Material-process-property relationships of 17-4 stainless steel fabricated by laser-powder bed fusion followed by hot isostatic pressing.

Harish Irrinki

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MATERIAL-PROCESS-PROPERTY RELATIONSHIPS OF
17-4 PH STAINLESS STEEL FABRICATED BY LASER-POWDER BED FUSION
FOLLOWED BY HOT ISOSTATIC PRESSING

By
Harish Irrinki

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MATERIAL-PROCESS-PROPERTY RELATIONSHIPS OF
17-4 PH STAINLESS STEEL FABRICATED BY LASER-POWDER BED FUSION
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By

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A Dissertation Approved on

July 20, 2018

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I dedicate this thesis to

my parents Sri Rama Murthy and Dhana Lakshmi Irrinki and grandfather Sri Gopala Krishna Murthy Sanku who have supported through the various obstacles in my life with their incessant generosity, support, and love.
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ABSTRACT

MATERIAL-PROCESS-PROPERTY RELATIONSHIPS OF
17-4 PH STAINLESS STEEL FABRICATED BY LASER-POWDER BED FUSION
FOLLOWED BY HOT ISOSTATIC PRESSING

Harish Irrinki
July 20, 2018

17-4 PH stainless steel is commonly used in medical, tooling, automotive, chemical and aerospace industries due to its excellent strength and corrosion properties. Additive manufacturing processes such as laser-powder bed fusion (L-PBF) have gained attention and importance due to the potential to produce complex-shaped three-dimensional parts for various industries. In order to manufacture three-dimensional components from 17-4 PH stainless steel powder using L-PBF, it is critical for design and manufacturing engineers to have an awareness of various material options and corresponding processing and post-processing conditions to obtain useful mechanical properties from the process. The goal of this dissertation is to establish a fundamental understanding of the material-process-property relationships of 17-4 PH stainless steel processed by laser-powder bed fusion (L-PBF). The investigation was carried out to understand the effects of post-processing
treatment cycles such as hot isostatic pressing (HIP) on the densification, mechanical properties, corrosion properties and microstructures of L-PBF parts fabricated at various energy densities using 17-4 PH stainless steel gas- and water-atomized powders. The microstructure formation and its impact on mechanical and corrosion properties due to different HIP treatment cycles were studied for 17-4 PH stainless steel gas and water-atomized L-PBF parts at various energy densities. The most significant aspect of this work is that density, microstructures, mechanical and corrosion properties of 17-4 PH stainless steel can be vastly improved and reliable properties can be achieved irrespective of starting powder attributes and L-PBF process conditions by using HIP treatment. Further, the properties obtained after the HIP treatment was less sensitive to variations in powder characteristics (size distribution and shape) and energy density during processing and were superior to known reported properties of 17-4 PH stainless steel fabricated by L-PBF, powder metallurgy, metal injection molding, or wrought samples.
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CHAPTER 1
INTRODUCTION

Laser powder bed fusion (L-PBF), popularly known as selective laser melting (SLM) process uses a focused laser beam to melt metal powders layer-by-layer into three-dimensional parts, based on CAD file [1]–[4]. To fabricate three-dimensional components using the L-PBF process, a design engineer needs understand the metal powder-laser interactions during the L-PBF process and their impact on the mechanical properties and microstructures of the L-PBF parts. In L-PBF, it is widely recognized that laser-powder interactions are affected by processing conditions (e.g. laser power, scan speed, hatch spacing and layer thickness) and powder characteristics (e.g. size, shape, and purity) for the successful fabrication of parts with desired properties [5]–[9]. However, there remains a wide gap in the understanding material-process-property relationships affected by powder attributes and processing conditions in L-PBF.

Typically, gas-atomized powders with a spherical shape and narrow particle size distribution (5 to 45 µm) have been preferred for fabricating parts using L-PBF [2], [3], [6], [8], [10]–[12]. However, such powders are inherently expensive as they use large amounts of inert process gas as well as only a small fraction (typically 30 ± 15 µm) of the production lot in each atomization run [2], [3], [6], [13].
On the other hand, water-atomized powders are relatively less expensive but are irregular in shape with variation in particle size distribution. The ability to fabricate high-density L-PBF parts from irregular water-atomized powders with properties competitive to parts fabricated using gas-atomized powders has not been convincingly demonstrated in the open literature [2], [3], [6], [13]. In this regard, previous studies reported by our group have shown the feasibility of fabricating L-PBF parts using 17-4 PH stainless steel fine water-atomized powders and achieving the properties similar or higher than 17-4 PH stainless steel gas-atomized parts in certain processing conditions, e.g. at an energy density of 104 J/mm$^3$ as shown in Figure 1.1 [1]–[4]. However, some porosity and variation in phase content in the microstructure still existed in water-atomized powders that impacted the mechanical properties of the 17-4 PH stainless steel L-PBF parts, especially at lower energy density, e.g. at an energy density of 64 J/mm$^3$ (Figure 1.2). The results obtained from these research studies were published in three peer-reviewed journal papers [1], [2], [14].
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In powder metallurgy, hot isostatic pressing (HIP) is routinely performed to eliminate porosity and anisotropy in the metal parts. Several HIP studies have emerged in recent times to improve the densification and mechanical properties of L-PBF parts using various materials such as Hastelloy, 316L stainless steel, CoCr and Ti-6Al-4V alloys [15]–[24].
However, no study in the literature studies the effect of HIP on the densification, mechanical properties and microstructures of 17-4 PH stainless steel L-PBF parts. In order to address the above gap, the current thesis builds off our earlier work [25] focuses on the effects of HIP and heat treatment on the densification, mechanical properties, corrosion properties and microstructures of the 17-4 PH stainless steel gas and water-atomized L-PBF parts when fabricated at various energy densities.

**Chapter 2** presents an understanding on the effects of a standard HIP cycle on the densification, mechanical properties and microstructures L-PBF parts fabricated from 17-4 PH stainless steel gas and water-atomized powders at various L-PBF processing conditions. The results from the current study will provide a better understanding of the effect of post-processing method such as HIP on the densification, mechanical properties and microstructure of L-PBF parts. The work presented in **Chapter 2** has been submitted to a peer-reviewed journal *Journal of the American Ceramic Society* in 2018 and is currently under review.

**Chapter 3** builds from the work presented in **Chapter 2** and the effects of modified HIP cycle is scaled to L-PBF parts fabricated from four different powders with varying shape and size attributes at different processing conditions. **Chapter 3** presents the effect of HIP on the densification behavior, mechanical properties, phase and microstructure development of the 17-4 PH stainless steel gas- and water-atomized parts processed by L-PBF. A part of the research presented in **Chapter 3** presented in “Additive Manufacturing-Powder Metallurgy Conference” 2018 and the entire work presented in **Chapter 3** is currently under preparation for submission to the *Additive Manufacturing* refereed journal.
Chapter 4 of the thesis examines the role of powder characteristics and processing conditions on the corrosion properties of 17-4 PH stainless steel fabricated using L-PBF. Our previous studies indicated that powder characteristics and processing conditions are known to strongly influence the densification and mechanical properties of parts fabricated by the L-PBF process. However, the influence of powder attributes and L-PBF process parameters on the corrosion performance of parts has not been studied extensively. In this regard, the work present in Chapter 4 compares the corrosion performance of the 17-4 PH stainless steel L-PBF parts fabricated from powders of different characteristics such as shape, size and the method utilized for atomization (gas-atomized or water-atomized) at various processing conditions. The corrosion performance of the L-PBF parts was evaluated using corrosion current, polarization resistance and corrosion rate values from the linear sweep voltammetry curves. The collected data from the Chapter 4 is expected to provide a better understanding of the effect of porosity, density and microstructures on the corrosion performance of 17-4 PH stainless steel L-PBF gas- and water-atomized parts. The work presented in Chapter 4 was published in the peer-reviewed journal Progress in Additive Manufacturing in 2018 [26].

Chapter 5 of this thesis builds form the wrok presented in Chapters 3 and 4 and studies the effects of modified hot isostatic pressing on the corrosion performance of 17-4 PH stainless steel gas- and water-atomized L-PBF parts. The entire work presented in Chapter 2 is currently under preparation for submission to the Journal of the Minerals, Metals and Materials Society referred journal.
The work presented in Appendix A is a collaboration work with Prof. Somayeh Pasebani’s group at the Oregon State University. The work focuses on the effects of post-processing (solution annealing and aging) on mechanical properties and microstructures of 17-4 PH stainless steel gas- and water-atomized L-PBF parts when fabricated at various energy densities. The work was published in peer-reviewed journal Additive Manufacturing in 2018 [3].

Appendix B reports the mechanical properties and microstructures data of the L-PBF parts fabricated from 17-4 PH stainless steel gas- and water-atomized powders when they are post-processed using various heat-treatment cycles. The work presented in Appendix B is an expansion of the work in Appendix A and will be the subject of future work to scale it into a refereed journal paper.

The desire to improve the sustainability and efficiency of the L-PBF process drives the requirement for recycling of starting powders that are unused during part fabrication. The data presented in Appendix C provides an initial understanding (qualitatively and quantitively) on how the powder shape and size varies as a function of starting powder characteristics and L-PBF processing conditions.

Appendix D provides data on the applications that have been developed using L-PBF process for aerospace, healthcare and automotive industries. The applications were developed based on materials-design-process relationship work presented in this thesis. A part of this work was published in “Powder Metallurgy Review” magazine in 2018 [27].
CHAPTER 2
EFFECTS OF HOT ISOSTATIC PRESSING ON THE DENSIFICATION, MECHANICAL PROPERTIES AND MICROSTRUCTURES OF 17-4 PH STAINLESS STEEL FABRICATED BY LASER-POWDER BED FUSION

2.1 INTRODUCTION

Laser powder bed fusion (L-PBF), alternately known as selective laser melting (SLM) is a powder-based process that uses focused laser energy to melt the metallic powders into solid parts. In the L-PBF process, the laser-powder interactions are largely affected by processing conditions such as laser power, scan speed, hatch spacing and layer thickness [28], [9], [29]–[34], [7], [35]. In general, the above processing conditions along with powder characteristics such as size, shape, and purity, significantly determine the density, microstructures and properties obtained from L-PBF parts [6], [32]–[34], [36]–[41]. In our former work [1], [2], a comprehensive study was performed to understand the role powder characteristics such as powder type, shape and size along with L-PBF processing conditions on densification, mechanical properties and microstructure of 17-4 PH stainless steel L-PBF parts. It was found that the % theoretical density, ultimate tensile strength, hardness of L-PBF parts are sensitive to L-PBF processing conditions and starting powder shape, size and type. An
important result of that study was a large variation in density, mechanical properties and microstructures were observed L-PBF parts printed with different powder attributes. Most of the research studies indicated that defects and anisotropy in the mechanical properties of L-PBF parts could be eliminated using post-processing methods such as hot isostatic pressing (HIP). Tillmann et al [20] discussed the efficiency of the HIP in decreasing the porosity and enhancing the mechanical properties of IN718 L-PBF parts. Many research groups performed HIP studies on various materials such as HastealloyX [15], [18], [42], 316L stainless steel [16], [17], CoCr [19] and Ti-6Al-4V alloys [23], [24]. However, there has been limited work performed on the elimination of the defects in 17-4 PH stainless steel L-PBF parts using HIP process [10], [22], [43].

In this regard, the present study was performed to understand the effect of HIP on the densification, mechanical properties and microstructures L-PBF parts fabricated from different powder size and types at various L-PBF processing conditions. The results from the current study will provide a better understanding of the effect of post-processing method such as HIP on the densification, mechanical properties and microstructure of L-PBF parts.

2.2 EXPERIMENTAL MATERIALS AND METHODS
The 17-4 PH stainless steel gas-atomized powder used for this study was supplied by 3D Systems and the 17-4 PH stainless steel water-atomized powder was produced by North American Hoganas. The particle size distributions of the 17-4 PH stainless steel powders used in this study are measured using a Microtrac S3000 particle size analyzer. A high-resolution FEI Quanta 600F scanning electron microscope (SEM) and a Bruker D8
DISCOVER X-ray diffraction (XRD) spectroscopy were used for morphology analysis of
the powders. L-PBF experiments were performed on a 3D Systems ProX 200 machine
under argon. The process parameters used in the L-PBF experiments consisted of laser
power, scan speed, layer thickness and hatch spacing as given in Table 2.1.

Table 2.1. L-PBF processing conditions and the corresponding energy densities used in
this study

<table>
<thead>
<tr>
<th>Laser power (W)</th>
<th>Scan speed (mm/s)</th>
<th>Hatch spacing (µm)</th>
<th>Layer thickness (µm)</th>
<th>Energy density (J/mm$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150</td>
<td>1550</td>
<td>50</td>
<td>30</td>
<td>64</td>
</tr>
<tr>
<td>150</td>
<td>1250</td>
<td>50</td>
<td>30</td>
<td>80</td>
</tr>
<tr>
<td>195</td>
<td>1550</td>
<td>50</td>
<td>30</td>
<td>84</td>
</tr>
<tr>
<td>195</td>
<td>1250</td>
<td>50</td>
<td>30</td>
<td>104</td>
</tr>
</tbody>
</table>

The laser power was varied between 150 and 195 W and scan speed was varied between
1250 and 1550 mm/s. The layer thickness and hatch spacing were kept constant at 30 µm
and 50 µm to fabricate tensile parts using the L-PBF process (Table 2.1). The set of process
parameters considered for the L-PBF experiments were further used to calculate laser
energy density using Equation 2.1.

$$E_p = \frac{P}{h \cdot v \cdot t}$$

where, $E_p$ is energy density (J/mm$^3$), P is laser power (W), v is scan speed (mm/s), t is layer
thickness (mm), and h is hatch spacing (mm).

Sixteen samples per type of powder were fabricated for each process condition. Thus, a
total of 256 samples were fabricated during the study. Of the 256 parts, 4 parts of each
powder type and process condition were selected for Archimedes density, mechanical
testing and microstructure analysis. The tensile samples were thermally stress relieved at
1200\(^0\) F for 1 hour in the air before their removal (electrical discharge machining) from the build plate. The tensile samples were cut by wire electrical discharge machining (EDM) into samples that were 0.68 m × 0.13 m × 0.318 m.

Standard HIP post-treatment was subsequently applied to the tensile samples using a commercial service provided by Quintus Technologies, USA. The HIP treatment was conducted for 2 h under a temperature of 1120 \(^0\)C and pressure of 105 MPa applied through Ar gas, followed by rapid cooling to 200 \(^0\)C at a rate of 100 \(^0\)C/min. The HIP treated tensile samples were analyzed for their mechanical properties as per ASTM E8M standard and physical properties as well as microstructures. Hardness testing was performed using a Rockwell ‘C’ hardness tester at 150 kg load. Tensile testing was performed using an Instron 5982 dual column machine.

Density measurements were conducted using a method based on Archimedes law on a Mettler Toledo XS104 weighing balance. The density values of the L-PBF parts reported in this paper were expressed as a percentage of the density of a 17-4 PH stainless steel cast part. The 17-4 PH stainless steel cast part was purchased from McMaster-Carr and density of the cast part was measured using the Archimedes method. The porosity of L-PBF parts was calculated using Equation 2.2.

\[
\phi = 1 - \frac{\rho (l)}{\rho (t)}
\]  

(3.2)

Where, \(\phi\) is the porosity of the L-PBF parts, \(\rho (l)\) is the density of the L-PBF parts, \(\rho (t)\) is the density of 17-4 PH stainless cast part.
XRD of the L-PBF parts was performed perpendicular to the build direction on the BRUKER D8 machine. Rietveld analysis was performed to quantify the phases in the XRD data. Microstructures of the L-PBF parts were analyzed using an Olympus BX53 microscope. Metallographic specimen preparation was carried out following standard procedures for microstructure characterization. The carpenter's etchant was used as an etchant to reveal the austenite and martensitic phases typically found 17-4PH stainless steel parts [26], [30]. The tensile testing of the L-PBF parts, microstructural characterization of the polished and etched L-PBF parts was performed in the horizontal plane perpendicular to the build direction.

2.3 RESULTS AND DISCUSSION

2.3.1 POWDER CHARACTERISTICS

The chemical compositions of the 17-4 PH stainless steel gas- and water- atomized powders are presented in Table 2.2.

<table>
<thead>
<tr>
<th>Powder type</th>
<th>C</th>
<th>Cr</th>
<th>Cu</th>
<th>Mn</th>
<th>Ni</th>
<th>P</th>
<th>S</th>
<th>Si</th>
<th>Nb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized</td>
<td>0.03</td>
<td>15</td>
<td>3.5</td>
<td>0.5</td>
<td>4.6</td>
<td>0.04</td>
<td>0.03</td>
<td>1</td>
<td>0.2</td>
</tr>
<tr>
<td>Water-atomized</td>
<td>0.2</td>
<td>18</td>
<td>4</td>
<td>0.1</td>
<td>3.5</td>
<td>0.01</td>
<td>0.01</td>
<td>0.3</td>
<td>0.3</td>
</tr>
</tbody>
</table>

The gas-atomized powder used in this study had a bimodal particle size distribution with a median particle size of 13 μm as shown in Figure 2.1. The water-atomized powder particle size distribution is monomodal and had median particle size 43 μm (Figure 2.1). Cumulative values of 10, 50, and 90% particle diameters are represented by $D_{10}$, $D_{50}$, $D_{90}$ values, respectively and are presented in Table 2.3. The morphology of the powders was
characterized using SEM and represented in Figure 2.2, showing the gas-atomized powders to be spherical while the water-atomized powders are irregular in shape.

![Figure 0.1](image)

Figure 0.1. Particle size distributions of the 17-4 PH stainless steel powders used in this study (a) gas-atomized powder $D_{50} = 13\, \mu m$ (b) water-atomized powder $D_{50} = 43\, \mu m$

Table 2.3. Particle characteristics of the 17-4 PH stainless steel gas- and water-atomized powders

<table>
<thead>
<tr>
<th>Powder</th>
<th>Particle distribution</th>
<th>Size width 2.56</th>
<th>Tap density (g/cc)</th>
<th>Apparent density (g/cc)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized (G)</td>
<td>$D_{10}$ 5 $D_{50}$ 13 $D_{90}$ 27</td>
<td>3.5</td>
<td>4 ± 0.05</td>
<td>3 ± 0.002</td>
</tr>
<tr>
<td>Water-atomized (W)</td>
<td>$D_{10}$ 26 $D_{50}$ 43 $D_{90}$ 67</td>
<td>6.2</td>
<td>3.7 ± 0.05</td>
<td>2.8 ± 0.05</td>
</tr>
</tbody>
</table>
Figure 0.2. SEM images of the 17-4 PH stainless steel powders used in this study (a) gas-atomized powder (D50 = 13 µm) (b) water-atomized powder (D50 = 43 µm)

Figure 2.3 shows the XRD patterns of gas- and water-atomized powders. The XRD pattern showed that martensite and austenite phases are present in both gas- and water-atomized powders. Rietveld analysis showed that the gas-atomized powder consisted of 70% martensite phase and 30% austenite phase whereas the water-atomized powders consisted of 80% austenite phase and 20% martensite phase. The phase difference between the gas- and water-atomized powders could be due to the difference in the atomization methods and chemistry of the starting powders. The cooling rate for water-atomized powders is 10 to 100 times higher than gas-atomized powders [13]. This variation in solidification rate resulted in high volume fraction of austenite phase in the water-atomized powders [33]. Furthermore, the austenite phase in gas-atomized powders could be due to the nitrogen gas which is used as atomizing media during the powder production process [33].

The porosity, phases, microstructure and properties of the L-PBF parts fabricated using 17-4 PH stainless steel gas and water-atomized powders were correlated with process
conditions using the energy density parameter calculated using Equation 2.2. The results are discussed in turn based on four values of increasing energy density.

