Recrystallization in non-conventional microstructures of 316L stainless steel produced via laser powder-bed fusion: effect of particle coarsening kinetics

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ABSTRACT
Alloys processed by laser powder-bed fusion show distinct microstructures composed of dislocation cells, dispersed nanoparticles, and columnar grains. Upon post-build annealing, such alloys show sluggish recrystallization kinetics compared to the conventionally processed counterpart. To understand this behavior, AISI 316L stainless steel samples were constructed using the island scan strategy. Rhodonite-like (MnSiO3) nanoparticles and dislocation cells are found within weakly-textured grains in the as-built condition. Upon isothermal annealing at 1150 °C (up to 2880 min), the nucleation of recrystallization occurs along the center of the melt pool, where nuclei sites, high stored elastic energy, and local large misorientation are found in the as-built condition. The low value of the Avrami coefficient (n = 1.16) can be explained based on the non-random distribution of nucleation sites. The local interaction of the recrystallization front with nanoparticles speeds up their coarsening causing the decrease of the Zener-Smith pinning force. This allows the progression of recrystallization in LPBF alloys, although sluggish. These results allow us to understand the progress of recrystallization in LPBF 316L stainless steel, shedding light on the nucleation mechanisms and on the competition between driving and dragging pressures in non-conventional microstructures. They also help to understand the most relevant microstructural aspects applicable for tuning microstructures and designing new LPBF alloys.
Introduction

Additive manufacturing (AM) consists of the repeated deposition and melting of a feedstock in a layer-by-layer iterative process until the final part is achieved. In this process, a broad range of metallic materials has been studied and applied in the industry. Among the existing AM techniques for such materials, laser powder-bed fusion (LPBF) has been attracting great interest due to the possibility of printing near-net-shape parts with complex geometries [1–3]. By adjusting the laser beam power [4], the powder layer thickness and the scan strategy [5, 6], microstructures can be tailored to achieve enhanced properties.

The engineered microstructures of LPBF materials are unique [7–9] compared to their wrought counterparts. At the mesoscopic level, they consist of several melt pools with a size of tens of micrometers containing columnar grains where regular dislocation cell structures are embedded. Another general feature is the presence of an unintended dispersion of nanoparticles in the matrix, most oxides, as reported in several metal systems [10–14].

An interesting phenomenon to further tailor LPBF microstructures is recrystallization, which is obtained by post-processing annealing. The driving force for recrystallization is the elastic energy stored in the dislocation cell walls formed due to the elevated thermal strains imposed by the high cooling rates and the subsequent alternating heating and cooling cycles [3, 15–22]. Post-build annealing behavior of LPBF-processed materials has been reported in the literature [3, 14, 23–38], where the final microstructure or the modification of mechanical or electrochemical properties following annealing have been investigated.

While texture evolution is fairly well understood during LPBF processing [39–41], only a few works have addressed the mesoscale texture evolution [26, 42–44] and the texture evolution [29, 42, 45] in...
LPBF-processed materials upon annealing. Therefore, key aspects such as those determining the underlying nucleation mechanisms, the competition between driving and dragging pressures, and recrystallization kinetics have been rarely reported. Most works cited above refer to the AISI 316L stainless steel, while the remaining ones refer to Ni-based superalloys. Only two works focus on the recrystallization mechanisms in AISI 316L and IN718 alloys in detail [45, 46], where their sluggish kinetics have been assigned to the interaction of high angle boundaries with oxide/silicate nanoparticles. Therefore, further investigations are required to understand whether the microstructure evolution and the recrystallization mechanisms are similar between samples from different alloys or obtained under different LPBF processing conditions.

In the present work, the aforementioned gaps in the literature were addressed to understand the recrystallization kinetics in LPBF alloys. In particular, this work quantifies nanoparticle coarsening due to their interaction with the recrystallization front and its effect on the sluggish recrystallization kinetics in LPBF AISI 316L stainless steel. To this end, the classical Johnson–Mehl–Avrami-Kolmogorov (JMAK) model was applied to estimate the recrystallization kinetics (Avrami exponent). The underlying recrystallization mechanisms in metallic parts produced by LPBF additive manufacturing are revealed, allowing to tune the microstructure during or after the building process.

**Experimental**

**Material**

The starting material is an AISI 316L austenitic stainless-steel CL-20 reused powder batch with spherical particles produced by Concept Laser GmbH. The mean particle size was 26 μm. The chemical composition of AISI 316L steel in the as-built condition is shown in Table 1. Bulk samples were manufactured as cylinders (φ15 mm × 50 mm) using a Mlab Cusing R machine under nitrogen atmosphere. The parts were produced using the island scan strategy where the layer is divided into smaller squares of size 5 × 5 mm² and the laser scanning direction rotates 90° between both every island and layer, as depicted in Fig. 1. The main processing parameters used in LPBF were the following: laser power (P) of 90 W, layer thickness (t) of 30 μm, hatch distance (dH) of 90 μm, and scanning velocity (v) 1500 mm/s. Further details about this manufacturing protocol can be found in a preceding work [47].

