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Article

Effect of Heat Treatments and Related Microstructural Modifications on High-Cycle Fatigue Behavior of Powder Bed Fusion–Laser Beam-Fabricated Ti-6Al-2Sn-4Zr-6Mo Alloy

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Abstract

The study investigates the influence of microstructures on fatigue behavior and failure mechanisms of the α - β titanium alloy Ti6246, fabricated via Powder Bed Fusion-Laser Beam (PBF-LB). In particular, the investigation assesses the effect of two post-processing heat treatments, namely α - β annealing at 875 °C (AN875) and solution treatment at 825 °C followed by aging at 500 °C (STA825), on the alloy's rotating and bending fatigue behavior. The results indicate that the STA825 condition provides superior fatigue resistance (+25%) compared to AN875, due to the presence of a finer bilamellar microstructure, characterized by thinner primary α lamellae (α_p) and a more homogeneous distribution of secondary α lamellae (α_s) within the β matrix. Additionally, an investigation conducted using the Kitagawa–Takahashi (KT) approach and the El-Haddad model, based on the relationship between the fatigue limit and defect sensitivity, revealed improved crack propagation resistance from pre-existing defects (ΔK_{th}) for the STA825 condition compared to AN875. Notably, the presence of fine α_s after aging for STA825 is effective in delaying crack nucleation and propagation at early stages, while refined α_p contributes to hindering macrocrack growth. The fatigue behavior of the STA825-treated Ti6246 alloy was even superior to that of the PBF-LB-processed Ti64, representing a viable alternative for the production of high-performance components in the automotive and aerospace sectors.

Keywords: Powder Bed Fusion–Laser Beam; titanium alloys; heat treatment; microstructures; high cycle fatigue; defects



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1. Introduction

The properties of titanium alloys, including their high strength-to-weight ratio, optimal fracture toughness, and excellent high-temperature and oxidation resistance, have led to their widespread adoption in sectors such as aerospace, motorsport, and automotive [1–4]. The growing demand for high-performance components, combined with the need for complex geometries unachievable through conventional machining technologies [5], has driven the adoption of additive manufacturing (AM) [6,7]. Among AM technologies, Powder Bed Fusion–Laser Beam (PBF-LB) is the most extensively studied and widely used, as it enables the fabrication of strong, lightweight, and geometrically complex titanium components, including those with internal channels, directly from Computer-Aided models.

Currently, the α - β titanium alloy Ti-6Al-4V (Ti64) is the most widely used in AM due to the following: (i) its long-standing application in the aerospace and automotive industries; (ii) its well-documented excellent mechanical performance; (iii) its availability on the market. For these reasons, PBF-LB Ti64 has been the subject of extensive research [3,8], particularly aimed at maximizing fatigue strength, as fatigue represents the primary cause of failure in automotive and aerospace components (e.g., airframe components, connecting rods, etc.).

However, the growing demand for enhanced performance in AM-produced components has sparked significant interest in alternative α - β titanium alloys, such as Ti-6Al-2Sn-4Zr-6Mo (Ti6246), which exhibits a higher β -phase fraction and enhanced mechanical potential compared to Ti64.

For conventionally processed Ti6246, the optimization of tailored heat treatments enables the achievement of high tensile properties [9,10]. In particular, these properties are attained through solution treatment below the β transus temperature, followed by aging, which promotes the formation of a fine bimodal microstructure consisting of a globular primary α phase (α_p) in a fine α - β lamellar matrix [11]. Beyond static properties, Peters et al. [12] reported that Ti6246 exhibits superior fatigue strength compared to Ti64 in both high-cycle fatigue (HCF) and very-high-cycle fatigue (VHCF) regimes. This improvement is primarily attributed to the higher α phase fraction present in Ti64 compared to Ti6246 under similar heat treatment conditions, resulting from its higher content of α -stabilizing elements. In wrought titanium alloys, which are typically free of macroscopic defects, fatigue crack nucleation preferentially occurs within the α_p , with subsequent propagation occurring either transgranularly within the α phase (especially when coarse) or along the α/β interfaces [13,14].

To date, no specific studies have been carried out on AM processed Ti6246. Nevertheless, it is well established that the unique microstructural features induced by PBF-LB in α - β Ti alloys significantly affect both tensile and fatigue behavior [15,16]. Many researchers have observed that high cooling rates typical of these processes can lead to the formation of metastable phases, such as α' [17] and α'' martensite [6], depending on the β -stabilizer content. These metastable microstructures can negatively affect their reliability during service [18,19]. Although optimized heat treatment can increase the tensile properties of the alloys, making them comparable and even superior to those of conventionally processed alloys, similar improvements in fatigue resistance remain challenging. In fact, the high surface roughness, gas porosity, and lack of fusion (LoF) defects of PBF-LB parts generate localized stress concentrations that promote crack nucleation, significantly reducing fatigue resistance [20–22].

Leuders et al. [18] observed that for a Ti64 alloy produced by PBF-LB, α + β annealing treatment improved crack propagation resistance compared to the as-built state, due to the transformation of α' martensite into a lamellar α + β structure. However, the heat treatment had no significant effect on crack nucleation resistance, which primarily occurred near pores. Yu et al. [15] extensively studied the effect of Solution Treating and Aging (STA) heat treatment within the α + β phase field on a Ti64 alloy fabricated via PBF-LB. Their findings revealed that this treatment promotes the formation of a distinct bilamellar microstructure, consisting of α_p lamellae within a fine matrix of secondary α (α_s) lamellae and a β phase, which exhibited enhanced fatigue resistance compared to the lamellar structure resulting from α + β annealing. This highlights the critical role of heat treatment optimization in improving fatigue performance.

Although the influence of microstructures on the fatigue behavior of α - β Ti64 alloys has been widely investigated, to the best of the authors' knowledge, a research gap exists regarding the effect of heat treatment on the fatigue strength of PBF-LB Ti6246 alloys

that requires further investigations. In previous works [23,24], the authors optimized the heat treatment parameters for PBF-LB Ti6246 and established correlations between its microstructural features and tensile properties. Building on these findings, the present study aims to investigate the effects of two heat treatments, STA and $\alpha+\beta$ annealing, on the fatigue behavior of PBF-LB Ti6246. Moreover, the correlation between fatigue failure and defects in the PBF-LB Ti6246 was investigated using the Kitagawa–Takahashi (KT) approach and the El-Haddad model [25,26], aiming to assess the potential for enhancing defect tolerance in this alloy. Indeed, enhancing defect tolerance could enable the elimination of the costly and time-consuming Hot Isostatic Pressing (HIP) process, which is commonly applied to titanium PBF-LB components to reduce internal defects and improve fatigue performance.