![XRD analysis for the 17-4 PH stainless steel gas- and water-atomized powders](image)

Figure 0.3. XRD analysis for the 17-4 PH stainless steel gas- and water-atomized powders showing the presence of α and γ phases

2.3.2 DENSITY

The variation of % theoretical density of gas- and water-atomized L-PBF parts before and after HIP treatment with energy density is shown in Figure 2.4. The % theoretical density of water-atomized L-PBF parts after HIP treatment was found to increase with increased energy density while the % theoretical density of gas-atomized L-PBF parts remained constant (99 ± 0.5 %) with the increased energy density (Figure 2.4). Among all energy densities, the post-densification after HIP treatment was significant for water-atomized L-
PBF parts when compared to gas-atomized L-PBF parts (Figures 2.5 and 2.6). A comparison of Figure 2.5a and 2.5b, which respectively show the cross-section images of the water-atomized L-PBF parts before and after HIP treatments at energy densities 64 and 80 J/mm$^3$, reveals that the HIP treatment closed large irregular pores in the as-printed L-PBF parts. In comparison to % theoretical density of as-printed L-PBF parts, the % theoretical density of water-atomized L-PBF parts after HIP treatment had increased between 3 to 6 % whereas the % increase in gas-atomized L-PBF parts were 2 %. The HIP treatment provides the high temperature and hydrostatic pressure, which evidently eliminated the pores in the L-PBF parts according to creep and diffusion densification law [28]. However, 3 to 4 % of porosity still exists in water-atomized L-PBF parts after the HIP treatment at energy densities 64 and 80 J/mm$^3$ as shown in Figure 2.5a and 2.5b. Furthermore, HIP experiments with increased temperature and hold time on various water-atomized L-PBF parts are currently being performed in our group and results will be reported elsewhere.

A notable aspect of this data is the highest % theoretical density (99 ± 0.5 %) after HIP treatment was observed for both gas- and water-atomized L-PBF parts at energy densities of 84 and 104 J/mm$^3$. Tillmann et al.[20] suggested that practically it might be impossible to achieve a 100 % dense L-PBF parts by HIP treatment. The possible reason for this being the open surface porosity and entrapped gas pores in the as-printed L-PBF parts hinder the post-densification during HIP Figures 2.5c, 2.5d and 2.6. During the HIP process, the heat treatment leads to an increase in pressure and expansion of these pores. This increase in inner pressure counteracts the densification during HIP process, thereby resulting in micro pores [15], [20], [23], [24], [47], [48].
Nevertheless, the results of the present study show that HIP treatment is effective in achieving high dense (99 ± 0.5 %) gas- and water-atomized L-PBF parts. These results indicate that using same processing and post-processing conditions, L-PBF parts with similar density can be achieved using inexpensive coarser water-atomized powders compared to relatively expensive finer gas-atomized powders that are typically used for L-PBF.
Figure 0.5. Optical micrographs of HIP and as-printed water-atomized $D_{50} = 43\mu m$

Figure 0.6. Optical micrographs of HIP and as-printed gas-atomized $D_{50} = 13\mu m$

2.3.3 MECHANICAL PROPERTIES AND MICROSTRUCTURES

The variation in the ultimate tensile strength of the L-PBF gas- and water-atomized parts before and after HIP treatment with energy density is shown in Figure 2.7. The ultimate
tensile strength of the L-PBF water-atomized parts after HIP treatment increased with increase in energy density and varied between 700 and 1050 MPa. Furthermore, at energy densities of 64, 80 and 84 J/mm$^3$, there was 20 to 50 % increase in the ultimate tensile strength of the L-PBF water-atomized parts after HIP treatment % when compared to the tensile strength of water-atomized L-PBF parts before the HIP treatment (Figure 2.7). This significant increase in the ultimate tensile strength of the water-atomized L-PBF parts after the HIP treatment could be attributed to the increase in density of L-PBF parts (3 to 6 %) (Figure 2.4). This relation between the ultimate tensile strength and the porosity of the L-PBF parts after the HIP treatment was observed in previous studies [15], [16], [18], [19], [21], [23], [24], [48].

Figure 0.7. Variation of the ultimate tensile strength of 17-4 PH stainless gas- and water-atomized L-PBF parts before and after HIP treatment as a function of energy density

However, the gas-atomized L-PBF parts after the HIP treatment showed significantly higher tensile strength (~1100 MPa) when compared to the tensile strength water-atomized L-PBF parts before and after the HIP treatment as shown in Figure 2.7. However, at the high energy density of 104 J/mm$^3$, the ultimate tensile strength of water-atomized L-PBF
parts (~950 MPa) after the HIP treatment was higher than the tensile strength as-printed gas-atomized L-PBF parts (~950 MPa) but lower than the as-printed water-atomized L-PBF parts. Furthermore, at high energy densities of 84 and 104 J/mm³, high dense (99 ± 0.5 %) L-PBF parts were produced after the HIP treatment but the variation in tensile strength ranged between 950 and 1150 MPa (20 % variation). To understand this variation in the ultimate tensile strength of the high dense (99 ± 0.5 %) L-PBF parts, X-ray diffraction (XRD) analysis was performed on the L-PBF parts (perpendicular to the build direction) as shown in Figure 8a and 8b (water-atomized L-PBF parts) and 9a and 9b (gas-atomized L-PBF parts).

Figure 0.8. XRD patterns of HIP and as-printed 17-4 PH stainless water-atomized L-PBF parts perpendicular to the build direction produced at energy densities 64, 80, 84 and 104 J/mm³.
Figure 0.9. XRD patterns of HIP and as-printed 17-4 PH stainless gas-atomized L-PBF parts perpendicular to the build direction produced at energy densities 64, 80, 84 and 104 J/mm$^3$.

The XRD patterns of gas- and water-atomized L-PBF parts at high energy densities of 84 and 104 J/mm$^3$ were completely different before and after the HIP treatment Figure 2.8a and 2.8b and Figure 2.9a and 2.9b. After the HIP treatment, the gas-atomized L-PBF parts at high energy densities of 84 and 104 J/mm$^3$, consisted of 95% martensite phase and 5% austenite phase (Rietveld analysis) whereas the water-atomized L-PBF parts consisted of 45% martensite phase and 55% austenite phase. The high-volume fraction of retained austenite (55%) in the water-atomized L-PBF parts after the HIP treatment could have attributed to 20% decrease in the ultimate tensile strength of water-atomized L-PBF parts when compared to highly martensitic gas-atomized powder L-PBF parts. The gas-atomized L-PBF parts with 95% martensite phase after the HIP treatment showed significantly higher tensile strength when compared to the tensile strength of as-printed gas- and water-atomized L-PBF parts and HIP water-atomized L-PBF parts.
Figure 2.8 and 2.9 also show that the XRD patterns of water-atomized L-PBF parts after the HIP treatment were significantly different from the XRD patterns of L-PBF parts before the HIP treatment (Figure 2.8a) and starting powder (Figure 2.3). The quantitative analysis showed that the volume fraction of retained austenite phases in water-atomized L-PBF parts before the HIP treatment varied between 10 and 30 \% whereas after the HIP treatment the volume fraction of retained austenite varied between 45 and 55 \%. This increase in volume percentage of austenite phase in water-atomized L-PBF parts could be attributed to the cooling rates during the HIP treatment. However, the formation of retained austenite in gas-atomized L-PBF parts after the HIP treatment was minimal (5\%) owing to its predominantly high martensite content in as-printed L-PBF parts (Figure 2.9a) and the starting powder (Figure 2.3). On-going uniform rapid quenching (URQ) experiments are expected to favor the formation of martensite in water-atomized L-PBF parts because of the rapid cooling rates during the HIP process. The interesting result from this data is both starting powder characteristics and L-PBF processing conditions effects the XRD phases in as-printed and HIP L-PBF parts.
Figure 0.10 shows the hardness of HIP and as-printed 17-4 PH stainless gas- and water-atomized L-PBF parts produced at energy densities 64, 80, 84, 104 J/mm³.

Figure 0.10 shows the after the HIP treatment, hardness values of the gas-atomized L-PBF parts (~35 HRC) were higher than hardness values of the water-atomized L-PBF parts (~25 HRC). The predominant percentage (95%) of martensite in HIP treated gas-atomized L-PBF parts could have contributed to the high hardness when compared to HIP treated water-atomized L-PBF parts. However, the hardness values of the gas- and water-atomized L-PBF parts did not change significantly with the increase in the energy density when subjected to HIP treatment. This behavior remains in consistent with the XRD phases present in the respective parts. Furthermore, the 10 to 30% decrease in hardness of the HIP treated water-atomized L-PBF parts when compared to hardness as-printed parts (~25 HRC) with the increase in energy density could be correlated to the 45 to 55% decrease in martensite phase in HIP treated L-PBF parts.

Figure 2.11 shows the variation of the yield strength of L-PBF parts before and after HIP treatment with energy density is similar to the variation of tensile strengths of L-PBF parts.
The yield strength of both gas- and water-atomized L-PBF parts after HIP treatment increased with the energy density and are higher than the yield strength of as-printed L-PBF parts. After the HIP treatment, the yield strength of water-atomized L-PBF parts varied between 400 and 800 MPa whereas the yield strength of gas-atomized L-PBF parts remained constant at 850 ± 15 MPa. According to Hall-Petch relationship [49], the yield strength of the L-PBF parts can be significantly affected by grain size in their microstructures. Microstructure analysis was performed on the L-PBF parts before and after the HIP treatment to understand the variation in grain strength with energy density as shown in Figure 2.12 and 2.13. The average grain intercept (AGI) method was used to quantify the grain size in the L-PBF parts.

Figure 0.11. The yield strength of HIP and as-printed 17-4 PH stainless gas- and water-atomized L-PBF parts produced at energy densities 64, 80, 84, 104 J/mm$^3$. 

[Graph showing yield strength vs. energy density]
The microstructures of gas- and water-atomized L-PBF parts at low energy densities 64 and 80 J/mm$^3$ were significantly different when subjected to HIP treatment as shown in Figures 2.12 and 2.13. Figure 2.12a and 2.12b show that at energy densities 64 and 80 J/mm$^3$, the columnar grains in the as-printed L-PBF parts did not change significantly after the HIP treatment whereas the columnar grains present in as-printed gas-atomized L-PBF parts...
parts completely disappeared when subjected to HIP treatment (Figure 2.13a and 2.13b). At energy densities of 64 and 80 J/mm³, the fine equiaxed grains of 15 ± 1 µm in size were observed in gas-atomized L-PBF parts after the HIP treatment whereas large columnar grains of 20 ± 5 µm in size was observed in HIP treated water-atomized L-PBF parts. This change in grains and grain size could explain the variation in the yield strength of L-PBF parts when fabricated from different powders at energy densities 64 and 80 J/mm³. Furthermore, the increase in yield strength of HIP treated water-atomized L-PBF parts when compared to the yield strength of as-printed L-PBF parts could be attributed to the decrease in porosity after the HIP treatment rather than on the grain size. Further analysis will be performed to understand the variation in grain diffusion as a function of starting powder and results will be reported elsewhere.

However, at high energy densities of 84 and 104 J/mm³, the columnar grains in as-printed L-PBF gas-and water-atomized L-PBF parts were changed to equiaxed grains and showed a higher level of homogenization when subjected to HIP treatment (Figure 2.12 and 2.13). The equiaxed grains of 5 ~ 10 µm and 5 µm in size were observed in the HIP-treated water- and gas-atomized L-PBF parts, respectively. According to Hall-Petch strengthening mechanism [49], the decrease in the grain size could have contributed to the increase in yield strength of water- and gas-atomized L-PBF parts after the HIP treatment.

Figure 2.14 shows the elongation of L-PBF gas- and water-atomized parts decreased with energy density when subjected to HIP treatment. These decrease in elongation of the HIP treated water-and gas-atomized L-PBF parts could be correlated to the decrease in the grain size (Figure 2.12 and 2.13). The HIP treated water-atomized L-PBF parts with long
columnar grains showed higher elongation (15 to 17 %) at energy densities 64 and 80 J/mm³.

![Graph showing % elongation vs. energy density](image)

Figure 0.14. The % elongation of HIP and as-printed 17-4 PH stainless gas- and water-atomized L-PBF parts produced at energy densities 64, 80, 84, 104 J/mm³.

### 2.4 CONCLUSIONS

This work presents a comprehensive study of the densification behavior and microstructure development of 17-4 PH stainless steel gas- and water-atomized L-PBF parts when subject to HIP treatment. The important result from this data is that both starting powder characteristics and L-PBF processing conditions affect the density, mechanical properties and microstructures of as-printed and HIP L-PBF parts. At low energy densities of 64 and 80 J/mm³, after the HIP treatment, large irregular pores in as-printed water-atomized L-PBF parts were eliminated and density increased from 90 to 97 %. For all the energy densities, the gas-atomized L-PBF parts after the HIP treatment showed significantly higher tensile strength, yield strength and hardness when compared to water-atomized L-PBF parts properties. The high-volume fraction of retained austenite (55 %) in the water-atomized L-PBF parts after the HIP treatment could have attributed to decrease in the
ultimate tensile strength, the hardness of water-atomized L-PBF parts when compared to highly martensitic gas-atomized powder L-PBF parts.

At low energy densities of 64 and 80 J/mm³, the columnar grains in the as-printed L-PBF parts did not change significantly after the HIP treatment whereas the columnar grains present in as-printed gas-atomized L-PBF parts completely disappeared when subjected to HIP treatment. However, at high energy densities of 84 and 104 J/mm³, long columnar grains in as-printed L-PBF gas- and water-atomized L-PBF parts were changed to equiaxed grains and showed a higher level of homogenization when subjected to HIP treatment. This variation in grains and grain size had significantly affected the yield strength and elongation of HIP treated gas- and water-atomized L-PBF parts. Furthermore, HIP experiments with increased temperature and hold time on various water-atomized L-PBF parts are currently being performed in our group and results will be reported elsewhere.
CHAPTER 3
PROPERTY GAINS FROM HOT ISOSTATIC PRESSING OF 17-4 PH STAINLESS STEEL GAS AND WATER-ATOMIZED PARTS FABRICATED BY LASER-POWDER BED FUSION

3.1 INTRODUCTION

Laser powder bed fusion (L-PBF) process, popularly known as selective laser melting (SLM) has received a lot of interest for fabricating complex three-dimensional components for automotive, aerospace, medical and tooling industries [1]–[3], [7], [26]. In the L-PBF process, three-dimensional components were fabricated using a fine focused laser beam as an energy source to selectively melt powders layer-by-layer based on CAD file [50]–[55]. Many research studies have been carried out in the past decade on iron, steel, nickel, aluminum, titanium and cobalt alloys to fabricate defect-free parts using the L-PBF process [6], [8], [15], [32], [50], [51], [56]–[68]. Most of the studies indicated that defects and porosities still exist in L-PBF parts when they were fabricated using powders of different shape and size attributes [6], [8], [15], [32], [50], [51], [56]–[68].

Typically, gas-atomized powders (<50µm) with a spherical shape and narrow size distribution were preferred for fabricating defect-free parts using L-PBF process [2], [3], [6], [8], [10]–[12]. However, such powders are inherently expensive as they use large
amounts of inert process gas as well as a small fraction of the production lot in each atomization run [2], [3], [6], [13]. On the other hand, water-atomized powders are relatively less expensive but the ability to fabricate high density L-PBF parts from water-atomized powders with properties competitive to parts using gas-atomized powders has not been convincingly demonstrated in the open literature [2], [3], [6], [13]. In this regard, previous studies conducted within our group have shown the feasibility of fabricating 17-4 PH stainless steel L-PBF parts using fine water-atomized powders and achieving the properties similar or higher than gas-atomized parts in certain processing conditions [1]–[4]. However, some porosity still existed in coarser size fractions of water-atomized L-PBF parts.

In powder metallurgy, hot isostatic pressing (HIP) is routinely performed to eliminate porosity and anisotropy. Several HIP studies have emerged in recent times to improve the densification and mechanical properties of L-PBF parts using various materials such as Hastelloy, 316L stainless steel, CoCr and Ti-6Al-4V alloys [15]–[24]. For example, Lavery et al [16] and Tillmann et al [20] discussed the effect of the HIP on decreasing the porosity and its impact on the mechanical properties for 316L stainless steel and Inconel 718 L-PBF parts, respectively. However, there are no reported studies in the literature that demonstrated fundamental understanding on effect of HIP on minimizing porosity and maximizing the properties of 17-4 PH stainless steel L-PBF parts.

In this regard, the present study was performed to understand the effect of the HIP on the densification, mechanical properties and microstructures of 17-4 PH stainless steel gas and
water-atomized as-printed L-PBF parts when they are fabricated at various energy densities. The current study builds from our initial work [69] on the effect of HIP on gas and water-atomized L-PBF parts.

3.2 EXPERIMENTAL

17-4 PH stainless steel water-atomized powders of median particle size 17, 24 and 43 µm and a gas-atomized powder of median particle size 13 µm were used as starting powders.

Figure 0.1. SEM micrographs of the four 17-4 PH stainless steel powders (a) gas-atomized powder D$_{50}$ = 13 µm (b) water-atomized powder D$_{50}$ = 17 µm, (c) water-atomized powder D$_{50}$ = 24 µm, (d) water-atomized powder D$_{50}$ = 43 µm
The morphology of the powders was characterized using an FEI Quanta 600F scanning electron microscope (SEM). The morphology of the gas and water-atomized powders used in the present study is shown in Figure 3.1. The SEM micrographs show typical spherical and irregular morphology for gas and water-atomized powders, respectively. The particle size measurement data, tap and apparent densities of 17-4 PH stainless steel water and gas-atomized powders used in the study are listed in previous journal papers published by the authors [1], [2].

L-PBF experiments using the gas and water-atomized 17-4 PH stainless steel powders were carried out using a 3D Systems ProX 320 machine in Ar atmosphere. Four different energy densities 64, 80, 84 and 104 J/mm³ were used to fabricate the L-PBF parts using gas and water-atomized powders used in this study. Detailed information about processing conditions used for calculating the energy densities for fabricating the L-PBF parts has reported previously [1]–[3], [26]. The tensile parts of dimensions 0.68 m × 0.13 m × 0.318 m was cut by wire electrical discharge machining (EDM) to test for mechanical properties as per ASTM E8M standard.

Three tensile parts of each water-atomized powder type at energy densities 64, 80, 84 and 104 J/mm³ and three tensile parts of gas- water-atomized powder type at energy densities 84 and 104 J/mm³ were selected for HIP treatment. The HIP cycle used in this experiment is shown in Figure 3.2. The density of the HIP treated parts was measured using Mettler Toledo XS104 weighing balance based on Archimedes density principle. The mechanical properties of the HIP treated L-PBF parts were measured with an MTS hydraulic dual
column testing system equipped with a 100 kN load cell at a strain rate of 0.001 s⁻¹. The hardness of the HIP treated parts parallel to the build direction was measured using a Rockwell ‘C’ hardness scale at 150 kg load. The phases present in the L-PBF parts perpendicular to the build direction were estimated using a BRUKER D8 X-ray diffraction (XRD) machine. Metallographic specimens were prepared perpendicular to the build direction and analyzed using an Olympus BX53 microscope. Carpenter's etchant was used to reveal the grain size and phases in the microstructures of as-printed and HIP treated 17-4PH stainless steel parts [26], [30].

![Graph showing hot-isostatic pressing processing conditions](image.png)

**Figure 0.2.** Hot-isostatic pressing processing conditions (temperature, pressure and time) used for post-processing as-printed 17-4 PH stainless steel gas-and water-atomized L-PBF parts.

3.3 RESULTS AND DISCUSSION

3.3.1 MECHANICAL PROPERTIES

The density of HIP treated samples fabricated using 17-4 PH gas and water-atomized powders at various energy densities is shown in **Figure 3.3**. The density was found to
increase after HIP treatment when compared to the density of as-printed parts for four 17-4 PH stainless steel powders at various energy densities. The density of HIP treated gas and water-atomized L-PBF parts under all energy densities ranged around \(\sim 99 \pm 0.5\%\) whereas the density of as-printed L-PBF parts using the four powders varied between 89 \(\pm 0.5\%\) and 98 \(\pm 0.5\%\) [1], [2].