**Annealing and microstructural characterization**

The LPBF samples were isothermally annealed in vacuum-sealed quartz tubes at 1150 °C from 5 min up to 2880 min (48 h), followed by water cooling. Thermodynamic simulations were performed using the software Thermo-Calc™ coupled with the TCFE-9 database to determine the phase stability diagram of this steel, thus helping us to understand the formation of silicate/oxide nanoparticles from the liquid and to select the optimal annealing temperature within the austenite single-phase field.

Metallographic sections taken parallel and perpendicular to the build direction (BD) were prepared by conventional metallography. Electrolytic etching was performed in an aqueous 10% oxalic acid solution at 5.0 V for 90 s for the acquisition of images in a Leica DM-IRM light optical microscope. Electron channeling contrast imaging was performed in a Zeiss Merlin scanning electron microscope (SEM) operating at 30 kV. EBSD maps were acquired on both the build plane (SD1 × SD2) and along a section parallel to the BD of the as-built samples, whereas only the SD1 × SD2 plane was analyzed on annealed samples. Here, SD stands for the scanning direction, as shown in Fig. 1. For this purpose, we used a JEOL JSM-6500F SEM, coupled with a DigiView EBSD detector from EDAX/Ametek with a step size of 700 nm and an accelerating voltage of 15 kV at a working distance of 18 mm. All pixels with confidence index (CI) below 0.1 were removed from the dataset. Grain orientation spread (GOS) maps were used to determine the recrystallized grains assuming

| Table 1 Chemical composition of AISI 316L stainless steel cylinders processed by LPBF in the present investigation (in wt%) |
|---|---|---|---|---|---|---|---|
| C  | Cr  | Ni  | Mo  | Mn  | Si  | N  |
| 0.0233 | 16.5819 | 10.066 | 2.0846 | 1.3028 | 0.4965 | 0.09 |
| O  | V  | W  | Nb  | Cu  | Co  | Fe  |
| 0.07 | 0.0724 | 0.0592 | 0.0191 | 0.3065 | 0.1772 | Bal. |
a GOS ≤ 1° criterion, considering grain boundaries as interfaces with misorientation higher than 5° and a minimum recrystallized grain size of 2 μm. Kernel average misorientation (KAM) maps were constructed using the 3rd nearest neighbor with a 5° tolerance angle to reveal the misorientation gradients developed within the microstructures after the LPBF process as well as their changes upon annealing. Contoured pole figures (PF) were calculated using a discrete binning method.

Vickers hardness testing was performed using a load of 200 g for 30 s parallel to the scanning direction (SD1 x SD2) in a total number of 15 individual measurements. X-ray diffraction (XRD) measurements were conducted in the as-built specimen on the SD1 x SD2 section using a Seifert Type ID3003 diffractometer equipped with a Co-Kα radiation source and operated at 40 kV and 30 mA. The sample was measured under rotation around the build direction (BD) under an angular speed of π s⁻¹. The diffracted data were acquired in the 2θ range between 30 and 120° with angular steps of 0.03° and a dwell time of 10 s. The obtained data were analyzed by means of a Rietveld-based refinement methodology using the software MAUD (Materials Analysis Using Diffraction) [48, 49]. The Popa model for size-strain analysis was also used to evaluate the effective crystallite size ($D_{\text{eff}}$) and the accumulated microstrain $<\varepsilon_{hh}^2>$ in austenite [50]. The Popa method is implemented in the software MAUD and the reliability of the refinements was established by the goodness of fit (GoF) parameter, which must converge to the unity for an optimal fit [51, 52].

The overall dislocation density ($\rho$) was estimated according to the procedure proposed by Williamson and Smallman [53] using Eq. 1:

$$\rho = \left(\frac{3 \cdot K \cdot <\varepsilon_{hh}^2>}{D_{\text{eff}}^2 \cdot b^2}\right)^{1/2} \tag{1}$$

where $K = 16.6$ for austenite (fcc) [53]; $b$ is the magnitude of the Burgers vector, which is equivalent to $a/\sqrt{2}$ along the <110> directions of the fcc structure, where $a$ is the lattice parameter. $D_{\text{eff}}$ and $<\varepsilon_{hh}^2>$ are, respectively, the effective crystallite size and the accumulated microstrain.

The size distribution of the nanoparticles was determined using an image analysis routine based on SEM micrographs and the Image J software. A total number of about 25,000 particles was considered to determine their size and volume fraction. Porosity was determined using the same image analysis procedure for light optical micrographs collected from the unetched material.

**Results and discussion**

**As-built condition**

Samples are dense and have a porosity of 0.28 ± 0.15%. We employed a combinatorial approach using the stability phase diagram calculation and
microstructural observations to understand the solidification path of LBPF 316L stainless steel and the resulting microstructure. Special focus was placed on the particle formation.