2. Materials and Methods

2.1. Sample Production

The PBF-LB samples were produced by gas-atomized Ti6246 powders supplied by Eckart TLS (Hartenstein, Zwickau, Germany). Glow Discharge Optical Emission Spectroscopy (GD-OES, GDA-650, Spectruma Analytik GmbH, Hof, Germany) was employed to verify that the chemical composition of the samples met the required standards. The chemical composition of the supplied powders, the printed samples, and the standard UNS R56260 [27] is reported in Table 1.

Table 1. Ti6246 alloy powders' chemical composition (wt.%), as declared by the supplier and reported in the UNS R56260 standard.

Elements [% wt.]	Ti	Al	Sn	Zr	Mo	Fe	N	O	H	Others
Supplier composition	Bal.	5.96	1.85	3.67	5.89	0.03	0.01	0.12	0.002	Total max < 0.30
Sample GD-OES	Bal.	6.36 ± 0.13	2.06 ± 0.08	4.21 ± 0.09	5.85 ± 0.14	0.14 ± 0.01	-	-	-	Total: 0.20 ± 0.06
Standard composition	Bal.	5.5 ÷ 6.5	1.75 ÷ 2.25	3.5 ÷ 4.5	5.5 ÷ 6.5	max 0.15	max. 0.04	max. 0.15	max. 0.0125	Each max 0.10 Total max 0.40

An analysis of the powder's particle size distribution was conducted, showing D10, D50, and D90 values of 24.1, 38.0, and 54.7 μm , respectively. Cylindrical samples (120 mm in height and 12 mm in diameter) were made using an EOS M270 Dual Mode (EOS GmbH, Krailling, Germany) 3D metal laser sintering system under a high-purity argon atmosphere. The system is equipped with a continuous Nd:YAG fiber laser capable of delivering up to 200 W of power, with a laser spot diameter of 100 μm . The PBF-LB process parameters, summarized in Table 2, were defined based on a previous study [6].

Table 2. PBF-LB scanning parameters for Ti6246 samples production.

Power [W]	Scanning Speed [mm/s]	Hatch Distance [μm]	Layer Thickness [μm]	Scanning Strategy	Platform Temperature [$^{\circ}\text{C}$]
190	1250	100	30	Standard 67° EOS strategy	100

The axisymmetric samples were fabricated with their axes oriented perpendicular to the pre-heated build platform, an industrial practice that minimizes the required support structure. To prevent distortions associated with mechanical removal, cylindrical specimens were detached from the build plate using electrical discharge machining. Based on previous

studies, the specimens underwent two distinct heat treatments. The first treatment, optimized by Carrozza et al. [24], was an annealing process conducted in a Pro.Ba VF8000/S high-vacuum furnace (TAV Vacuum Furnaces S.p.A, Caravaggio, Bergamo, Italy) within the $\alpha+\beta$ phase field at 875 °C for 2 h, followed by slow furnace cooling at a controlled rate of 5 °C/min. This treatment aimed to enhance the alloy's ductility and toughness. The second treatment, optimized by the authors [23], was an STA process performed in a Nabertherm LT 9/14 muffle furnace (Nabertherm GmbH, Lilienthal, Germany). It involved solution treatment at 825 °C for 2 h, rapid water quenching, and subsequent aging at 500 °C for 3 h to achieve high strength. For clarity, these heat treatments will hereafter be referred to as AN875 and STA825, respectively (Figure 1). Table 3 summarizes the tensile test data, in terms of Yield Strength (YS), ultimate tensile strength (UTS), and elongation to failure (EL), carried out on the alloy and reported in [23]. These data were averaged over three samples for each condition.

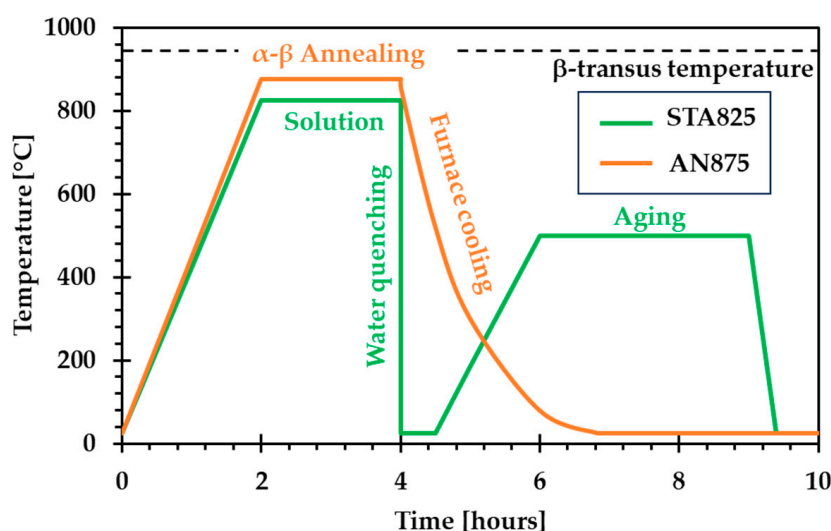


Figure 1. Representation of STA825 (green) and AN875 (orange) thermal cycles.

Table 3. A summary of tensile properties for machined tensile specimens subjected to AN875 and STA825 treatments (Adapted from ref. [23]).

	YS [MPa]	UTS [MPa]	EL [%]
STA825	1118 ± 12	1327 ± 1	10.1 ± 0.4
AN875	936 ± 9	1085 ± 1	22.3 ± 0.5

The cylindrical specimens were machined and polished to produce 4-point bend fatigue samples according to BS ISO 1143:2021 [28], with a $\Phi 10$ mm grip diameter and a $\Phi 5$ mm gauge section. Surface roughness measurements were carried out on the gauge section of the fatigue samples using a JR25 Nanovea (Irvine, CA, USA) optical profilometer to verify compliance with the specifications. The samples showed a smooth and regular surface with average Ra values of 0.32 ± 0.02 μm . Figure 2 illustrates the geometry and dimensions of the samples used in this study.

