Oxide Evolution During the Solidification of 316L Stainless Steel from Additive Manufacturing Powders with Different Oxygen Contents

XINLIANG YANG, FENGZAI TANG, XINJIANG HAO, and ZUSHU LI

The oxide evolution during the solidification of 316L stainless steel from additive manufacturing powders with different oxygen contents is studied by in situ observation of the melting and solidification of the powder materials, advanced characterization of the solidified materials, and non-equilibrium thermodynamic analysis. An oxide evolution map is established for the 316L powders with different oxygen contents. It reveals the relationship between the surface oxidation in the reused powder and its expected oxide species and morphology in the as-solidified component. For the 316L powder with oxygen content higher than ~0.039 pct, the liquid oxide formed first from the steel melt and then crystallized to certain oxide phases during solidification, while for the powder with lower oxygen, oxide phases are suggested to directly form from the steel melt. The oxide species in the as-solidified sample was predicted by the Scheil–Gulliver cooling calculation and verified by the TEM-based phase identification. The oxides formed in the melt of low O 316L alloy (0.0355 pct O) are predicted to be (Mn, Cr)Cr₂O₄ spinel and SiO₂ oxide. In the high O (0.4814 pct O) 316L melt solidification, the final oxides formed are (Mn, Cr)Cr₂O₄ spinel, SiO₂ oxide, and Cr₂O₃ corundum. As an important characteristic of powder materials, the oxygen pick-up due to the powder surface oxidation significantly influences the inclusion evolution in the powder fusion process.

https://doi.org/10.1007/s11663-021-02191-w
© The Author(s) 2021

I. INTRODUCTION

The research of the inclusion evolution is one of the critical considerations in the steel metallurgy, which has extensively focused on the reaction between the intrinsic elements (e.g., O, N, H, C) and the metallic elements, and the sulfides and phosphide impurities in the initial steelmaking and subsequent thermomechanical processes.[1] By the extended application of the powder formed steel materials via the powder fusion (liquid–solid transition)-based process,[2] the fundamental understanding of inclusion formation and evolution was demanded due to its non-equilibrium nature[3] for the powder manufacturing and additive manufacturing (AM) technology marching into the structure component business.[4–6]

[1] Song et al.[11] have emphasized the importance of the protective gas purity on the oxide inclusion formation and reported that varying oxygen contents in the atmosphere of the laser solid forming process significantly changed the morphology and amount of the oxide inclusion in the as-processed samples. Eo et al.[12] studied the effect of processing parameters (laser power, scan speed, beam spot size and powder steel composition) on the size distribution and number density of the inclusion evolution in the laser powder deposition process and proposed a linear relationship between the inclusion number density and the tensile yield strength of the as-processed components. Recently, Deng et al.[13] reported a wide range of experimental results of the oxygen source on the formation of the oxide inclusion from the precursor powder, processing atmosphere to the moisture during the powder storage. The above-mentioned investigations nicely covered the processing
aspect of inclusion evolution, but limited information on the oxide evolution during the powder fusion process was reported from the material quality point of view.

With the rapid development of additive manufacturing technology, the significance of extending the powder service life to improve the process cost-effectiveness has drawn great attention. Researchers have studied the effect of the reused powder on the characteristics of powder flowability, packing density and resultant mechanical performance of the as-processed components. However, the effect of the reused powder with different oxygen pick-up contents on the oxide formation and evolution in the powder fusion is still not clear.

Therefore, this study aims to reveal the oxide formation and evolution in the non-equilibrium solidification of the 316L stainless steel from the additive manufacturing powders with different surface oxidation conditions. The steel powders with different oxygen contents were subjected to the in situ observation of melting and solidification under the high-temperature confocal laser scanning microscope (HT-CLSM), TEM-based phase identification and non-equilibrium thermodynamic analysis. A relationship between the powder oxygen pick-up and the oxide formation route during the solidification is proposed. The solidified oxide species from different powder conditions were analyzed using the Scheil–Gülder cooling function and confirmed by the crystallographic identification.