The fractional increase in densification of the L-PBF parts following HIP treatment was significant for coarser water-atomized L-PBF parts \((D_{50}=24\) and \(43\) \(\mu\)m) when fabricated at energy densities of 64 and 80 J/mm\(^3\). These results show that densification and reduction in porosity in the as-printed L-PBF parts can be achieved using HIP treatment and does not depend significantly on the powder shape and size or the energy density used to fabricate the parts. An another important result of this study is that the minimum starting density of L-PBF parts in order to achieve 99 \(\pm 0.5\%\) dense parts after HIP treatment, was 92 \(\pm 1\%\) (water-atomized \(D_{50}=43\) \(\mu\)m at energy density 80 J/mm\(^3\)). A similar increase in density due to HIP treatment was observed in previous studies for L-PBF parts fabricated from other compositions [15]–[24]. Further, the densities of gas- \((D_{50}=13\) \(\mu\)m) and water-atomized \((D_{50}=43\) \(\mu\)m) L-PBF parts obtained from the current study is higher than densities of L-PBF parts fabricated from the same powders when they were post-processed using a different HIP cycle [69]. The HIP cycle used in the current study had an additional step in the cycle (Figure 3.2), at a solutionizing temperature of 1050\(^0\)C and 70 MPa pressure for 20 minutes which could have accelerated the densification of the L-PBF parts [69]. Atkinson et al. [70] indicated that addition of solutionizing temperature step at high
pressures would lead to accelerated densification as per the plasticity/power-law creep for low-alloy steels.
Figure 0.3. The density of 17-4 PH stainless steel gas-and water-atomized L-PBF parts before and after HIP treatment as a function of energy density.
The variation in the ultimate tensile strength of the HIP treated L-PBF parts with energy density for four 17-4 PH stainless steel powders is shown in Figure 3.4. The ultimate tensile strength of the L-PBF parts was found to increase due to HIP treatment. The increase in the ultimate tensile strength of the L-PBF parts can be attributed to the increase in density following HIP treatment (Figure 3.3). The HIP treated L-PBF parts fabricated using fine gas- (D$_{50}$=13µm) and water-atomized (D$_{50}$=17µm) powders showed significantly higher tensile strength (1450 ± 50 MPa) compared to as-printed L-PBF parts at all energy densities. Similar ultimate tensile strength values (1450 ± 50 MPa) were observed for HIP treated coarser water-atomized powder (D$_{50}$=43µm) for the L-PBF parts fabricated at energy densities 84 and 104 J/mm$^3$. However, the ultimate tensile strength of L-PBF parts fabricated using one of the water-atomized powder lots (D$_{50}$=24µm) was slightly lower; ranging between 1300 ± 50 MPa and 1400 ± 20 MPa after HIP treatment but still higher than the corresponding value for as-printed L-PBF parts. The fractional increase in ultimate tensile strength due to HIP treatment is significant for as-printed water-atomized L-PBF parts when compared to gas-atomized L-PBF parts (Figure 3.3). Further, the ultimate tensile strength of gas (D$_{50}$=13µm) and water-atomized (D$_{50}$=43µm) L-PBF parts after HIP treatment used in the current study is higher than the tensile strength of the L-PBF parts obtained after the HIP cycle used in our previous study [69].

The data obtained from the current study shows the effectiveness of HIP treatment on improving the ultimate tensile strength of the gas and water-atomized L-PBF parts. Overall, the data clearly shows the after using the current HIP treatment cycle, similar ultimate tensile strength values can be obtained using both water-atomized and gas-atomized L-PBF
powders. Further, the ultimate tensile strength of L-PBF parts after HIP treatment obtained in the current study is higher than the ultimate tensile strength of the L-PBF parts when they were heat treated as shown in previous studies [3], [62], [71], [72]. For reference, the ultimate tensile strength of 17-4 PH stainless steel L-PBF parts in various studies after solutionizing and aging treatment ranged around 1275 ± 75 MPa [3], [62], [71], [72].

The hardness of gas and water-atomized L-PBF parts increased following HIP treatment and was found to be higher than the hardness of as-printed L-PBF parts at all energy densities, as shown in Figure 3.4. The hardness of the L-PBF parts after HIP treatment varied between 36 ± 3 and 40 ± 3 HRC and was higher than as-printed, and comparable to MIM and wrought values [2], [3], [13]. For reference, Murr et al. [45], [73] and Kumar et al. [74] reported hardness values of 35 ± 2 HRC for 17-4 PH stainless gas-atomized L-PBF parts. For further comparison, the hardness of 17-4PH stainless steel in the wrought state was reported to be 38 ± 2 HRC while parts fabricated using casting and MIM were reported to be 37 ± 2 HRC after heat treatment [75].

Figure 3.5 shows the effect of HIP treatment on the yield strength of gas and water-atomized L-PBF parts fabricated at various energy densities. It can be seen that the yield strength of the L-PBF parts increased due to the HIP treatment and higher than as-printed properties. The trends observed in Figure 3.5 are similar to the trends observed for ultimate tensile strength and hardness (Figure 3.3 and 3.4). The yield strength of the L-PBF parts fabricated using the fine gas (D₅₀=13µm) and water-atomized (D₅₀=17µm) powders after HIP treatment was 1200 ± 30 MPa irrespective of variation in energy density. These yield
strength values are higher than the yield of coarser water-atomized parts at energy densities 64 and 80 J/mm\(^3\). However, no difference in yield strength was observed between gas and water-atomized parts after HIP treatment at an energy density of 104 J/mm\(^3\). Furthermore, the yield strength of the L-PBF parts after HIP treatment obtained in the current study is higher than the yield strength of L-PBF parts when they heat treated data reported in the literature [3], [62], [71], [72].

The elongation of gas and water-atomized L-PBF parts decreased due to HIP treatment and was lower than the elongation of the corresponding as-printed L-PBF parts at all energy densities, as shown in Figure 3.6. The trends observed in Figure 3.6 contrasted with the trends observed in the ultimate tensile strength, yield strength and hardness trends in this study. The elongation of the gas and water-atomized L-PBF parts after HIP treatment ranged between 5-11 % but was higher than the elongation of heat-treated 17-4 PH stainless steel L-PBF parts reported in previous studies [3], [62], [71], [72]. For comparison, the elongation of 17-4 PH stainless steel L-PBF parts in various studies after solutionizing and aging ranged between 2-5 % [3], [62], [71], [72].

Overall, the current data suggest after HIP treatment that similar densities and mechanical properties can be achieved using a wide number of particle sizes for water-atomized powders compared to gas-atomized powders. Further, the values of HIP treated properties in this study were higher than values reported in literature for L-PBF 17-4 PH stainless steel using several heat treatment cycles. In the next section, results from x-ray diffraction
and microstructure analysis performed on the HIP treated and as-printed L-PBF parts were to understand the origins of mechanical properties.

Figure 0.4. The variation in ultimate tensile strength with energy density for 17-4 PH stainless steel gas- and water-atomized L-PBF parts before and after HIP treatment.
Figure 0.5. The variation in hardness with energy density for 17-4 PH stainless steel gas- and water-atomized L-PBF parts before and after HIP treatment.
Figure 0.6. The variation in yield strength with energy density for 17-4 PH stainless steel gas- and water-atomized L-PBF parts before and after HIP treatment.
Figure 0.7. The variation in elongation with energy density for 17-4 PH stainless steel gas-and water-atomized L-PBF parts before and after HIP treatment.
3.3.2 MICROSTRUCTURES

Figure 3.8 shows the XRD data on the effects of HIP treatment on gas and water-atomized L-PBF parts at all energy densities. HIP treatment resulted in predominantly forming the martensite phase in the L-PBF parts irrespective of the initial phases present in the as-printed parts. Further, the phases in the HIP treated L-PBF parts were not significantly sensitive to the size and shape of the powder. It is expected that the martensite phase present in the HIP treated L-PBF parts would have also contributed (in addition to density increase) to the increase in ultimate tensile strength as shown in Figure 3.3 when compared to as-printed parts. The phase change is significant for water-atomized L-PBF parts after the HIP treatment when compared to gas-atomized L-PBF parts. The increase in martensite content in the L-PBF parts may be attributed to the rapid cooling rates during the HIP treatment [21 -30]. According to continuous cooling transformation (CCT) curves, martensite forms in the L-PBF parts when the austenite phase in the starting condition is rapidly cooled to room temperature [32].

The microstructures of the L-PBF parts before and after the HIP treatment at energy densities 84 and 104 J/mm³ for gas-atomized powders and at energy densities 64 and 104 J/mm³ for water-atomized powders were shown in Figure 3.9. The following symbols were used in Figure 3.9 to distinguish the microstructures (G-13: gas-atomized D₅₀=13µm; W-17: water-atomized D₅₀=17µm; W-24: water-atomized D₅₀=24µm; W-43: water-atomized D₅₀=43µm and 64 and 104 J/mm³ represent energy densities).
Figure 0.8. The XRD phases in gas and water-atomized L-PBF parts before and after HIP treatment for parts fabricated at energy densities of 64, 80, 84 and 104 J/mm$^3$
Figure 0.9. The microstructures of gas and water-atomized L-PBF parts before and after HIP treatment for parts fabricated at energy densities of 64 and 104 J/mm$^3$.

It can be clearly seen that the microstructural features present in the as-printed parts were completely different from the HIP’ed gas and water-atomized L-PBF parts as shown in Figure 3.9. The microstructure formation trends which were observed in the L-PBF parts after the HIP treatment is a contrast to the trends observed in the as-printed L-PBF parts microstructures. The long columnar grains of average size 25 ± 5 µm and 20 ± 5 µm present in as-printed gas and water-atomized L-PBF parts, respectively, were translated to homogeneous equiaxed grains 18 ± 1 µm and 15 ± 1 µm, respectively, after the HIP
treatment. This variation in grain shape and size could have also influenced the ultimate tensile strength, yield, and elongation of the L-PBF parts as shown in Figures 3.4, 3.6 and 3.7. During HIP treatment, the fluid at high temperature exhibits an isotropic pressure on the L-PBF parts, resulting in creep and diffusion phenomena that could cause the densification and microstructure variation in the L-PBF parts [70]. It is surmised that the creep and diffusion phenomena could result in strengthening grain boundaries and increasing dislocations along the grain boundaries. During tensile testing, larger stresses are needed to move these dislocations resulting in an increase in the yield strength of the material [70]. Previous studies by Tillman et al. [20], Lavery et al. [16] and Wu et al. [24] showed the transformation of long columnar grains to equiaxed grain structure for various materials after the HIP treatment.

A notable aspect of the current study is the achievement of similar densities, mechanical properties and microstructures in the L-PBF parts from both gas and water-atomized powders after the HIP treatment. The density, mechanical properties and microstructures variation data for the four 17-4 PH stainless steel L-PBF parts are summarized in Table 3.1.
Table 3.1. The density, mechanical properties and microstructure variation in 17-4 PH stainless steel gas-and water-atomized L-PBF parts following HIP treatment.

<table>
<thead>
<tr>
<th>17-4 PH stainless steel L-PBF parts from:</th>
<th>Effects on properties following HIP treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized powder D$_{50}$=13µm</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>Density: 98 ± 1 to 99.5 ± 0.2 %,</td>
</tr>
<tr>
<td></td>
<td>Ultimate tensile strength: 950 ± 50 to 1450 ± 20 MPa,</td>
</tr>
<tr>
<td></td>
<td>Hardness: 30 ± 1 to 40 ± 1 HRC,</td>
</tr>
<tr>
<td></td>
<td>Yield strength: 800 ± 50 to 1200 ± 20 MPa</td>
</tr>
<tr>
<td></td>
<td>Other changes</td>
</tr>
<tr>
<td></td>
<td>Elongation: 22 ± 1 to 11 ± 1 %</td>
</tr>
<tr>
<td></td>
<td>Martensite phase did not change</td>
</tr>
<tr>
<td></td>
<td>Long columnar grains (25 ± 5 µm) transformed to equiaxed grains (18 ± 1 µm)</td>
</tr>
<tr>
<td>Water-atomized powder D$_{50}$=17µm</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>Density: 96 ± 1 to 99.5 ± 0.2 %,</td>
</tr>
<tr>
<td></td>
<td>Ultimate tensile strength: 500 ± 50 to 1450 ± 20 MPa,</td>
</tr>
<tr>
<td></td>
<td>Hardness: 27 ± 1 to 40 ± 1 HRC,</td>
</tr>
<tr>
<td></td>
<td>Yield strength: 400 ± 50 to 1200 ± 20 MPa</td>
</tr>
<tr>
<td></td>
<td>Other changes</td>
</tr>
<tr>
<td></td>
<td>Elongation: 14 ± 1 to 8 ± 1 %</td>
</tr>
<tr>
<td></td>
<td>Austenite phase transformed to the martensite phase</td>
</tr>
<tr>
<td></td>
<td>Long columnar grains (20 ± 5 µm) transformed to equiaxed grains (15 ± 1 µm)</td>
</tr>
<tr>
<td>Water-atomized powder D$_{50}$=24µm</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>Density: 88 ± 1 to 99 ± 0.2 %,</td>
</tr>
<tr>
<td></td>
<td>Ultimate tensile strength: 480 ± 30 to 1400 ± 20 MPa,</td>
</tr>
<tr>
<td></td>
<td>Hardness: 25 ± 1 to 40 ± 1 HRC,</td>
</tr>
<tr>
<td></td>
<td>Yield strength: 400 ± 50 to 1100 ± 20 MPa</td>
</tr>
<tr>
<td></td>
<td>Other changes</td>
</tr>
<tr>
<td></td>
<td>Elongation: 12 ± 1 to 6 ± 1 %</td>
</tr>
<tr>
<td></td>
<td>Austenite phase transformed to the martensite phase</td>
</tr>
<tr>
<td></td>
<td>Long columnar grains (20 ± 5 µm) transformed to equiaxed grains (14 ± 1 µm)</td>
</tr>
<tr>
<td>Water-atomized powder D$_{50}$=43µm</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>Density: 90 ± 1 to 99 ± 0.2 %,</td>
</tr>
<tr>
<td></td>
<td>Ultimate tensile strength: 490 ± 30 to 1440 ± 20 MPa,</td>
</tr>
<tr>
<td></td>
<td>Hardness: 27 ± 1 to 40 ± 1 HRC,</td>
</tr>
<tr>
<td></td>
<td>Yield strength: 400 ± 50 to 1180 ± 20 MPa</td>
</tr>
<tr>
<td></td>
<td>Other changes</td>
</tr>
<tr>
<td></td>
<td>Elongation: 14 ± 1 to 7 ± 1 %</td>
</tr>
<tr>
<td></td>
<td>Austenite phase transformed to the martensite phase</td>
</tr>
<tr>
<td></td>
<td>Long columnar grains (20 ± 5 µm) transformed to equiaxed grains (16 ± 1 µm)</td>
</tr>
</tbody>
</table>
3.4 CONCLUSIONS

A comprehensive study was performed to understand the effects of HIP treatment on the densification, mechanical properties and microstructures of 17-4 PH stainless steel gas and water-atomized L-PBF parts at various energy densities. The results from the current study show the strong dependence of densification, mechanical properties and microstructures on temperature, pressure and time during the HIP cycle. The density, ultimate tensile strength, hardness and yield strength of gas and water-atomized L-PBF parts increased due to HIP treatment and were higher than as-printed properties. Following HIP treatment, 99.5 ± 0.2 % dense L-PBF parts with ultimate tensile strength: 1450 ± 20 MPa, hardness: 40 ± 1 HRC, yield strength: 1180 ± 20 MPa and elongation: 10 ± 1 % were achieved using both water and gas-atomized powders. The density, martensite content and equiaxed microstructure in L-PBF parts after the HIP treatment were consistent with the trends in mechanical properties of the L-PBF parts. The austenite phase present in the as-printed water-atomized parts transformed to martensite phase after HIP treatment and contributed to the increase ultimate tensile strength of the L-PBF parts at all energy densities. Further, the long columnar grains present in the as-printed gas and water-atomized L-PBF parts (20 ± 5 µm) translated to homogeneous equiaxed grains (15 ± 1 µm) in HIP treated L-PBF parts at all energy densities. This variation in grain structure and grain size significantly affected the yield strength, ultimate tensile strength and elongation of HIP treated gas and water-atomized L-PBF parts.
CHAPTER 4
EFFECTS OF POWDER CHARACTERISTICS AND PROCESSING CONDITIONS ON THE CORROSION PERFORMANCE OF 17-4 PH STAINLESS STEEL FABRICATED BY LASER-POWDER BED FUSION

4.1 INTRODUCTION

Stainless steel alloys have been used in many applications due to their excellent strength and corrosion properties [1–4]. The laser-powder bed fusion (L-PBF) process has gained attention and importance due to its potential to produce complex-shaped parts for medical, tooling, automotive and aerospace industries [1, 5–8]. The L-PBF process is a layer by laser melting or fusion process which uses a laser beam to melt the starting powder for fabricating a desired 3D component [1–4]. Among stainless steels, 17-4 PH stainless steel is desirable for fabrication of parts using L-PBF process in transportation (engine components, gears, fuel injectors) and injection molding industries (injection mold with conformal cooling channels) within our research group as shown in Figure 4.1.
Many research studies have been carried out to understand the effects of powder and processing conditions on the densification behavior and structural attributes of components fabricated by the L-PBF process [1,3,7,9–15]. Most of the prior studies indicated that quality of parts produced using the L-PBF process depends on many factors, the most important being processing conditions and powder characteristics. The effect of processing conditions such as laser power, scan speed, layer thickness, scan spacing on the densification of powders has been extensively studied in recent years [1-4]. On the other hand, in-depth experimental studies on the influence of powder characteristics such as powder shape, size on the densification behavior have been reported only in the recent past [2, 5]. The variation in densification phenomena of the powder as a function of powder type and processing conditions could lead to porosities in the parts fabricated using L-PBF process [1,3,7,9–15].
The porosity in L-PBF parts fabricated using stainless steel powders could not only affect the mechanical properties but also influence the sensitivity to localized corrosion attack, especially in the corrosion environments [16–19]. However, the corrosion properties of L-PBF components fabricated from powders with different characteristics and in different processing conditions have not received much attention in the literature. The corrosion resistance of stainless steel parts results from a thin and protective passive film chromium oxide (Cr$_2$O$_3$) or iron oxide (Fe$_2$O$_3$) based passive film. The chromium or iron oxide passive film acts as a partial barrier between the metal and the corrosive species present in the environment [16,18,20–23]. The stability of the passive film when exposed to aggressive corrosive species such as chloride ions is important in preventing corrosion [16,18, 20, 22–25]. However, the porosity in L-PBF parts can be expected to compromise the passive film layer stability and could result in enhanced corrosion of the L-PBF parts [16–19,26]. Therefore, it is important to understand the corrosion behavior of 17-4 PH stainless steel components built using L-PBF process in natural service conditions.

Independent corrosion studies in various electrolytes have been reported on 17-4 PH stainless steel L-PBF parts fabricated using gas-atomized powders [16, 18, 20, 23, 25, 27]. However, few studies have been reported that compare the corrosion resistance of the 17-4 PH stainless steel L-PBF parts fabricated from powders of different characteristics such as shape, size and the method utilized for atomization (gas-atomized or water-atomized) [16,18]. In order to address this gap in the literature, the present study was carried out to understand the effect of powder characteristics and processing conditions on the corrosion performance of 17-4 PH stainless steel L-PBF parts. The results from the current study will
provide a better understanding of the effect of porosity, density and microstructures on the corrosion performance of 17-4 PH stainless steel L-PBF gas- and water-atomized parts.

4.2 EXPERIMENTAL MATERIALS AND METHODS

One 17-4 PH stainless steel gas-atomized powder and three water-atomized powders were used in this study. The gas-atomized 17-4 PH stainless steel powder utilized for this study was supplied by 3D Systems. The water-atomized 17-4 PH stainless steel powders were obtained from North American Höganäs. A high-resolution FEI Quanta 600F scanning electron microscope (SEM) was used for morphology analysis of the powders. The L-PBF experiments were performed on a 3D Systems ProX 200 machine in an argon environment. The tensile geometries were fabricated using four energy densities to melt the powders, i.e. 64, 80, 84 and 104 J/mm³. The energy densities were calculated using the laser processing parameters like laser power, scan speed, hatch spacing and layer thickness. In this study, two laser powers (150 and 195 W) and two scan speeds (1250 and 1550 mm/s) were used to fabricate the tensile geometries while layer thickness and hatch spacing were kept constant at 30 µm and 50 µm, respectively.

A total of 64 parts from four powders in four processing conditions were fabricated during the study. Of the 64 parts, four parts fabricated from each powder type and in one processing condition were selected for Archimedes density and corrosion performance analysis. Density measurements were conducted using the Archimedes method on a Mettler Toledo XS104 weighing balance. The density values of the 17-4 PH stainless steel gas- and water-atomized L-PBF parts were expressed as a percentage relative to the density
of the 17-4 PH stainless steel cast part (7.87 g/cm³). The porosity of L-PBF parts was calculated using **Equation 4.1**.

\[ \phi = 1 - \frac{\rho (I)}{\rho (t)} \]  

(1)

Where, \( \phi \) is the porosity of the L-PBF parts, \( \rho (I) \) is the density of the L-PBF parts, \( \rho (t) \) is the density of 17-4 PH stainless cast part.