The stability phase diagram shown in Fig. 2a reveals a miscibility gap between two liquids at high temperatures; i.e., a Fe-enriched portion (namely LIQUID in Fig. 2a) that solidifies into austenite at 1450 °C, and a second one that is enriched in Si and O (namely LIQUID #2 in Fig. 2a), which solidifies as manganese silicate nanoparticles at 1600 °C. The formation of these amorphous manganese silicate in the powder feedstock occurs in the precursor gas atomization process [54]. In the LPBF process, these nanoparticles dissociate and re-nucleate during the fast solidification, further reducing their size.

The formation of these silicate nanoparticles in the precursor powder occurs due to the high affinity of silicon and manganese with oxygen. The oxygen concentration is further increased upon LPBF due to the residual concentration of oxygen in the processing chamber [54]. The fast spinodal decomposition kinetics associated with these liquids allows their separation, even under the high cooling/solidification rates imposed by the gas atomization/LPBF process. However, the degree of separation of these liquids is dependent on the cooling rate [55]. In our case, the Si–O-enriched liquid solidifies first into spherical particles whose morphology is explained by the decrease of surface area prior to solidification. The high solidification rates in gas atomization process and LPBF lead to short time for liquid separation, yielding amorphous and nanosized particles [23, 56–58]. Thus, under such solidification conditions, one may expect that the equilibrium manganese chromate spinel (MnCr2O4) predicted in Fig. 2a would not be readily formed. Instead, the non-equilibrium amorphous rhodonite-like silicate (MnSiO3) is the most plausible stoichiometry for the particles visualized in our steel [14]. The Fe-enriched liquid portion undergoes dynamic supercooling due to the high cooling rates [49], leading its solidification to start and finish within the temperatures of the austenite stability field. Other minor phases such as M23C6, Cr7N3, and sigma (σ) are also predicted to occur in the equilibrium phase diagram of Fig. 2. However, despite being thermodynamically predicted, they were not experimentally observed in this work because of the high cooling rates employed during the process, thus hindering their formation. MnS is a detrimental inclusion observed in conventionally processed AISI 316L [59]. They are neither thermodynamically predicted nor experimentally observed in this study and they were also not reported in the work by Kong et al. [36].
To assess the relative chemical composition of the nanoparticles in our steel, we monitored their stability by tracking their qualitative chemical composition changes after isothermal annealing for 48 h at 1150 °C using EDS as reported in Fig. 2b. For this purpose, a coarse particle was chosen to minimize the iteration of the primary electron beam with the matrix during EDS mapping. Figure 2b shows that the interior of this particle is enriched in Mn and Si, resembling the rhodonite (MnSiO$_3$) constituent predicted in Fig. 2a, similarly to the reported in the literature for the as-built 316L steel [11, 23, 56, 57, 60, 61]. Yet, Fig. 2b reveals a diffusion front enriched in Cr and O that partially consumes the original manganese silicate from its surface towards the interior. This result suggests the ongoing transformation of the amorphous MnSiO$_3$ into the corresponding stable crystalline spinel (MnCr$_2$O$_4$) variant, as predicted in Fig. 2a. However, the formation of other phases cannot be discarded, and only high-resolution transmission electron microscopy (HRTEM) can clarify this issue. Therefore, we have chosen to refer to these second-phase particles as (nano)particles when discussing about the annealed microstructure, as we are unable to unambiguously identify their structure. The particle volume percentage was determined as 0.25 ± 0.11% using quantitative image analysis and fairly agrees with both, the predicted percentage of 0.4% shown in Fig. 2b and the results reported by Chao et al. [62].

Figure 3a–d show the as-built microstructures visualized in different construction sections, i.e., parallel and perpendicular to the BD. The top view (SD1 × SD2) is shown in Fig. 3a–b, where a cross-like structure (chessboard pattern) is noticed due to the 90° rotation after every built layer [63–65]. In contrast, stacks of melt pools resembling “fish scales” [10, 66] are observed on the BD × SD1 section (Fig. 3c, d). The schematic sketch depicted in Fig. 3e shows how this peculiar microstructure forms in top and side views.

Coarser columnar grains grow inward from the sides of the melt pool, while finer columnar grains grow vertically in the melt pool center via epitaxial growth, due to the thermal gradient direction [40]. These square-shaped coarse-grained regions lying in between the path of the laser tracks are called “pockets” hereafter. The white circles in Fig. 3b and d highlight the presence of a cellular substructure. Figure 4 shows an enlarged view of the microstructure.