II. MATERIALS AND METHODS

A. The 316L Powder Materials Under Different Oxygen Pick-up Conditions

Three 316L powders with different oxygen contents (Table I), i.e., low O powder, medium O powder and high O powder were used in this study, representing three oxygen pick-up conditions of the steel powders in their service life in the powder fusion-based AM process. The low O powder was the as-received 316L powder produced by inert gas atomization with limited oxygen pick-up, and the medium O powder was the rejected powder in the powder fusion process of the as-received 316L powder. The high O powder representing excessive-high oxygen pick-up in process was produced by oxidizing the as-received 316L powder at 800 °C under compressed air with 0.2 L/min flow rate for 20 minutes to simulate the oxygen pick-up level reported in the research community.

The chemical compositions of the three types of powder with different surface oxidation conditions are listed in Table I. The chemical composition of metallic elements was determined by the inductively coupled plasma mass spectrometry (ICP-MS) method. Carbon, sulfur and phosphorus contents were obtained by combustion analysis. Oxygen content was measured by inert gas fusion. Morphology of these powders is shown in Figure 1 with the feature of the small powders adhering over large ones, which is commonly seen in powder manufacturing due to the impingement of solidified fine powders over the coarse molten/semi-molten powders in the powder atomization process. A clean surface is observed in the low O powders in Figures 1(a) and (d). The oxygen pick-up in the Medium O powder led to the contamination formed on the powder surface (Figure 1(e)), while the excessive oxygen pick-up in the high O powder resulted in the surface oxides covering the individual steel powders in Figures 1(c) and (f). According to the previous studies, the Si, Mn and Cr, as oxygen getter elements in the 316L stainless steel, preferentially participated in the powder surface oxidation. With further extended oxidation conditions, Fe could also be involved in forming iron oxides.

B. Characterization of Powders and Solidified Droplets

To study the effect of 316L powder with different oxygen pick-up contents on the inclusion evolution during the powder fusion, the Yonekura VL2000DX-SVF17SP high-temperature confocal laser scanning microscope (HT-CLSM) was utilized to in situ observe the oxide/metal interaction with higher morphological contrast comparing to high energy synchrotron X-ray approach. This can simulate a non-equilibrium solidification condition which can be captured by the laser optical system using suitable cooling rate, and exclude the local ionization and powder spattering due to the complicated physical phenomenon of laser beam/powder interaction. The powder sample was heated to 1600 °C at a heating rate of 10 K/s. After 20 seconds holding at 1600 °C, the melt was then rapidly solidified at the cooling rate of 10 K/s in a non-equilibrium state. The Ar atmosphere (N5.0, BOC) at the flow rate of 0.2 L/min was applied in the programed thermo-cycles. 0.25 g powder dose was used to simulate the interaction volume of a single beam spot with powder material without the interference of the powder spreadability. For each powder conditions, seven sets of thermo-cycle were conducted for in situ observation and as-solidified droplet preparation.

The post-mortem characterization of the formed oxide phase in the as-solidified droplet was conducted by the morphological, chemical and crystallographic analysis. A scanning electron microscope (SEM, JEOL 7800F) and a dual-beam focused ion beam (FIB) SEM (Versa FEI, now part of Thermo Fisher Scientific) were employed to study the surface morphology of the as-solidified droplet, during which both secondary electron (SE) and backscattered electron (BSE) images were acquired. Chemical compositions of the samples were examined using the equipped energy-dispersive X-ray spectroscopy (EDS, Oxford Instruments) at 10 to 15 kV. Cross-sectional TEM (transmission electron microscopy) samples were sectioned from the oxide layer of the as-solidified 316L droplet using the standard in situ lift-out technique. Scanning TEM (STEM) analysis was conducted using a Talos F200X microscope (FEI, now part of Thermo Fisher Scientific) operating at 200 kV with a Super-X EDS system.
C. Non-equilibrium Thermodynamic Calculation

The thermodynamic calculation was processed with FactSage 7.3 software package (GTT-Technologies). The Scheil–Gulliver cooling calculation was processed for the oxide formation in the non-equilibrium solidification\(^{[31]}\) of 316L stainless steel from its powder form with different oxygen pick-up conditions. The Scheil–Gulliver model\(^{[32]}\) assumes infinitely diffusion in the liquid phase, no diffusion in the solid phase, and local equilibrium at the solid/liquid interface. The gas–liquid and gas–solid reactions were excluded in the calculation due to the limited duration of the solidification process. Due to the limited mobility data of the element diffusion in oxide materials, the back diffusion from the solidified phase is not considered in the current calculation. As a reference, the equilibrium cooling calculation was also conducted to compare the oxide species prediction. The input data included the powder chemical composition and the different oxygen content for the given conditions. The database sets of FactPS, FToxid and FSstel were used. The slag liquid, monoxide, spinel, corundum, and rhodonite were selected as oxide solution candidates. The liquid, fcc austenite and bcc ferrite were selected as steel solution candidates.