Potentiostatic polarization experiments were used to compare the electrochemical or corrosion behavior of the test coupons. All polarization experiments were conducted in an electrochemical glass cell shown in **Figure 2.2**. Metrohm Autolab PGSTAT100N was used to perform three electrode electrochemical measurement. Ag/AgCl was used as a reference electrode and the platinum was used as a counter electrode. The polarization experiments were carried out on L-PBF parts by varying the potential from 1 V/Ag/AgCl to -1 V/Ag/AgCl. The surface area of 1 cm² of each L-PBF part, which was finished with 1200 grit SiC paper, was exposed to the 0.5 M NaCl solution in ambient conditions. All experiments were carried out at room temperatures and were repeated 3 times for each specimen.

![Figure 0.2.Electrochemical corrosion studies setup used in this study.](image)

Figure 0.2.Electrochemical corrosion studies setup used in this study.
A summary of 17-4 PH stainless steel particle characteristics used in this study is reported in Table 4.1 [28, 29]. The morphology (shape and size) of the powders was characterized using SEM and shown in Figure 4.3, showing the gas-atomized powders to be spherical while the water-atomized powders were irregular in shape.

**Table 4.1.** Particle characteristics of the 17-4 PH stainless steel gas- and water-atomized powders.

<table>
<thead>
<tr>
<th>Powder</th>
<th>Particle distribution</th>
<th>Tap density (g/cc)</th>
<th>Apparent density (g/cc)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized (G)</td>
<td>D&lt;sub&gt;10&lt;/sub&gt; (µm) 13</td>
<td>4 ± 0.05</td>
<td>3 ± 0.002</td>
</tr>
<tr>
<td>Water-atomized (W) 1</td>
<td>D&lt;sub&gt;50&lt;/sub&gt; (µm) 27</td>
<td>4.3 ± 0.05</td>
<td>3 ± 0.05</td>
</tr>
<tr>
<td>Water-atomized (W) 2</td>
<td>D&lt;sub&gt;90&lt;/sub&gt; (µm) 37</td>
<td>3.7 ± 0.01</td>
<td>2.8 ± 0.05</td>
</tr>
<tr>
<td>Water-atomized (W) 3</td>
<td>D&lt;sub&gt;10&lt;/sub&gt; (µm) 26</td>
<td>3.7 ± 0.05</td>
<td>2.8 ± 0.05</td>
</tr>
</tbody>
</table>
4.3 RESULTS AND DISCUSSION

The variation of density as a function of the energy density used to fabricate L-PBF parts using four 17-4 PH stainless steel powders are reported in Table 4.2.
Table 4.2. Densities (expressed as % theoretical) of the 17-4 PH stainless steel gas- and water-atomized L-PBF parts fabricated under energy densities ranging from 64 to 104 J/mm³.

<table>
<thead>
<tr>
<th>L-PBF parts from</th>
<th>Energy density 64 J/mm³</th>
<th>Energy density 80 J/mm³</th>
<th>Energy density 84 J/mm³</th>
<th>Energy density 104 J/mm³</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized D₅₀ = 13 µm</td>
<td>97 ± 0.5 %</td>
<td>97.5 ± 0.5 %</td>
<td>97.5 ± 0.5 %</td>
<td>97.5 ± 0.5 %</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 17 µm</td>
<td>96 ± 1 %</td>
<td>97 ± 0.5 %</td>
<td>97 ± 0.5 %</td>
<td>97 ± 0.5 %</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 24 µm</td>
<td>87 ± 0.5 %</td>
<td>91 ± 0.5 %</td>
<td>96 ± 1 %</td>
<td>97 ± 0.5 %</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 43 µm</td>
<td>89 ± 0.5 %</td>
<td>95 ± 0.5 %</td>
<td>97 ± 0.5 %</td>
<td>97 ± 0.5 %</td>
</tr>
</tbody>
</table>

The densities of L-PBF parts produced using coarser water-atomized powders (D₅₀=24 µm and 43 µm) were lower when compared to the density of L-PBF parts produced using the finer gas-atomized (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) powders (97± 0.5 %) at low energy densities of 64 and 80 J/mm³. The high density of the L-PBF parts (97± 0.5 %) fabricated using finer gas-atomized (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) powders can be attributed to their high tap (4 ± 0.05 g/cc) and apparent density (3± 0.05 g/cc) (Table 4.2). Lee et al. [30] showed that high apparent density of powders could result in better packing density of the powder bed. The high packing density of the powder bed presumably results in better melt pool hydrodynamic stabilities ensuing enhance densification of the powders during the L-PBF process [1], [96]–[99]. However, the density of three 17-4 PH stainless steel water-atomized powders is similar (97 ± 0.5 %) and comparable to 17-4 PH stainless steel gas-atomized powders when processed at high energy densities 84 and 104 J/mm³. At higher energy densities, a large amount of molten
pool with low viscosity and better wettability properties is likely to be formed in the powder bed producing high dense L-PBF parts[28, 30–33].

Figure 0.4. Tafel plots of the components fabricated from four 17-4 PH stainless steel powders fabricated at (a) energy density 64 J/mm³ (b) energy density 80 J/mm³ (c) energy density 84 J/mm³ and (d) energy density 104 J/mm³.

The variation in Tafel plots curves as a function of the energy density used to fabricate L-PBF parts from the four 17-4 PH stainless steel powders is shown in Figure 4.4. The Tafel plots curves shown in Figure 4.4 qualitatively indicated different anode and cathode curve behavior in 17-4 PH stainless steel gas- and water-atomized L-PBF parts when fabricated at different energy densities. The quantification of the Tafel plots behavior was
subsequently performed to further understand the effect of density on the corrosion performance of four 17-4 PH stainless steel powders. Using the Tafel plot quantification method [84], [86], the 17-4 PH stainless steel L-PBF parts Tafel plots were analyzed as shown in Figure 4.5 for calculating the corrosion current, \(i_{\text{corr}}\) and Tafel constants. The corrosion current, \(i_{\text{corr}}\) and Tafel constants were used to determine the polarization resistance \(R_p\) and corrosion rate from Equation 4.2 and 4.3, respectively.

\[
Polarization\ resistance, \ R_p = \frac{1}{i_{\text{corr}}} \left( \frac{\beta_a \beta_c}{\beta_a + \beta_c} \right) \quad [4.2]
\]

Where the Tafel constants (\(\beta_a\) and \(\beta_c\)) represent the anodic and cathodic slopes, respectively.

\[
Corrosion\ rate, \ CR = \frac{i_{\text{corr}}}{dA} (k \times EW) \quad [4.3]
\]

Where, \(d\) is the density of the material (Kg/m\(^3\)), \(A\) is exposed surface area of the material to corrosion (m\(^2\)), \(k\) is a constant (3.272 m/year), and EW is the equivalent weight of the material.
Figure 0.5. Schematic diagram showing the extracting of corrosion current and constants from a Tafel plot.

The variation in corrosion current with the density of the L-PBF parts fabricated using finer gas-atomized (D<sub>50</sub>=13 µm) and water-atomized (D<sub>50</sub>=17 µm) powders are shown in Figure 4.6a. High density and low corrosion current were observed for the 17-4 PH stainless steel L-PBF parts irrespective of the atomization technique to produce starting powders and the energy density of these two systems. The corrosion current of the gas-atomized (D<sub>50</sub>=13 µm) and water-atomized (D<sub>50</sub>=17 µm) L-PBF parts ranged between 0.1 and 1.0 µA, which is lower than the wrought sample (0.9 ± 0.1 µA). It can also be seen from Figure 4.6a that the corrosion current of the L-PBF parts was not significantly varied for finer gas (D<sub>50</sub>=13 µm) and water-atomized (D<sub>50</sub>=17 µm) powders when fabricated at different energy densities. Previous studies by Frankel et al. [100] showed that the low corrosion current could be an indication that the rate of electrochemical reaction is slow either because of the thick oxide film or low porosity on the surface of the L-PBF parts.
Further, no systematic variation in the corrosion current was observed for high dense (97 ± 0.5 %) gas-atomized ($D_{50}=13 \ \mu m$) and water-atomized ($D_{50}=17 \ \mu m$) L-PBF parts when fabricated at different energy densities.

Figure 0.6. Variation in corrosion current with % theoretical density and energy density for samples fabricated using (a) gas-atomized ($D_{50}=13 \ \mu m$) and water-atomized ($D_{50}=17 \ \mu m$) powders (b) water-atomized ($D_{50}=24 \ \mu m$ and 43 $\mu m$) powders. The symbols represent the four powders and the colors represent the four energy densities used in the fabrication.

Figure 4.6b shows that the density and consequently corrosion current significantly varied as a function of energy density for coarser water-atomized powders ($D_{50}=24 \ \mu m$ and 43 $\mu m$) powders when compared to gas ($D_{50}=13 \ \mu m$) and water-atomized ($D_{50}=17 \ \mu m$)
powders (Figure 4.6a). The corrosion current decreased with increased density of the water-atomized (D$_{50}$=24 µm and 43 µm) L-PBF parts. At a high energy density of 104 J/mm$^3$ and with densities higher than 97 %, the corrosion current of the water-atomized (D$_{50}$ = 24 µm and 43 µm) L-PBF parts was found to be lower than the wrought samples and similar to the corrosion current of gas (D$_{50}$=13 µm) and water-atomized (D$_{50}$=17 µm) L-PBF parts (Figure 4.6). The corrosion currents of the 17-4 PH stainless gas- and water-atomized L-PBF parts with densities higher than 97 % varied between 0.1 and 0.8 µA (Figure 4.6b) whereas the corrosion current value wrought sample is 0.9 ± 0.1 µA. Similar corrosion current values for high dense 17-4 PH stainless steel parts fabricated with gas-atomized powders have been reported in previous L-PBF research studies [84]–[87], [94]. For example, Stoudt et al. [16] carried out potentiostatic polarization studies on L-PBF parts fabricated using 17-4 PH stainless steel gas-atomized powders in 0.5 M NaCl solution and reported that corrosion potential of the nearly fully dense (99 ± 0.5 %) varied between 0.5 and 0.8 µA. Furthermore, at a high energy density of 104 J/mm$^3$, the corrosion current is independent of starting powder characteristics (shape and size) and atomization method (gas or water-atomization) used to produce these powders (Figure 4.6).

In Figure 4.7, the polarization resistance is plotted as a function of the density of L-PBF parts when fabricated using four 17-4 PH stainless powders at various energy densities. It was seen that the polarization resistance of the high dense (99 ± 0.5 %) L-PBF parts fabricated using finer gas-atomized (D$_{50}$=13 µm) and water-atomized (D$_{50}$=17 µm) powders showed high polarization resistance (26,000 ± 2,000 Ω) when fabricated at different energy densities (Figure 4.7a). For comparison, these values are higher than
wrought sample \((25,000 \pm 1,000 \, \Omega)\) and similar to the polarization resistance values \((25,000-28,000 \, \Omega)\) of 17-4 PH stainless steel L-PBF parts fabricated using gas-atomized powders as reported in the literature [14]. The high polarization resistance value of the L-PBF parts indicates that the resistance of the samples in the corrosion environment is high [27,40,41].

The density and consequently, the polarization resistance of coarser water-atomized powders \((D_{50}=24 \, \mu m \text{ and } 43 \, \mu m)\) powders were found to be strongly dependent on energy density used to fabricate the samples (Figure 4.7b). The polarization resistance of coarser water-atomized powders \((D_{50}=24 \, \mu m \text{ and } 43 \, \mu m)\) L-PBF parts increased with an increase in energy density and density of the L-PBF parts. At low energy densities of 64 to 84 J/mm\(^3\), the polarization resistance samples fabricated using coarser water-atomized powders \((D_{50}=24 \, \mu m \text{ and } 43 \, \mu m)\) powders is 11,000 ± 3,000 Ω and 15,000 ± 4,000 Ω, respectively. These polarization resistance values are lower than that of the finer gas-\((D_{50}=13 \, \mu m)\) and water-atomized \((D_{50}=17 \, \mu m)\) L-PBF parts as shown in Figure 4.7.

Figure 4.7b also indicate that at high energy density of 104 J/mm\(^3\), samples fabricated using coarser water-atomized powders \((D_{50}= 43 \, \mu m)\) exhibited a slightly higher polarization resistance \((28,000 \pm 500 \, \Omega)\) than samples fabricated using gas-atomized \((D_{50}=13 \, \mu m)\) powders \((23,000 \pm 1000 \, \Omega)\) and wrought samples \((25,000 \pm 1,000 \, \Omega)\). Thus, the corrosion performance of the 17-4 PH stainless steel L-PBF parts is strongly dependent on their density, which in turn depends on the powder characteristics and L-PBF processing conditions.
Figure 0.7. Variation in polarization resistance as a function of % theoretical density and energy density for samples fabricated using (a) gas-atomized ($D_{50} = 13 \, \mu m$) and water-atomized ($D_{50} = 17 \, \mu m$) powders (b) water-atomized ($D_{50} = 24 \, \mu m$ and $43 \, \mu m$) powders. The symbol represent the four powders and the colors represent the four energy densities used in the fabrication.

Figure 4.8 shows the variation in corrosion rate as a function of the L-PBF parts density when fabricated at various energy densities using the four 17-4 PH stainless steel powders. The L-PBF parts fabricated using the finer gas ($D_{50} = 13 \, \mu m$) and water-atomized ($D_{50} = 17 \, \mu m$) powders ranged between 1 to 8 $\mu m$/year at different energy densities as shown in the Figure 4.8a. For further comparison, the corrosion rates of finer gas ($D_{50} = 13 \, \mu m$) and water-atomized ($D_{50} = 17 \, \mu m$) L-PBF parts were lower than that of the wrought 17-4 PH
stainless steel is 3 ± 0.5 µm/year except at an energy density 64 J/mm³. However, the corrosion rate for samples fabricated using the coarser water-atomized (D₅₀=24 µm and 43 µm) powders decreased with an increase in energy density (Figure 4.8b). At energy densities of 64 and 80 J/mm³, the corrosion rate of the coarser water-atomized (D₅₀=24 µm and 43 µm) L-PBF parts varied between 20 to 45 µm/year. These corrosion rate values are higher than that of the finer gas (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) L-PBF parts fabricated at the same energy densities as well as the wrought samples. However, at an energy density of 104 J/mm³, the corrosion rates were similar for L-PBF parts fabricated using all four powders (Figure 4.8).

Figure 0.8. Variation in corrosion rate as a function of % theoretical density and energy density for samples fabricated using (a) gas-atomized (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) powders (b) water-atomized (D₅₀=24 µm and 43 µm) powders. The symbols represent the four powders and the colors represent the four energy densities used in the fabrication.
**Figure 4.9** shows the optical microscopy images of the surfaces of samples fabricated at 64 J/mm³ obtained before and after conducting potentiostatic polarization experiments. It can be found that there is a significant difference in the images before and after the corrosion tests for the coarser water-atomized (D₅₀=24 µm and 43 µm) L-PBF parts at energy density 64 J/mm³ as evidenced by an increase in a number of pores. Large pores are seen in the coarser water-atomized (D₅₀=24 µm and 43 µm) L-PBF parts images whereas small pores can be found in finer gas (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) L-PBF parts images before and after the potentiostatic polarization experiments. The presence of minor pores in the finer gas (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) L-PBF parts after the corrosion tests could indicate the high resistance of the L-PBF parts to the corrosion phenomena [20,22,34–36]. The pore formation in the L-PBF parts after the corrosion tests is consistent with the corrosion performance data when fabricated at energy density 64 J/mm³ (**Figure 4.6, 4.7 and 4.8**). The corrosion performance of the L-PBF parts fabricated using finer gas (D₅₀=13 µm) and water-atomized (D₅₀=17 µm) powders are better than the water-atomized powders (D₅₀=17 µm and 43 µm) at energy density 64 J/mm³.

**Figure 4.10** shows the optical microscopy images of the surfaces of samples fabricated at 104 J/mm³ obtained before and after conducting potentiostatic polarization experiments. Compared to the images presented in **Figure 4.9**, only minor pitting-induced pores can be found in the images before and after the corrosion tests for all L-PBF parts. Further, the minor pits seen in all L-PBF parts at energy density 104 J/mm³ is consistent with the
corrosion performance data extracted from the Tafel plots (Figure 4.6, 4.7 and 4.8). The corrosion performance of the L-PBF parts fabricated using water-atomized powders ($D_{50}$=17 µm and 43 µm) at energy density 104 J/mm$^3$ is better than the wrought sample.

The porosity formed during the L-PBF process play a critical role in the initiation of pitting corrosion in the L-PBF parts. The pore geometry at low densities appears to be favorable for pitting corrosion since the pores would lead to stagnation of the sodium chloride solution in part [20,22,34–36]. The stagnation of chloride solution could cause a breakdown the passive layer ensuing favorable sites for initiation of pitting corrosion and increases the rate of corrosion [20,22,34–36]. The L-PBF parts with densities higher than 97%, available pore sites desirable for pitting corrosion were less and could have decreased the rate of corrosion in the L-PBF parts [20,37–39]. The rate of electrochemical corrosion reaction can be expressed as corrosion current in the L-PBF parts and decreases for the dense parts of the thick oxide layer. The decrease in the rate of electrochemical corrosion reactions could decrease the corrosion current of the L-PBF parts densities higher than 97% [20,37–39].
Figure 0.9. Optical microscopy of samples fabricated at energy density 64 J/mm$^3$ using four 17-4 PH stainless steel powders acquired before and after conducting potentiostatic polarization experiments in 0.5 M NaCl solution.

Figure 0.10. Optical microscopy of samples fabricated at energy density 104 J/mm$^3$ using four 17-4 PH stainless steel powders acquired before and after conducting potentiostatic polarization experiments in 0.5 M NaCl solution.

Future experiments on densification and corrosion performance of 17-4 PH stainless steel L-PBF parts are currently being performed using hot isostatic pressing (HIP). Additionally,
impedance tests will be performed to compare the corrosion performance of the L-PBF parts with the experimental data reported in this study. Further, surface analysis and impedance tests will be performed on the gas-and water-atomized L-PBF parts to understand the mechanisms of pit formation and corrosion performance as a function of microstructure and the results will be reported elsewhere.

4.4 CONCLUSIONS

The following conclusions were drawn from this study to understand the effects of processing conditions (energy density) and powder characteristics (shape and size) on the corrosion performance of 17-4 PH stainless steel parts produced by laser-powder bed fusion (L-PBF):

1) The corrosion performance of the 17-4 PH stainless steel L-PBF parts fabricated using gas-and water-atomized powders strongly depend on their density. The density and consequently the corrosion current, polarization resistance, corrosion rate of the L-PBF parts did not significantly vary for high dense (97 ± 0.5 %) finer gas (D_{50}=13 \, \mu m) and water-atomized (D_{50}=17 \, \mu m) powders when fabricated using energy densities in the range from 64-104 J/mm^3. Furthermore, the corrosion performance of L-PBF parts of high density (97 ± 0.5%) showed slightly higher corrosion performance compared to the wrought sample.
2) The density and consequently the corrosion performance of coarser water-atomized powders (D$_{50}$=24 µm and 43 µm) of L-PBF parts increased with energy density when fabricated in the same range of energy densities 64-104 J/mm$^3$.

3) At energy densities of 64, 80, 84 J/mm$^3$, L-PBF parts fabricated using the finer gas-atomized (D$_{50}$=13 µm) and water-atomized (D$_{50}$=17 µm) powders showed significantly better corrosion performance compared to coarser water-atomized powders (D$_{50}$=24 µm and 43 µm) L-PBF parts.

4) At the energy density of 104 J/mm$^3$, the corrosion performance is independent of starting powder characteristics (shape and size) and atomization method (gas or water-atomization) used to produce these powders.

5) At the energy densities of 104 J/mm$^3$, the L-PBF parts fabricated using water-atomized powders (D$_{50}$=17 µm and 43 µm) powders exhibited higher polarization resistance (28000 ± 500 Ω) than wrought samples (25000 ± 1000 Ω) in the 0.5 M NaCl environment.
CHAPTER 5
EFFECTS OF HOT ISOSTATIC PRESSING ON THE CORROSION PERFORMANCE OF 17-4 PH STAINLESS STEEL GAS-AND WATER-ATOMIZED PARTS FABRICATED USING LASER-POWDER BED FUSION

5.1 INTRODUCTION

Laser powder bed fusion (L-PBF) process has received a lot of interest for fabricating complex three-dimensional components for automotive, aerospace, medical and tooling industries using various materials [1]–[3], [7], [26]. Among engineering alloys, 17-4 PH stainless steel is used in many applications that take advantage of its high mechanical properties in combination with its excellent corrosion resistance [26], [84], [90], [101], [104], [107]. Several research studies have been carried in our group as well as literature to fabricate defect-free L-PBF parts from 17-4 PH stainless steel and enhance the resulting mechanical properties using the post-processing operations such as hot isostatic pressing (HIP) and heat-treatment [2], [3], [6], [8], [15], [32], [50], [51], [56]–[69].