Transmission electron microscopy studies showed that these dislocation cells have typical misorientations ranging from 0.5 to 1° in 316L stainless steel [10] and 0.2–0.8° in Hastelloy X [67] LPBF alloys. The formation of such solidification substructure is facilitated by ultra-high cooling rates that vary between 10$^5$ and 10$^6$ K/s [68, 69] together with high-temperature gradients (e.g., in the order of 10$^6$ K/m) [66]. The presence of a large number of nanoparticles dispersed in the matrix of this material is readily observed as indicated by the yellow arrows in Fig. 4. Fifteen ECCI-SEM images were used to calculate the average size of the cells and the particle size distribution in the as-built specimen. The average dislocation cell size found was 383 ± 40 nm and the mean rhodonite-like particle size in this condition is 37 ± 19 nm. The dislocation density of the as-built steel was estimated, according to the protocol described in Section “Annealing and microstructural characterization”, as being ~1.5 × 10$^{14}$ m$^{-2}$ (see Table 2). The literature reports that a rough estimate of $\rho \sim 4 \times 10^{14}$ m$^{-2}$ can be assumed for LPBF materials having dislocation cells with sizes of ~400 nm, which has a good agreement with our findings [70].

Figure 5a–c show the EBSD top view maps (SD1 × SD2) of the sample, i.e., the maps were acquired from sections perpendicular to the build direction. Just as a pictorial remark, these engineered microstructures depicted as EBSD inverse pole figure (IPF) maps have geometric forms that very much resemble those found in some of the colorful paintings portrayed by F. Hundertwasser (1928–2000) and W. W. Kandinsky (1866–1944). The inverse pole figure (IPF) presented in Fig. 5a shows the grain orientations along the tracks and in the interior of the pockets. Small grains with sizes below 10 μm are found in the tracks, whereas in the pockets they are much larger (50–60 μm). The difference in grain size between tracks and pockets is also observed in Fig. 3b, where fine grains are observed at the center of the laser tracks, while coarser square-shaped grains are observed between the laser tracks. The resulting chessboard pattern is created from the superposition of two perpendicular linear patterns (90° rotation), where epitaxial growth ensures the partial growth of grains from the previous layer onto the next one [45]. Figure 5b shows that high angle
boundaries (black lines) and low angle boundaries (red lines) are not evenly distributed in the microstructure. There is a much larger fraction of low angle boundaries along the laser tracks (about 0.35) and much less in the interior of the pockets (about 0.05). High angle boundaries respond for 0.6 of the total number of boundaries. From the side cross-section in Fig. 6a, one can observe that, although exhibited as equiaxed grains in the SD1 x SD2 cross-section, almost no equiaxed grains are visible in a 3D perspective. Rather, grains are columnar and grow toward the BD.

KAM maps are useful to estimate the elastic stored energy in the material in a qualitative manner [71]. Low-misorientation dislocation cells like those depicted in Fig. 4 can be found throughout the microstructure (tracks and pockets); however, low angle boundaries with larger misorientations (2°–15° interval) are more numerous along the tracks, leading to a local increase of stored energy (Fig. 5c).
Regions with locally fairly low dislocation densities inside of heterogeneously deformed microstructures can also provide regions that may act as potential recrystallization nuclei. This means that such nuclei may appear both, at low or at high dislocation density regions, depending on several factors such as the processing parameters, the studied material, and the degree of heterogeneity of the deformation substructure [72]. In all cases, such nuclei must be surrounded by at least one segment of a mobile high angle boundary [72, 73], where the adjacent deformed microstructure provides a higher stored elastic energy to drive the reaction in a net direction and sustain their growth. Therefore, it is important to identify, investigate and classify the nucleation sites, their spatial distribution, and their adjacent deformation substructure. A visualization of the possible recrystallization nucleation sites in a large section of the LPBF-manufactured steel is given in Fig. 6. The former melt pool zones can be clearly recognized, and only high angle boundaries are indicated by black lines (Fig. 6c).

When applying the GOS ≤ 1° criterion, i.e., marking grains with high inner local crystalline perfection and negligible lattice curvature, possible recrystallization nucleation sites can be distinguished (Fig. 6b). However, a careful evaluation must be conducted when identifying recrystallized grains based solely on the GOS criterion, as smaller grains are known to intrinsically exhibit lower orientation spread values [73]. It must be stated that such grains are not recrystallized, as they have dislocation cells, although with a likely lower dislocation density. These local regions with lower dislocation density, yet surrounded by mobile high angle boundaries, possess the required conditions to serve as

**Table 2** Parameters obtained via both Rietveld-based refinement and Popa model for the as-built LPBF-processed AISI 316L steel. \(< D_{\text{eff}} >\) and \(< e^2_{hh} >^{1/2}\) are, respectively, the effective crystallite size and the accumulated microstrain. \(\rho\) stands for dislocation density. The goodness of fit (GoF) parameter is also displayed in this table.

| Condition | Lattice parameter (nm) | \(< D_{\text{eff}} >\) (nm) | Microstrain \(< e^2_{hh} >^{1/2}\) | \(\rho\) (m\(^{-2}\)) | GoF |
|-----------|------------------------|----------------------------|-------------------------|-----------------|-----|
| as-built  | 0.35899                | 190                       | 0.00100                 | \(1.5 \times 10^{14}\) | 1.38 |
suitable nucleation sites upon annealing. Most of these nuclei are fairly elongated towards the build direction. Their spatial distribution is also non-uniform throughout the microstructure. This fact is related to the different levels of straining in the microstructure (misorientation gradients) from the bottom to the top, as revealed by the KAM map shown in Fig. 6c. This difference in straining along BD may occur due to the variation of process conditions during building.