### Table I. The Chemical Compositions (Wt Pct) of Three 316L Powders With Different Oxygen Pick-up Conditions

|        | Cr    | Ni    | Mo    | Mn    | Si   | C   | S   | P   | O   | Fe  |
|--------|-------|-------|-------|-------|------|-----|-----|-----|-----|-----|
| Low O  | 15.77 | 10.22 | 2.12  | 1.26  | 0.63 | 0.0147 | 0.0142 | 0.038 | 0.0355 bal. |
| Medium O | 15.74 | 10.23 | 2.12  | 1.25  | 0.51 | 0.0145 | 0.0144 | 0.037 | 0.1575 bal. |
| High O | 15.72 | 10.22 | 2.12  | 1.23  | 0.62 | 0.0148 | 0.0142 | 0.035 | 0.4814 bal. |

The Upper Limit of the Oxygen in the Commercial 316L Stainless Steel Powder
carpenter additive 0.10
Renishaw 0.10

---

III. RESULTS AND DISCUSSION

The experimental characterization and thermodynamic analysis are detailed in the following sections to reveal the oxide evolution of the droplet samples solidified from 316L powders with different oxygen pick-up conditions.

A. Oxide Evolution During the Solidification of 316L Stainless Steel

The snapshots of the \textit{in situ} observation of the powder melting and solidification in HT-CLSM are presented in Figure 2. For the case of the low O powder, the gas bubble floating was observed from the molten liquid (Figure 2(a)) and then a clean melt surface was maintained (Figure 2(b)). The initiation of the dendritic structure was observed at 1437 °C (Figure 2(c)). A fully developed dendrite could be seen in Figure 2(d) with decreasing temperature. The bubbling phenomenon suggested that the gas in between the powder materials was wrapped into the liquid pool in the melting stage, and then rapidly driven out of the melt pool by the buoyant force. In the case of the medium O powder, the oxides in the form of a porous network occurred on the
top of the melt pool (Figure 2(e)) when the solid powders were transformed into an integrated liquid pool. With the assistance of interfacial energy and melt convection, the discrete oxide clusters were agglomerated as the oxide network. A significant oxide gathering was captured between Figures 2(f) and (g). The solidification of 316L stainless steel from the high O powder in Figure 2(i) showed a similar oxide clustering phenomenon, but a thicker oxide layer compared to the medium O powder case. This is attributed to the high surface oxides formed in the powder oxidation treatment. During the floating up of the powder surface oxide with increased volume, the majority of the large oxide clusters has been integrated as a large oxide film shown on the top of the melt. A fringe feature was observed in Figure 2(j) between the melt and the oxide film at 1497°C, which disappeared with further cooling (Figure 2(k)).

The observation in the HT-CLSM experiment shows that the powder oxygen pick-up due to the surface oxidation can influence the oxide evolution in the small volume powder fusion. Pre-existing surface oxide could release during the powder melting and gathered and agglomerated during solidification. Such agglomeration behavior assists the formation of continuous oxide films which can be entrapped in the as-processed component and potentially deteriorate the component mechanical performance. Eo et al.\cite{12} reported that the oxide layer covered the melt pool of 316L stainless steel during the laser powder deposition process. When a new layer of powders is deposited, the formed oxide layer can either melt or float up to the newly formed melt pool surface. Similar existence of the large oxide film was also reported in the Inconel 718 alloy.\cite{33}
Marangoni convection may disperse the clustered oxide. One may suggest that the temperature-induced tension of the crystallized (solid) oxide and the liquid steel melt and the oxide liquid, as well as the low surface force of interfacial energy difference between immiscible phases in a wide temperature range due to the driving path promotes the agglomeration of the non-metallic non-equilibrium solidification (temperature decreasing), total oxygen content in the steel is more than the oxygen powder and above 2000°C for high O powder (e.g., medium O powder and high O powder). With the increasing oxygen content, the oxide agglomeration may well exist in the fusion process and the internal convention. However, Brownian motion is a time-dependent phenomenon, and the Marangoni flow characterization, the turbulence kinetic energy is ~ 0.35 m²/s² by assuming turbulence intensity is 50 pct of the mean flow rate, which is still lower than the energy required to disperse the oxides agglomerate smaller than 100 μm (0.75 m²/s²). The oxide agglomeration may well exist in the fusion process from the steel powder with excessive oxygen pick-up during its service life. It can either pre-exist or melt and solidify to form the oxide liquid and then crystallize.

