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Article Type: Full length article

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Effect of powder characteristics and oxygen content on modifications to the microstructural topology during hot isostatic pressing of an austenitic steel

S. Irukuvarghula *, H. Hassanin², C. Cayron³, M. Aristizabel⁴, M. M. Attallah⁵, and M. Preuss¹

¹School of Materials, University of Manchester, U.K, M1 3BB
²School of Mechanical and Automotive Engineering, Kingston University, London, KT1 2EE, UK
³Ecole Polytechnique Fédérale de Lausanne (EPFL), Rue de la Maladière 71b, 2000 Neuchâtel, Switzerland
⁴Ceit-IK4, PO Manuel Lardizabal 15, 20018 Donostia-San Sebastian, Spain
⁵School of Metallurgy and Materials, University of Birmingham, Edgbaston, B15 2TT, UK

Abstract

The effect of powder size distribution and oxygen content on the extent of multiple twinning and spatial distribution of oxide inclusions in hot isostatic pressed (HIPed) 316L steels was investigated using powders with different characteristics. Modifications to, and differences in their microstructural topology, were tracked quantitatively by evaluating the metrics related to twin related domains (TRDs) on specimens produced by interrupting the HIPing process at various points in time. Results revealed that powder size distribution has a strong effect on the extent of multiple twinning in the fully HIPed microstructure, with specimens produced using narrow distribution showing better statistics (i.e., homogeneously recrystallized) than the ones produced using broad size distribution. The oxide inclusion density in fully HIPed microstructures increased with the amount of oxygen content in the

*Corresponding author: sandeep.irkuvarghula@twi.co.uk
powders while prior particle boundaries (PPBs) were only observed in the specimens that were HIPed using broad powder distribution. More importantly, results clearly revealed that the spatial distribution of the inclusions was strongly affected by the homogeneity of recrystallization. Implications of the results are further discussed in a broader context, emphasizing the importance of utilizing the occurrence of solid state phase transformations during HIPing for controlling the microstructure evolution.

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### 1 Introduction

Powder hot isostatic pressing (HIPing) is a manufacturing process that is used to produce near net shape components with fine grain size, chemical homogeneity, and improved inspectability [1]. Used in many industries, including oil & gas and aerospace, it is a thermomechanical process in which powder encapsulated in a canister is consolidated to theoretical density by the application of temperature and pressure. The HIPing conditions, chosen according to the material (see for e.g., Table. 1 in [1]), ensure complete densification of the powder compact by the end of the HIPing cycle. While the densification kinetics during HIPing for various alloys have been studied to considerable detail [2, 3, 4, 5, 6, 7], the effect of powder characteristics on the microstructure development during HIPing, and on fully HIPed microstructure, still remains unclear, and is of significant interest.

Albeit the applied pressure during HIPing is isostatic, powders deform inhomogeneously depending on their size; i.e., small particles deform more than large particles [5, 6, 8, 9, 10, 11]. In powder HIPed 316L austenitic stainless steels, it has previously been demonstrated that deformation of the powder is a prerequisite for recrystallization to occur during HIPing [11]. This has been linked to the absence of annealing twins (Σ3 boundaries in coincidence site lattice formalism) and associated twin chains in powder particles that do not undergo plastic deformation; these non-deformed powders can be discerned by their prior particle boundaries (PPBs) in the fully consolidated microstructure [11]. This observation has a more general applicability to low-medium stacking fault energy (SFE) materials processed by powder HIPing, and is not limited to 316L steels (see for e.g., [12, 13] for microstructures of powder HIPed Inconel 718).

It is known that inclusions and second phase particles act as void nucleation sites during ductile fracture, and strongly affect material toughness [14, 15, 16]. Compared to conventionally processed materials, powders contain an order of magnitude more oxygen, which, depending on its solubility in the material, can be in the form of oxide inclusions
(or oxycarbides, depending on the chemistry of the alloy) in the microstructure of a fully HIPed component. It has previously been demonstrated that oxide inclusions lower the impact toughness of powder HIPed 316L and 304L steels [17, 18]. In powder HIPed nickel-base superalloys, presence of inclusions at PPBs in the as-HIPed and heat treated state has been linked to the reduction in their ductility and stress rupture properties [12, 13, 19], although post-HIP hot working has been shown to “break” the PPB networks and randomize the spatial distribution of inclusions [20, 21, 22, 23, 24]. In other words, as a result of recrystallization during post-HIP thermomechanical processing, inclusions at PPBs are pushed by moving grain boundaries, and therefore, are homogeneously distributed in the bulk. More specifically, spatial redistribution of the inclusions, which are initially present at PPBs, is promoted by the formation and propagation of twin chains (i.e., recrystallization) during hot working.

Since the microstructure in low-medium SFE materials evolves by recrystallization during HIPing (provided there is enough stored energy from particle deformation) [11], understanding the effect of powder characteristics on microstructure development, and more specifically, on the extent and homogeneity of recrystallization – and its effect on the spatial distribution of inclusions, will give insights for the development of high value manufacturing of near net shape HIPed components (e.g., Nickel-base superalloys and austenitic steels for aerospace, oil & gas, and nuclear industries). Therefore, in the present study, by HIPing 316L powders (a representative low-medium SFE material) with different size distributions and oxygen content, we aim to address (a) the effect of powder size distribution on recrystallization and the extent of multiple twinning during HIPing (b) the effect of oxygen content in the powders on the inclusion size and density in the fully HIPed microstructure and (c) the effect of recrystallization on the spatial distribution of oxide inclusions during HIPing.