The enhanced densification, reduction in porosity, increased martensite phase content, as well as changes in grain size and shape resulted in an increase in mechanical properties for 17-4 PH stainless steel L-PBF parts when they were HIP treated at various conditions [69], [108]. These results are consistent with previous powder metallurgy studies on stainless
steel, magnesium, aluminum alloys on the effects of post-processing on microstructural features such as phase variation and grain refinement [16], [18]–[20], [24], [109]–[111]. Other studies have indicated how microstructural features can also affect the corrosion properties of post-processed [32], [40]–[42] L-PBF parts [87], [109], [112], [113]. For example, Ralston and Birbilis [114] reviewed the effect of grain size and phase variation on the corrosion performance of Co-Cr, steel, Al and Mg alloys. They have suggested that grain refinement and phase variation in the parts could change the electrochemical behavior and corrosion performance of the parts in different corrosion environments. They also indicated that grain refinement which occurs due to post-processing treatment could decrease or increase the pitting corrosion rate in the parts depending on passive layer formation in the corrosion environment.

To the best of our knowledge, there are no studies in the open literature on the effects of grain size and phase content on corrosion performance of 17-4 PH stainless steel L-PBF parts following HIP treatment. In order to address this gap, the present study was performed to understand the effects of HIP treatment on the corrosion resistance of 17-4 PH stainless steel parts fabricated by L-PBF using gas and water-atomized powders. The current study builds from our initial work on the role of powder attributes and processing conditions on the corrosion performance of as-printed 17-4 PH stainless steel L-PBF parts.

5.2 EXPERIMENTAL

L-PBF parts were fabricated from 17-4 PH stainless steel water-atomized powders of median particle size 17, 24 and 43 µm and a gas-atomized powder of median particle size
13 µm as shown in Figure 5.1 at energy densities 64 and 104 J/mm³. The detailed information about powder characteristics and L-PBF processing conditions used in the study were listed in previous journal papers published by the authors [1], [2].

HIP post-treatment was performed on as-printed L-PBF parts using a commercial service provided by Quintus Technologies, USA. The HIP treatment was conducted for 2.5 h under a temperature of 1120 °C and pressure of 105 MPa and solutionized at a temperature of 1050 °C and pressure of 70 MPa for 20 minutes applied through Argon gas, followed by
rapid cooling to room temperature at a rate of 200 °C/min [108]. Three L-PBF parts from each powder type were used for linear-sweep voltammetry experiments in 0.5 M NaCl solution at ambient conditions, to evaluate their electrochemical corrosion behavior. The electrochemical environment was exposed to the surface of the L-PBF parts parallel to the build direction. The exposed surface area of each HIP treated L-PBF part was 1 cm² and was finished by diamond polishing using a 1µm grade medium. Detailed information about the experimental setup was discussed in our previous paper [26]. The optical images of the HIP treated L-PBF parts before and after corrosion were analyzed using an Olympus BX53 microscope.

5.3 RESULTS AND DISCUSSION

The properties such as density, martensite content and average grain size of 17-4 PH stainless steel L-PBF parts before and after HIP treatment are represented in Table 5.1. It is clear from the Table 5.1 that the density, hardness of the L-PBF parts increased following HIP treatment when compared to the properties of as-printed parts. The density and hardness of all the HIP treated L-PBF parts was ~ 99.5 ± 0.2 % and ~ 40 ± 1 HRC, respectively, whereas the density and hardness of as-printed L-PBF parts slightly varied (97 ± 0.2 % and 32 ± 2 HRC) depending on the starting powder type. However, there was no measurable effect of starting powder type and size observed on the density and hardness of HIP treated L-PBF parts. Further, the increase in density and hardness in the HIP treated L-PBF parts was accompanied by a decrease in grain size when compared to as-printed L-PBF parts. Figure 5.2 shows the optical microstructures of the HIP treated and as-printed L-PBF parts parallel to the build direction which is also the surface exposed to corrosion.
environment in electrochemical corrosion tests. The microstructures of the HIP treated gas- and water-atomized L-PBF parts were noticeably different when compared to the microstructure of as-printed L-PBF parts. The scan patterns which were clearly observed in the as-printed L-PBF parts were transformed to equiaxed grains following HIP treatment. Further, the grain size in the L-PBF parts decreased following HIP treatment when compared to as-printed L-PBF parts. A similar transformation in microstructure was observed in Inconel 718 parts following HIP treatment [20]. Atkinson et al [70] indicated that the post-treatment of parts at high temperature and pressure followed by rapid cooling resulted in grain refinement.
Table 5.1. The variation in density, hardness, martensite content and average grain size of fabricated from various powders by L-PBF at an energy density 104 J/mm³, before and after HIP treatment.

<table>
<thead>
<tr>
<th>Condition</th>
<th>Property</th>
<th>Gas-atomized powder D₅₀=13µm</th>
<th>Water-atomized powder D₅₀=17µm</th>
<th>Water-atomized powder D₅₀=24µm</th>
<th>Water-atomized powder D₅₀=43µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIP</td>
<td>Density (%)</td>
<td>99.5 ± 0.2</td>
<td>99.5 ± 0.2</td>
<td>99.5 ± 0.2</td>
<td>99.5 ± 0.2</td>
</tr>
<tr>
<td>As-Printed</td>
<td>Density (%)</td>
<td>97 ± 0.5</td>
<td>97.5 ± 0.5</td>
<td>97 ± 0.2</td>
<td>97.5 ± 0.5</td>
</tr>
<tr>
<td>HIP</td>
<td>Hardness (HRC)</td>
<td>40 ± 1</td>
<td>40 ± 1</td>
<td>40 ± 1</td>
<td>40 ± 1</td>
</tr>
<tr>
<td>As-Printed</td>
<td>Hardness (HRC)</td>
<td>31 ± 1</td>
<td>34 ± 1</td>
<td>32 ± 1</td>
<td>32 ± 1</td>
</tr>
<tr>
<td>HIP</td>
<td>Martensite (%)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>As-Printed</td>
<td>Martensite (%)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>HIP</td>
<td>Grain size (µm)</td>
<td>15 ± 3</td>
<td>15 ± 2</td>
<td>15 ± 2</td>
<td>15 ± 3</td>
</tr>
<tr>
<td>As-Printed</td>
<td>Grain size (µm)</td>
<td>35 ± 2</td>
<td>25 ± 3</td>
<td>28 ± 3</td>
<td>25 ± 2</td>
</tr>
</tbody>
</table>

The average grain size of HIP treated and as-printed gas and water-atomized parts are also shown in Table 5.1. The average grain intersect method (AGI) [115] was used for quantifying the grain size present in the optical microstructures (Figure 5.2). The average grain size of the HIP treated L-PBF parts was ~ 15 ± 3 µm whereas the grain size of the as-printed L-PBF parts was dependent on starting powder type, ranging between 25 ± 2 µm and 35 ± 2 µm. A notable aspect of the study is the increase in mechanical properties of the HIP treated L-PBF parts was accompanied by not only an increase in density but
also a decrease in grain size. Therefore, it is essential to understand the variation in grain size and density on the corrosion performance of the HIP treated L-PBF parts.

Potentiostatic polarization tests were performed on HIP treated L-PBF parts in 0.5 M NaCl solution and the Tafel plots of were shown in Figure 5.3. Figure 5.3 shows that the anodic and cathodic behavior of HIP treated L-PBF parts were noticeably different from the as-printed L-PBF parts. The Tafel plots of the HIP treated L-PBF parts slightly moved to positive direction on X-axis (potential) and negative direction on Y-axis (corrosion current density) when compared to as-printed Tafel plots of both gas and water-atomized powders. This indicates that the corrosion current density of the L-PBF parts decreased and corrosion potential increased following HIP treatment. Further, the passive region in the L-PBF parts increased following HIP treatment. Previous electrochemical corrosion study by Stoudt et al [113] on 17-4 PH stainless steel gas-atomized L-PBF parts have shown the similar cathodic behavior in Tafel plots when they are heat-treated.
Figure 0.3. Tafel plots of 17-4 PH stainless steel parts fabricated with various powder by L-PBF at an energy density 104 J/mm$^3$, before and after HIP treatment.
Table 5.2. The variation in corrosion current and corrosion potential of 17-4 PH stainless steel parts fabricated by L-PBF at an energy density 104 J/mm$^3$, before and after HIP treatment.

<table>
<thead>
<tr>
<th>Property</th>
<th>Corrosion current (A)</th>
<th>Corrosion potential (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HIP</td>
<td>As-printed</td>
</tr>
<tr>
<td>Gas-atomized $D_{50} = 13 \mu m$</td>
<td>0.008 ± 0.001</td>
<td>0.35 ± 0.05</td>
</tr>
<tr>
<td>Water-atomized $D_{50} = 17 \mu m$</td>
<td>0.008 ± 0.001</td>
<td>0.3 ± 0.04</td>
</tr>
<tr>
<td>Water-atomized $D_{50} = 24 \mu m$</td>
<td>0.008 ± 0.001</td>
<td>0.6 ± 0.07</td>
</tr>
<tr>
<td>Water-atomized $D_{50} = 43 \mu m$</td>
<td>0.008 ± 0.001</td>
<td>0.3 ± 0.05</td>
</tr>
</tbody>
</table>

The corrosion current of L-PBF parts decreased due to HIP treatment when compared to as-printed L-PBF parts and remained constant at 0.008 ± 0.001 µA. However, the corrosion current of the as-printed L-PBF parts slightly varied between 0.03 ± 0.005 and 0.6 ± 0.07 µA depending on the starting powder type. Further, an increase in the corrosion potential was observed for the HIP treated L-PBF parts (-150 ± 5 mV) when compared to corrosion potential of the as-printed L-PBF parts (-380 ± 20 mV). The increase in corrosion potential and a decrease in the corrosion current in the HIP treated L-PBF parts can be attributed to the reduced grain size (15 ± 3 µm) and increased density (99 ± 0.5 %). Previous studies by Gollapudi et al [116] and Obayi et al [117] have reported a similar behavior for annealed ferrous parts. For further comparison, the corrosion current and corrosion potential of the wrought 17-4 PH stainless steel is 0.9 ± 0.1 µA and -400 ± 20 mV, respectively.

The corrosion current values were used for the calculation of polarization resistance and corrosion rate of the L-PBF parts in 0.5 M NaCl solution as shown in Table 5.2. The equations used for calculation of the above-mentioned parameters have been listed in previous papers published by authors [26].
Table 5.3. The variation in polarization resistance and corrosion rate of 17-4 PH stainless steel parts fabricated by L-PBF at an energy density 104 J/mm³, before and after HIP treatment.

<table>
<thead>
<tr>
<th>Property</th>
<th>Polarization resistance (Ω cm²)</th>
<th>Corrosion rate (µm/year)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HIP</td>
<td>As-printed</td>
</tr>
<tr>
<td>Gas-atomized D₅₀ = 13 µm</td>
<td>40000 ± 500</td>
<td>23500 ± 500</td>
</tr>
<tr>
<td></td>
<td>HIP</td>
<td>1 ± 0.3</td>
</tr>
<tr>
<td></td>
<td>As-printed</td>
<td>3.5 ± 0.5</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 17 µm</td>
<td>40500 ± 200</td>
<td>27500 ± 500</td>
</tr>
<tr>
<td></td>
<td>HIP</td>
<td>1 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>As-printed</td>
<td>1.8 ± 0.5</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 24 µm</td>
<td>40000 ± 700</td>
<td>23000 ± 500</td>
</tr>
<tr>
<td></td>
<td>HIP</td>
<td>1.2 ± 0.1</td>
</tr>
<tr>
<td></td>
<td>As-printed</td>
<td>7 ± 1</td>
</tr>
<tr>
<td>Water-atomized D₅₀ = 43 µm</td>
<td>40200 ± 500</td>
<td>27500 ± 200</td>
</tr>
<tr>
<td></td>
<td>HIP</td>
<td>1 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>As-printed</td>
<td>1.7 ± 0.5</td>
</tr>
</tbody>
</table>

The polarization resistance of the L-PBF parts increased following HIP treatment when compared to the polarization resistance of as-printed L-PBF parts. The polarization resistance of the HIP treated gas- and water-atomized L-PBF parts was ~ 40,000 ± 500 Ω cm² whereas the polarization resistance of the as-printed gas- and water-atomized L-PBF parts ranged between 23,000 ± 500 Ω cm² and 27,500 ± 500 Ω cm². However, the corrosion rate of the HIP treated samples was significantly lower than the corrosion rate of as-printed L-PBF parts. The corrosion rate of the HIP treated gas- and water-atomized L-PBF parts was found to be ~ 1 ± 0.3 µm/year whereas the corrosion rate of the as-printed gas- and water-atomized L-PBF parts ranged between 1.7 ± 0.5 µm/year and 7 ± 1 µm/year, depending on powder type used for the parts fabrication. For further comparison, the polarization rate and corrosion rate of wrought 17-4 PH stainless steel is 25,000 ± 1,000 Ω cm² and 3 ± 0.5 µm/year, respectively.

The low corrosion rate in the HIP treated L-PBF parts could be correlated to their optical images after corrosion testing as shown in Figure 5.4. The pits observed in the HIP treated L-PBF parts for all powders after the corrosion experiment was lower than for as-printed parts, which was consistent with the quantitative Tafel parameters presented in Table 5.3.
This high polarization resistance and low corrosion rate in the L-PBF parts after HIP treatment is consistent with the increased density as well as reduced grain size in the microstructures. An important result of this study is the corrosion resistance of 17-4 PH stainless steel L-PBF parts were enhanced due to HIP treatment and did not significantly depend on the powder type used for the L-PBF parts fabrication, in contrast to the corrosion trends in as-printed parts. These results also demonstrated the feasibility of achieving similar corrosion performance from both gas and water-atomized L-PBF parts following HIP treatment.

Figure 0.4. Optical micrographs following corrosion tests of HIP treated and as-printed 17-4 PH stainless steel parts fabricated by L-PBF at energy density 104 J/mm$^3$ using gas and water-atomized powders.

HIP treated gas and water-atomized L-PBF parts have a smaller grain size compared to as-printed L-PBF parts which will result in a high volume of grain boundaries. It is probable that the high volume grain boundaries would enhance the chemical activity and dissolution rate in the corrosion environment. This increase in chemical activity could favor the formation of a thick and stable oxide layer on the surface. The thick and stable oxide layer could, in turn, inhibit the adsorption of the Cl$^-$ ions on the surface of HIP treated L-PBF parts [114], [116]–[118].
5.4 CONCLUSIONS

A study was performed to understand the effect of hot isostatic pressing on the corrosion performance of 17-4 PH stainless steel gas- and water-atomized parts produced by laser-powder bed fusion (L-PBF). The corrosion resistance of the L-PBF parts was evaluated using linear-sweep voltammetry curves (Tafel plots) obtained from the electrochemical corrosion tests. The anodic and cathodic behavior as well as the location of passive region in Tafel plots HIP treated L-PBF parts were significantly different from Tafel plots of as-printed L-PBF parts. The density (99.5 ± 0.2 %), corrosion potential (-150 ± 5 mV), and polarization resistance (40000 ± 500 Ω cm²) of the L-PBF parts increased due to HIP treatment and higher than as-printed properties. Further, the corrosion rate of the HIP treated L-PBF parts was lower than the as-printed L-PBF parts irrespective of starting powder type. The superior corrosion performance of the HIP treated L-PBF parts could be correlated to the decrease in grain size when compared to as-printed L-PBF parts. A similar corrosion performance was observed for both expensive gas-atomized and inexpensive water-atomized L-PBF parts when they are HIP treated.
CHAPTER 6
CONCLUSIONS AND FUTURE WORK

6.1 CONCLUSIONS
The goal of this dissertation is to establish a fundamental understanding of the material-process-property relationships of 17-4 PH stainless steel processed by laser-powder bed fusion (L-PBF). The investigation was carried out to understand the effects of post-processing treatment cycles such as hot isostatic pressing (HIP) (Chapters 2 and 3) on the densification, mechanical properties, corrosion properties (Chapters 4 and 5) and microstructures of L-PBF parts fabricated using 17-4 PH stainless steel gas- and water-atomized powders at various energy densities. The microstructure formation and its impact on mechanical and corrosion properties due to different HIP treatment cycles were studied for 17-4 PH stainless steel gas and water-atomized L-PBF parts at various energy densities. The most significant aspect of this work is that density, microstructures, mechanical and corrosion properties of 17-4 PH stainless steel can be vastly improved irrespective of starting powder attributes and L-PBF process conditions by using HIP treatment and are superior to known reported properties of 17-4 PH stainless steel fabricated by L-PBF or wrought samples. The conclusions emerging from this dissertation are summarized in the following sections.
6.1.1 MECHANICAL PROPERTIES AND MICROSTRUCTURES

A detailed study was performed using different HIP treatment cycles on L-PBF parts fabricated at various energy densities using 17-4 PH stainless steel gas and water-atomized powders. The results from this work show the strong dependence of densification, mechanical properties and microstructures on temperature, pressure and time during the HIP cycle.

- The mechanical properties of both gas and water-atomized L-PBF parts improved following HIP treatment.

Figure 6.1. Ultimate tensile strength of HIP treated gas and water-atomized L-PBF parts when compared to other approaches

- The ultimate tensile strength of both gas and water-atomized L-PBF parts increased due to HIP treatment and less sensitive to powder and process variations compared to other approaches as shown in Figure 6.1.

- Similar trends were observed when yield strength and hardness values of HIP treated gas and water-atomized L-PBF parts were compared to other approaches.

- Following HIP treatment, 99.5 ± 0.2 % dense L-PBF parts with ultimate tensile strength: 1450 ± 20 MPa, hardness: 40 ± 1 HRC, yield strength: 1180 ± 20 MPa and elongation: 10 ± 1 % were achieved using both water and gas-atomized
powders. For reference, the as-printed properties ranged as follows: Density: 88 ± 1 to 97.5 ± 0.2 %, Ultimate tensile strength: 480 ± 30 to 1070 ± 50 MPa, Hardness: 27 ± 1 to 36 ± 1 HRC, Yield strength: 400 ± 50 to 800 ± 50 MPa.

- The density, martensite content and equiaxed microstructure in L-PBF parts after HIP treatment were consistent with the trends in the mechanical properties of the L-PBF parts.
- Martensite phase and equiaxed microstructures after HIP treatment was less sensitive to powder/process variations compared to other approaches.
- Robust and reliable properties were obtained after the HIP treatment from both gas and water-atomized powders at various energy densities. The properties obtained after the HIP treatment were less sensitive to variations in powder characteristics (size distribution and shape) and energy density during processing and were superior to known reported properties of 17-4 PH stainless steel fabricated by L-PBF, powder metallurgy, metal injection molding, or wrought samples.

6.1.2 CORROSION PROPERTIES

The following conclusions were drawn from the corrosion performance evaluation studies performed on as-printed (Chapter 4) and HIP treated L-PBF parts (Chapter 5):

6.1.2.1 AS-PRINTED L-PBF PARTS

Corrosion studies were conducted to understand the effects of processing conditions (energy density) and powder characteristics (shape and size distribution) on the corrosion
performance of as-printed 17-4 PH stainless steel parts produced by laser-powder bed fusion (L-PBF):

- The corrosion performance of the as-printed 17-4 PH stainless steel L-PBF parts fabricated using gas and water-atomized powders strongly depend on their density. The density and consequently the corrosion current, polarization resistance, and corrosion rate of the as-printed L-PBF parts did not vary significantly for highly dense (97 ± 0.5 %), finer gas (D_{50}=13 µm) and water-atomized (D_{50}=17 µm) powders when fabricated using energy densities in the range from 64-104 J/mm³. Furthermore, the corrosion performance of as-printed L-PBF parts of high density (97 ± 0.5%) showed slightly higher corrosion performance compared to the wrought sample.

- The density and consequently the corrosion performance of as-printed L-PBF parts fabricated using the coarser water-atomized powders (D_{50}=24 µm and 43 µm) increased with energy density when fabricated in the same range of energy densities 64-104 J/mm³.

- At energy densities of 64, 80, and 84 J/mm³, as-printed L-PBF parts fabricated using the finer gas-atomized (D_{50}=13 µm) and water-atomized (D_{50}=17 µm) powders showed significantly better corrosion performance compared to coarser water-atomized powders (D_{50}=24 µm and 43 µm) L-PBF parts.

