research of Tucho et al. [65] showed strong influences of E density on the porosity and they reported the lowest (176 ± 9 HV5) and highest (213 ± 3 HV5) hardness values for the 316LSS produced (using SLM 280 HL) at 50 J·mm−3 and 80 J·mm−3, respectively. Recently, Eliasu et al. [83] investigated the AM of 316LSS on an EOS M280 system (hL = 30 µm) by varying the E density from 20.4 to 142.9 J·mm−3 to examine the influence of porosity on hardness and reported maximum values ranging from 230 HV0.3 to 240 HV0.3 for the highest relative densities (91–93%) realized between 44.44 and 111.11 J·mm−3. Thus, the role of the E density on the part density and, in turn, the hardness of 316LSS that was observed in the present study, concurs reasonably well with previously reported findings for SLM and LPB processes. Furthermore, the similar Vickers hardness of near-full-density 316LSS (226.1 HV0.5 to 235.6 HV0.5) in the TD and (227.8 HV0.5 to 234.7 HV0.5) in the BD produced by A/SM with the Matsuura LUMEX Avance-25 technology also agrees with the values and low anisotropy reported for the stand-alone LPB AM of 316LSS, as tabulated in Table 5. One noteworthy exception is the reported highest value by Saedi et al. [64] of 325 HV1, which was attributed to a fine cellular size and high amounts of amorphous nano-scale silicate inclusions in their 316LSS builds [84,85].
Table 5. As-built, room temperature properties of 316LSS using stand-alone LPB AM.

| AM Systems | Hardness  | YS (MPa)  | UTS (MPa) | EL (%) | YM (GPa) |
|------------|-----------|-----------|-----------|--------|----------|
| Renishaw, h₂=50 µm Datasheet [86] | 198 ± 8 HV₀,₅(H) | 547 ± 3 (H) | 676 ± 2 (H) | 43 ± 2 (H) | 197 ± 4 (H) |
| EOS M270, h₂=50 µm [56] | 228 ± 4 HV₁ (H) | 494 ± 14 (V) | 624 ± 17 (V) | 35 ± 8 (V) | 190 ± 10 (V) |
| Renishaw AM250, h₂=50 µm [39] | 219–239 ± 5 HV₁ (V) | 487 ± 3 (V) | 594 ± 4 (V) | 49 ± 4 (V) | |
| Renishaw AM250, h₂=50 µm [39] | 554 ± 5 (H) | N/A (V) | 685 ± 5 (H) | 36 ± 3.2 (H) | 25.7 ± 12.2 (V) |
| Renishaw AM125, h₂=50 µm [87] | 215 ± 10 HV₁ (H) | 539 ± 3 (H) | 600 ± 3 (H) | 28 ± 0.5 (H) | |
| EOS M100, h₂=20 µm FlexLine [88] | N/A | 535 (H) | 650 (H) | 35 (H) | 45 (V) |
| EOSINT M280, h₂=20 µm [89] | 85 HRB | 530 ± 60 (H) | 640 ± 50 (H) | 40 ± 15 (H) | 185 (H) |
| EOS M290, TRL 8 h₂=20 µm [89,90] | 530 ± 20 (H) | 640 ± 20 (H) | 540 ± 20 (H) | 54 (V) | |
| EOS M290, TRL 7 FlexLine, h₂=40 µm [90] | 540 ± 50 (H) | 640 ± 50 (H) | 570 ± 50 (H) | 51 ± 9 (V) | |
| EOS M400-4, TRL 7 FlexLine, h₂=40 µm [90,91] | 90 HRB | 550 ± 10 (H) | 650 ± 10 (H) | 40 ± 2 (H) | 45 ± 8 (V) |
| EOS M270, h₂=20 µm [64] | 325 HV₁ | 456 ± 17 | 703 ± 8 | ~48 | |
| EOS M270, h₂=20 µm [92] | 493 ± 11 (H) | 636 ± 15 (H) | 55.8 ± 3.0 (45°) | 39.2 ± 15.8 (V) | |
| EOS M290 [93] | ~625 (H) | ~750 (H) | ~54% (H) | |
| SLM Solutions, h₂=30 µm [94] | 209 ± 2 HV₁₀ | 519 ± 25 | 633 ± 28 | 31 ± 6 | 184 ± 20 |
| SLM Solutions 125 HL, h₂=30 µm [62] | ~229 HV₃ (H) | ~490 ± 5 | ~640 ± 10 | ~25 ± 2 | |
| SLM 250 Realizer, h₂=30 µm [81] | 669 ± 12 (H) | 667 ± 12 (H) | 30.0 ± 2.9 (H) | 43.4 ± 1.5 (V) | |
| SLM 250 Realizer, h₂=60 µm [95] | 534 ± 5.7 (H₆) | 653 ± 3.4 (H₆) | 16.2 ± 0.8 * (H₆) | 16.6 ± 0.4 * (H₆) | |
| SLM Solutions 280 HL, h₂=30 µm [96] | 528 ± 5 (H) | 639 ± 5 (H) | 38.0 ± 0.7 (H) | 179 ± 25 (H) | |
| Concept Laser Datasheet [97] | 20 HRC | 234 ± 5 (V) | 640 ± 7 (45°) | 63 ± 5 (45°) | |
| Concept Laser Mlab-Cusing, h₂=25 µm [7] | 532 ± 3 (H) | 573 ± 6 (H) | 63 ± 5 (H) | |
| Concept Laser Mlab-Cusing, h₂=30 µm [60] | N/A | 512 ± 14 (SM-V) | 62 ± 12 (SM-V) | 20.4 ± 3 (SM-V) | |
| Concept Laser M2 [98] | 536 ± 4 (CB-V) | 668 ± 5 (CB-V) | 24.7 ± 2 (CB-V) | |
| Concept Laser M2 [98] | 488 ± 20 (CB-H) | 528 ± 7 (CB-H) | 11.6 ± 1 (CB-H) | |
| Fraunhofer open architecture [98] | 450 ± 10 (V,H) | 640 (V,H) | ~87% (V,H) | |
| Sisma MYSINT100 [63] | 210–240 HV₁ | ~505–525 (45°) | ~650 (45°) | ~40 (45°) | |
| SYNDAYA Dimetal-100, h₂=30 µm [66] | 281.6 HV₀,₁ | ~590 (H) | ~21.1 (H) | |