Modifications to the microstructural topology of different powders during HIPing were investigated by quantifying multiple twinning using the metrics related to twin related domains (TRDs). The notion of TRDs as a characteristic microstructural dimension in low-medium SFE materials that are prone to recrystallization was first proposed by Gertsman and Henager [25], and later developed by Reed et al. [26, 27] and Cayron [28, 29]. Specifically, TRDs represent clusters of grains that are related by $\Sigma3$ misorientations, and connected by chains of $\Sigma3$ boundaries [25]. Therefore, the entire recrystallized microstructure consists of multiple twinned clusters, i.e., TRDs. Since $\Sigma3$ boundaries are generally immune to percolation phenomena (such as crack propagation), and because all grains within TRDs are connected by $\Sigma3$ boundaries, it was suggested that the material performance is controlled at the length scale of TRDs [25]. Reed et al. [30] and Barr et al. [31], by providing experimental evidence, clearly demonstrated the existence of a correlation between TRD size and material response towards intergranular phenomena.
Therefore, in this study, data analysis was focused towards identifying TRDs and quantifying multiple twinning. Using the metrics related to TRDs, we analyze the data from microstructures produced by interrupting the HIP process at various points in time. Inclusion density (i.e., #/unit area) is measured in the fully HIPed microstructures and is linked back to the oxygen content in the as-received powders. Connections between powder size distribution, PPBs, oxygen content, and the spatial distribution of inclusions on recrystallization are made, and are shown to be associated with the powder deformation during HIPing. Suggestions in terms of powder size distribution and oxygen content are put forward, that produce homogeneously recrystallized microstructures with low volume fraction and random spatial distribution of inclusions (i.e., not on PPBs) in a near net shape component, potentially eliminating the need for post-HIP thermomechanical processing.

2 Materials and methods

2.1 Experimental

![Figure 1: Particle size distribution of various powders](image)

Figure 1: Particle size distribution of various powders (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Six powders of 316L austenitic steel with different characteristics (chemical and physical) were used in this study. Specifically, four nitrogen atomized powders were obtained from commercial vendors (designated as C, D, E, and F) while two powders were gas atomised at TLS technik, Germany (atomised using argon and nitrogen; designated as A

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1Data for one of the six powders (powder C) were presented in our earlier study [11], and we show them here along with the data of five powders to demonstrate better, the correlation between powder characteristics and their microstructure evolution during HIP consolidation.
and B, respectively), from a forged plate supplied by Rolls-Royce. Particle size distribution of the six powders is shown in Fig. 1. It can be seen that powders C and D have much narrower size distribution while others have much broader distribution. Chemical composition of the powders and some of their properties are shown in Tables 1 and 2, respectively. The oxygen content in the powders varies from 110 ppm to 600 ppm and the tap density from 4.9 to 5.7 g/cc. For interrupted HIP runs, powders were first filled in 25 mm diameter, 30 mm height, and 2 mm thick mild steel canisters, vibrated and vacuum degassed at room temperature before hot crimping the evacuation tube. The canisters were then HIPed using simultaneous application of temperature and pressure, at 950 °C, 1000 °C, 1050 °C, 1120 °C, without any dwell time, at 103 MPa.

Additionally, powders were filled in mild steel canisters with 76 mm outside diameter, 2.3 mm wall thickness, 200 mm height and HIPed at 1160°C and 103 MPa, and held at those conditions for 4 hours. The canisters were then solution annealed at 1050°C for 1.5 hours and water quenched. These are designated as fully HIPed samples (or fully consolidated state) in this study. For microstructural examination, specimens were sectioned, ground and polished using standard metallographic procedures. Final polishing was performed on a vibratory polisher using colloidal silica. Backscattered electron (BSE), secondary electron (SE) images, energy dispersive spectroscopy (EDS) data for chemical mapping, and Electron Backscatter Diffraction (EBSD) data were acquired on a Field Emission Gun Scanning Electron Microscope. EBSD maps were acquired to study the differences in the evolution of grain boundary network in all six specimens during the HIPing process. A step size of 0.5 µm and 1 µm were used for partially HIPed and fully HIPed specimens, respectively. For statistical analysis of the data, EBSD maps from 5 random locations were acquired.