- At the energy density of 104 J/mm³, the corrosion performance does not vary significantly as a function of starting powder characteristics (shape and size) and atomization method (gas or water-atomization) used to produce these powders.
• At the energy densities of 104 J/mm$^3$, the as-printed L-PBF parts fabricated using water-atomized powders ($D_{50}$=17 µm and 43 µm) powders exhibited a slightly higher polarization resistance (28000 ± 500 Ω) compared to wrought samples (25000 ± 1000 Ω) in 0.5 M NaCl.

6.1.2.1 HIP TREATED L-PBF PARTS

A study was performed to understand the effect of hot isostatic pressing on the corrosion performance of 17-4 PH stainless steel gas and water-atomized parts produced by laser-powder bed fusion (L-PBF) at an energy density 104 J/mm$^3$.

• The anodic and cathodic behavior as well as the breakdown of the passive region in Tafel plots HIP treated L-PBF parts were significantly different from the corresponding attributes found in Tafel plots of as-printed L-PBF parts.

• The corrosion potential (-150 ± 5 mV), polarization resistance (40000 ± 500 Ω cm$^{-2}$) of the L-PBF parts increased following HIP treatment and was higher than the corresponding as-printed properties, consistent with the increase in density (99.5 ± 0.2 %).

• The corrosion rate (1 ± 0.3 µm/year) of the HIP treated L-PBF parts was lower than the as-printed L-PBF parts (1.7 ± 0.5 µm/year to 7 ± 1 µm/year) irrespective of starting powder type.

• The superior corrosion performance of the HIP treated L-PBF parts could also be correlated to the decrease in grain size when compared to as-printed L-PBF parts.
• A similar corrosion performance (corrosion current: 0.008 ± 0.001 µA, corrosion potential: -150 ± 5 mV, polarization resistance: 40000 ± 500 Ω cm-2, corrosion rate: 1 ± 0.3 µm/year) was observed for both gas and water-atomized L-PBF parts following HIP treatment

6.2 FUTURE WORK

This dissertation was undertaken to understand several new aspects affecting material-process-property-microstructure relationships when 17-4 PH stainless steel gas and water-atomized powders were used for fabricating L-PBF parts. The findings of this work have identified several areas for further work to build from the studies presented in this dissertation. They are:

• An understanding of the fatigue life of 17-4 PH stainless steel L-PBF parts as a function of powder attributes, L-PBF processing conditions and post-processing operations such as HIP and heat treatment needs to be undertaken

• A three-dimensional porosity, phase and grain distribution analysis of 17-4 PH stainless steel L-PBF parts can provide further insight into the formation as well as influence of pore, phase and grain distributions on mechanical and corrosion properties

• Understanding the effects of part build orientation on the mechanical and corrosion properties as well as microstructures can provide an insight on the impact of HIP treatment on anisotropic L-PBF parts
• The experimental results from the present study need to help refine modeling and simulation to further understand the effects of powder packing, flow and densification as a function of initial powder attributes (shape, chemistry and size) to improve process reliability, revise methodologies to evenly distribute powders to a high packing density in powder bed process and enhance design process tools for L-PBF.

• The results from this work on 17-4 PH stainless steel should be extended to other alloy systems as well as revised water-atomization procedures that enhance powder packing and flow attributes by modifying shape and size distributions.
REFERENCES


APPENDIX A

EFFECTS OF ATOMIZING MEDIA AND POST PROCESSING ON MECHANICAL PROPERTIES OF 17-4 PH STAINLESS STEEL MANUFACTURED VIA SELECTIVE LASER MELTING

A.1 INTRODUCTION

17-4 PH stainless steel is known as AISI 630 stainless steel and is a lath martensitic stainless steel hardened by precipitation of Cu-rich spherical particles in the martensitic matrix. 17-4 PH stainless steels have good mechanical properties and corrosion resistance at typical service temperatures below 300 °C [1,2] and are used in marine environments, power plants (light-water and pressurized water reactors) and powder metal injection molding industries. With solution annealing of 17-4 PH stainless steel at 1050 °C and aging at 482 °C, Cu-rich phase precipitate as coherent spherical bcc structures that would result in increased tensile strength and toughness [3]. Coherent (bcc) Cu-rich precipitates usually transform to non-coherent fcc-Cu-rich particles upon extended aging at 400 °C [4]. Reversion of martensite to the austenite during aging is explained by the concentration of austenite stabilizing elements led by diffusion and lower martensitic transformation start (Ms) temperature [5].
Selective Selective laser melting (SLM) is an additive manufacturing (AM) technology to produce complex three-dimensional parts through solidifying successive layers of powder materials on the basis of a 3D computer aided design (CAD) model. SLM is associated with complete melting of the powder material [6,7]. Because the weldability of 17-4 PH martensitic stainless steel is good for the usual arc welding processes, this class of steels is a promising alloy for SLM method [8,9]. Researchers have manufactured 17-4 PH components via SLM technique and have reported formation of metastable austenite and martensite phase as a result [10,11]. Murr et al. [12] explored the role of nitrogen and argon as the atmosphere of SLM chamber on formation of austenitic structure and ferritic/martensitic, respectively. Co-existence of both martensite and retained austenite in the SLM manufactured parts produced in nitrogen atmosphere clearly indicates that the nitrogen atmosphere has influenced the stabilization of austenite [12]. Facchini et al. [11] analysis showed that the as-fabricated 17-4 PH stainless steel has 70% mass fraction of metastable austenite, which transformed to the martensite phase during tensile deformation [13,14].

Farshidianfar et al. [15] has carried experiments to control the microstructure of Laser Additive Manufacturing (LAM) parts by controlling the cooling rate. Yadollahi et al. and Luecke [8,16] investigated the effect of build rotation during SLM on tensile and fatigue properties of 17-4 PH steel parts and found un-melted regions caused by lack of fusion significantly reduced their tensile and fatigue strength. Rafi et al. [17] investigated the mechanical properties and microstructure of 17-4 PH manufactured by SLM and found that columnar grains with smaller diameters (< 2 μm) which appear in the melt pool had a
combination of martensite and retained austenite phases. They found that the processing environment, grain size and chemistry of powder had a great influence on the phase content of sample.

Recently, powder characteristic such as size, distribution and morphology of particles got special concern in SLM process [18–20]. In all of the SLM process of 17-4 PH SS, gas atomized 17-4 PH was used as feedstock [8,12,17,21]. However, the production cost of per kilogram of water atomized powder is relatively lower than that of gas atomized powder [22]. Considering 17-4 PH water-atomized powder that has much less nitrogen (0.02 wt.%) at a cheaper production rate is a novel approach that will mitigate the aforementioned issues of cost and high nitrogen content. Few studies have evaluated the final properties of water-atomized 316L stainless steel and H13 tool steel powder equivalent to those produced with gas-atomized powder. Comparing those made with gas-atomized and water-atomized powder, the water-atomized samples shows superior surface finish, uniformity in deposition and bonding between layers. [23–25]. In our former study [18], gas-atomized 17-4 PH powders when processed at lower energy densities (80–84 J/mm3) showed better densification and mechanical properties compare to high energy density (104 J/mm3). Using high energy density of 104 J/mm3 for water-atomized powder, made the relative density and mechanical properties, comparable to the gas atomized powder. Only as printed parts were characterized in our former study [18], and no data were provided on the post-processing of 17-4 PH components manufactured by SLM process.
The objective of this study is to investigate the effects of atomizing media (water atomization versus gas atomization) as well as heat treatment (solution annealing and aging) temperature on phase transformation and mechanical properties of laser melted 17-4 PH parts. Our results (as shown later) will demonstrate the practicality of using relatively cheaper water-atomized 17-4 PH powders as feedstock strongly depends on designing a non-standard and effective thermal post processing.

A.2 MATERIALS AND METHODS

Gas-atomized and water-atomized 17-4 PH SS powders procured from Sandvik and North American Hoganas (NAH), respectively. The chemical composition of gas-atomized and water-atomized powder, which is provided by vendors, is given in Table A.1. Water-atomized 17-4 PH powder had higher carbon content than gas-atomized powder because it was designed for conventional powder metallurgy (pressing and sintering) and metal injection molding. It is because higher carbon content of water-atomized 17-4 PH would result in higher green strength required in conventional powder metallurgy [22]. Water-atomized 17-4 PH powder with chemical composition similar to that of gas-atomized powder is not available.

Table A.1. Chemical composition of 17-4 PH stainless steel for gas-atomized and water-atomized powders (wt.%).

<table>
<thead>
<tr>
<th>Powder</th>
<th>C</th>
<th>Cr</th>
<th>Cu</th>
<th>Mn</th>
<th>Ni</th>
<th>P</th>
<th>S</th>
<th>Si</th>
<th>Nb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized</td>
<td>0.03</td>
<td>15-17.5</td>
<td>3-5</td>
<td>1</td>
<td>3-5</td>
<td>0.04</td>
<td>0.03</td>
<td>1</td>
<td>0.25</td>
</tr>
<tr>
<td>Water-atomized</td>
<td>0.208</td>
<td>17.74</td>
<td>3.94</td>
<td>0.13</td>
<td>3.54</td>
<td>0.013</td>
<td>0.008</td>
<td>0.3</td>
<td>0.35</td>
</tr>
</tbody>
</table>

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A 3D Systems ProX 200 machine with an yttrium fiber laser system with maximum power of 300W was used to manufacture the tensile specimens under argon atmosphere as per specifications of ASTM E8 standard. The specimens were printed using two different laser energy densities (64 and 104 J/mm$^3$). Two different post processing regimes were used in this study; (1) specimens were solutionized at 1051 °C for 1h in H2 atmosphere and then aged at 482 °C in N$_2$ atmosphere for 1h (H900) and (2) specimens were solutionized at 1315 °C for 45 min in H$_2$ atmosphere and then aged at 482 °C for 1 h in N$_2$ atmosphere. The specimens were then polished using a standard polishing procedure and electro-etched in a solution of (NaOH 33% Electrolyte at 5 V for 2 min). An Instron 5982 dual column testing system utilized with a 100 kN force load cell and strain rate of 10$^{-3}$ s$^{-1}$ was used for tensile tests.

Hardness was measured using the Rockwell C hardness scale (150kg load). Density of specimens were measured by Archimedes method using a Metter Toledo XS104 weighing balance equipped with a density measurement kit. Leica optical microscope was utilized to study the microstructure of printed components. A Bruker D8 discover X-ray diffraction instrument was used to analyze the phase structure. Scheil model for non-equilibrium solidification and prediction of phase fraction during SLM process were performed using Thermo-Calc software integrated with TCFE8 database. The simulation were performed for a step size of 1 °C from liquid phase up to remaining liquid fraction reach the 0.1. Microstructure of heat treated samples were studied by transmission electron microscopy (TEM) using FEI Titan TEM/STEM with ChemiSTEM capabilities. FEI Quanta 3D field emission dual beam scanning electron microscope (SEM/FIB) was used to prepare samples for TEM.
A.3 RESULTS AND DISCUSSION

The powder feedstock characteristics such as particle size distribution (PSD), apparent density (AD), tap density (TD) and Hausner ratio values were measured as shown in Table A.2. The AD of powder is defined by mass per unit volume of loose powder which provides a measure of the “fluffiness” of a powder [26].

<table>
<thead>
<tr>
<th>Powder</th>
<th>Particle distribution</th>
<th>Tap density (g/cc)</th>
<th>Apparent density (g/cc)</th>
<th>Hausner Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized (G)</td>
<td>$D_{10}$ (µm): 5</td>
<td>$D_{50}$ (µm): 13</td>
<td>$D_{90}$ (µm): 27</td>
<td>4 ± 0.05</td>
</tr>
<tr>
<td>Water-atomized (W)</td>
<td>$D_{10}$ (µm): 26</td>
<td>$D_{50}$ (µm): 43</td>
<td>$D_{90}$ (µm): 67</td>
<td>3.7 ± 0.05</td>
</tr>
</tbody>
</table>

The median particle size of gas atomized and water atomized powder were 13 and 43 µm, respectively. PSD selected here was similar to PSD used by Rafi et al. [17] with the mean particle size of 39 µm and PSD used by Murr et al. [27] with two different mean particles sizes of 19 µm and 25 µm. In our former study [16], four different PSD were used with mean particle size of 13, 17, 24 and 43 µm, however our primary focus of this study is on mean particle size of 13 and 43 µm. The density of manufactured part is related to the density of powder bed. Many factors influence the density of powder bed such as particle shape, particle size, powder surface morphology and flowability of the powder [28]. Therefore, powder flowability is one of the topics of concern when producing parts in SLM process. If the powder does not spread in a consistent manner, the density of manufactured part varies from region to other region and the porosity emerged in the manufactured parts.
and reduces the mechanical properties. The Hausner ratio is a number which is correlated to the powder flowability and is a useful measure of cohesion [26]. It is calculated using the equation shown below [11]:

\[ H = \frac{\rho_T}{\rho_A} \]  

(A.1)

Where \( \rho_T \) is tap density and \( \rho_A \) is apparent density. Hausner ratio of water-atomized (1.301) was less than gas-atomized (1.324). Many factors play rolls in flowability of powders such as the shape and size of particles. In general, gas atomized powders, because of having spherical shape, show better flowability rather than water-atomized powders which have irregular shape [28]. However, in this study, because water-atomized powders have bigger particle size, the interlocking forces between the particles reduce and eliminate the roll of particle shape, therefore, the flowability of water-atomized powders is comparable to gas-atomized powder. **Figure A.1** shows the SEM micrographs of the two powders used in this study. Gas-atomized powders shown in **Figure A.1a** are spherical particles with \( D_{50}=13 \) μm. Water-atomized powder shown in **Figure A.1b** had a mixture of semi-spherical and rounded shapes with relatively clean particles without satellites and \( D_{50}=43 \) μm.
Figure A.0.1. Morphology of 17-4 PH SS powder: (a) gas atomized powder and (b) water atomized powder [29].

Figure A.2a and b shows the XRD-patterns of gas-atomized and water-atomized powders, respectively. While gas-atomized and water-atomized powders contain both austenite and martensite phases, austenitic phase of water-atomized is more dominant in volume fraction. On the other hand, gas-atomized powder has a higher volume fraction of martensitic phase. The austenite phase in gas-atomized powder is because of using nitrogen as atomizing media for atomization of the melt with relatively a slower cooling rate [12]. As Table A.1 shows, the carbon content of water-atomized powder (0.208 wt.%) is much higher than gas-atomized carbon content (0.03 wt.%). Furthermore, the cooling rate for water-atomized powder are 10–100 times larger than gas atomized powders [30]. Therefore, combined effect of higher carbon as an austenite stabilizer [11] and faster cooling rate in water atomization led to the presence of higher amount of austenite phase in the powder feedstock. Furthermore, nitrogen can be picked up during gas atomization process and along with martensitic crystal lattice expansion can lead to deformation, smaller grains and broader peaks [31] as observed in Figure A.2a. In case of water-atomized powder, grain size of austenite is coarser than that of gas-atomized. The average crystallite size of gas
and water-atomized powder were calculated to be 235 nm and 552 nm, respectively by using Williamson-Hall method [32]. The microstructure of as printed 17-4 PH gas-atomized powder perpendicular to the build direction at energy density of 64 J/mm³ and 104 J/mm³ are shown in Figure A.3a and b, respectively. The microstructure printed at 64 J/mm³ exhibited coarser lath martensitic structure as shown in Figure A.3a, and ultrafine martensitic structure was observed in Figure A.3b. Single martensitic phase was observed in samples manufactured from gas-atomized powder independent of energy density. Microstructure of water-atomized powder printed at energy density of 64 J/mm³ and 104 J/mm³ are shown in Figure A.3c and d, respectively. Similar lath martensitic structure located in islands of retained austenite were observed in Figure A.3c and d. Water-atomized powder printed at 64 J/mm³, showed formation of a continuous network of precipitates along the grain boundaries (as shown in Figure A.3c) that are likely enriched in carbon. As shown in Figure A.3c, significant amount of large porosity (larger than 10 μm) can be due to lack of fusion at lower energy density of 64 J/mm³. In Figure A.3d, the precipitates formation was not observed along the grain boundaries likely due to dissolution at higher laser power and slower scan rate (larger melt pool region with slower solidification rate). Both austenite and martensite phases were present in samples manufactured from water-atomized powder, because of combined effect of dissolution of carbon that can stabilize formation of austenite and rapid solidification during SLM that can lead to formation of martensite [17].

The XRD-patterns for both water and gas-atomized specimens at both the laser energy densities (64 J/mm³ and 104 J/mm³) are shown in Figure A.4a and b, respectively. Printed
gas-atomized powder showed only martensitic phase. The phases for gas-atomized remained unchanged after increasing energy density from 64 to 104 J/mm$^3$ (Figure A.4a). The as printed water-atomized powder samples dual austenitic and martensitic structure at both energy density of 64 and 104 J/mm$^3$. At energy density of 64 J/mm$^3$, the laser power and scan speed was at 150W and 1550 mm/s, respectively. At energy density of 104 J/mm$^3$, the laser power and scan speed was at 195W and 1250 mm/s, respectively. At higher energy density of 104 J/mm$^3$, the fraction of martensitic phase increased. The presence of retained austenite phase after printing of water-atomized powder was found to be independent of laser energy density (Figure A.4b). This could likely due to the higher carbon content in water-atomized powder feedstock. This could imply that the phase transformation during SLM is highly dependent to initial feedstock chemistry. Therefore, feedstock chemistry and cooling rate can have a significant role on phase transformation upon rapid solidification during SLM process [33]. Water-atomized powder used in this study was not commercially manufactured and modified for SLM process. This study is only a preliminary evaluation of water-atomized powder capability to be used as a cheaper alternative feedstock for SLM.

Figure A.5 shows cross section of gas-atomized and water-atomized samples, parallel to the build direction. Gas-atomized powder printed at low energy density (Figure A.5a) shows a pattern of repeated rectangular with the mean height of about 50 μm, which is the same as the thickness of each layer of powder bed. Saeidi et al. [2] reported similar pattern in SLM of duplex stainless steel. On the other hand, water-atomized sample printed at energy density of 64 J/mm$^3$ (Figure A.5b) did not show such pattern. This may be
attributed to the lower apparent density of water-atomized powder (2.813 g/cm$^3$) compare to gas-atomized powder (3.040 g/cm$^3$). The particles in water-atomized powders after melting may collapse into the empty spaces between the particles, so the melting pool boundaries can change and became very hard to distinguish the melting pool boundaries. With increasing the energy density to 104 J/mm$^3$ (Figure A.5c and d), the melt pool boundaries are no longer distinct. Therefore, the temperature is high enough for melting and mixing all the particles to make a fully austenite liquid that could transform to martensite.

![XRD patterns](image)

Figure A.0.2. The XRD pattern of 17-4 PH powders (a) gas-atomized and (b) water-atomized.

The optical micrographs, perpendicular to the build direction, obtained from gas-atomized powder printed at 64 and 104 J/mm$^3$ and solutionized at 1051 °C and aged at 482 °C are shown in Figure A.6a and b, respectively. Microstructure of gas atomized samples printed at 64 and 104 J/mm$^3$, revealed martensitic structure. Furthermore, fine martensitic structure transformed to coarser martensitic structure at energy density of 104 J/mm$^3$ after heat treatment. The microstructure of heat treated water-atomized powder revealed single martensitic phase at energy density of 64 J/mm$^3$ and dual martensitic and austenitic
structure at energy density of 104 J/mm$^3$ as presented in Figure A.6c and d, respectively. After heat treatment, the microstructure became uniform and predominately a martensitic structure with no trace of the melt pools for 64 J/mm$^3$ sample for both water-atomized and gas-atomized powder. Presence of austenite in Figure A.6d could be due to reversion austenite phenomena. Reversion austenite was also observed in selective laser melting of 17-4 PH [5]. Process of austenite reversion has been generally attributed to the chemical stabilization by the diffusion of alloying element constituents, commonly toward lath boundaries. More specifically, as Cu diffuses to form precipitates, the local $M_f$ temperature is effectively lowered at the sites of their formation and thereby permits austenite chemical stabilization at room temperature [5].