For the oxygen content in the powder lower than ~ 390 ppm (e.g., low O powder), oxide liquid is not formed during non-equilibrium solidification but solid oxides uniformly precipitate in the liquid steel under preferred thermal conditions. The contact of the neighboring oxide precipitates can be determined by the Brownian motion, the sedimentation by density difference and the internal convention. Brownian motion is a time-dependent phenomenon, and the drag force of the oxide sedimentation is limited by the particle size (micron to sub-micron). Therefore, limited clustering/agglomeration in powder fusion due to the un-favored kinetics condition is expected, which is consistent with the experimental observation of a clean melt surface in the solidification of 316L stainless steel from low O powder.

The oxide formation and evolution path are of great importance in the control of the clustering/agglomeration defect. Ideally, the oxygen content of the 316L powder material should be below ~ 390 ppm for the uniformly precipitated oxide particles in the steel melt. With the increasing oxygen content, the oxide agglomeration becomes severe as seen in the medium O case with 0.1575 wt pct oxygen and even worse in the high O powder with 0.4814 wt pct oxygen content.

### B. Oxide Composition of the As-solidified 316L Stainless Steel Droplet

The as-solidified droplets from the melting and solidification experiments of the three powders in HT-CLSM are taken to analyze the inherent influence of the powder oxygen pick-up on its oxide evolution during the solidification. Three distinctive morphologies of inclusions are exhibited on the surface of the as-solidified droplets from the three different powders. The droplet made by the low O powder shows a smooth surface curved by fine dendrite tips in Figures 4(a) and (d) and its detailed microstructure (Figure 4(g)). The Si-rich oxides (gray contrast) at the size of several microns and finer chromium-rich oxides (bright contrast) in sub-micron scale were identified by the EDS chemical analysis. According to Figure 3, the oxide precipitates directly in the steel melt during the solidification of the 316L melt from the low O powder (0.035 wt pct O). The formed oxide phase is predicted by FactSage calculation to be (Mn, Cr)Cr₂O₄ spinel at 1607.37°C and then SiO₂ oxides.

Figures 4(b), (e), and (h) show the surface morphology of the droplet solidified from the melt of the medium O powder in the HT-CLSM experiment. The droplet sample is partially covered with the oxides observed in the Figures 2(e) through (h), and the steel surface presents a pattern of directional dendritic structure with severe shrinkage. The RoI 1 and RoI 2 are two representative oxide morphologies illustrated in Figure 4(b). The RoI 1 (Figure 4(e)) shows the silica particles and chromium oxides uniformly formed on the droplet surface similar to the case of the low O powder. The RoI 2 area (Figure 4(h)) corresponds to the porous oxide network observed during the melting and solidification, which was inherited from the medium O powder surface.

The TEM characterization of the FIB sliced oxide crystal is presented in Figure 5. The well-developed facet structure was observed in the High-angle annular dark-field (HAADF) image (Figure 5(a)) with the compounds entrapped within the oxide crystal. The selected area diffraction (SAD) pattern (Figure 5(b)) of the same crystal shows a cubic crystal structure with an incident beam parallel to the [111] zone axis. Together with the EDS mapping and quantitative analysis of Figures 5(d) through (g), the oxide observed in Figure 4(h) is confirmed to be MnCr₂O₄ spinel oxide. The entrapped compound indicated in Figure 5(a) is confirmed as amorphous silicon oxide by the bright-field (BF) image with the insertion of Fast Fourier transform (FFT) in Figure 5(c) and EDS mapping in Figures 5(g) and (h). An iron particle is also observed in the oxide crystal as shown in Figure 5(d), which could be the entrapped liquid steel in the oxide crystallization.