Table 1: Chemical composition (in wt%) of 316L stainless steel powders determined using inductively coupled plasma mass spectrometry and intert gas fusion.

| Powder | Cr | Mn | Mo | Ni | P  | Si | C  | S  | N  | O  | Fe  |
|--------|----|----|----|----|----|----|----|----|----|----|-----|
| A      | 17.93 | 1.83 | 2.10 | 10.09 | 0.036 | 0.35 | 0.028 | 0.001 | 0.079 | 0.029 | Bal. |
| B      | 17.84 | 1.85 | 2.08 | 10.07 | 0.035 | 0.39 | 0.029 | 0.001 | 0.129 | 0.011 | Bal. |
| C      | 16.44 | 1.32 | 2.08 | 10.14 | 0.023 | 0.57 | 0.018 | 0.002 | 0.098 | 0.021 | Bal. |
| D      | 16.94 | 1.48 | 2.41 | 10.45 | 0.017 | 0.69 | 0.016 | 0.007 | 0.118 | 0.061 | Bal. |
| E      | 17.9  | 1.84 | 2.44 | 11.78 | 0.009 | 0.73 | 0.02  | 0.007 | 0.061 | 0.022 | Bal. |
| F      | 17.6  | 0.64 | 2.2  | 11.3  | 0.012 | 0.69 | 0.012 | 0.010 | 0.05  | 0.0149 | Bal. |
Table 2: Physical properties of the powders.

| Property            | A  | B  | C  | D  | E  | F  |
|---------------------|----|----|----|----|----|----|
| Apparent density (g/cc) | 5.4 | 5.3 | 4.5 | 4.9 | 5.2 | 5.1 |
| Tap density (g/cc)    | 5.7 | 5.5 | 4.9 | 5.2 | 5.5 | 5.6 |
| Flowability (FFC)     | 17 | 23 | 10 | 19 | 18 | 16 |
| Satellites           | Yes | Yes | Yes | Yes | Yes | Yes |

2.2 Data analysis

Differences in the microstructural topology of the fully HIPed specimens were initially studied by quantifying the distribution of the triple junctions (TJs) associated with Σ3 boundaries. Specifically, three types of TJs were identified: $J_0$ type triple junction containing three boundaries which are not Σ3, $J_1$ type containing one Σ3 boundary, and $J_2$ type containing two Σ3 boundaries. TJs were identified using scripts written in MATLAB, and executed as part of the freely available MTEX package [32]. This approach allows qualitative understanding of the extent of twinning. Identification of higher order twins, the evolution of TRDs and the associated metrics for intermediate HIP states for all specimens was performed using ARPGE software developed by Cayron [33]. More specifically, average values of size of TRDs, number of grains per TRD ($\langle N_g \rangle$), length of longest chain ($\langle LLC \rangle$), polysynthetism ($\langle p \rangle$), and twinning anisotropy ($\langle a \rangle$) were analyzed to better quantify multiple twinning.

Detailed description and their interpretation are discussed in [29]. Briefly, length of longest chain and polysynthetism represent the longest chain of Σ3 twins in each grain and the tendency to form twin chains of type A-B-A-B-A... where the grains A and B are linked by Σ3 boundaries in a TRD, respectively. Twinning anisotropy, which depends on the choice of the primary grain used for reconstructing the TRD, represents the structure of the TRD. In other words, it describes the morphology of the TRD from its twinning tree. For statistical analysis of the oxide inclusions, several BSE images of the specimens were acquired and processed using imageJ software [34]. The average size of inclusions, their number density (#/unit area), and the nearest neighbour distance were computed with built-in plugins available in imageJ.

3 Results

Figure 2 shows representative images highlighting general microstructural aspects of as-received powder. Figure 2a shows morphology of the powder; it is seen that the powder particles are spherical and contain smaller particles (referred to as satellites) on their surfaces. Such a morphology was observed for all powders, with powder C having the
Figure 2: Representative images showing (a) the morphology of as-received powder D (b) small particles adhering to the powder surface (c) SE image and corresponding chemical maps of nickel and molybdenum (d) the grain boundary misorientations in a powder particle using electron backscatter diffraction, along with its legend. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
most number of satellites. Presence of satellites on powder surfaces reduces flowability and affects powder packing. Figure 2b highlights the surface features of the powder. Small particles, < 1 µm, are seen to adhere the powder surface, presumably from the gas-atomization process. Figure 2c shows the SE image along with the chemical maps of nickel and molybdenum obtained on polished surface of a powder particle. Nickel, and to a lesser extent, molybdenum, are seen to be segregated in what appears to be a cellular microstructure. Figure 2d shows the grain boundary misorientations of a particle from the as-received powder. The misorientations are colour coded according to the scheme proposed by Patala et al. [35, 36], which allows complete misorientation representation (axis and angle) of grain boundaries. It is seen that the boundaries have a jagged appearance and are predominantly high angle.

While the grain boundaries in as-received powder principally are random high angle, the fully HIPed specimens on the other hand contain a large fraction of annealing twins (Fig. 3), indicating that the powders undergo recrystallization during consolidation by HIPing. Qualitatively, Fig. 3 also reveals differences in the microstructures of specimens HIPed with different powders. Specifically, it can be seen that the specimens consolidated using powders A, B, and F show an inhomogeneously recrystallized microstructure with large grains surrounded by several small recrystallized grains (Fig. 3a, Fig. 3b, and Fig. 3f, respectively) while the specimens HIPed using powders C, D, and E are more homogeneously recrystallized (Fig. 3c, Fig. 3d, and Fig. 3e, respectively).