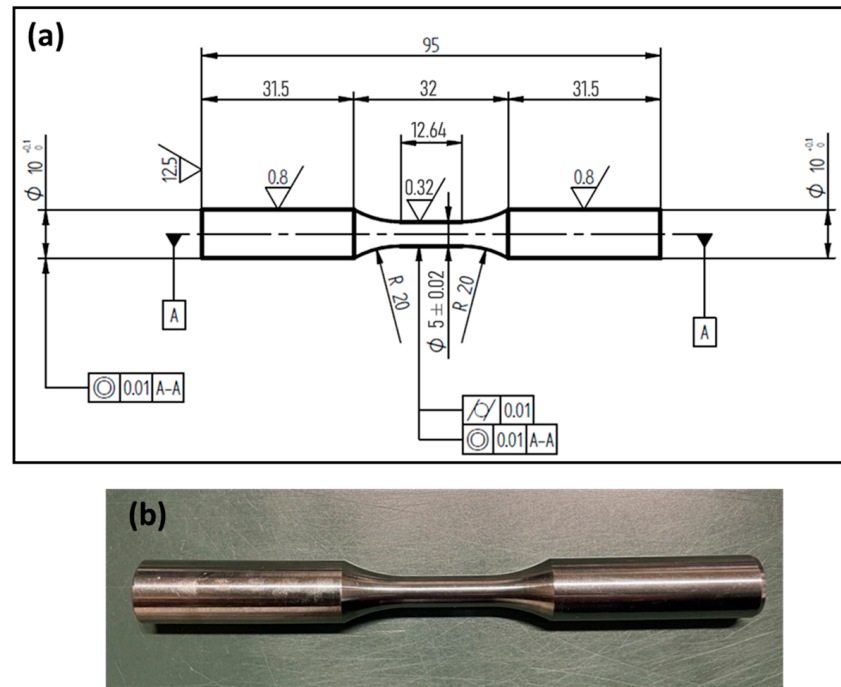


Figure 2. The geometry and dimensions [mm] (a) and an image of the fatigue samples (b) according to BS ISO 1143:2021.

2.2. Microstructural Characterization

Microstructural characterization was performed on the longitudinal sections of the cylindrical specimens. Three samples for each heat treatment condition were analyzed. One cylinder with a diameter of 10 mm and a length of 10 mm was extracted from each grip end, hot-mounted in conductive resin, ground with abrasive papers up to P1200 grit, and polished with diamond suspensions (9 μm and 3 μm), followed by final polishing with a water-based colloidal silica and hydrogen peroxide solution to achieve a mirror finish [29]. Polished cross-sections were examined using a Zeiss Axio Imager A.1M optical microscope (OM, Oberkochen, Germany) to identify process-related defects present in the samples. For detailed microstructural analysis, samples were etched with Kroll's reagent (93% H_2O , 5% HNO_3 , 2% HF) and imaged at low magnification with OM and at high magnification using a Tescan Mira 3 Field Emission Gun–Scanning Electron Microscope (FEG-SEM, Brno, Czech Republic). Image analysis was performed using ImageJ v1.52a [30] to assess sample density, defect content and size, as well as key microstructural features, including the α_p lamellae content and thickness. The α_p content was defined as the α_p (dark area in Figure 3a) area fraction relative to the total analyzed area. α_p lamellae were isolated in the microstructure using the “Outlines” function of the ImageJ software v1.52a (Figure 3b), which then aggregates the contributions from the delineated areas. The “Ellipse” ImageJ tool was instead applied to approximate the morphology of the α_p lamellae via elliptical fitting (Figure 3c). The arithmetic mean of the minor axes of the fitted ellipses was defined as the characteristic lamellar thickness, as described in [22]. Microstructural analyses were conducted on two samples for each heat treatment condition, with five images for each sample. Images were taken with varying magnifications for each condition (5kx for AN875 and 30kx for STA825), due to their differing microstructural fineness.

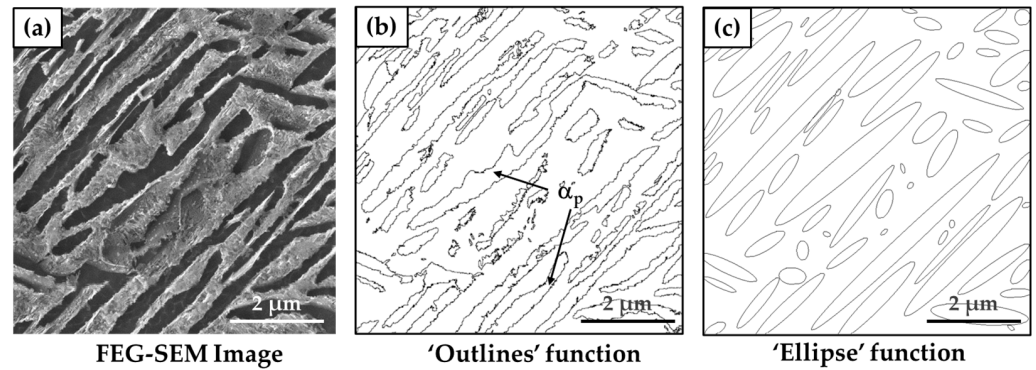


Figure 3. FEG-SEM microstructure (a) for image analysis made by using the ‘Outlines’ Function (b) to determine the α_p content and the ‘Ellipse’ Function (c) to quantify the α_p lamella thickness.

2.3. Fatigue, Hardness Tests, and Fractographic Analyses

High-cycle fatigue strength was evaluated through fully reversed ($R = -1$) rotating–bending tests conducted in a four-point configuration machine (RB35, Italsigma, Forlì, Italy), according to the ISO 1143:2021 standard. The tests were performed at a frequency of 50 Hz, with a run-out limit of 2×10^6 cycles. Fatigue strength at a 50% probability of failure ($\sigma_{d,50\%}$) and the associated standard deviation (σ_γ) were determined using the staircase method (ISO 12107:2012 [31]). Based on preliminary tests, a stress increment of 20 MPa (approximately 5% of the estimated fatigue strength) was selected, and 15 specimens were tested for each heat treatment condition. Vickers hardness measurements were performed on the grip sections of the fatigue specimens using a 30 kg load and a 15 s dwell time to ensure consistency in hardness across all the samples. For both treatment conditions, the values were obtained by averaging 5 measurements performed on two different samples.