The thick and large patches of inclusions (oxides) are observed on the surface of the as-solidified droplet from the high O powder during the HT-CLSM experiment in Figure 4(c) and the two RoIs (Figures 4(f) and (i)). In RoI 1, three sections of the droplet surface have been separated by dash lines, which are the steel surface.
intermediate phase and the thick layer of oxides corresponding to the liquid metal, the fringe feature and the oxides layer in the HT-CSLM experiment (Figure 2(j)). The thick oxide layer in RoI 2 (Figure 4(i)) could be a mixture of silicon oxides and Cr, Mn containing oxides from the EDS analysis. The silicon oxides present a bright contrast, while the dark contrast is the Cr, Mn-containing oxides. In Figures 6(a) and (b), the thick oxide layer in RoI 2 (Figure 4(i)) is confirmed to be a mixture of dark facet crystals and bright binder compound. The high-resolution TEM-BF images (Figures 6(c) and (d)) showed a cubic crystal structure with (111) plane $d$-spacing of 0.4873 nm for the dark facet crystal, while an amorphous feature for bright binder compound (Figure 6(e)). Together with the chemical analysis in Figures 6(g) through (j), the dark facet crystal is MnCr$_2$O$_4$ spinel$^{[38]}$ and the bright compound is the amorphous silica. The intermediate phase observed in Figure 4(f) was analyzed using EDS through FIB sectioning due to its limited layer thickness (Figures 7(a) and (b)). The element mapping in Figures 7(e) through (h) suggests that the intermediate phase was mainly composed of a high fraction of the Cr and Mn containing oxide with a small volume of silicon oxides in between.

To understand the phase evolution in the oxide agglomeration on the medium O and high O droplets, the oxide phase calculations using the Scheil–Gulliver cooling and the equilibrium cooling functions are presented in Figure 8 by setting the oxide liquid as the cooling solution phase. In the equilibrium cooling, both cases follow the similar solidification route, i.e., Liquid (steel) + Oxide liquid $\rightarrow$ Spinel, by keeping the total mass balance constant. The temperature range of the above reaction is between 1633.71 $^\circ$C to 1597.52 $^\circ$C for the medium O melt, and 1697.56 $^\circ$C to 1568.41 $^\circ$C for the high O melt. However, once the solid phase has been removed from the total mass balance of the target phase in the Scheil–Gulliver cooling, the solidification routes significantly deviated from the equilibrium state. In the medium O case (Figure 8(a)), the (Mn, Cr)Cr$_2$O$_4$ spinel first formed at 1736.26 $^\circ$C and consumed the majority of the oxide liquid, with further cooling, the SiO$_2$ oxide occurred at 1482.98 $^\circ$C. This is consistent with the oxide phase identified in Figure 5. However, the SiO$_2$ was presented in an amorphous state rather than the tridymite crystal structure due to the influence of the cooling rate subjected to the experiment. A good agreement is obtained between the oxide phase identified in the TEM characterization (Figure 6) and the
Fig. 5—The crystal structure identification and chemical analysis of the oxide shown in the RoI 2 in the medium O droplet (Fig. 4(h)), (a) HAADF image of the oxide crystal, (b) selected area diffraction (SAD) pattern of the same oxide, (c) the BF image of the entrapped compound RoI in (a) with the insertion of FFT image suggesting a feature of amorphous structure, and (d) through (h) the element maps of Fe, Mn, Cr, O, and Si to confirm the oxide from chemical analysis.

Fig. 6—The crystal structure identification and chemical analysis of the oxides shown in the RoI 2 in the high O droplet (Fig. 4(i)), (a) and (b) BF image of the oxide mixture, (c) the high-resolution TEM with the two FFT images (d) and (e) of the selected area, (f) the HAADF image of the mixture of the oxide layer, and (g) through (j) the element maps of O, Mn, Cr, and Si to confirm the oxide from chemical analysis.
The current work suggests that the oxide formation and the resultant species under the non-equilibrium condition deviated significantly from its equilibrium state in particular for the powders containing medium and high oxygen contents (> 390 ppm). This may improve the oxide inclusion prediction in rapid solidification processes, such as the powder fusion-based additive manufacturing, where the inclusion engineering is lack of consideration at the moment, however, could play a critical role, due to its inherent high oxygen content in the metal powder, in the properties of the as-built components. For the low O powder, it favors the powder fusion-based AM process due to its much lower oxide formation temperature and eliminated liquid oxide phase in the non-equilibrium solidification, which leads to the fine inclusions precipitating uniformly in the
as-built component. The medium and high O powders may not be suitable for the conventional AM component due to the excessive oxide inclusion. The unique feature of metallic powder coated with oxide layer may enable them into the functional grade materials, e.g., to improve the stiffness of the component made by retained oxide network in between the steel matrix uniformly.