SM = Single-melt laser scanning; CB = checker board laser scanning; TRL = Technology readiness level; V = Vertical and H = Horizontal samples/direction. * Uniform percent elongation.
The nominal engineering stress–strain curves of the as-built 316LSS tensile samples, as given in Figure 12, were analyzed to evaluate the mechanical properties—namely YS, UTS, EL, and YM—for the four laser powers (Table 6). The mechanical response of the as-built 316LSS produced at 160 W consisted of a linear elastic region characterized by a YM of 175.1 ± 10.4 GPa with the onset of early yielding at 493.8 ± 10.5 MPa, which was followed by limited plastic deformation to a maximum stress (UTS) of 603.9 ± 9.0 MPa with premature failure occurring before necking at a total EL (strain) of 17.3%. The observed premature failure of the 316LSS built at 160 W may be attributed to its low relative density (98%) and characteristics of the internal defects, namely the lack of fusion layers with large interconnected and irregular pores that can act as initiation sites for crack formation and subsequent propagation through the microstructure. The high degree of these defects (as seen in Figures 6a and 8a) thus played a crucial role in limiting the mechanical performance of the 316LSS built at 160 W, as corroborated by the lower YS (by ~10%), UTS (by ~16%), YM (by ~21%), and EL (by ~70%), relative to the 240 W condition.