All the run-out specimens were subsequently retested until failure. Fractographic analyses were conducted using the FEG-SEM to identify crack nucleation sites, dominant failure mechanisms, and the size and nature of the defect responsible for initiation, referred to as the “killer defect”. For each specimen tested, the stress range ($\Delta\sigma$)—a function of the maximum (σ_{max}) and minimum (σ_{min}) stress—and the stress intensity factor range (ΔK) at the crack tip were determined, calculated using Equations (1) and (2), respectively. Since the tests were conducted under symmetric alternating load conditions ($R = -1$), σ_{max} and σ_{min} are equal in magnitude and opposite in sign, and $\Delta\sigma$ is twice the alternating stress (σ_a). The stress intensity factor is a function of the stress range, the critical defect size (in terms of $\sqrt{\text{area}}$, also known as the Murakami parameter), and the defect’s location (where coefficient C is 0.5 for internal defects or 0.65 for surface defects) [26,32,33].

$$\Delta\sigma = \sigma_{max} - \sigma_{min} = 2\sigma_a, \quad (1)$$

$$\Delta K = C \cdot \Delta\sigma \cdot \sqrt{\pi \cdot \sqrt{\text{area}}}, \quad (2)$$

KT diagrams were developed based on the fracture surface analysis data, in accordance with established methodologies for additively manufactured alloys [26]. These diagrams plot $\Delta\sigma$ and ΔK as functions of the $\sqrt{\text{area}}$, establishing a correlation between high-cycle fatigue strength and the critical defect size for the Ti6246 alloy. The diagrams include both run-out conditions, marked with “O”, and failure conditions, indicated by “X”. The data points on the KT diagram were then fitted using the El-Haddad model [34]. This model generates threshold curves that identify regions of failure (above the curve) from non-failure (below the curve). Specifically, the obtained curves are described by Equations (3)–(5),

which represent the relationship between the fatigue threshold stress range ($\Delta\sigma_{th}$), the threshold crack propagation resistance range (ΔK_{th}), and the critical defect size (\sqrt{area}):

$$\Delta\sigma_{th}(\sqrt{area}) = \Delta\sigma_{th,0} \sqrt{\frac{\sqrt{area_0}}{\sqrt{area} + \sqrt{area_0}}}, \quad (3)$$

$$\Delta K_{th}(\sqrt{area}) = \Delta K_{th,lc} \sqrt{\frac{\sqrt{area}}{\sqrt{area} + \sqrt{area_0}}}, \quad (4)$$

$$\sqrt{area_0} = \frac{1}{\pi} \cdot \left(\frac{\Delta K_{th,lc}}{Y \cdot \Delta\sigma_{th,0}} \right)^2 \quad (5)$$

$\Delta\sigma_{th,0}$ represents the limiting stress range for defect-free specimens; $\Delta K_{th,lc}$ represents the resistance limit to long crack (macrocrack) propagation; and $\sqrt{area_0}$ is a modeling parameter proportional to the critical crack length, distinguishing short from long cracks, and it is dependent on $\Delta\sigma_{th,0}$ and $\Delta K_{th,lc}$, which best fit the experimental data.

3. Results and Discussion

3.1. Microstructural Characterization

Figure 4a presents a representative OM image showing small, circular defects indicative of gas porosity (marked by arrows) and a larger, irregular defect characteristic of a LoF, further detailed in the FEG-SEM image in Figure 4b. Porosity analyses for the AN875 and STA825 samples revealed no significant deviations from the previously reported values in [23]. The average density measurements are consistent with prior results, with densities of $99.81 \pm 0.05\%$ and $99.87 \pm 0.09\%$, respectively.

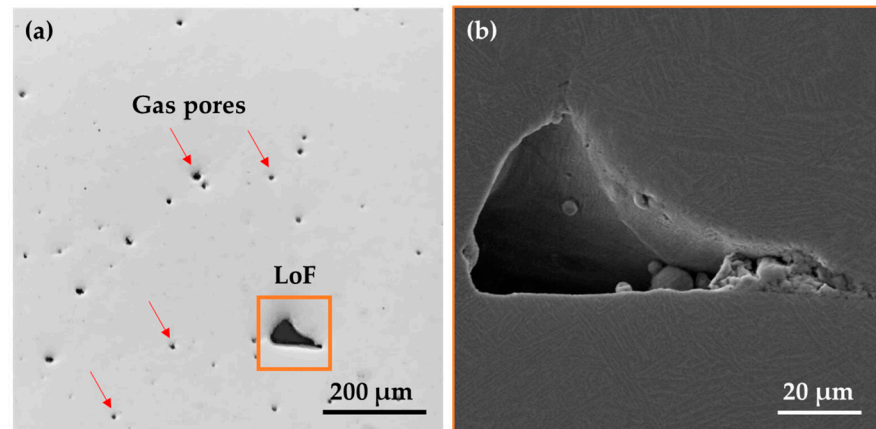


Figure 4. An OM image of a Ti6246 sample showing gas porosities (indicated by red arrows) and a larger LoF defect (a), with a corresponding high-magnification FEG-SEM image (b).

The defect size distribution, derived from image analysis of OM micrographs, is illustrated in Figure 5a. The data represents the number of defects per unit area as a function of the square root of the defect area. The accompanying table reports the 50th (D_{50}) and 90th (D_{90}) percentiles of defect size and the maximum defect size (D_{max}). In the AN875 heat-treated alloy, a lower relative frequency of defects is observed, accompanied by a higher proportion of larger defects compared to the STA825 condition. This behavior is attributed to the higher solution treatment temperature used in the AN875 process, which may promote slight growth of gas-induced porosities, as evidenced in previous works on AM-processed alloys [35–37]. Nonetheless, the variation in defect size is limited to just a few microns and is considered negligible in terms of its impact on the mechanical performance of the material, particularly its fatigue behavior.

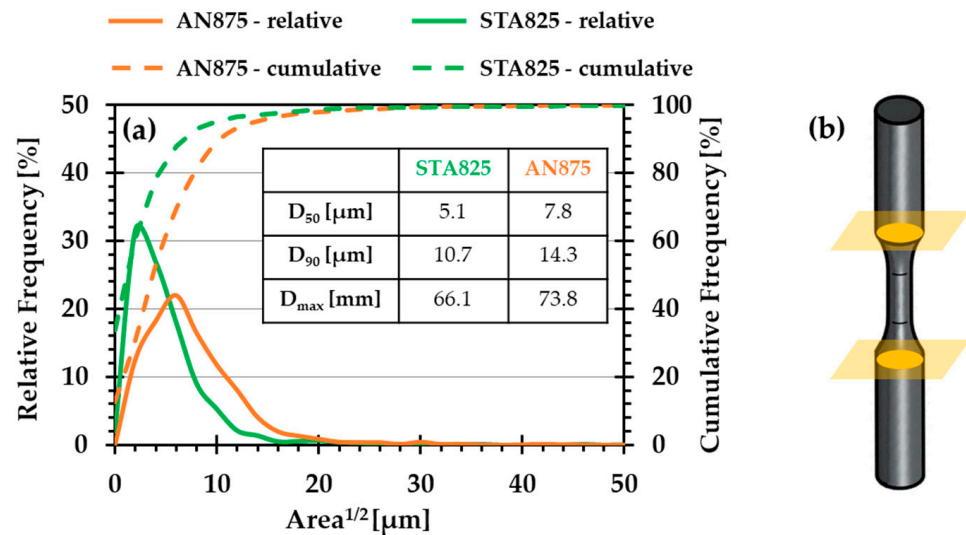


Figure 5. The relative and cumulative frequency distributions of defect sizes (in terms of the square root of the area) (a) obtained from image analysis on OM micrographs of polished cross sections (b).