**Figure 5** EBSD maps of the build plane (SD1 x SD2) of AISI 316L steel manufactured by LPBF process in the as-built condition showing: a Inverse pole figure, b Image quality map indicating high angle boundaries marked by black lines and low angle boundaries in red; c KAM maps of the corresponding IPF map. EBSD: electron backscatter diffraction; LPBF: laser powder-bed fusion; KAM: kernel average misorientation; IPF: inverse pole figure.

**Figure 6** EBSD maps of LPBF 316L steel in the as-built condition in the view parallel to the build direction showing: a IPF of the entire mapped area; b GOS map for regions displaying GOS ≤ 1°, constructed to highlight microstructure sites that could act as possible recrystallization nuclei; c KAM maps of the corresponding IPF map. LPBF: laser powder-bed fusion; IPF: inverse pole figure; GOS: grain orientation spread; KAM: kernel average misorientation; BD: building direction; SD1: scanning direction 1.
Annealing behavior of as-built samples

Samples were annealed at 1150 °C for several time intervals. Based on the thermodynamic simulations displayed in Fig. 2a and on previous experiments (data not shown), this annealing temperature was chosen because only austenite and spinel (MnCr$_2$O$_4$) should be present in the microstructure in equilibrium. Lower temperatures would make the progress of recrystallization sluggish, as the grain boundary mobility follows an Arrhenius-type thermal activation behavior, whereas higher temperatures would allow fast particle coarsening (see Section “Annealing behavior of as-built samples”), grain growth, or the likely formation of δ-ferrite.

Figure 7a–d show the IPF maps of as-built and annealed samples at 1150 °C for 5, 15, and 60 min, respectively. Significant microstructural changes become noticeable only after 60 min of annealing. The initial chessboard-like pattern is replaced to some extent by a new microstructure, formed by equiaxed grains and annealing twins, as recrystallization progresses. After this period of time, only a small portion of the heat-treated microstructure preserves the initial features of the as-built condition.

The evolution of recrystallization and the identification of potential nucleation sites were investigated by constructing GOS maps from the EBSD data, as depicted in Fig. 8a–d. In these figures, the recrystallized grains and possible recrystallization nuclei were considered as those with GOS ≤ 1° [74–76] and are displayed in white along their volume percentage. We distinguish possible recrystallization nuclei and recrystallized grains based on the presence of annealing twins, as both exhibit GOS ≤ 1°. The former is present up to 5 min annealing at 1150 °C, but such grains do not exhibit annealing twins nor morphology of recrystallized grains, differently from the recrystallized grains observed at 15 min annealing (1150 °C) onwards. Enlarged views of the samples depicted in Fig. 8b and c were chosen to highlight the nucleation sites, as reproduced in Fig. 9a–d. These results clearly indicate that nucleation occurs along the laser beam tracks (Fig. 9c). These regions hold the higher local stored energy and larger misorientations (Figs. 5 and 6) and the possible recrystallization nuclei (Fig. 6b). For the same reasons, at longer annealing times, the recrystallization front progresses along the tracks (higher stored energy) and then into the pockets (Fig. 9c and d) driven by the stored energy from dislocation cells and LABs formed due to static recovery.

According to the Johnson–Mehl–Avrami-Kolmogorov (JMAK) model [72], the volume fraction of recrystallized grains ($X_v$) varies with time according to Eq. 2, where $B$ is a constant and $n$ is the Avrami exponent. This exponent depends on the material (stacking fault energy), dimensionality, and...
nucleation process (site saturation or constant nucleation rate).

\[ X_V = 1 - \exp(-Bt^n) \]  \hspace{1cm} (2)

Taking into account the values of \( X_V \) obtained from the EBSD data, and using Eq. (2), a value of \( n = 1.16 \) was found (Fig. 10). Assuming that nucleation sites are randomly distributed, that all of them are available at \( t = 0 \) s (site-saturation) and that isotropic growth occurs, the Avrami coefficient is expected to be 3. Low \( n \) values result from a non-random nucleation of recrystallization [72]. Compared to 304L grade, 316L stainless steel is less prone to undergo static recovery due to the high thermal stability resulting from Mo segregation at cell walls [21] and low-to-medium stacking fault energy of 33 mJ/m\(^2\) [77]. Therefore, the low Avrami exponent of \( n = 1.16 \), herein observed, can be explained by the preferential nucleation and growth along the build tracks, restraining their growth along the transverse/building direction, as the build tracks exhibit higher dislocation density and higher local misorientation [45]. As a result, elongated recrystallized grains are observed in Fig. 9d. The most equiaxed grains are found at the pockets’ corners, or at the regions where two perpendicular laser tracks meet. This occurs due to the fact that they are allowed to grow in 3D by consuming the build tracks along all directions.