Differences in their microstructures were evaluated first by analyzing the distribution of \( J_0, J_1, \) and \( J_2 \) type triple junctions. Figure 4 shows a representative grain boundary misorientation map of the fully HIPed specimen (using powder D) overlayed with different types of triple junctions. It is seen that most of the triple junctions are of \( J_1 \) type, followed by \( J_0 \) and \( J_2 \), respectively. The distribution of types of triple junctions in fully HIPed specimens is tabulated in Table 3. In all specimens, \( J_1 \) type dominates, followed by \( J_0 \) and \( J_2 \). The statistics also confirm the qualitative observations made from the BSE images (Fig. 3). Specifically, compared to A, B, and F, specimens C, D, and E have lower \( J_0 \) type junctions and higher fraction of \( J_1 \) triple junctions. In other words, specimens A, B, and F have lower number fraction of \( \Sigma 3 \) boundaries compared to C, D and E; it then follows that they also contain more grains which have not undergone multiple twinning. This further suggests that the observed differences arise from their contrasting powder characteristics (see Fig. 1, and Tables 1 and 2).

It is emphasized that, owing to the crystallographic constraint at triple junctions [37], the third boundary in a \( J_2 \) type triple junction can either be \( \Sigma 9 \) or \( \Sigma 1 \) boundary (i.e., \( \Sigma 3^2 \) and \( \Sigma 3^0 \), respectively), while for the \( J_1 \) type junction, the other two boundaries will be higher order twins (i.e., according to the following rule: \( \Sigma 3^n \Sigma 3^m = \Sigma 3^{n+m-2i} \), where \( i \) is an integer between 0 and \( n \) [26, 28]. It is also possible that the \( J_0 \) type junctions contain
Figure 3: BSE images of fully consolidated specimens. (a) specimen HIPed using powder A, (b) to B, and so on.
Figure 4: Representative map showing grain boundary misorientations colour coded according to the legend in Fig. 2b, overlayed with various triple junction types. Here, triple junctions consisting of three non-$\Sigma$3 boundaries (i.e., $J_0$) are marked by $\blacktriangle$; one $\Sigma$3 boundary (i.e., $J_1$) by $\blacktriangle$; two $\Sigma$3 boundaries (i.e., $J_2$) by $\blacktriangle$ (Colour online).
higher order twins (e.g., \(\Sigma 9-\Sigma 9-\Sigma 9\)) and therefore in that respect, the above analysis only gives a rudimentary picture regarding the differences in grain boundary network topology between the specimens. Therefore, in order to identify higher order twins and to better understand the topological differences, we go beyond the quantification of triple junctions associated with \(\Sigma 3\) boundaries, and evaluate advanced metrics associated with twin related domains.

A representative image with TRDs for specimen E reconstructed using ARPGE is shown in Figure 5a. Here, the grain boundaries can be identified using the legend shown in Figure 5b, with the numbers representing \(n\) in \(\Sigma 3^n\). Figure 5c is the largest TRD highlighted in Figure 5a (with an arrow). All grains in this TRD are related by \(\Sigma 3^n\) misorientations, and are connected by \(\Sigma 3\) boundaries; they are shown on a twinning tree for this TRD using a fractal representation (Figure 5d). All parameters associated with the TRDs are automatically calculated using ARPGE and then averaged.

Table 3: Triple junction distribution in the fully consolidated specimens.

| Specimen | \(J_0\)  | \(J_1\)  | \(J_2\)  |
|----------|----------|----------|----------|
| A        | 36 (+3)  | 56 (+2)  | 8 (+1)   |
| B        | 36 (+3)  | 55 (+4)  | 9 (+1)   |
| C        | 31 (+2)  | 62 (+2)  | 7 (+1)   |
| D        | 29 (+3)  | 62 (+2)  | 9 (+1)   |
| E        | 28 (+1)  | 61 (+1)  | 11 (+1)  |
| F        | 36 (+3)  | 56 (+3)  | 8 (+1)   |

Figures 6a-f show the metrics (i.e., Length fraction of \(\Sigma 3\), \(\langle TRD \rangle\), \(\langle N_g \rangle\), \(\langle LLC \rangle\), \(\langle p \rangle\), and \(\langle a \rangle\)) quantifying the extent of multiple twinning for the evolutionary microstructural states of all specimens, and very clearly reveal the differences between them. It can be seen that the statistics for all powders HIPed at 950\(^\circ\)C are very similar, but begin to diverge as the HIPing temperature increases (Figures 6a-f). This is because, there is incipient plastic deformation of powders at 950\(^\circ\)C, and the corresponding microstructures are similar to those of the as-received powders. Figure 6a shows the evolution of length fraction of \(\Sigma 3\) boundaries as a function of HIPing temperature. While the length fraction in all specimens increases with HIPing temperature, the rate of increase (with temperature and hence, time) is different. In the fully HIPed condition, \(\Sigma 3\) boundary length fraction of C, D, and E is similar (\(~\sim 51\%) and higher than those for A, B, and F (\(~\sim 45\%)). Specimen C has the largest \(\langle TRD \rangle\) size (46 \(\mu\)m), followed by D, E (\(~\sim 31 \mu\)m) then by F (23 \(\mu\)m), and A and B (19 \(\mu\)m), (Fig: 6b).