Figure 5b shows a schematic representation of the cross-sections analyzed for each sample.

Figure 6 presents FEG-SEM micrographs of chemically etched samples following heat treatment. Both conditions exhibit columnar prior β grains, generated from PBF-LB printing, and a bilamellar microstructure characterized by the formation of α_s within the β phase, located between α_p lamellae. However, the STA825 condition exhibits a significantly finer arrangement of both α_p and α_s lamellae compared to the AN875 condition. These microstructural differences arise from the distinct solution treatment parameters and cooling rates applied in each heat treatment, which influence phase transformation kinetics and lamellar refinement. As reported by Galarraga et al. [38], the growth of the α phase is primarily governed by two factors: solution temperature and cooling rate. Higher solution temperatures, approaching the β -transus, combined with slower cooling rates, promote coarsening of α_p lamellae. In the AN875 condition, the microstructure consists of large α_p lamellae with rounded tips, along with finer α_s lamellae that nucleate within the β phase ($\alpha_s + \beta$) separating the α_p lamellae (Figure 6d). However, the α_s fraction is relatively low and inhomogeneously distributed. During isothermal holding at 875 °C, accelerated diffusion processes drive the coarsening of α_p . This effect is further amplified by slow furnace cooling, which promotes additional coarsening of α_p and the formation and consequent coarsening of α_s lamellae [39]. Conversely, the STA825 condition exhibits a finer bilamellar microstructure with a significantly reduced size and volume fraction of α_p lamellae, primarily influenced by the solution temperature (825 °C) and holding time. Additionally, α_s lamellae are extremely fine and uniformly distributed within the β phase ($\alpha_s + \beta$). Rapid water quenching, in fact, inhibits diffusive transformations, preserving the α_p amount achieved at 825 °C and leading to the formation of metastable β phase, as discussed by the authors in [23]. The subsequent aging process induces transformation of the metastable β phase, leading to the formation of a finely distributed α_s lamellar structure within the β phase, as indicated in Figure 6b, and the precipitation of very fine strengthening α_2 (Ti_3Al) within α_p , in agreement with the results of the XRD analysis reported in our previous study [23]. Similar findings for α - β alloys containing a minimum of 6% Al after STA heat treatment are also described in [38,40–42].

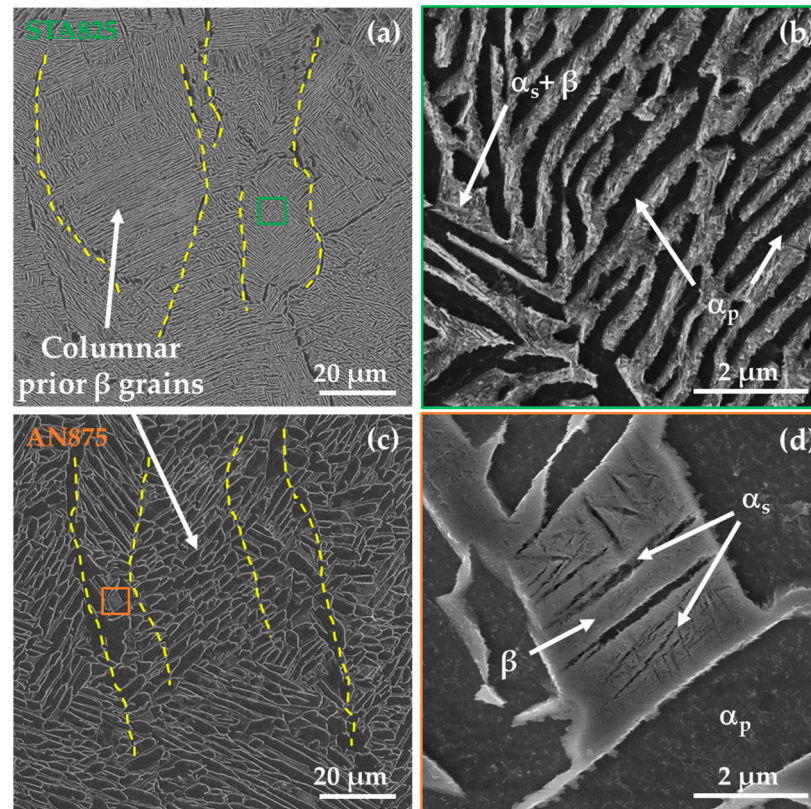


Figure 6. Low- and high-magnification FEG-SEM images of STA825 (a,b) and AN875 microstructures (c,d). The yellow dotted lines and the white arrows highlight, respectively, the columnar prior β -grain boundaries and the principal microstructural features.

Table 4 summarizes the key microstructural characteristics, including the α_p lamellae content and thickness, for each heat treatment condition. These values agree with those obtained during the heat treatment optimization phase [23].

Table 4. Summary of microstructural features (α_p lamellae content and thickness) for STA825 (measured after aging) and AN875 specimens.

	α_p Content [%]	α_p Thickness [μm]
STA825	39.32 ± 0.69	0.29 ± 0.02
AN875	69.99 ± 1.28	1.46 ± 0.10

3.2. Fatigue Behavior

The results of the fatigue tests are reported in Table 5. The STA825 samples ($HV_{30} = 425 \pm 5$) exhibited higher fatigue strength compared to the AN875 ones ($HV_{30} = 351 \pm 9$), with the 50% probability of failure ($\sigma_{d,50\%}$) being approximately 25% higher (474 ± 15 MPa vs. 382 ± 15 MPa). For both heat treatment conditions, $\sigma_{d,50\%}$ was found to be approximately 35–36% of the ultimate tensile strength (UTS) (Table 5), highlighting a correlation between the static and fatigue properties of the alloy, regardless of the heat treatment conditions.

Table 5. A summary of fatigue strength at the 50% failure probability ($\sigma_{d,50\%}$) and standard deviation (σ_γ), and the $\sigma_{d,50\%}/\text{UTS}$ ratio of the investigated heat treatment conditions.

	$\sigma_{d,50\%}$ [MPa]	σ_γ [MPa]	$\frac{\sigma_{d,50\%}}{\text{UTS}}$ [%]
STA825	474	15	35.7
AN875	382	15	35.2

This observation confirms the soundness of the design strategy employed for the STA825 heat treatment. Specifically, the STA825 treatment offers enhanced fatigue resistance over the conventional AN875 condition due to the following: (i) enhanced static strength, resulting from the synergistic strengthening effect of a refined and homogeneously distributed bilamellar architecture (comprising α_s lamellae and a retained β phase within the interlamellar regions) and the precipitation of Ti_3Al intermetallics during the aging stage; (ii) improved resistance to early-stage fatigue crack initiation and propagation, particularly along $\alpha_p/(\alpha_s + \beta)$ interfaces, due to the presence of fine α_s lamellae, as confirmed by previous studies [14,43]; and (iii) an increased capability to deflect or arrest macrocrack propagation, attributed to the presence of fine and hardened primary α lamellae, which outperform the coarser counterparts typically formed under the AN875 regime.