![Figure 12. Average tensile stress–strain curves for as-built 316LSS produced at laser powers of 160 W, 240 W, 320 W, and 380 W.](image)

**Table 6.** Average as-built static tensile properties of 316LSS.

| Power (W) | Infill E density (J·mm⁻³) | UTS (MPa) | STD | YS (MPa) | STD | EL (%) | STD | YM (GPa) | STD |
|-----------|---------------------------|-----------|------|----------|------|--------|------|-----------|------|
| 160       | 38.1                      | 603.9     | 9.0  | 493.8    | 10.5 | 17.3   | 1.9  | 175.1     | 10.4 |
| 240       | 57.1                      | 715.2     | 3.1  | 551.3    | 5.6  | 56.9   | 0.5  | 222.9     | 8.8  |
| 320       | 76.2                      | 714.0     | 2.0  | 554.9    | 2.4  | 53.4   | 1.0  | 223.8     | 10.4 |
| 380       | 90.5                      | 714.8     | 2.5  | 559.1    | 2.8  | 52.5   | 1.8  | 224.4     | 41.4 |

By contrast, the as-built 316LSS produced at the other three laser powers exhibited more ductile stress–strain behaviors with relatively higher values for the YS (551.3–559.1 MPa), UTS (714.0–715.2), EL (52.5%–56.9%), and YM (222.9–224.4 GPa). Considering the measured standard deviations in Table 6, the average values of these tensile properties were statistically comparable (from a t-test with an alpha level of 0.05) for the 316LSS produced at 240 W, 320 W, and 380 W (lowest p-value of 0.4629). This finding agrees with their comparable densities (Table 2), as well as their similar porosity characteristics of small, spherical, and isolated pores within the as-built 316LSS microstructure (Figure 6b–d). How-
ever, it is worth mentioning that the relative density (or porosity percentage) differences between all four laser power conditions were quite small (at $\leq 2\%$), which alludes to the more significant role of the porosity features on defining the mechanical behavior and properties of as-built 316LSS. In regards to this, it is interesting to note the slightly better UTS and EL properties of the as-built 316LSS produced at 240 W, which had the finest pores (Figure 6b), relative to the 320 W and 380 W conditions. The coarsening of the isolated pores in the 316LSS microstructure was observed to have the greatest and most significant impact on the ductility with reductions of 6% and 8%, respectively, in the EL for the 320 W and 380 W laser powers relative to the 240 W condition.

The tensile properties of the near-full-density samples produced in the present study at laser powers of 240 W, 320 W, and 380 W more than adequately met the minimum specifications stipulated in ASTM F3184–16 [76] for LPB-processed 316LSS, as given in Table 4. Moreover, the mechanical properties measured in this current work for the near-full-density samples showed good agreement with the vast tensile property data in the open literature, as tabulated in Table 5 from the results of previous research studies on the LPB AM of 316LSS using different stand-alone 3D printers. In this comparison, it is noteworthy that the overall tensile properties measured for the near-full-density 316LSS manufactured using the Matsuura Lumex Avance-25 hybrid LPB process were on the high side of the reported ranges of properties from conventional stand-alone LPB technologies. The observed differences can be attributed to the additive system, LPB process parameters ($h_L$, $h_S$, $\nu$, $d$, $P$, etc.), scan strategy, orientation, powder composition, etc. Finally, in comparing the tensile properties of 316LSS produced using hybrid (LPB) manufacturing or stand-alone LPB technologies to the equivalent wrought or cast alloys, the main issue appeared to be the ductility. As supported by the EL values in Table 5, the ductility of as-built 316LSS can fall substantially below the minimum value of 40% for the wrought equivalent alloy [71,72] and even the minimum of 30% stipulated for cast [74,75] and LPB [76] products. This is because the ductility of LPB-processed 316LSS was found to be highly sensitive to remnant porosity [99,100], which is unlike the high-strength (YS and UTS) properties that have been previously ascribed to a Hall–Petch (grain-boundary) strengthening effect of the fine cellular structure [62,95]. This corroborates the EL findings in the present study that showed a strong dependence of the ductility on the pore volume, morphology, and distribution; this points to the importance of process optimization and developing an understanding of the MPSP linkages for advancing the A/SM of 316LSS.