The average number of grains in the TRDs (i.e., \(\langle N_g \rangle\)) are lowest for A, B, and F (\(~\sim 3\) per TRD), while those for C, D, and E are much higher (3.5, 4, and 4.5, respectively), Figure 6c. The average length of longest chain \(\langle LLC \rangle\) that represents the twinning
Figure 5: (a) Representative map showing the reconstructed TRDs for specimen E (b) the legend for identifying various boundaries, with the numbers representing $n$ in $\Sigma 3^n$ (c) largest TRD identified in the map along with grain numbers (d) fractal representation of twin chains in the TRD identified by the arrow in (c); here the grains numbers occupy red circles while the squares show the operators that connect the grains. $\Sigma 3$ corresponds to 1a, $\Sigma 9$ corresponds to 2a and so on (Colour online).
Figure 6: Evolution of various parameters associated with TRDs for the specimens as a function of HIPing temperature: (a) length fraction of Σ3 grain boundaries (b) average sizes of TRDs, (c) number of grains in a TRD, (d) length of longest chain, (e) twinning polysynthetism, and (f) twinning anisotropy. Lines joining the data points are only a guide to the eye. Here FC stands for ‘Fully Consolidated’.
order in a TRD, is greatest for E (1.62), followed by D (1.55) and C (1.43), and then by F, A, and B (0.98, 0.9, 0.88, respectively), Figure 6d. The twins in E, D, and C are more polysynthetic ($\langle p \rangle$ values are 1.29, 1.27, 1.26, respectively) than F, A, and B (1.16, 1.15, 1.13, respectively), Figure 6e. The average values for twinning anisotropy ($\langle a \rangle$) are in the following order: E, C, D (1.83, 1.66, 1.55, respectively), and F, A, and B (1.44, 1.39, 1.39, respectively), Figure 6f. Figures 6a-f also reveal that the propagation of twin chains is the most during the dwell time of HIPing cycle.

As mentioned previously, powder processed materials usually contain an order of magnitude more oxygen than cast and forged components. Our results show the specimens containing numerous oxide inclusions around 0.5 $\mu$m in diameter, enriched in manganese, although their precise chemical composition was not evaluated (Figure 7). Most of the inclusions were dislodged during grinding and polishing, leaving “holes” in the matrix, suggesting that they are very weakly bonded to the matrix. In order to estimate their fraction in the specimens, the “holes” were treated as oxide inclusions. Their statistical analysis is shown in Table 4. Their density and average size increase with the oxygen content in the powders, while the nearest neighbour (NN) distance decreases. Although the oxygen content in specimens B and F are very low compared to others (110 ppm and 150 ppm, respectively), the errors associated with inclusion density and NN distance in them are high. This was observed to be the consequence of their presence along prior particle boundaries (PPBs). Specifically, it was observed that if the PPBs were present in the specimens, inclusions decorated them, and were more closely spaced along them. Moreover, PPBs were only observed in specimens with broad powder size distributions (A, B, and F) and virtually nonexistent in specimens C, D, and E. Interestingly, specimen D which has the highest oxygen content (600 ppm) and a narrow powder size distribution, was homogeneously recrystallized and did not contain any PPBs. In other words,
the inclusions were randomly distributed in the bulk. Representative images for specimen F (110 ppm) and D (600 ppm) are shown in Figure 8. PPBs are seen in Figure 8a which contains fewer annealing twins than in Figure 8b. The results thus indicate a correlation between (a) powder size distribution and the extent multiple twinning (i.e., recrystallization) (b) oxygen content in the powder and the inclusion density, and (c) spatial distribution of oxide inclusions and recrystallization.

Table 4: Inclusion distribution in the fully HIPed microstructure.

|                   | A    | B     | C     | D     | E     | F     |
|-------------------|------|-------|-------|-------|-------|-------|
| Oxygen content    | 290 ppm | 110 ppm | 200 ppm | 600 ppm | 220 ppm | 150 ppm |
| Inclusion density (x10^9/m²) | 16 (±3) | 7.5 (±5.5) | 8.9 (±1) | 21.5 (±3.6) | 12.1 (±2.8) | 20 (±10.7) |
| Nearest neighbour distance (µm) | 3.64 (±0.39) | 4.82 (±3.9) | 4.23 (±0.52) | 3.23 (±0.38) | 4.18 (±0.58) | 2.11 (±0.9) |
| Average size (µm) | 0.46 (±0.06) | 0.43 (±0.09) | 0.49 (±0.05) | 0.5 (±0.04) | 0.41 (±0.07) | 0.36 (±0.1) |

Figure 8: Secondary electron showing the effect of recrystallization on inclusion distribution in fully HIPed specimen of powder F (a) and D (b).