![Figure A.0.3 Optical micrographs of as printed 17-4 PH SS powder perpendicular to the build direction: (a) gas-atomized at 64 J/mm$^3$, (b) gas-atomized at 104 J/mm$^3$, (c) water-atomized at 64 J/mm$^3$ and (d) water-atomized at 104 J/mm$^3$.](image-url)
The XRD patterns for gas-atomized and water-atomized powder after printing at energy density of 64 and 104 J/mm$^3$ and solution annealing at 1051 °C and aging at 482 °C are shown in Figure A.7a and Figure A.7b, respectively. In gas-atomized powder, single martensite phase was observed. In water-atomized powder, after thermal aging, single martensite phase was observed at energy density of 64 J/mm$^3$. This is likely because of the austenite formed during solutionizing transforms to martensite while quenching through the Ms temperature. However, at energy density of 104 J/mm$^3$ dual martensite and reverse austenite phases were observed as shown in Figure A.6d. Reversion or local stabilization of austenite by the diffusion of Cu and Ni in water-atomized powder during aging could lead to austenite concentrations at energy density of 104 J/mm$^3$. The activation energy for forming of reverse austenite is 240 kJ/mol which is very close to the activation energy needed for diffusion of Ni in martensitic matrix (245 kJ/mol) [34]. Higher carbon content in water-atomized powder and dissolutions of these carbon at higher energy density of 104 J/mm$^3$ (higher laser powder 195 W and slower scan rate 1250 mms) would lead in to local enrichment of carbon, which is an austenite stabilizer elements, as shown in Figure A.3d. Increasing localized concentration of austenite stabilizing elements would locally lower the martensitic transformation start (Ms) temperature below room temperature and increase the amount of austenite [5]. This phenomenon became significant after aging and as a result, led to significant reverse austenite formation in water-atomized powder printed at 104 J/mm$^3$ as can be seen in Figure A.6d and in XRD plot of Figure A.7b.
While 17-4 PH water-atomized powder has higher content of carbides (MX and M23C6), these carbides can potentially get dissolved at higher energy density (104 J/mm$^3$) due to higher power and lower scanning speed. However, at lower energy density (64 J/mm$^3$), lower input energy and higher scanning speed does not lead to dissolution of carbides. Relationship between energy density and dissolution of carbides can be explained due to more fusion and larger melt pool at higher energy density and higher diffusion rate of Ni and Cu in the matrix.

According to Ziewiec et al. [35] in non-equilibrium solidification conditions of the weld 17-4PH, segregation and the diffusion of copper and the elements stabilizing the austenite (such as carbon and nitrogen) cause the occurrence of the reverse transformation of the martensite into austenite as fast as just 1 h at 620 °C. The increase of copper-carbides solubility in the austenite causes dissolution of the fcc-Cu precipitates and carbides would lead to segregation and local enrichment of austenite stabilizer elements. In the regions where segregation of the austenite stabilizer elements occurred, a stable austenite is formed.
as a result of $\alpha \rightarrow \gamma$ transformation. Similarly, after SLM and upon solutionizing, if the carbides are dissolved (at 104 J/mm$^3$), the combined effect of carbon segregation and Cu enrichment would likely lead to austenite stabilization, and this will facilitate formation of reverse austenite. Such austenite is stable after cooling down to the ambient temperature. But if carbides are not dissolved (at 64 J/mm$^3$), then carbon enrichment will not occur and reverse austenite does not form. Therefore, in water-atomized powder, reverse austenite is more plausible at higher energy density and less likely at lower energy. Gas-atomized powder did not exhibit any reverse austenite due to much lower carbon content. Yield stress (YS), ultimate tensile strength (UTS), hardness values, elongation (%) and density after solution annealing at 1051 °C and aging at 482 °C are shown in Table A.3. For comparison, the properties of wrought 17-4 PH alloy is also provided. The hardness values of gas atomized powder showed similar hardness values (42-45HRC) at energy density of 64 and 104 J/mm$^3$, which was found to be higher than hardness of wrought alloy (39 HRC). This is attributed to the rapid melting and solidification of laser additive manufacturing process which resulted to finer grain size and increase in hardness value and potentially higher dislocation density. On the other hand, hardness value of water-atomized powder was significantly less than gas-atomized and wrought alloy, but increased from 18 to 24 HRC with increasing energy density from 64 to 104 J/mm$^3$. This hardness increase could be attributed to the higher relative density of manufactured part at higher energy density (Table A.3). Similar results were observed for yield strength (YS) and ultimate tensile strength (UTS) of gas-atomized powder and water-atomized powder. While YS and UTS values of gas-atomized powder were comparable with wrought values, these values were much lower for water-atomized powder.
Figure A.0.5. Optical micrographs, parallel to the build direction, of as printed 17-4 PH SS powder at 64 J/mm$^3$ (a) gas-atomized, (b) water-atomized and at 104 J/mm$^3$ (c) gas-atomized, (d) water-atomized.
Figure A.0.6. The optical micrographs of heat treated parts (solutionized at 1051 °C and aged at 482 °C) perpendicular to the build direction: (1) gas-atomized at 64 J/mm$^3$, (b) gas-atomized at 104 J/mm$^3$, (c) water-atomized at 64 J/mm$^3$ and (d) water-atomized at 104 J/mm$^3$.

Figure A.0.7. The XRD pattern of (a) gas-atomized and (b) water-atomized printed at 64 and 104 J/mm$^3$ and heat treated (solutionized at 1051 °C and aged at 482 °C).
Table A.3. Mechanical properties at energy density of 64 and 104 J/mm$^3$ - solutionized at 1051 °C and aged at 482 °C (wrought values are taken from literature [36]).

<table>
<thead>
<tr>
<th>Powders</th>
<th>Energy density (J/mm$^3$)</th>
<th>Yield stress (MPa)</th>
<th>UTS (MPa)</th>
<th>Hardness (HRC)</th>
<th>Elongation (%)</th>
<th>Density (g/cm$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized D$_{50}$ = 13µm</td>
<td>64</td>
<td>1116</td>
<td>1358</td>
<td>45</td>
<td>5.1</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>1200</td>
<td>1368</td>
<td>42</td>
<td>2.6</td>
<td>7.7</td>
</tr>
<tr>
<td>Water-atomized D$_{50}$ = 43µm</td>
<td>64</td>
<td>365</td>
<td>510</td>
<td>18</td>
<td>1</td>
<td>7.3</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>500</td>
<td>990</td>
<td>24</td>
<td>3.3</td>
<td>7.5</td>
</tr>
<tr>
<td>Wrought</td>
<td>-</td>
<td>1170</td>
<td>1310</td>
<td>39</td>
<td>5</td>
<td>7.9</td>
</tr>
</tbody>
</table>

The relative density of manufactured part after heat treatment for gas-atomized powder at both low and high energy densities were 97%. Comparing these data with relative density of parts before heat treatment (94% for energy density of 64 J/mm$^3$ and 95% for energy density of 104 J/mm$^3$) [18] showed slight increase in density by elimination of porosities [22]. The relative density of water-atomized powder at energy density of 104 J/mm$^3$ was approximately the same as 95%, before and after the heat treatment. The YS and UTS were lower than expected for water-atomized powder at energy density of 64 J/mm$^3$, due to porosities and unmelted particles, as shown in Fig. 5b. Elongation values of water-atomized powder were higher than gas-atomized powder at energy density of 104 J/mm$^3$, consistent with the presence of reversed austenite in parts printed at higher energy density. To reduce the retained austenite in water-atomized powder, the samples were solutionized at a higher temperature of 1315 °C in H2 atmosphere for 45 min followed by 1h of aging at 482 °C. The microstructure of heat treated gas-atomized powder are fully martensitic as shown in
Figure A.8a and b. Martensite lath were found to be coarser at energy density of 104 J/mm$^3$ due to higher laser energy and slower cooling rate. Similarly, microstructure of water-atomized powder after heat treatment are shown in Figure A.8c and d for energy density of 64 and 104 J/mm$^3$, respectively. Fully martensitic microstructure was observed at 64 and 104 J/mm$^3$ energy densities with finer lath martensite at 104 J/mm$^3$.

![Micrographs](image)

Figure A.8. The optical micrographs of heat treated parts (solutionized at 1315 °C and aged at 482 °C) perpendicular to the build direction: (1) gas-atomized at 64 J/mm$^3$, (b) G at 104 J/mm$^3$, (c) W at 64 J/mm$^3$ and (d) W at 104 J/mm$^3$.

Table A.4 shows the YS, UTS, hardness, elongation and density values for two different laser energy densities after being solutionized at 1315 °C and aged at 482 °C. Hardness for water atomized powder at energy density of 104 J/mm$^3$ is 40 HRC, which is comparable
to hardness values for gas-atomized and wrought alloy, reported 43 and 39 HRC, respectively. The YS and UTS values for water-atomized powder at low energy density are relatively lower, because of low relative density (92% due to presence of porosity and unmelted or partially melted particles) that would cause weak metallurgical bonds. The elongation for water-atomized powder at 104 J/mm$^3$ shows considerably higher magnitude (5.5%) compare to gas-atomized (2%) and slightly higher than wrought alloy (5%). The YS and UTS values for gas atomized at both energy densities didn’t show a considerable change with solutionizing at the temperature of 1315 °C and the density values remained the same (7.7 g/cm$^3$). The UTS values for water-atomized powder (1261 MPa) at 104 J/mm$^3$ is nearly the same as gas-atomized powder and wrought alloy, 1300 and 1310 MPa, respectively.

**Table A.4.** Variation in mechanical properties at energy density of 64 and 104 J/mm$^3$ - solutionized at 1315 °C and aged at 482 °C.

<table>
<thead>
<tr>
<th>Powders</th>
<th>Energy density (J/mm$^3$)</th>
<th>Yield stress (MPa)</th>
<th>UTS (MPa)</th>
<th>Hardness (HRC)</th>
<th>Elongation (%)</th>
<th>Density (g/cm$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-atomized D$_{50}$ = 13µm</td>
<td>64</td>
<td>1186</td>
<td>1308</td>
<td>45</td>
<td>2.6</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>1255</td>
<td>1300</td>
<td>43</td>
<td>2</td>
<td>7.7</td>
</tr>
<tr>
<td>Water-atomized D$_{50}$ = 43µm</td>
<td>64</td>
<td>650</td>
<td>780</td>
<td>24</td>
<td>0.7</td>
<td>7.3</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>1000</td>
<td>1261</td>
<td>40</td>
<td>5.5</td>
<td>7.5</td>
</tr>
<tr>
<td>Wrought</td>
<td>-</td>
<td>1170</td>
<td>1310</td>
<td>39</td>
<td>5</td>
<td>7.9</td>
</tr>
</tbody>
</table>

Comparing the relative densities of parts for water-atomized powder at two different heat treatments showed that there isn’t a considerable change in the densities due to higher
temperature heat treatment, however, at high energy density (104 J/mm$^3$), a considerable increase (500 MPa in YS and 271 MPa in UTS) happened after solutionizing at 1315 °C comparing to solutionizing at lower temperature (1051 °C). Furthermore, the water atomized powder at energy density of 64 J/ mm$^3$ showed an increase of 285 MPa in YS and 270 MPa in UTS. Therefore, this is likely due to finer martensitic microstructure achieved in water atomized samples after solutionizing at 1315 °C. Furthermore, carbides and other second phase particles were dissolved and then reprecipitated homogenously during aging with furnace cooling. This could also lead to higher elongation in water-atomized powder. In order to understand phase transformation during solutionizing at 1315 °C followed by aging at 482 °C, Thermo-Calc simulation was used. High cooling rate in SLM process may result in micro segregation features, therefore, it is reasonable to assume no diffusion occurs in solid state. In this case, we can use Scheil model for solidification [37]. Figure A.9 shows the mole fraction of solid for different phase with temperature through cooling from liquid phase to solid phase for water atomized and gas atomized powders. The simulation for gas-atomized powder (Figure A.9a) shows that the first phase which forms from liquid is ferrite- BBC and the second phase which forms during cooling is austenite-FCC. The phases which form at the end of solidification process are ferrite-BCC + austenite-FCC (FCC#1) + Cu-rich (FCC#2) + Laves Phase + MnS phases. The simulation for water-atomized powder (Figure A.9b) shows the first phase is ferrite-BCC followed by ferrite-BCC + austenite-FCC.

At the last stage of solidification process, the simulation shows ferrite-BCC + austenite-FCC (FCC#1) + Cu-rich (FCC#2) + M23C6 (a Nb-rich carbide) phases. The presence of
M23C6 is attributed to the higher carbon content of water-atomized powder (0.208 wt.%) compare to gas atomized (0.03 wt.%) which doesn't show any carbide formation in Thermo-Calc simulation (Figure A.9a). With increasing temperature above 1075 °C, the M23C6 phase gets dissolved in the austenitic matrix. During aging, carbide phase in martensitic stainless steel can act as nuclei for precipitation of M23C6 and hence accelerate the coarsening and spheroidization of M23C6 due to Ostwald ripening [38]. Agglomeration of carbide would increase the mobility of boundaries, because the large particles are no longer effective like small particles to pin the boundaries [39]. Solutionizing of water-atomized powder at the temperature of 1315 °C completely dissolve the M23C6 and lead to local enrichment of the carbide-forming elements such as Cr and Nb in martensitic matrix. The Thermo-Calc analysis shows that after quenching and heat treatment at 482 °C, the M23C6 precipitate in martensitic matrix and cooperate with Cu-rich precipitation in pinning the grain boundaries. This cooperation could result to increase the strength of water-atomized samples which was solutionized at the high temperature of 1315 °C compare to low temperature solutionizing at 1051 °C.

Microstructure of gas-atomized and water-atomized powder printed at energy density of 104 J/mm³, solutionized at 1315 °C and aged at 482 °C were studied using TEM. Figure A.10a and c shows bright field (BF) TEM micrograph, high angle annular dark field (HAADF) scanning transmission electron microscope (STEM) micrograph and energy dispersive spectroscopy (EDS) mapping corresponding to gas-atomized powder. In Figure A.10a and b, second phase precipitates with different size varying from approximately 30
nm to 100 nm and larger were observed. These fine precipitates could form at the initial stage of aging with coherent structure and no obvious strain contrast. These coherent bcc clusters would then nucleate and grow in the supersaturated bcc matrix and lose coherency after exceeding a certain critical size and become incoherent precipitates with fcc structure [40]. The particle shown in Figure A.10b are found to be Cu-enriched according to EDS map shown in Figure A.10c. It is clear that they pose a rather small volume fraction in the microstructure and are not uniformly distributed in the matrix. The large precipitates would preferably form at the grain boundaries as shown in Figure A.10a and b.
Figure A.0.9. Scheil nonequilibrium solidification of (a) gas-atomized and (b) water-atomized powder simulation show solid phase during solidification.
Figure A.0.10. TEM micrographs from gas-atomized 17-4 PH powder after SLM at energy density of 104 J/mm$^3$ – followed by solutionizing at 1315 °C and aging at 482 °C: (a) TEM BF micrograph, (b) HAADF STEM micrograph showing coarse and fine precipitates, (c) EDS mapping of Cu-enriched precipitates.

**Figure A.11a and c** shows microstructure and EDS map corresponding to water-atomized powder after heat treatment. Narrow distribution of size and number density of second phase precipitates were present in the microstructure as shown in Figure A.11a and b. These precipitates are in the uniform size ranging from 20 to 35 nm. There are nanoporosity in the microstructure and some of them could serve as nucleation sites for formation of larger Cu-enriched precipitates ($> 100$ nm) as shown in Fig. 11c. Absence of
large precipitates could lead to higher ductility in water-atomized powder compared to gas-atomized powder, but detailed investigation deems necessary.

Figure A.0.11. TEM micrographs from water-atomized 17-4 PH powder after SLM at energy density of 104 J/mm$^3$ – followed by solutionizing at 1315 °C and aging at 482 °C: (a) bright field micrograph, (b) HAADF STEM micrograph showing fine precipitates, (c) EDS mapping of Cu-enriched precipitates.
4. CONCLUSIONS

The effect of two different atomizing media (nitrogen atomized versus water atomized) and two different SLM post processing (solutionizing at 1050 °C and aging at 482 °C versus solutionizing at 1315 °C followed by aging at 482 °C) on mechanical properties and microstructure of 17-4 PH SS, were investigated. Following results are concluded:

- Both as-printed water-atomized and gas-atomized samples of additively manufactured 17-4 PH SS consisted of mixture of dual austenitic phase and lath martensitic phase.
- Gas-atomized powder, after low temperature solutionizing (1051 °C) and aging (482 °C), revealed single martensitic phase. However, water-atomized powder, at energy density of 104 J/mm$^3$, showed dual martensite and reversed austenite phase due to local carbon enrichment and higher diffusion rate and segregation of Cu and Ni into the matrix.
- After high temperature solutionizing (1315 °C) and aging (482 °C), both gas-atomized and water-atomized components exhibited fully martensitic structure (regardless of energy density values).
- Mechanical properties of water-atomized powder printed at energy density of 104 J/mm$^3$ and post processed at 1315 °C and aged at 482 °C was significantly improved and found to be comparable to UTS and YS values of gas-atomized and wrought alloy.
- Thermo-Calc simulation showed the dissolution of M23C6 at 1315 °C for water-atomized powder with higher carbon content. After aging, M23C6 precipitate along
with fine Cu-enriched precipitate could result in pining the grain boundaries and improving the mechanical properties and ductility.

- Microstructure of gas-atomized powder after heat treatment at 1315 °C and aged at 482 °C revealed small (∼30–40 nm) and large (> 100 nm) Cu-enriched precipitates whereas water-atomized powder revealed presence of fine (∼30 nm) Cu-enriched precipitates uniformly distributed in BCC martensitic structure.

A. 5 ACKNOWLEDGEMENTS

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A. 6 REFERENCES


APPENDIX B
HEAT TREATMENT STUDIES ON 17-4 PH STAINLESS STEEL PARTS
FABRICATED BY LASER-POWDER BED FUSION

B.1 INTRODUCTION

Laser-powder bed fusion (L-PBF), is an additive manufacturing (AM) process that uses a focused laser beam as an energy source to melt fine layers of powders to yield a solid three-dimensional part [1 - 5]. The L-PBF process has received attention to produce complex three-dimensional parts for functional applications such as tooling for injection molding, customized implants for medical industries, and lightweight components for aerospace industries [1 - 5]. Many independent investigations have been carried out in the past decade on different materials and indicated that defect-free L-PBF parts can be produced if appropriate conditions and materials are used [6 - 8]. The porosity and microstructures in the L-PBF manufactured parts have a direct effect on the resulting mechanical properties. For instance, micro-flaws and grain dislocations would serve as stress raisers for crack initiation and compromise the mechanical performance of the L-PBF parts [1 - 12]. Most of the studies indicated that L-PBF as-built parts fabricated from powders of different powder shape and size produce different porosities and microstructures under various processing conditions [1 - 12]. Studies have also shown that in the L-PBF process, the material is subjected to repeated cycles of rapid heating, melting, solidification and cooling.
Due to this phenomenon, the microstructures and mechanical properties of the as-built samples are different for various processing conditions and powder characteristics such as powder shape and size. Heat-treatment is routinely used to improve the microstructure and enhance mechanical properties of metal components [2, 12]. However, there is limited fundamental understanding of the mechanical properties and microstructures of the 17-4 PH stainless steel heat-treated L-PBF parts when properties of starting powders such as powder shape and size are varied [2, 12].