The softening behavior and the corresponding JMAK plot are shown in Fig. 10. The hardness of the as-built specimen is 246 ± 10 HV-0.2. Similar values were also reported in refs. [14, 69, 78]. These values are considerably higher than the ones found in wrought 316L stainless steel, namely, about 165 HV [69]. This difference is consistent with the complex dislocation cell structure observed in the present specimens and the presence of nanoparticles dispersed in the matrix [14]. The results also show that most of the softening occur during the first 240 min. For longer annealing times, the hardness levels out indicating that recrystallization is completed at less than 4 h at this temperature. Compared to our preceding work [45], where the recrystallization kinetics did not follow the JMAK behavior, the present study exhibits faster recrystallization kinetics (incomplete recrystallization after annealing at 1150 °C for 8 h in our previous study). The likely reason for such differences is the higher number density of recrystallized grains in the present study.

The discontinuous character of recrystallization is confirmed by clear nucleation and growth processes (Figs. 7 and 9). Contrastingly, continuous recrystallization does not exhibit such recognizable and consecutive steps (nucleation and growth) and is usually observed in highly strained materials, mainly at high annealing temperatures, where an increased fraction
of high angle boundaries are present [72]. Upon annealing, the microstructure slightly changes in continuous recrystallization, frequently resulting in a fine-grained material. Such behavior is completely different from the response observed upon annealing for the current AISI 316L steel.

The population evolution of $\Sigma 3$-type boundaries (annealing twins), high angle (HAB), and low angle boundaries (LAB) is shown in Fig. 11. During primary recrystallization, the number of LABs decreases as the fraction of HABs steadily increases. $\Sigma 3$-type special boundaries (twins) are absent in the as-built condition. Only after 15 min, larger fractions of twins become noticeable in the microstructure, as depicted in Fig. 9c, d and Fig. 11. Although the line fraction of annealing twins evidenced is relatively large (about 48%), it did not vary in a significant manner between 240 and 2880 min. This fraction of $\Sigma 3$ boundaries after long-term annealing is significantly lower than the one reported for a grain-boundary engineered 316L stainless steel [79], where a fraction of 79% was present after 5% thickness reduction and annealing at 1027 °C for 30 min. It is known that grain-boundary-engineered alloys are subjected to low cold deformation (usually around 5% strain) followed by high-temperature annealing. In our alloy, the dislocation density is similar to a 20% tensile strained wrought-processed counterpart, which explains the reduced
Thus, the \( \Sigma 3 \) boundary fraction may be increased by controlled post-thermomechanical processing [44, 81].

**Driving and dragging pressures in recrystallization**

The driving pressure for static recrystallization (\( P_D \)) can be approximated by using Eq. 3 [82]:

\[
P_D \approx 0.5 \rho Gb^2
\]

where \( \rho \) is the dislocation density \( (1.5 \times 10^{14} \text{ m}^{-2}) \), here obtained from Table 2, \( G \) is the shear modulus and \( b \) is the magnitude of the Burgers vector. Assuming \( b = 0.2541 \text{ nm} \) for austenite and \( G = 77 \text{ GPa} \) [83], the driving pressure for recrystallization in the as-built material (Eq. 2) can be calculated as \( 3.7 \times 10^5 \text{ J/m}^3 \) or 0.37 MPa. The drag force due to Zener-Smith pinning (\( P_Z \)) for the same initial condition can be estimated using Eq. 4 [84, 85]:

\[
P_Z = -3f \gamma_{GB} / 2r
\]

where \( f \) is the particle volume fraction, \( \gamma_{GB} \) is the grain boundary energy, and \( r \) is the particle radius. Taking a mean particle size \( 2r = 37 \pm 19 \text{ nm} \), \( f = 0.25 \pm 0.11\% \), and \( \gamma_{GB} = 668 \text{ mJ/m}^2 \) [86], the calculated pinning pressure in the as-built condition is approximately 6.8 MPa. This number drops to 0.9 MPa when considering the particle size after 1-h annealing at 1150°C (viz. 170 ± 137 nm). These general estimates assume that both driving pressure (stored energy) and drag forces (particle distribution) are uniformly distributed in the microstructure. However, as the microstructure mapping revealed, this is not the present case. In qualitative terms, the stored energy was observed to be much larger along the building tracks. A rough estimate based on the
misorientation distribution data from EBSD maps shows that about 40% of the total length of boundaries have low-angle character (2°–15°) (Figs. 5 and 11). In turn, these LABs are formed by geometrically necessary dislocations (GNDs). Since over 90% of the LABs are located along the melt tracks and about less than 10% are found within the pockets, the dislocation density and, thus, the stored energy for recrystallization is much larger in these regions, Fig. 9. Furthermore, nanoparticles are not evenly dispersed in the matrix making boundary pinning uneven in the microstructure.