4 Discussion

4.1 Effect of powder size distribution on recrystallization during HIPing

Depending on the powder size distribution, the extent to which the powder compact is plastically strained during early stages of HIPing varies. Specifically, powders with low tap density experience more strain as they can accommodate more plastic flow than powders with high tap density. In fact, Li and Funkenbusch, in their HIP model, have demonstrated that the deformation of powder in a monosize distribution could be almost
twice of that in a bimodal distribution [6]. Since the stored energy increases with the
amount of plastic strain experienced, so will the driving force for recrystallization during
HIPing. In other words, powder size distribution has a strong effect on the extent of
multiple twinning during HIPing. From Fig. 1 and Table 2, it can be seen that the powders
considered in this study have different size distributions and consequently, varied tap
densities (i.e., in the following order: C < D < E < B < F < A). Therefore, the metrics
quantifying multiple twinning can be expected to follow a similar trend, although it
must be noted that locally, powders experience strains that will be different from the
macroscopic strain due to the shrinkage of canisters during HIPing. In other words, while
the differences in tap density/powder size distribution give an indication of the relative
strains, the development of multiple twinned microstructure depends on the energy stored
locally. More specifically, it is analogous to the dependence of applied strain on the
differences in the extent of multiple twinning observed in the grain boundary engineering
studies of low-medium SFE materials (e.g., [38, 39, 40]).

From Table 3, which shows the distribution of various triple junction types, it can be
seen that $J_1$ type junctions are higher in specimens C, D, and E compared to A, B, and F.
In other words, the number of annealing twins is higher in C, D, and E than in A, B, F.
The average number of grains, $\langle N_g \rangle$, within TRDs is highest in E, followed by D and C,
while they are lowest for A, B, and F. It suggests that the twin chains in specimens C, D, E
have propagated more than those in A, B, and F. This is further confirmed if we compare
$\langle LLC \rangle$, $\langle p \rangle$, and $\langle a \rangle$. It is seen that these parameters are nicely grouped for C, D, E (a
shade higher), and A, B, F (comparitively lower), clearly revealing that broad powder
size distributions, owing to higher tap densities (and consequently, experiencing lower
plastic strains during HIPing), have lower driving force for recrystallization compared
to powders with lower tap densities (Table. 2). In summary, a narrow powder size
distribution with low tap density produces a homogeneously recrystallized microstructure.
Such a microstructure is beneficial from the point of view of enhanced resistance towards
material degradation mechanisms that propagate intergranularly (e.g., stress corrosion
cracking).

4.2 Mechanism of formation of oxide inclusions

As previously pointed out, during the early stages of HIPing, plastic deformation of
powders is initiated at their contact points with other powders. The temperature and
pressure at which the deformation starts and proceeds depends on the elevated tempera-
ture strength of the material and the powder size distribution. Hedberg et al. have
shown that gas atomised 316L powder surface consists of a homogeneous layer (< 10 nm)
of iron/manganese/chromium oxide [41]. Because of the presence of a continuous oxide
film, complete bonding between powders during their incipient deformation at elevated
temperature (i.e., at \( \sim 950 \)°C in this study) cannot occur. The oxide layer is disrupted during HIPing and metal to metal contacts are established. In order to reduce the interfacial energy between the metal and oxide layer, the disrupted oxide layer coalesces to form inclusions (\( \sim 0.2 \) µm) at the inter-particle boundaries, Fig. 9a.

![Representative BSE images highlighting the effect of recrystallization on the randomisation of inclusions in the microstructure (HIPed specimens from powder D). Large dark regions between the powders in (a) and (b) correspond to the porosity resulting from partial consolidation.](image)

With increasing temperature, powders experience larger strains and simultaneously begin to recrystallize, Fig. 9b, where the presence of annealing twins can be seen. However, the inclusions are still present at the inter-particle boundaries. At 1050°C, depending on the local plastic strain experienced by the powders, the recrystallized grains begin to grow past the inter-particle boundaries with the inclusions still arranged on them, Fig. 9c. However, after the full HIP cycle (i.e., HIPed at 1160°C, 103 MPa, 4 hours dwell time), the inclusions are distributed in the bulk, depending on the homogeneity of recrystallization, albeit there are few regions where they are situated at the grain
boundaries (Fig. 9d), indicating that inclusions are dragged by moving grain boundaries during recrystallization. Moreover, their average size has also increased to around 0.5 µm (refer Table 4), further suggesting that the coarsening of inclusions is diffusion and/or coalescence induced. We note that the growth of oxide inclusions during recrystallization has previously been reported for Ni-Cr and Cu-Si alloys, respectively [42, 43]. Once nucleated, the oxide inclusions must be quite stable at HIPing temperatures (i.e., 1160°C). This is because, if they were to be in solid solution, there is no a priori reason for their re-nucleation exactly at PPBs during the cooling step of HIPing process. We further note the presence of sub-micron sized cellular microstructure in the partially HIPed specimens (evident in Fig. 9a-c), which is the consequence of rapid solidification of the melt during gas atomisation process (i.e., powder production process). However, the microstructure homogenizes during the “dwell time” of the HIPing cycle.

4.2.1 Effect of oxygen content on recrystallization

The powders considered in the present study had oxygen content ranging from 110 ppm (powder B) to 610 ppm (powder D), Table 1. We now compare the metrics for specimens B and D (in other words, the lowest and highest oxygen content specimens, respectively) to specifically understand the effect of oxygen content in the powder on recrystallization during HIPing. Firstly, the microstructures qualitatively reveal that specimen B has recrystallized inhomogeneously while specimen D has a more homogeneous microstructure. The triple junction distributions (Table. 3) quantitatively indicate a higher fraction of Σ3 boundaries in D compared to B (fraction of \( J_0 \) is lower while \( J_1 \) is higher in D). In addition, the advanced metrics related to TRDs (i.e., \( \langle TRD \rangle \), \( \langle N_g \rangle \), \( \langle LLC \rangle \), \( \langle p \rangle \), and \( \langle a \rangle \)) show that the extent of multiple twinning is much larger in specimen D than in B, clearly indicating that the oxygen content has no effect on recrystallization. On the other hand, the powder size distribution of B is much broader than D (tap densities of 5.5 g/cc and 5.2 g/cc, respectively) due to which the plastic strain (and hence stored energy) in the compact of powder B during HIPing will be lower than powder D. This in turn results in larger driving force for recrystallization in D and hence, better twinning statistics than in B.