Figure 7a,b shows low-magnification images of fracture surfaces of the different fatigue samples, highlighting regions characterized by stable fatigue crack propagation and final overload fractures. For both treatment conditions, nucleation was observed to occur superficially or sub-superficially, mainly from Lof defects, as visible in high-magnification images (Figure 7c,d).

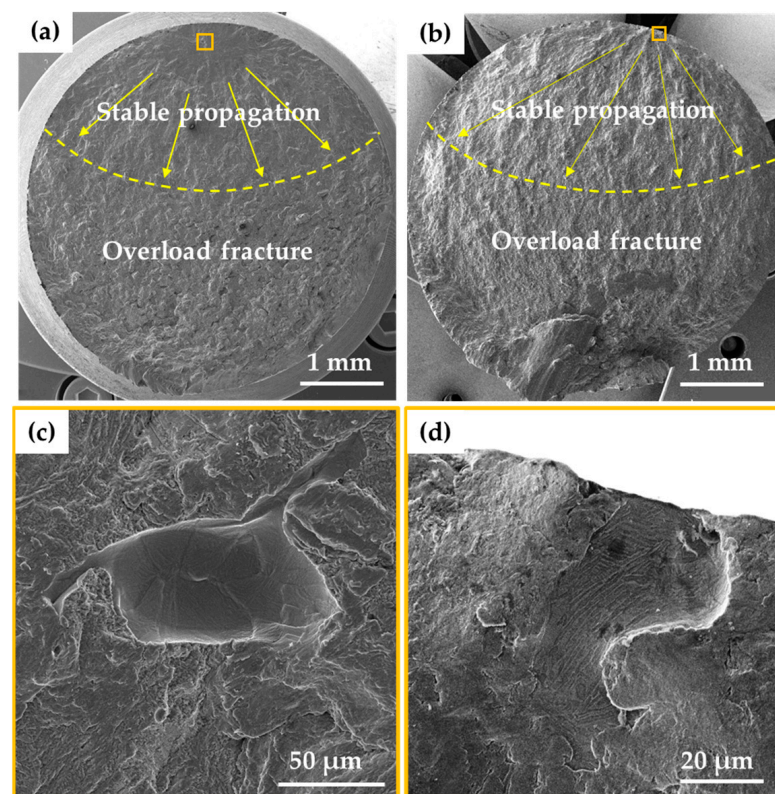


Figure 7. Low-magnification (a,b) FEG-SEM images of different fracture surfaces with a focus on crack nucleation sites originating from sub-superficial (c) or superficial Lof (d). The yellow arrows and dotted lines highlight, respectively, the stable propagation direction and the end of the stable propagation zone.

When nucleation occurs from subsurface killer defects, a low-speed crack propagation zone (LSP) and a high-speed crack propagation zone (HSP) can be identified (Figure 8a). The LSP zone (Figure 8b) can be associated with fish-eye-like crack propagation, as it exhibits an alternating distribution of parallel bright strips and dark grooves, a phenomenon termed the “mackerel pattern”. This pattern was identified by Gao et al. for Ti64 [44] and Ti6.5Al-3.5Mo-1.5Zr-0.3Si (TC11) [45] alloys. The mackerel pattern can be attributed to the difficulty encountered by the crack during its early growth stages in propagating along

the fine $\alpha_p/(\alpha_s + \beta)$ interface [45]. This effect is much more readily identifiable in the STA825 sample because the crack propagates with greater difficulty due to the presence of uniformly distributed α_s lamellae within the β phase. According to Lütjering et al. [40], α_s lamellae are difficult to deform plastically and can be considered undeformable. Therefore, their presence induces strong resistance to crack propagation, resulting in the formation of grooves and strips within the LSP [44]. In contrast, the HSP in Figure 8c zone exhibits a different morphology, characterized by the presence of tearing ridges and quasi-cleavage facets. This phenomenon is not observed when the crack nucleates and propagates from surface defects, as shown in Figure 8d. In such cases, only HSP zones can be detected, exhibiting quasi-cleavage propagation, characterized by tearing ridges and the presence of microcracks both at the crack tip (Figure 8e) and further along the propagation path (Figure 8f), in which the fatigue striations typical of a fracture under cyclic loading are detectable also. These observations agree with the findings of Li et al. [46], who attributed these peculiar fracture surface morphologies to differences in the ΔK values associated with fracture initiation at the surface versus the subsurface.