**Particle-boundary interaction**

As recrystallization proceeds, nanoparticles interact with high angle grain boundaries making their migration sluggish. The interaction between moving boundaries and particles enables a fast coarsening of the latter, which increases particle interspacing and causes a progressive decrease of the Zener-Smith pinning force (Eq. 4), as demonstrated in Fig. 12a, b. It is worth mentioning that these parameters (particle size and interspacing) are mean values taken from the global microstructure as a function of the annealing time.

To demonstrate that the interaction between high angle boundaries and particles speeds up Ostwald ripening, we measured the particle sizes in different regions (recrystallized grain interior, recrystallization front, and non-recrystallized grains). Figure 13a–c show SEM images depicting the evolution of the particle size with the progress of recrystallization. In Fig. 13a, Ostwald ripening [87] of nanoparticles lying at the grain boundaries of the recrystallized grains is nearly twice as large as those in non-recrystallized regions after 5-min annealing. In general, particles coarsen, but the coarsening kinetics at grain boundaries or within recrystallized grains is faster than in recovered regions (Fig. 13b–d).

This difference in terms of particle coarsening kinetics can be mostly attributed to the much higher grain boundary diffusivity (D_b) compared to the bulk one (D). Determining D_b is a very complex experimental task and far from the scope of this work, since it depends on the local boundary width (δ) and on the boundary misorientation. Assuming δ ≈ 0.5 nm, one can estimate that D_b is about four to eight orders of magnitude higher than D. Solute elements possessing high solubility in austenite speed up Ostwald ripening [88]. Thermokinetics calculations allowed to determine the diffusion coefficients and the solubility in austenite of five solute elements (Mn, Si, O, Cr, and Mo) over a wide range of temperature (900–1200 °C).

For the sake of simplicity, we only focus on the bulk diffusivity of these species. This is justified because even in non-recrystallized regions, where bulk diffusivity is active, significant particle coarsening occurs. Figure 14a shows the changes of D as a function of temperature for several alloying elements in the austenite. Oxygen is an interstitial solute in austenite and it is the fastest-diffusing species among these five elements followed by Si, Mo, Mn, and Cr, in this order. Its bulk diffusivity is about 4 orders of magnitude higher than the other species at 1150 °C.
These numbers support the experimental results concerning particle coarsening in this metal system.

Particle coarsening can be controlled by either bulk or grain boundary diffusion. According to Kirchner [89], the particle radius is proportional to $t^{1/4}$ for particles sitting at the (high-angle) grain boundaries. For low-angle grain boundaries (subgrains), where pipe diffusion is the rate-controlling transport mechanism, the dependence drops to $t^{1/5}$. Taking the experimental points depicted in Fig. 12a, our data fit the dependence on $t^{1/4}$, in good agreement with the microstructural observations, i.e., enhanced particle coarsening due to grain boundary diffusion. One straight implication of particle coarsening is the change in corrosion properties upon annealing. An improvement of corrosion properties is observed due to the formation of a thicker and more stable protective layer after annealing between 800 and 1050 °C [36, 90]. Contrastingly, a more negative corrosion potential is observed after annealing at 1200 °C up to 2 h, while a lower pitting potential was observed [26]. The reason is the formation of a less effective passivation film due to the presence of larger pores [26], likely created by the dissolution of particles during Ostwald ripening.

**Texture evolution**

The crystallographic texture of LPBF materials depends on the heat flow and local nucleation conditions [91] and it can be tailored by adjusting the main process parameters, such as the laser beam power, the adopted scanning strategy, and the temperature of the build plate [92]. Figure 15a–e show...
the contoured pole figures calculated from the EBSD mapping of the as-built condition and after annealing at 1150 °C for 5, 15, and 60 min, respectively. The scale bars are shown on the right-hand side of the pole figures for each condition. The texture intensity is expressed in terms of multiples of a random distribution (mrd). The material in the as-built condition has a weak texture (2.292 mrd) related to the Goss \{011\}% component. When either possible recrystallization nuclei or recrystallized grains (GOS \{110\}%/C176) are considered, their starting texture is also rather weak, as shown in Fig. 15e. As annealing proceeds, texture becomes random for practical purposes. The weak-to-absent texture in the as-built condition can be attributed to the low laser power (90 W) used in the manufacturing process of our samples [83], which favors nucleation of finer grains rather than strong epitaxial growth.