Rao et al. studied the effect of oxygen content on the microstructure of powder HIPed Inconel 718 and concluded that the effect of oxygen content in the powders is to retard recrystallization [12]. Specifically, using three powders with different oxygen contents (275 ppm, 180 ppm, and 140 ppm) they observed (qualitatively, from the micrographs) that the extent of recrystallization decreased with an increase in oxygen content, although the effect of powder size distribution was ignored in their analysis (refer Figures 2c and 3c in [12]). More specifically, the tap density was highest for the powder with 275 ppm oxygen (broad size distribution) and lowest for 140 ppm (narrow size distribution), which clearly
suggests that the driving force for recrystallization for the powder with lowest tap density will be greater than that for highest tap density. We believe that the results of Rao et al. tacitly demonstrated the dependence of powder size distribution on recrystallization, but were interpreted otherwise. In summary, our results clearly demonstrate that the oxygen content in the powders has no effect on the degree of plastic strain experienced during initial stages of HIPing.

### 4.2.2 Spatial distribution of oxide inclusions during consolidation and the effect of recrystallization

While powder HIPed components can be manufactured in near net shape, one of the challenges in using them in high integrity structural applications is their lower toughness values (attributed to the presence of oxide inclusions) compared to cast and forged ones, although it is emphasized that the toughness values in many cases are still above the ones stipulated in ASME codes. See for e.g., [17, 18] for the case of powder HIPed 316L and 304L. The size, volume fraction, coherency with the matrix, and spatial distribution of inclusions have a strong effect on the material toughness. Figure 8 revealed the effect of recrystallization on the spatial distribution of inclusions. Specifically, Fig. 8a, which corresponds to powder F (highest tap density and lowest oxygen content), showed a particular region where the inclusions are decorated at PPBs while Fig. 8b, which corresponds to powder D (lowest tap density and highest oxygen content), showed a random distribution of the inclusions. The triple junction distribution and twinning statistics reveal that the scale and homogeneity of recrystallization in specimen D is much better than specimen F, clearly demonstrating that recrystallization strongly affects the spatial distribution of the inclusions.

During powder HIPing, a general view regarding the effect of inclusions at PPBs is that they act as pinning centres for moving grain boundaries [12, 19, 44, 45, 46]. Indeed, one of the ways suggested to promote the movement of grain boundaries past them is to perform HIPing at temperatures where the driving force for the boundary movement is greater than the pinning force of the inclusions at PPBs, although it is recognized that it results in increased grain size [13, 19]. However, pushing of particles by moving grain boundaries during solid state phase transformations (e.g., recrystallization, allotropic transformation), and diffusional movement of inert particles have previously been reported [42, 43, 47, 48, 49, 50, 51, 52]. These studies focused on the effect of recrystallization in a material with homogeneous dispersion of inclusions. However, in powder HIPing, the opposite holds; i.e., the inclusions nucleate at inter-particle boundaries (or PPBs) where the contact stresses are the highest (provided there is room for plastic flow), and are then pushed away from PPBs as a result of recrystallization during the dwell time of the HIPing cycle.
While it is known that, in low-medium SFE materials, recrystallization occurs during HIPing, previous studies have not explicitly correlated the spatial distribution of inclusions to multiple twinning. In fact, results from this study have clearly demonstrated that, although the inclusions nucleate at the inter-particle boundaries, their spatial distribution after the HIPing cycle (i.e., after complete consolidation) is strongly affected by the extent of recrystallization, which is governed by the powder size distribution. Therefore, in a much broader context, the naturally occurring mechanisms of phase transformation and related phenomena during powder processing of certain alloy systems (e.g., recrystallization in low-medium SFE materials, transformation mismatch plasticity in titanium alloys and martensitic/bainitic steels; see for e.g., [53, 54]) can be used for optimizing the manufacturing process and controlling the microstructural evolution.

4.3 Implications of the present study for powder-HIPed components with low-medium SFE

One of the main results originating from the present study is that, the homogeneity of recrystallization during HIPing is governed by the powder size distribution and not by the amount of oxygen content. More importantly, the spatial distribution of oxide inclusions and their volume fraction are governed by the extent of recrystallization and the amount of oxygen content in the powders, respectively. We believe that these observations are very important and are of high value, and can be used to potentially manufacture near net shape components for critical applications only by powder-HIPing. As an example, we identify its applicability to powder processed nickel-base superalloys.