3.5. Fractography

The fractured surfaces of select tensile samples were observed by SEM to distinguish the microscopic features and mode of failure, as shown in Figure 13 for each laser power condition. The fractography of the 160 W sample surfaces revealed, at low magnification, layered lack of fusion defects consisting of unmelted powder particles, as well as large irregularly sized pores that were often interconnected (Figure 13a); these observations corroborate well with the macrostructural findings illustrated in Figure 6a and Figure 6a. By contrast, the bonded regions between these lack of fusion layers revealed, at higher magnification, a dimpled texture consisting of micro-voids (Figure 13b), which is indicative of ductile failure occurring in local regions where the fracture progressed through the cellular/grain boundary structure in the as-built 316LSS. These features (dimples and defects) on the fracture surface point to the potential of the as-built 316LSS for higher strength and ductility, which were limited in the 160 W tensile samples by the overriding influence of the layered defects that caused premature failure (through the layers with interconnected pores and unmelted powder) and the accompanying low mechanical performance.
Unlike the as-built 316LSS produced at 160 W, the lack of fusion defects were absent from the tensile fracture surfaces of the 240 W (Figure 13c–d), 320 W (Figure 13e–f), and 380 W (Figure 13g–h) sample conditions (due to the adequate $E_{\text{density}}$). At low magnification, the fracture surfaces of the near-full-density tensile samples produced at 240 W, 320 W, and 380 W showed ductile tear ridges with small rounded pores (Figure 13c,e,g) and, at high magnification, there was the predominant presence of small equiaxed dimples (Figure 13d,f,h), indicating that the main fracture mode was through nucleation, growth, and the coalescence of micro-voids. Though the overall features observed on the tensile fracture surfaces of the near-full-density samples were comparable, the 240 W sample exhibited the finest spherical pores relative to the 320 W and 380 W conditions, which agrees reasonably with the $\mu$CT (Figure 6) findings, macrostructural observations (Figure 8b-d), and the highest EL values. These fractographic observations in the present study for the as-built 316LSS produced using hybrid (LPB) processing corroborate well with previously reported findings for near-full-density samples built with stand-alone LPB AM [100]; specifically, low levels of remnant spherical porosity ($\leq 1\%$) bring about a ductile failure mode with EL values comparable to the wrought equivalent alloy, while remnant layered porosity (interconnected irregular pores with unmelted powder) at high levels ($\geq 2\%$) leads to premature failure with significantly lower EL. Moreover, from a microscale perspective, though it is difficult to directly relate the dimple size with ductility, the predominant presence of a healthy population of fine, uniformly distributed dimples on the tensile fracture surfaces of the near-full density 316LSS samples is typically indicative of high microstructural integrity and high plasticity [101].

4. Discussion

Due to the newness of A/SM, the research so far has been considerably limited on in-envelope LPB processing with the integrated micro-machining of 316LSS, relative to the large volume of studies and data on stand-alone LPB technologies, as highlighted by the tabulated property data in Table 5. Thus, the present study strived to advance the body of knowledge on the A/SM of 316LSS by investigating the influence of in-envelope processing (LPB processing and dry micro-machining) on the geometry, surface texture/finish, density, porosity attributes, microstructure, mechanical properties, and tensile fracture/failure modes.
For the first time, the characteristics of the manufacturer’s reference 316LSS powder were evaluated and this newly generated data provides benchmarking of the feedstock material to increase build-to-build reproducibility through appropriate monitoring and quality control. Previously, a couple of studies applied the Matsuura Lumex Avance-25 A/SM system to build 316LSS samples but explored micro-machining out-of-envelope [20,21]. Through the present research, the in-envelope machining of 316LSS was investigated for the first time and the measured surface roughness (Ra between 0.33 µm and 0.65 µm) was equivalent to stand-alone micro-milling after LPB printing (minimized Ra between 0.55 µm and 0.65 µm) [21], which is encouraging for advancing a synergistic manufacturing approach that takes advantage of the complementary elements of additive and subtractive processing.