The laser power is the main processing parameter controlling the texture in LPBF alloys, as high laser powers lead to strong Goss \{110\}% < 100 > [BD] < SD > textures [40, 45] or even single crystal-like parts [41]. Additionally, the scanning strategy plays a key role, as the rotation between layers weakens epitaxial growth, triggering nucleation of grains with different crystallographic orientations [93] and, thus, giving rise to overall weaker crystallographic textures. Upon annealing, the recrystallization weakens the texture further through the nucleation of strain-free grains, where twinning aids in creating a higher orientation variety of the crystallographic variants [94]. Texture strengthening occurs upon annealing only when abnormal grain growth occurs, where a few grains have a growth advantage and overgrow the matrix grains [95], which is not the behavior observed in this study. As a result, the weak texture of the as-built samples is randomized upon annealing. Texture intensities between as-built and annealed LPBF alloys are directly related. A similar AISI 316L with stronger texture (11.402 mrd vs. 2.292 mrd for the present as-built sample) in the as-built condition exhibits stronger texture after annealing at 1150 °C for 1 h (3.171 mrd) [45] when compared to the annealed samples in the present investigation (1.9 mrd). These results indicate that using the island scan strategy (with rotation between layers) combined with a low laser power followed by post-build recrystallization annealing gives rise to fairly isotropic texture and microstructures.

**Comparison of recrystallization kinetics between LPBF and conventionally manufactured materials**

A straightforward comparison between the recrystallization kinetics of LPBF- and wrought-processed 316L stainless steel is not simple. Many microstructural parameters differ from one route to another depending on the processing conditions and might impact the recrystallization kinetics, such as starting grain size, texture, residual amounts of delta ferrite, strain-induced martensite (SIM) formation, chemical segregation, and so on. However, a general trend is observed when considering all the differences: the temperature to initiate the recrystallization in LPBF 316L is higher and the kinetics is more sluggish compared to the wrought manufactured counterpart. For example, recrystallization takes place at 700 °C for 1 h following 95% cold rolling reduction [94]. Complete recrystallization after 1000 °C for 30 min following 13% cold drawing is observed in another study [96]. For the steel herein investigated, incomplete recrystallization persists after annealing at 1150 °C for 4 h.

Considering all the microstructural features in wrought and LPBF 316L stainless steels, the weak texture and low strain found in the latter are apparently not the major contributors for the difference in their recrystallization kinetics, as seen in ref. [96]. Another point is the solute drag caused by
molybdenum segregation at cell walls in the as-built LPBF AISI 316L, which guarantees enhanced thermal stability [21]. Despite this factor, molybdenum segregation was not investigated in detail, since the matrix is chemically homogenized before 10 s at the investigated temperature, as observed by thermokinetic simulations using the DICTRA software [45]. Therefore, our results suggest that, in LPBF-316L stainless steel, recrystallization is mainly controlled by particle coarsening and heterogeneous nuclei distribution. Thus, LPBF alloys, which frequently contain a fine dispersion of nanoparticles, exhibit a sluggish recrystallization kinetics when compared to a wrough counterpart (for other alloys as well [46]) due to the interaction of nanoparticles with migrating grain boundaries.
Our study reports the in-depth characterization of the recrystallization kinetics and the importance of the as-built LPBF microstructure, especially nanoparticles, on this phenomenon. We quantitatively show that recrystallization is mainly controlled by particle coarsening with annealing and the concurrent decrease of the Zener-Smith pinning force. These results confirm that the yield stress and other mechanical properties can be tuned by controlling a given recrystallized volume fraction and their nucleation sites by a combination of proper process parameters (scanning strategy and laser beam power) and post-processing recrystallization annealing.

Summary and conclusions

The post-build annealing behavior of texture-free AISI 316L austenitic stainless-steel cylinders manufactured by LPBF additive manufacturing was investigated at 1150 °C. The main microstructural changes upon annealing were followed by means of several characterization techniques such as strain analysis from EBSD data (GOS and KAM maps), XRD, and quantitative metallography. Based on the main findings reported in this work, the following conclusions can be drawn:

(a) A larger fraction of low angle boundaries can be noticed along the melt tracks (center of the melt pools) compared to the pockets (sides of the melt pools). In the as-built condition, potential recrystallization nuclei lie fairly parallel to the build direction at the intersections of the melt tracks of layers $n$ and $n + 1$, i.e., regions with higher stored elastic energy.

(b) Recrystallization starts at the melt tracks because of their larger stored energy. Recrystallization proceeds along these tracks and after consuming them, migrate towards the coarse-grained areas (pockets).

(c) Recrystallization kinetics is most affected by particle coarsening. The Zener-Smith pinning pressure exerted by nanoparticles progressively decreases upon annealing due to coarsening. When particles reach a critical size, the recrystallization front is released since the driving pressure overcomes the dragging one.

(d) The interaction between the second-phase particles and the recrystallization front, where grain boundary diffusivity is the most active mass transport mechanism, allows fast Ostwald ripening. The mean particle size increases from $37 \pm 19$ nm to $170 \pm 137$ nm after 1-h annealing. Longer annealing times transform rhodonite ($\text{MnSiO}_3$) into spinel-type manganese chromate ($\text{MnCr}_2\text{O}_4$) particles, the predicted equilibrium phase.

(e) The recrystallization kinetics is particularly sluggish due to the non-random distribution of nuclei, as suggested by the low value of the Avrami exponent $n = 1.16$ and concurrent pinning effects.

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Declarations

Conflict of interest The authors declare that no conflict of interests or competing interests exists that could have influenced the work reported in the paper.

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