Highly alloyed nickel-base superalloys which are of interest for demanding environments, are processed using powder metallurgy techniques to avoid casting induced segregation of alloying elements. However, presence of PPBs in the as-HIPed condition has been a perennial problem, and has necessitated post-HIP thermomechanical processing of these alloys, which not only increases the manufacturing cost, but also restricts the efficient use of HIPing as a process for manufacturing near net shape components. Specifically, processing involves canning of gas-atomized powders and HIPing, followed by hot extrusion and/or isothermal forging [20, 21]. Post-HIP thermomechanical processing is performed in order to “break” the PPB network present in the as-HIPed products (i.e., via recrystallization). Hot-working of these alloys after HIPing has been shown to be effective not only in enhancing the fraction of Σ3 boundaries, but also in breaking the PPB networks [22, 55, 56], clearly suggesting a direct correlation between the extent and homogeneity of recrystallization and the absence of PPBs.

In our study on powder HIPed 316L, by considering different powder size distributions, and by just HIPing, we clearly demonstrated a correlation between the extent and
homogeneity of recrystallization, and the absence of PPBs (Fig. 3, Table. 3, Fig. 6, and Fig. 8). More specifically, a narrow powder size distribution has produced a more homogeneously recrystallized microstructure free from PPBs than the broadly distributed ones. While a broad powder size distribution (in other words, distribution with high tap density) provides better control over the shape change of the component after HIPing\(^2\), modelling can be used to predict the shape change so that the initial canister can be designed in order to obtain near net shape after HIPing even with narrow powder size distribution. The benefit of uniform recrystallization during HIPing is two-fold. First, the increased fraction of Σ3 grain boundaries due to multiple twinning during HIPing can improve high cycle fatigue crack propagation behaviour [57]. Second, the inclusions that nucleate at inter-particle boundaries will more likely be dispersed homogeneously after HIPing (because of the absence of PPBs), potentially improving the ductility, although it must be noted that any improvement in ductility is dependent on the volume fraction of inclusions.

While it can be argued that the propensity for twinning depends on the SFE of the material, and that there would be differences in microstructural topology between 316L and Ni-base superalloys, we emphasize that hot working has a strong effect on grain boundary network topology, and we also underscore that HIPing is essentially a thermomechanical process. For example, within the context of twinning induced grain boundary engineering in castings, the effect of process variables (e.g., strain, annealing temperature and time, or strain rate and hot deformation temperature) on grain boundary connectivity can be inferred by referring to [58] for a general overview and [59, 60, 61] specifically for Ni-base superalloys. In the case of powder HIPing, the process variables are powder size distribution, temperature, pressure, and time.

The effectiveness of HIPing as a thermomechanical process can therefore be taken advantage of, to reduce/potentially eliminate the problem of PPBs by proper choice of powder size distribution, chemistry, and HIPing parameters. More specifically, the inclusion density can be reduced by using powder with low oxygen content while their spatial distribution can be randomized by optimizing the HIP process variables to promote homogeneous recrystallization. That is, in addition to using narrow powder size distribution, the stored energy in the deformed powders can be tuned by adjusting the way in which pressure and temperature are applied in order to promote multiple twinning and eliminate PPBs during HIPing. Such an optimized process reduces the number of steps and overall cost of manufacturing components for critical applications, such as hot sections of gas turbines used in aerospace industry.

\(^2\)During HIPing, shrinkage of the canister is more uniform for a broad powder size distribution compared to a narrow size because of the lesser extent of plastic flow offered.
Conclusions

The aim of the present study was to understand the effect of powder characteristics and oxygen content on the microstructural evolution during hot isostatic pressing of 316L austenitic steel. The main findings are summarized as follows:

- The principal mechanism by which the microstructure evolves during HIPing is recrystallization, the homogeneity of which, is strongly dependent on the powder size distribution.

- Fully HIPed specimens with narrow powder size distribution (i.e., lower tap density) were more homogeneously recrystallized than the ones produced using broad size distribution (i.e., higher tap density). Prior particle boundaries (PPBs) were more frequently observed in specimens HIPed with broad powder size distribution than the ones HIPed with narrow distribution.

- The density of oxide inclusions in the fully HIPed specimens increased with the oxygen content in the powders, with their spatial distribution strongly dependent on the homogeneity of recrystallization.

- The oxide inclusions rich in manganese nucleated on inter-particle boundaries (in other words, PPBs) during early stages of HIPing, and depending on whether or not there was sufficient stored energy, were pushed by the moving grain boundaries during recrystallization.

Finally, as demonstrated here, homogeneous recrystallization during HIPing results in random distribution of oxide inclusions and a microstructure free from PPBs. This is highly relevant from the point of view of manufacturing PPB free near net shape components for critical applications, since it would eliminate the need for post-HIP thermo-mechanical processing (in order to break the PPBs) and reduce manufacturing costs.

Acknowledgements

The authors would like to thank the EPSRC for funding through New Nuclear Manufacturing (NNUMAN): EP/J021172/1, and Manufacture using Advanced Powder Processes (MAPP): EP/P006566/1. Rolls-Royce is also thanked for supplying the 316L forging for this study.

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Graphical Abstract

Broad powder size distribution (PSD1)  Narrow powder size distribution (PSD2)

Average values of length of longest chain, polysynthetism, and twinning anisotropy for PSD1 and PSD2.