The present study was carried out to understand the effect of the heat-treatment on the mechanical properties and microstructures of the L-PBF parts fabricated from 17-4 PH stainless steels powders varied in shape (gas- and water-atomized), size-distribution at various L-PBF processing conditions. Two different post processing regimes were used in this study; (1) **Heat treatment-1**: specimens were solutionized at 1051 °C for 1h in H₂ atmosphere and then aged at 482 °C in N₂ atmosphere for 1h (H900) and (2) **Heat treatment-2**: specimens were solutionized at 1315 °C for 45 min in H₂ atmosphere and then aged at 482 °C for 1 h in N₂ atmosphere. The work presented in this study is an expansion of the work in Appendix A and will be the subject of future work to scale it into a refereed journal paper.
B.2 RESULTS

**Table B.1** The variation in density (g/cm$^3$) of 17-4 PH stainless steel L-PBF parts due to heat treatment at energy density 64 J/mm$^3$

<table>
<thead>
<tr>
<th>Condition</th>
<th>As printed</th>
<th>Heat treatment-1</th>
<th>Heat treatment-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy density (J/mm$^3$)</td>
<td>64</td>
<td>64</td>
<td>64</td>
</tr>
<tr>
<td>Gas atomized D$_{50} = 13$ µm</td>
<td>7.6 ± 0.05</td>
<td>7.6 ± 0.05</td>
<td>7.8 ± 0.05</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 24$ µm</td>
<td>6.9 ± 0.05</td>
<td>6.9 ± 0.05</td>
<td>7.5 ± 0.05</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 17$ µm</td>
<td>7.6 ± 0.05</td>
<td>7.6 ± 0.05</td>
<td>7.7 ± 0.1</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 43$ µm</td>
<td>7.0 ± 0.05</td>
<td>7 ± 0.05</td>
<td>7.6 ± 0.05</td>
</tr>
</tbody>
</table>

**Table B.2** The variation in density (g/cm$^3$) of 17-4 PH stainless steel L-PBF parts due to heat treatment at energy density 104 J/mm$^3$

<table>
<thead>
<tr>
<th>Condition</th>
<th>As printed</th>
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<th>Heat treatment-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy density (J/mm$^3$)</td>
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<td>104</td>
<td>104</td>
</tr>
<tr>
<td>Gas atomized D$_{50} = 13$ µm</td>
<td>7.7 ± 0.05</td>
<td>7.7 ± 0.05</td>
<td>7.8 ± 0.05</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 24$ µm</td>
<td>7.6 ± 0.05</td>
<td>7.6 ± 0.05</td>
<td>7.6 ± 0.05</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 17$ µm</td>
<td>7.7 ± 0.05</td>
<td>7.7 ± 0.05</td>
<td>7.7 ± 0.05</td>
</tr>
<tr>
<td>Water atomized D$_{50} = 43$ µm</td>
<td>7.5 ± 0.05</td>
<td>7.5 ± 0.05</td>
<td>7.6 ± 0.05</td>
</tr>
</tbody>
</table>
Figure B.1. The variation in the ultimate tensile strength of 17-4 PH stainless steel gas-atomized D$_{50}$ = 13 µm L-PBF parts due to heat treatment at various energy densities.
Figure B.0.12. The variation in the ultimate tensile strength of 17-4 PH stainless steel water-atomized $D_{50} = 43 \, \mu m$ L-PBF parts due to heat treatment at various energy densities.

<table>
<thead>
<tr>
<th>Energy density ($J/mm^3$)</th>
<th>Gas-atomized $D_{50} = 13 , \mu m$</th>
<th>Water-atomized $D_{50} = 24 , \mu m$</th>
<th>Water-atomized $D_{50} = 17 , \mu m$</th>
<th>Water-atomized $D_{50} = 43 , \mu m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>64</td>
<td>![Image]</td>
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<td>80</td>
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<tr>
<td>84</td>
<td>![Image]</td>
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<td>![Image]</td>
<td>![Image]</td>
</tr>
<tr>
<td>104</td>
<td>![Image]</td>
<td>![Image]</td>
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<td>![Image]</td>
</tr>
</tbody>
</table>

Figure B.0.13. The microstructures of 17-4 PH stainless steel water-atomized $D_{50} = 43 \, \mu m$ L-PBF parts due to heat treatment-1 at various energy densities.
B.3 REFERENCES


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APPENDIX C

EFFECTS OF INITIAL POWDER ATTRIBUTES AND PROCESS CONDITIONS ON
THE RECYCLABILITY OF 17-4 PH STAINLESS STEEL POWDER IN
LASER-POWDER BED FUSION

C.1. INTRODUCTION

The desire to improve the sustainability and efficiency of the laser-powder bed fusion (L-PBF) process drives the requirement for recycling of starting powders that are recycled during the fabrication process. An initial study was carried out to understand the effect of L-PBF processing conditions and starting powder characteristics on the recyclability of the 17-4 PH stainless steel powders. To this effect, following a single L-PBF run, powder was collected in the vicinity of the part, potentially in the heat-affected zone after laser energy input. These powders were referred to as recycled powders or recycled powders, in contrast to the term, "initial powders" for the starting powders prior to use in an L-PBF run. The powders were characterized by SEM and XRD to detect any changes in morphological or chemistry.

Under all energy densities used in this study, no large changes in shape (qualitatively and quantitively) was observed in the 17-4 PH stainless steel gas and water-atomized powders after a single use. However, at a high energy density of 84 and 104 J/mm³, the percentage of large spherical agglomerates qualitatively increased while fine spherical particles
decreased in the used 17-4 PH stainless steel gas and water-atomized powders. The likely formation of sintered agglomerates in the recycled 17-4 PH stainless steel gas- and water-atomized powder will affect the powder flowability and density during the L-PBF process. XRD analysis also indicated minor changes possibly as a result of oxidation or volatilization of alloy constituents. These results pinpoint the need for detailed follow up to track the changes in microstructure and properties in parts printed with recycled powders following multiple uses. The work presented in this study will be the subject of future work to scale it into a refereed journal paper.

C.2 RESULTS

Figure C.1. SEM images of starting 17-4 PH stainless steel gas-and water-atomized powders
Figure C.2. SEM images of recycled 17-4 PH stainless steel gas-atomized powders at energy density 64 J/mm$^3$
Figure C.3. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50}=17 \mu m$) powder at energy density 64 J/mm$^3$.

Figure C.4. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50}=24 \mu m$) powder at energy density 64 J/mm$^3$. 
Figure C.5. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50} = 43\ \mu m$) powder at energy density 64 J/mm$^3$. 
Figure C.6. SEM images of recycled 17-4 PH stainless steel gas-atomized ($D_{50}= 13 \, \mu m$) powder at energy density 80 J/mm³
Figure C.7. SEM images of recycled 17-4 PH stainless steel water-atomized (D$_{50}$ = 17 µm) powder at energy density 80 J/mm$^3$
Figure C.8. SEM images of recycled 17-4 PH stainless steel water-atomized \((D_{50} = 24 \mu m)\) powder at energy density 80 J/mm³
Figure C.9. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50}=43$ µm) powder at energy density 80 J/mm$^3$. 
Figure C.10. SEM images of recycled 17-4 PH stainless steel gas-atomized (D$_{50}$= 13 µm) powder at energy density 84 J/mm$^3$
Figure C.11. SEM images of recycled 17-4 PH stainless steel water-atomized (D_{50}= 17 µm) powder at energy density 84 J/mm^3
Figure C.12. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50} = 24 \mu m$) powder at energy density $84 J/mm^3$. 
Figure C.13. SEM images of recycled 17-4 PH stainless steel water-atomized ($D_{50}=43 \ \mu m$) powder at energy density $84 \ J/mm^3$
Figure C.0.14. SEM images of recycled 17-4 PH stainless steel gas-atomized ($D_{50}=13$ µm) powder at energy density 104 J/mm$^3$.

Figure C.0.15. XRD pattern of the recycled 17-4 PH stainless steel gas-and water-atomized powders (left) and starting powders (right).
Figure C.0.16. XRD pattern of the recycled 17-4 PH stainless steel gas-and water-atomized powders (left) and corresponding L-PBF parts (right) processed at an energy density of 64 J/mm³.
APPENDIX D

UOF 3D PRINTING BUSINESS INCUBATOR: INTEGRATING PRODUCT DESIGN, PROCESSING AND MARKET ANALYSIS FOR MANUFACTURING WITH METAL POWDERS

D. 1 INTRODUCTION

Powder metallurgy and metal injection molding are proven net-shaping processes for the high volume fabrication of precision parts for a broad range of transportation, medical, electronics, industrial and consumer applications. However, the growth of applications and market size in these technologies depends to a good extent on design engineers being familiar with these prescribing parts based on processing metal powders. The advent of metal 3D printing provides an opportunity for design engineers to get acquainted with powder metallurgy and metal injection molding technologies at an early stage of their education and training. A new academic program at the University of Louisville has been addressing this very bottleneck during the last year.

The lead time to translate a new product design into manufacturing is time-consuming and expensive [1]–[10]. In each design cycle, the production moves through various stages involving physical prototype is created to test and validate the design. Implementing
additive manufacturing or 3D printing in the design cycle of a new product can disrupt the product design and supply chain of various aerospace, defense, automotive and healthcare industries [1]–[10]. However, understanding material-process-properties relationship is one current barrier to broader adoption of metal 3D printing to complement and integrate with the traditional process in industries [1]-[3].

The research performed at the Materials Innovation Guild, University of Louisville (3dmaterials.us) focuses on obtaining a fundamental understanding of metal 3D printing processing conditions and ensuing materials properties and microstructure through the integration of carefully designed experiments to enable optimization of the materials-design-process relationship as shown in Figure D.1. With an understanding of the materials-design-process relationships in metal 3D printing could emerge an opportunity that allows new design innovations and products in aerospace, defense, automotive and healthcare industries.

In this regard, University of Louisville developed a pilot project, UofL 3D Printing Business Incubator (uofl3dpbiz.com) that connected capstone student design teams with various industries to understand the techno-economic points through case studies.
A total of 20 design teams compromised of 100 engineering students shown in Figure D.2 were trained and educated in techno-economic aspects of 3D printing. During the 9 months pilot project period, students worked with academia and industry partners. Industry partners defined a real-world capstone project and whereas, academia provided them necessary research support. This support helped capstone students to redesign parts for 3D printing to produce and test them technical and business aspects (surface, geometric, mechanical and material requirements, cost and economic batch analysis).

In this article, one interesting case study is presented that shows the potential of 3D printing in automotive industries. In addition to the designing and manufacturing the prototypes, the business case is also developed by students to demonstrate how 3D printing can complement the traditional processes like powder metallurgy by saving time and costs.
Figure D.2. Mechanical engineering capstone students (Spring 2017) were involved in the UofL 3D Printing Business Incubator

D. 2 AUTOMOTIVE ENGINE GEARs

The automotive industry, among other manufacturing industries, is constantly driving towards a shorter lead time-to-market for its new products. New cars and trucks need to be designed, tested and released for production in a much shorter period [4], [5], [7], [8], [10]. The shorter time-to-market evolution has also influence on the gearbox manufacturers who must design and test their products in a much shorter period. This can also speed up the process by placing a high-volume production order of gears using powder metallurgy for various automotive industries. The critical aspect of this process is the lead time to create new gear prototypes, which is currently around 10 weeks. This is mostly caused by the need to develop custom hobbing and grinding tools, a process which is time-consuming and expensive [4], [8], [10]. This long lead time limits the number of designs which can be tested and makes it difficult to achieve the best tribological and acoustic performance. For this reason, gearbox manufacturers need a faster production technology such as 3D printing to create precision gears for prototyping purposes. The 3D printing process has the potential to significantly accelerate the process of prototyping new product designs and
validating them under real conditions. In this regard, a student startup team at the University of Louisville, Bluegrass Engineering Solutions (Figure D.3), worked with an automobile engine manufacturer to decrease the lead time to produce the gear prototypes using 3D printing process.

Figure D.3. The Bluegrass Engineering Solutions student startup team that worked on the automobile engine gear: Austin Anderson, Brad Gootee, Michael Kjelby, Devon Warman, Gunnar Wagoner

The primary challenge for design engineers for choosing a prototype production method for a specific product like automotive engine gear depends on batch size and cost of the production. The economic batch size and cost analysis of L-PBF process were performed using CES Edupack as shown in Figure D.4. It can be concluded that L-PBF process is the cost-effective and efficient low volume production process for the prototype fabrication.
The gear geometry was designed using commercially available CAD software (Figure D.5). The initial strength verification was performed using FEM techniques (ANSYS software) (Figure D.5). Through various testing and studies, it has been found that laser-powder bed fusion process can manufacture functional gear prototypes (Figure D.6) and can reduce the design and testing lead time for new products. The gear is redesigned as per the design principles of L-PBF process and fabricated in EOS M 290 machine using 17-4 PH stainless steel powder. 17-4 PH stainless steel is chosen because of its high strength and corrosion resistance which is widely used in automotive, medical and aerospace industries.
The geometric analysis on the 3D printed gear and powder metallurgy gear was performed and results are shown in Table D.1. The geometric dimensions of the 3D printed gear were comparable to the gear manufactured by powder metallurgy. However, to test the 3D printed gear experimentally requires additional surface finishing by machining. For comparison, Boquet et al.[4] compared machining time and surface finish of the spur gears manufactured using wire EDM, milling, L-PBF process. Boquet et al. concluded that L-PBF process is a potential technology to fabricate prototype gears, but additional finishing is required for further testing of the gears.
The time and cost analysis are performed using the cost models when prototype gears were produced by L-PBF process [1] -[10]. It is estimated that using 3D printing in the design phase could reduce the prototyping costs by 50% and design cycle time by 90% compared to the current methods used by the engine manufacturer. This study could provide the pathway for decreasing the turn around time in the automobile industry for sending the optimized designs to powder metallurgy companies for high volume production. Thus, a case of how 3D printing can help the powder metallurgy companies and automobile manufacturers by decreasing the lead time and cost of manufacturing was demonstrated in this article. Future work includes working with powder metallurgy alloys specific to powder metallurgy using binder jetting as well as incorporating functional features such as weight reduction and noise reduction. Further testing of the gear will be performed using test setup at the automotive engine manufacturer’s facility to validate the 3D printed part.

Figure D.6. Automotive engine gear re-designed and produced using the L-PBF process
Table D.1. Comparison of dimensions of 3D printed, powder metallurgy gears with CAD model

<table>
<thead>
<tr>
<th>Dimension</th>
<th>3D printed part (mm)</th>
<th>Powder metallurgy part (mm)</th>
<th>CAD model (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tooth thickness</td>
<td>9.75</td>
<td>9.8</td>
<td>9.8</td>
</tr>
<tr>
<td>Overall diameter</td>
<td>64.5</td>
<td>64.5</td>
<td>64.6</td>
</tr>
<tr>
<td>Base diameter</td>
<td>49.5</td>
<td>49.5</td>
<td>49.5</td>
</tr>
<tr>
<td>Overall gear</td>
<td>16.5</td>
<td>16.6</td>
<td>16.8</td>
</tr>
<tr>
<td>thickness</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shaft diameter</td>
<td>15.8</td>
<td>16</td>
<td>16</td>
</tr>
</tbody>
</table>

D. 3 CUSTOMIZED FUEL INJECTOR

In the automobile industry, fuel injector designs developed for traditional manufacturing have been constrained by processes that require the creation of separate components that are assembled to create the product. 3-D printing has the potential to create complex geometries at no additional cost. This provided freedom to a team of 5 students at the University of Louisville shown in Figure D.7 to create a consolidated fuel injector (design provided by automobile engine manufacturer) to decrease products weight and manufacturing steps. Furthermore, students have demonstrated efficient design for fuel injector as shown in Figure D.8 and Table D.2 with complex internal cooling channels. However, further analysis needs to perform to evaluate and validate this fuel injector for production.
Figure D.7. Students worked on automobile fuel injector case study: Ian Yockey, Kyle Swenson, Nick Bowling, Wil Johnson, Zach Winfield

Table D.2. Performance comparison of 3D printed fuel injector with and without cooling channels

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuel injector tip failure</td>
<td>662 °F</td>
</tr>
<tr>
<td>temperature</td>
<td></td>
</tr>
<tr>
<td>Maximum heat input rate with</td>
<td>35.3 W</td>
</tr>
<tr>
<td>cooling channels</td>
<td></td>
</tr>
<tr>
<td>Maximum heat input rate</td>
<td>40.1 W</td>
</tr>
<tr>
<td>without cooling channels</td>
<td></td>
</tr>
<tr>
<td>Percent increase in</td>
<td>14%</td>
</tr>
<tr>
<td>performance</td>
<td></td>
</tr>
</tbody>
</table>
D.4. TOOLING FOR INJECTION MOLDING

In plastic injection molding industry, the production of injection molding tools is one of the most expensive aspects of the production processes. It is expensive, time consuming and very technically demanding to use traditional processes to manufacture the tools for injection molding. A company which does plastic injection molding to produce various
commercial products indicated that it takes them 40 to 50 weeks and spends several thousands of dollars to manufacture a production grade tooling.

A team of 8 students shown in Figure D.9. identified this niche and worked with injection molding company to demonstrate the potential of 3D printing in manufacturing the tooling by decreasing the cost and time. Students redesigned the existing tool and integrated cooling channels in them to improve the performance of the tool when compared to traditional made tools shown in Figure D.10. The performance of the 3D printed tool when compared to traditional tools is shown in Table D.3. In addition, students demonstrated the business case and potential of 3D printing by reducing the tool cycle time by 80% and improving the performance of the tools 20%.

Figure D.9. Students worked on tooling case study: Nathan Westurn, Shri Patel, Aaron Hartpence, David Campbell, Robin Winebrenner, Chris Barnett, Bradley Ford, Joe Tobbe, Travis Wickizer
Figure D.10. Design and simulation of conformal cooling channels of a 3D printed mold

Table D.3. Performance evaluation of a 3D printed mold

<table>
<thead>
<tr>
<th>Type of cooling channel</th>
<th>Cycle time (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mold with no cooling channel</td>
<td>14.4</td>
</tr>
<tr>
<td>Mold with non-conformal cooling channel</td>
<td>13.5</td>
</tr>
<tr>
<td>Mold with conformal cooling channel</td>
<td>12.4</td>
</tr>
</tbody>
</table>
D.5. SUMMARY AND FUTURE OUTLOOK

A total of 20 design teams compromised of 100 engineering students were trained and educated in techno-economic aspects of 3D printing as a part of 3D Printing Business Incubator program at the University of Louisville. During the 9 months pilot project period, students worked with academia and industry partners on various case studies. These case studies demonstrated the feasibility of using 3D printing in multiple industries that support traditional manufacturing technologies by decreasing lead time and cost. Further, the academic program strongly suggests a pathway for powder metallurgy and metal injection molding industries to benefit from new products as well as fresh engineering talent. Efforts are underway to scale the program to achieve a regional economic impact in the form of an industry-government-academia collaboration, MADE IN KY: MAnufacturing and DEsign Innovation Network of Kentucky.
D.6 REFERENCES


CURRICULUM VITAE

Harish Irrinki
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Education

University of Louisville
Ph.D. Mechanical Engineering; GPA 4.0/4.0
Structure-process-property relationship of 17-4 PH stainless steel processed by laser-powder bed fusion
Advisor: Dr. Sundar Atre

University of Louisville
M.S. Mechanical Engineering; GPA 4.0/4.0
April 2016

Jawaharlal Nehru Technological University
B.Tech. Mechanical Engineering; GPA 3.8/4.0
Hyderabad, INDIA
June 2014

Selected Awards and Recognitions

Hsing Chuang Award for Excellence in Graduate Study -2018
NSF Student Travel Award, AMPM Conference- 2018
MIMA Student Travel Grant, MIM Conference-2017
NSF I-CORP Award -2017
Grosscurth Fellowship, University of Louisville -2016-18
AMPM Best Poster Award - 2016
CPMT Axel Madison Award - 2016
ASME Best Presentation Award -2016

Research Experience

Tesla
Fremont, CA

• Led the project to identify the additive manufacturing (AM) opportunities in automobile industry using a scientific model

• Developed a techno-economic model to demonstrate the techno-economic merits in resulted in adapting AM technology for fabricating the BIW of Tesla cars
Grosscurth Fellow and Graduate Research Assistant
June 2016 - present

- Experience in powder-based additive manufacturing design, process, materials and applications development

- Developed a fundamental understanding of the role of powder attributes, L-PBF process and post-process conditions on the densification, mechanical and electrochemical properties and microstructures of L-PBF parts

- Evaluated different polymer and metal 3D printers and responsible for the purchase of $300K metal 3D printer, $10K polymer 3D printers

- Developed the applications based on the understanding of relationship between: design to process, process to microstructure; microstructure to properties

- Designed and evaluated the performance of various 3D printed functional parts for various OEM

- Applications and process development for various OEM’s (Cummins, Walmart, Amaray, GE Concept Laser, HTS, Mirus, NAH, Sandvik Osprey) based on research, technical and business merits

- Communicated technical results with collaborators and business leaders at all levels through formal technical presentations and informal discussions

Materials: 17-4 PH, 420 and duplex stainless steel, bronze, niobium, tungsten, molybdenum-rhenium

3D Printing: Concept Laser Mlab, EOS M 290, 3DSYSTEM ProX 320, FDM

Design Skills: SolidWorks, Inspire, Catia 3D Experience, Materialise, Limitstate, MatLab, Ansys, C

Characterization: SEM, XRD, EDAX, hardness and tensile testing

Leadership Experience

Graduate Mentor-3D Printing Business Incubator
Aug. 2016 – Apr. 2017

- Planned and executed 4 mechanical engineering undergraduate projects that support technology-based business opportunities on metal additive manufacturing platforms
University of Louisville and Oregon State University  Sep. 2015 – Apr. 2017

Graduate Teaching Assistant

- Taught and helped students to design and fabrication techniques to develop various applications
- Topics mainly focused on CAD, ANSYS simulations, manufacturing and prototype development

Publications


Conference Talks and Proceedings