**IV. CONCLUSIONS**

In the current work, the oxide evolution in the solidification of 316L stainless steel from additive manufacturing powders with different oxygen contents was investigated through the in situ observation, TEM phase identification, and non-equilibrium thermodynamic analysis. The conclusions can be drawn below.

1. The oxygen pick-up on the powder materials through surface oxidation significantly influences the oxide formation/evolution, morphology, and species during the solidification of the 316L stainless steel from its powder form.
2. The Scheil–Gulliver cooling-based non-equilibrium thermodynamic calculation of the oxide evolution provides consistent oxide phase prediction to the crystallographic phase identification, and it deviates from the equilibrium cooling function.
3. For the powder with oxygen content higher than ~390 ppm (medium and high O powder), the formation of liquid oxide mixture from the steel melt and then crystallization to certain oxide phases are observed during melting and solidification, while for the powder with lower than ~390 ppm oxygen (low O powder), such oxide evolution behavior is not ob-
served and oxide phases are suggested to form from the steel melt uniformly during solidification.

4. The oxides formed in the melt of the low O 316L alloy (0.0355 pct O) are predicted to be (Mn, Cr)Cr2O4 spinel and SiO2 oxide. With increasing oxygen content, oxide liquid forms first, and then the spinel phase crystallizes from the oxide liquid at the temperature of 1736.26 °C and SiO2 at 1482.98 °C in the medium O case (0.1575 pct O). In the high O (0.4814 pct O) 316L melt solidification, the formation temperature of spinel is raised to 1792.68°C. The final oxides formed are (Mn, Cr)Cr2O4 spinel, SiO2 oxide, and Cr2O3 corundum.

ACKNOWLEDGMENTS

This work is financially supported by EPSRC with Grant No. EP/N011368/1. The authors would like to thank Dr. Sabrina Yan for the TEM sample preparation in WMG, the University of Warwick.