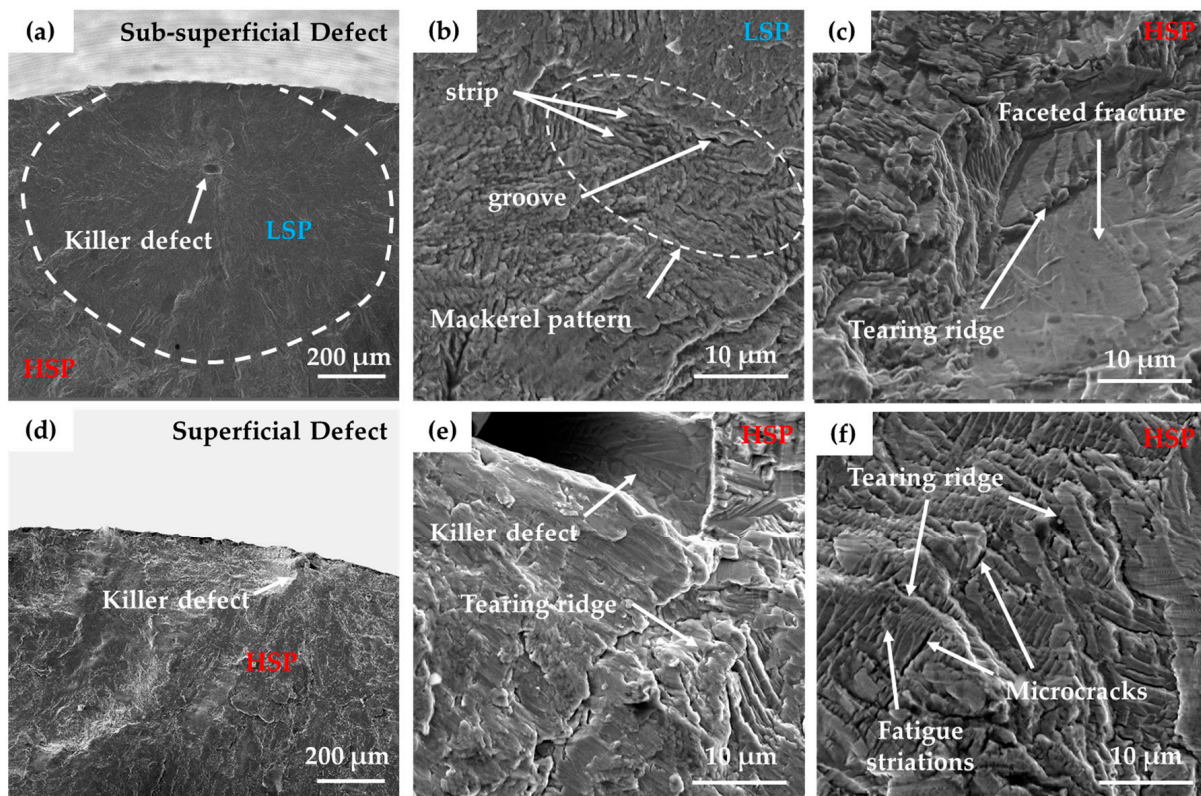


Figure 8. Crack propagation from subsurface defects (a), characterized by LSP (b) and HSP (c) zones, and from surface defects (d) with quasi-cleavage zones both at the crack tip (e) and in the propagation zone (f).

Micrographs of the overload fracture zones (Figure 9) reveal a ductile morphology, characterized by the presence of large dimples for the AN875 condition and shallow dimples for the STA825, consistent with the higher elongation to failure of the AN875 alloy than the STA825 alloy and the fracture surfaces observed in the tensile specimens documented in [23].

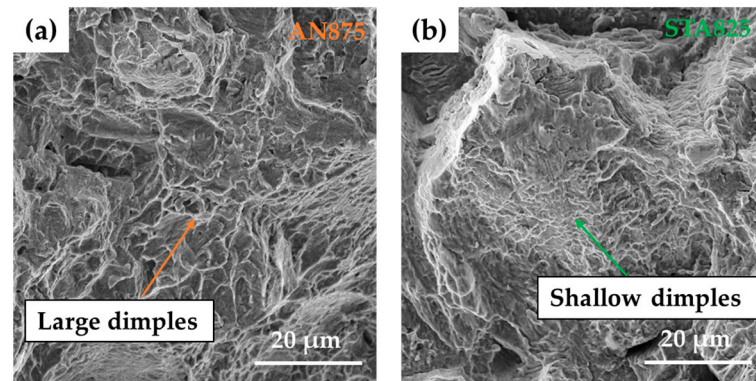


Figure 9. High-magnification images of the overload fracture zone of the specimens AN875 (a) and STA825 (b).

Figure 10a,b present the KT diagrams. These diagrams illustrate, respectively, the stress range as a function of the defect size ($\sqrt{\text{area}}$; $\Delta\sigma$) and the stress intensity factor range versus the defect size ($\sqrt{\text{area}}$; ΔK), complete with their respective El-Haddad best-fit curves. The best-fit parameters derived from the El-Haddad model for the two tested conditions are listed in Table 6.

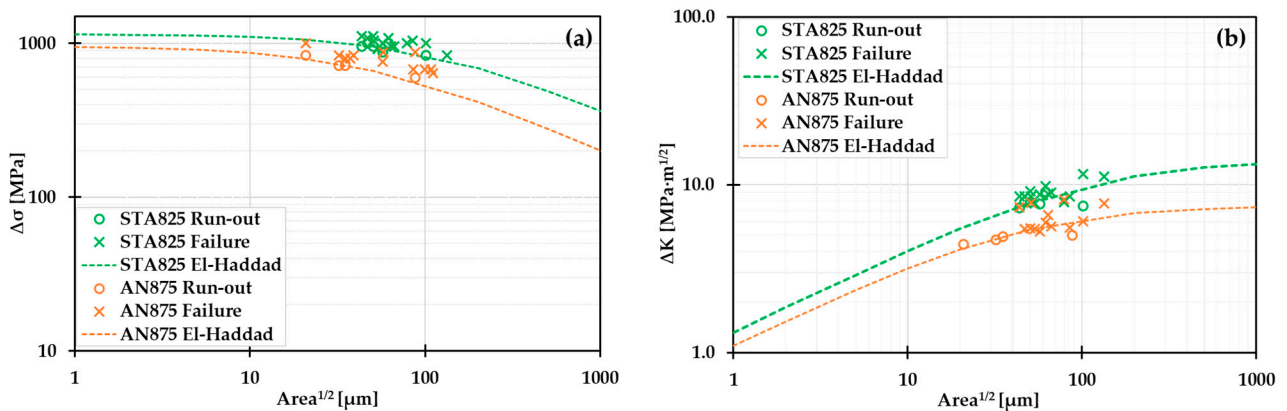


Figure 10. KT diagrams of the stress range ($\Delta\sigma$) (a) and the stress intensity factor (ΔK) (b) as a function of the defect size ($\sqrt{\text{area}}$). “X” indicates failed specimens, while “O” refers to run-out specimens.

Table 6. El-Haddad best-fit parameters for the analyzed conditions.

	$\Delta\sigma_{th,0}$ [MPa]	$\Delta K_{th,lc}$ [MPa $\sqrt{\text{m}}$]	$\sqrt{\text{area}_0}$ [μm]
STA825	1170	14.5	116
AN875	980	8.2	53

The results obtained from the KT diagrams revealed that the $\Delta\sigma_{th,0}$ and $\Delta K_{th,lc}$ resistances were 19 and 77% higher, respectively, for the STA825 sample compared to the AN875. For samples produced with additive manufacturing technologies, the latter value is particularly significant since crack nucleation almost always originates from pre-existing defects generated during the process. This confirms the positive effect of the fine bilamellar microstructure on the fatigue performance of the alloy, resulting from the synergistic combination of high tensile strength, good ductility, and enhanced resistance to crack propagation.

Although the heat-treated PBF-LB Ti6246 alloy can achieve a higher UTS compared to its wrought counterpart (1327 MPa [23] vs. 1279 MPa [10]), its fatigue performance is significantly lower due to the presence of defects, such as LoF and pores [14,21], which act as preferential sites for crack initiation. In contrast, in conventionally processed alloys, cracks

generally initiate within large α grains or α grain clusters, where localized microstructural features promote irreversible slip accumulation [43], rather than from manufacturing-induced defects. Peters et al. [12] reported that the STA-treated alloy with a fine bimodal microstructure exhibited HCF strength at stress levels up to 625 MPa, approximately 37 and 21% higher than the AN875 and STA825 conditions, respectively, due to the different crack initiation mechanisms.