Moreover, Mutua [23] studied the LPB processing of 316LSS with the Matsuura Lumex Avance-25 A/SM system and built samples using different conditions, namely \( p = 100–400 \text{ W} \), \( \nu = 400–1000 \text{ mm} \cdot \text{s}^{-1} \), \( h_S = 0.025–0.2 \text{ mm} \), and \( d = 0.05–0.3 \text{ mm} \). Their design of experiments (DOE) approach identified a process map [23] with a wide optimum window for the laser power (100 W to 320 W) at fixed values for \( \nu \) (700 mm \cdot s\(^{-1}\)), \( h_S \) (0.12 mm), \( d \) (0.2 mm), and \( h_L \) (0.05 mm). However, the robustness of this optimum window remained uncertain, as the basis for its validation rested on a single experimental point at 320 W [23]. In view of these findings, the present research study explored the robustness of the LPB process by systematically varying the laser power (from 160 W to 380 W) with all the other parameters fixed—\( \nu \) (700 mm \cdot s\(^{-1}\)), \( h_S \) (0.12 mm), \( d \) (0.2 mm), and \( h_L \) (0.05 mm)—to match the validation point from the study by Mutua [23]. The measured roughness values (Ra of 11.12 µm—11.73 µm) in the present study for the as-built surfaces of 316LSS corroborated well with the reported Ra values of 9.95 µm by Mutua [23] and 14.05 µm by Avegnon et al. [20]. It is noteworthy here that the roughness after the in-envelope machining of the as-built surfaces could be reduced by more than an order of magnitude, which provides flexibility for selecting LPB parameters that are suitable/robust for maximized performance. In this regard, the evolution in the density and porosity with increasing laser power clearly indicated that the 160 W (\( E_{\text{density}} = 38.1 \text{ J} \cdot \text{mm}^{-3} \)) condition was insufficient for achieving near-full density in 316LSS, and (for the first time) the impact on the mechanical performance was evaluated through tensile testing and fractographic analysis. At 160 W, the tensile properties of as-built 316LSS were compromised by the layered balling and lack of fusion defects (consisting of irregularly-shaped interconnected pores and unmelted powder) that caused premature failure. It is noteworthy here that the strength properties (YS = 493.8 ± 10.5 MPa and UTS = 603.9 ± 9.0 MPa) met the minimum specifications for the YS (205 MPa) and UTS (515 MPa) in ASTM F3184—16 [76], but the measured EL (17.3%) was far below the specified value of 30%. Thus, the lower limit for the laser power was identified to be 240 W for achieving near-full density (with only small spherical pores), a ductile mechanical response, and tensile properties that met all the minimum requirements in ASTM F3184—16 [76]. The upper limit may be defined by the coarsening of the remnant spherical pores in the near-full-density microstructure of the as-built 316LSS, which was noted for the 320 W condition and, more so, at 380 W, with the concomitant impact of statistically significant reductions in the EL by, respectively, 6% and 8%, relative to the 240 W condition. However, considering the tensile property specifications in ASTM F3184—16 [76], the 240 W, 320 W, and 380 W conditions more than adequately surpassed the minimum requirements for YS, UTS, and EL, though the finer pores for the 240 W laser power may prove to be more advantageous under cyclic loading, an important area for future research in the A/SM of 316LSS. Thus, based on the MPSP linkages explored in the present study, a more robust optimum process window could be identified for the A/SM of 316LSS with laser powers ranging from 240 W (\( E_{\text{density}} = 57.1 \text{ J} \cdot \text{mm}^{-3} \)) to 380 W (\( E_{\text{density}} = 90.5 \text{ J} \cdot \text{mm}^{-3} \)). The in-depth analyses performed showed (for the first time) the relationship between the laser power and density/porosity evolution for the A/SM of 316LSS powder, especially with regard to the understanding of their impacts on the mechanical response, failure mode, and material allowable (property) data.
5. Conclusions

The current study examined the characteristics and properties of 316LSS additively manufactured using laser powder bed (LPB) processing in-envelope with micro-machining. Based on the investigations conducted in this research, the following conclusions can be drawn:

1. The inspection of the part dimensions indicated that the overall deviations (with respect to the CAD model) were lower than 280 µm and 10 µm for the as-built and machined surfaces, respectively, which indicates the potential for high accuracy with in-envelope additive/subtractive processing.

2. The linear and areal surface roughness parameters on the as-built vertical side-walls had average Ra and Sa values of 11.4 µm and 14.9 µm, respectively. After in-envelope machining, the Ra (0.33–0.65 µm) and Sa (0.35–0.70 µm) values on the machined surfaces were more than an order of magnitude lower relative to the as-built surfaces.