OPEN ACCESS

This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

REFERENCES

1. J.F. Elliott and M. Gleiser: Thermochimica Acta: Steelmaking, Addison-Wesley Publishing Co., Reading, 1963.
2. D. Herzog, V. Seyda, E. Wycisk, and C. Emmelmann: Acta Mater., 2016, vol. 117, pp. 371-92.
3. J. Oliveira, T. Santos, and R. Miranda: Prog. Mater. Sci., 2020, vol. 107, p. 100590.
4. T. DebRoy, T. Mukherjee, J. Milewski, J. Elmer, B. Ribic, J. Blecher, and W. Zhang: Nat. Mater., 2019, vol. 18, pp. 1026-32.
5. S. Öztürk, K. Için, B. Öztürk, U. Topal, and H.K. Odabasi: Int. J. Mater. Sci. Appl., 2017, vol. 6, pp. 241-49.
6. S. Öztürk, S.E. Sünbül, and K. Için: Trans. Nonferrous Met. Soc. China, 2020, vol. 30, pp. 1169-82.
7. T. Lau, G. Weatherly, and A. McLean: Weld. J., 1986, vol. 65, pp. 343-47.
8. T. Kuwana, Y. Sato, and S. Kaneda: Weld. Int., 1993, vol. 7, pp. 365-70.
9. T. Koseki, S. Ohkita, and N. Yurioka: Sci. Technol. Weld. Join., 1997, vol. 2, pp. 65-69.
10. O. Grong, T. Siewert, G. Martins, and D. Olson: Metall. Mater. Trans. A, 1986, vol. 17A, pp. 1797-1807.
11. M. Song, X. Lin, F. Liu, H. Yang, and W. Huang: Mater. Des., 2016, vol. 90, pp. 459-67.
12. D. Eqo, S. Park, and J. Cho: Mater. Des., 2018, vol. 155, pp. 212-19.
13. P. Deng, M. Karadge, R.B. Rebak, V.K. Gupta, B.C. Prorok, and X. Lou: Addit. Manuf., 2020, vol. 35, p. 101334.
14. I.E. Anderson, E.M. White, and R. Dehoff: Curr. Opin. Solid State Mater. Sci., 2018, vol. 22, pp. 8-15.
15. E. Hryha, R. Shvab, H. Gruber, A. Leicht, and L. Nyborg: Metall. Ital., 2018, vol. 3, pp. 34-39.
16. S.E. Brika, M. Letenkee, C.A. Dion, and V. Brailoovski: Addit. Manuf., 2020, vol. 31, p. 100929.
17. A.B. Spierings, M. Voegtlín, T. Bauer, and K. Wegener: Prog. Addit. Manuf., 2016, vol. 1, pp. 9-20.
18. P. Quinn, S. O’Halloran, J. Lawlor, and R. RagHAVendra: Adv. Mater. Process. Technol., 2019, vol. 5, pp. 348-59.
19. Z. Dong, H. Kang, Y. Xie, and C. Chi: Mater. Lett., 2019, vol. 236, pp. 214-17.
20. M.J. Heiden, L.A. Deibler, J.M. Rodelas, J.R. Koepke, D.J. Tung, and D.J. Saiz: Addit. Manuf., 2019, vol. 25, pp. 84-103.
21. C.L.A. Leung, S. Marussi, M. Towrie, R.C. Atwood, P.J. Withers, and P.D. Lee: Acta Mater., 2019, vol. 166, pp. 294-305.
22. Carpenter Additive technical data sheet CT PowderRange 316L F. https://www.carpenteradditive.com/hubfs/carpenter_additive/image/Resources/Datasheets/CT%20PowderRange%20316L%20F.pdf. Accessed 10 Nov 2020.
23. Renishaw data sheet SS 316L-0407 powder for additive manufacturing. https://www.renishaw.com/media/pdf/en/8cbea72a843440d3bd8a09fd5021ad89.pdf. Accessed 10 Nov 2020.
24. S. Öztürk, S.E. Sünbül, A. Meçoğlu, S. Önal, and K. Için: Mater. Sci. Technol., 2020, vol. 36, pp. 1771-84.
25. H. Karlsson, L. Nyborg, and S. Berg: Powder Metall., 2005, vol. 48, pp. 51-58.
26. I. Nyborg, T. Tunberg, and P. Wang: Met. Powder Rep., 1990, vol. 45, pp. 750-53.
27. S.E. Sünbül, K. Için, F.Z. Seren, Ö. Sahin, D.D. Çakıl, R. Sezer, and S. Öztürk: Vacuum, 2021, vol. 187, p. 110072.
28. A. Erdogam, S.E. Sünbül, K. Için, and K.M. Doleker: Vacuum, 2021, vol. 187, p. 110143.
29. J. Zhou, H. Tsai, and P. Wang: Weld. J. Heat Transf., 2005, vol. 128, pp. 680-90.
30. C. Zhao, C. Kwakernaak, Y. Pan, I. Richardson, Z. Saldi, and S. Kenjeres: Acta Mater., 2010, vol. 58, pp. 6345-57.
31. D. Durinc, P.T. Jones, B. Blampain, P. Wollants, G. Mertens, and J. Elsen: J. Am. Ceram. Soc., 2007, vol. 90, pp. 1177-85.
32. A.D. Pelton, G. Eriksson, and C.W. Bale: Metall. Mater. Trans. A, 2017, vol. 48A, pp. 3113-29.
33. Y. Zhang, X. Cao, P. Wanjarja, and M. Medraj: Acta Mater., 2013, vol. 61, pp. 6562-76.
34. H. Duan, Y. Ren, B.G. Thomas, and L. Zhang: Metall. Mater. Trans. B, 2019, vol. 50B, pp. 36-41.
35. S. Clark, C. Leung, Y. Chen, L. Sinclair, S. Marussi, and P. Lee: IOP Conf. Ser. Mater. Sci. Eng., 2020, vol. 861, p. 012010.
36. P. Shima, J. Philip, and B. Raj: Appl. Phys. Lett., 2009, vol. 94, p. 223101.
37. J Bellot, V. Descotes, and A. Jardy: JOM, 2013, vol. 65, pp. 1164-72.
38. J. Hastings and L. Corliss: Phys. Rev., 1962, vol. 126, p. 556.

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.