Figure 11 provides a comparative analysis of the fatigue behavior of the Ti6246 alloy produced via PBF-LB in this study, the same alloy manufactured through conventional methods, and the Ti64 alloy fabricated with PBF-LB technology and subjected to different heat treatments. The reported data for the Ti64 alloy refer to tests conducted under a stress ratio of $R = -1$, using surface-polished specimens, consistent with the testing conditions applied to the Ti6246 alloy in this work. When comparing the PBF-LB STA825 Ti6246 with the PBF-LB Ti64 alloy, the STA825 Ti6246 shows superior fatigue performance regardless of the heat treatment conditions, as highlighted in Figure 10.

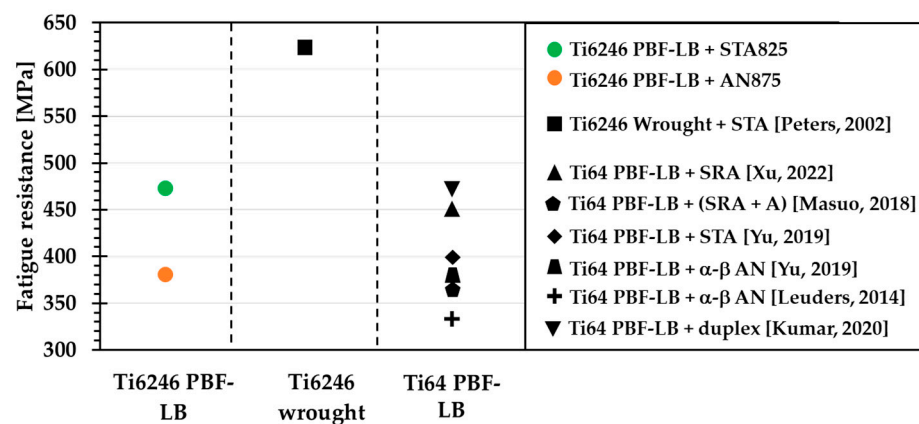


Figure 11. A comparison of the fatigue behavior of the following: (i) the Ti6246 alloy produced via PBF-LB in this study; (ii) the same alloy manufactured through conventional methods (Adapted from [12]); (iii) the Ti64 alloy fabricated with PBF-LB technology and subjected to various heat treatments, including Stress-Relieving Annealing (SRA) (Adapted from [16]), SRA and aging (SRA+A) (Adapted from [47]), α + β annealing (α - β AN) (Adapted from [15,48]), STA (Adapted from [15]), and double step annealing (duplex) (Adapted from [49]).

4. Conclusions

This study focused on assessing the effect of heat treatments and the resulting microstructures on the fatigue resistance of a Ti6246 alloy produced via PBF-LB. Two heat treatment conditions were investigated: α + β annealing at 875 °C for 2 h (AN875) and α + β solution treatment at 825 °C, followed by water quenching and aging at 500 °C for 3 h (STA825). Based on microstructural characterization, fatigue testing, and fractographic analysis, the following conclusions can be drawn:

- (i) The AN875-treated samples exhibited the lowest fatigue strength ($\sigma_{d,50\%} = 382 \pm 15$ MPa), primarily due to the low mechanical properties associated with significant microstructural coarsening caused by the high annealing temperature, slow furnace cooling rate, and the absence of strengthening precipitates. The low strength primarily affected crack initiation, while the coarse primary α (α_p) phase facilitated crack propagation.
- (ii) The STA825-treated samples demonstrated the highest fatigue strength ($\sigma_{d,50\%} = 474 \pm 15$ MPa), corresponding to an improvement of nearly 25% over the AN875 condition. This enhancement is attributed to the presence of strengthening precipitates, the refinement of α_p lamellae, and the formation of ultrafine, homogeneously

- distributed secondary α lamellae within the β phase ($\alpha_s + \beta$), which hinder fatigue crack nucleation and early-stage propagation.
- (iii) Both heat treatment conditions exhibited a similar $\sigma_{D,50\%}/UTS$ ratio, ranging between 35% and 36%, demonstrating a clear correlation between static and fatigue properties.
 - (iv) In both heat treatment conditions, fatigue crack nucleation consistently occurred at surface or subsurface lack of fusion defects. These process-induced defects represent the primary limiting factor in the fatigue performance of PBF-BF Ti6246 compared to its conventionally produced counterpart.
 - (v) Crack propagation behavior was strongly influenced by defect location. Subsurface defects triggered an initial low-speed propagation (LSP) region characterized by a “mackerel pattern”, likely due to the high microcrack growth resistance in the early propagation stage at $\alpha_p/(\alpha_s + \beta)$ interfaces. This was followed by a high-speed propagation (HSP) region exhibiting quasi-cleavage fracture. In contrast, surface-initiated cracks propagated directly through quasi-cleavage mechanisms from the crack tip. Fatigue striations were observed in the HSP zone.
 - (vi) El-Haddad’s best-fit parameters derived from the Kitagawa–Takahashi diagrams confirmed the beneficial role of the fine bilamellar microstructure of the STA825 condition in enhancing fatigue performance, with an increase of 19% in the intrinsic fatigue limit ($\Delta\sigma_{th,0}$) and 77% in the threshold stress intensity factor ($\Delta K_{th,lc}$).
 - (vii) The STA825-treated Ti6246 alloy exhibited superior fatigue strength compared to PBF-LB Ti-6Al-4V subjected to various heat treatments.

This study addresses a critical gap in the current literature concerning the influence of heat treatments on the microstructure and fatigue performance of Ti-6Al-2Sn-4Zr-6Mo (Ti6246) alloy processed via Powder Bed Fusion—Laser Beam (PBF-LB). Future investigations will focus on the role of surface finishing and Hot Isostatic Pressing (HIP) as potential post-processing strategies to mitigate the detrimental effects of process-induced defects and further improve fatigue resistance. Expanding the knowledge base on this alloy and optimizing its post-processing treatments are essential steps toward enabling its broader adoption in high-performance structural applications, such as fan blades, engine shafts, or connecting rods.

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Abbreviations

The following abbreviations are used in this manuscript:

AM	additive manufacturing
PBF-LB	Powder Bed Fusion–Laser Beam
HCF	high cycle fatigue
STA	solution treating and aging
LoF	lack of fusion
KT	Kitagawa–Takahashi
GD-OES	Glow Discharge Optical Emission Spectroscopy
OM	optical microscope
FEG-SEM	Field Emission Gun–Scanning Electron Microscope
LSP	low-speed propagation
HSP	high-speed propagation

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