3. The density and porosity characteristics—examined by Archimedes, helium gas pycnometry, and X-ray µCT methods—showed the presence of a lack of fusion layers and large interconnected pores with irregular morphologies for the 160 W condition, which exhibited the lowest relative density of 98.0%. Near-full density, with the porosity transformed to isolated spherical pores, was possible at the higher laser powers, and the measured relative densities for 240 W (99.5%), 320 W (99.3%), and 380 W (99.0%) were statistically equivalent. LPB additive manufacturing at 240 W and 320 W showed improved densification, with the latter showing coarser pores (up to 60 µm) relative to the former. Further increases in the laser power to 380 W increased the volume of coarser pores (up to 100 µm), likely due to excessive energy input that promoted evaporation, the development of gas bubbles, and the eventual coalescence of many small pores into fewer larger pores. The relative densities measured for 240 W, 320 W, and 380 W were statistically equivalent and ranged from 99.0% to 99.5%.

4. The as-built mechanical properties measured for the 160 W condition were low in hardness (11.0 ± 0.5 HRC and 206.9 ± 17.0 HV0.5) and, though the yield (493.8 MPa) and ultimate tensile (603.9 MPa) strengths met the minimum requirements of 205 MPa and 515 MPa, respectively, in ASTM F3184—16, the measured elongation (17.3%) was far below the specified value of 30%. The higher laser powers (240 W, 320 W, and 380 W) that led to near-full density also resulted in tensile properties that more than adequately met the requirements stipulated in ASTM F3184—16 and were comparable to 316LSS produced using stand-alone LPB processing.

5. Fractographic observations indicated a change in the fracture mode from failure through the layered lack of fusion defects and interconnected porous network (with unmelted powder particles) in the 160 W sample to ductile failure by micro-void formation, growth, and coalescence for the 240 W, 320 W, and 380 W samples.

6. This study of the hybrid additive/subtractive manufacturing of 316LSS using a Matsura LUMEX Avance-25 system has identified important linkages between the in-envelope processes (LPB and/or high-speed micro-machining) and the geometry, surface finish, density, porosity, microstructure, tensile properties, and fracture mode, which has led to the definition of a robust operating window for reaching high mechanical performance and near-full density, alongside typical surface finishes and tolerances of traditional subtractive manufacturing technologies.

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**Abbreviations**

| Symbol | Abbreviation | Definition |
|--------|--------------|------------|
| 2D     | Two-dimensional |            |
| 3D     | Three-dimensional |            |
| 316LSS | 316 L stainless steel |            |
| AM     | Additive manufacturing |            |
| A/SM   | Additive/subtractive manufacturing |            |
| BD     | Build direction |            |
| CAD    | Computer-aided design |            |
| CRM    | Critical raw material |            |
| CT     | Computed tomography |            |
| d      | Spot diameter |            |
| DOE    | Design of experiments |            |
| EDM    | Electro-discharge machining |            |
| EL     | Elongation |            |
| E\text{density} | Energy density |            |
| GD&T   | Geometric dimensioning & tolerancing |            |
| h\text{L} | Layer height |            |
| h\text{S} | Hatch spacing |            |
| HRC    | Hardness Rockwell C-scale |            |
| HV\text{0.5} | Vickers hardness at 500 g load |            |
| KGT    | Kurz–Giovanola–Trivedi (KGT) |            |
| LPB    | Laser powder bed |            |
| µ      | Micro |            |
| MPSP   | Material–process–structure–property |            |
| P      | Power |            |
| Ra     | Arithmetic average height (line) |            |
| RPM    | Revolutions per minute |            |
| Rq     | Root mean square height (line) |            |
| Rz     | Maximum height (line) |            |
| Sa     | Arithmetic average height (surface) |            |
| SEM    | Scanning electron microscope |            |
| SLM    | Selective laser melting |            |
| Sq     | Root mean square height (surface) |            |
| STD    | Standard deviation |            |
| Sz     | Maximum height (surface) |            |
| TD     | Transverse direction |            |
| UTS    | Ultimate tensile strength |            |
| ν      | Scanning speed |            |
| λ\text{c} | cut-off (wavelength) |            |
| XRD    | X-ray diffraction |            |
| YS     | Yield strength |            |
| YM     | Young’s modulus |            |
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