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Tailoring residual stress profile of Selective Laser Melted parts by Laser Shock Peening

Nikola Kalentics, Eric Boillat, Patrice Peyre, Snežana Ćirić-Kostić, Nebojša Bogojević, Roland E. Logé

Abstract

The paper describes a new approach in controlling and tailoring residual stress profile of parts made by Selective Laser Melting (SLM). SLM parts are well known for the high tensile stresses in the as-built state in the surface or subsurface region. These stresses have a detrimental effect on the mechanical properties and especially on the fatigue life. Laser Shock Peening (LSP) as a surface treatment method was applied on SLM parts and residual stress measurements with the hole-drilling method were performed. Two different grades of stainless steel were used: a martensitic 15-5 precipitation hardenable PH1 and an austenitic 316L. Different LSP parameters were used, varying laser energy, shot overlap, laser spot size and treatments with and without an ablative medium. For both materials the as-built (AB) residual stress state was changed to a more beneficial compressive state. The value and the depth of the compressive stress was analyzed and showed a clear dependence on the LSP processing parameters. Application of LSP on SLM parts showed promising results, and a novel method that would combine these two processes is proposed. The use of LSP during the building phase of SLM as a “3D LSP” method would possibly give the advantage of further increasing the depth and volume of compressive residual stresses, and selectively treating key areas of the part, thereby further increasing fatigue life.

Keywords:
Selective laser melting
Laser shock peening
3D Laser shock peening
Residual stress profile
15-5 PH stainless steel
316L stainless steel

1. Introduction

Selective laser melting (SLM) is a part of a large family of Additive Manufacturing (AM) processes. Over the last decades more than thirty different types of Additive Manufacturing processes have been developed [1,2], with SLM being one of the most researched over the past years.

However, although the mechanical properties have become close to those of bulk materials [3–14], SLM has some inherent drawbacks such as warping, cracking and detrimental tensile residual stresses (TRS). A large degree of shrinkage occurs during liquid – solid transformation, thus accumulating considerable tensile residual stresses on the surface of the SLM produced components. The complex residual stresses (RS) that arise during cooling are regarded as key factors responsible for the distortion and even delamination of the final parts [9,10,15–17]. These residual stresses may even cause process failure during the building phase [18].

The last melted layer shrinks during cooling while the layer underneath, already solidified constrains it and prevents further shrinking [15,16]. Since this mechanism occurs for each layer at each step of the SLM process, large tensile residual stresses accumulate inside the manufactured component which cause significant and detrimental anisotropy of the mechanical properties of produced parts [5,7,19–21] thus limiting their application.

Different methods have arisen to reduce residual stresses. In situ heating (e.g. build plate heating; reheating of the melt pool) is commonly used [15,22]. Adapting scanning strategies can also have a strong impact on residual stresses [15,23]. As a post treatment, annealing is widely used and has demonstrated in some cases a 70 percent reduction of residual stresses [24]. Although these methods have demonstrated certain improvements of the final residual stress state, they have shown to be unable to completely remove tensile residual stresses. In the case of annealing, as a post treatment it is limited to addressing this issue only after the building process is done, and as such it cannot address the issue of stress accumulation that can cause process failure.

In the present paper, a novel strategy is proposed for tailoring residual stresses in parts manufactured by SLM by using another
2. Experimental setup

2.1. Material

Materials used in this paper are two grades of stainless steel, PH1 and 316L. PH1 with a standard denomination 15-5 PH, is a martensitic precipitation hardenable stainless steel made by the SLM machine producer EOS (EOS GmbH, Germany). PH1 has an ultimate tensile strength (UTS) of 1150 MPa [43] and provides excellent mechanical properties, especially in the precipitation hardened state. It is widely used in medical, aerospace and other engineering applications requiring high hardness and strength. PH1 samples were produced on an EOSMINT M280 additive manufacturing machine, with a closed set of parameters and a fixed scanning strategy provided by the machine producer for this powder.

316L is a widely used austenitic stainless steel and has an UTS of 760 MPa [44]. The powder that was used was DIAMALLOY 1003 obtained from Sulzer Metco, Switzerland. These samples were produced on a Concept laser machine equipped with a fiber laser operated in continuous mode at a wavelength of 1070 nm and a spot size of 90 µm. The parameters laser power, scanning speed, hatch distance and powder layer thickness were 125 W, 600 mm/s, 0.105 mm and 0.03 mm, respectively. A bi-directional scanning strategy that was parallel to the part edges was used in every layer without a change in scanning direction between layers. This was done to deliberately create large tensile residual stresses and to show the ability of the LSP process to address even such stresses and to convert them to CRS.

The chemical composition of both PH1 and 316L stainless steel is shown in Table 1.

2.2. Laser shock peening

Laser shock peening (LSP) experiments were made at the PIMM laboratory at CNRS-ENSM Paristech [45]. The laser source used is a 7.1 ns at 532 nm Nd:YAG GALA – class laser from Thales Laser company. The beam spatial energy distribution is “top-hat” and the pulse shape is near – Gaussian. LSP processing parameters are shown in Table 2. Round laser spots of 1, 2 and 5 mm diameter were used with the laser energy of 0.4, 1.6 and 10 J respectively. This ratio of the spot size and the energy kept the power density constant at 7.2 GW/cm². The pressure at the surface of the treated part was estimated at 4.7 GPa using an empirical equation

\[ P(GPa) = 1.75 \sqrt{\frac{I(GW)}{I_0(cm^2)}} \]  

Pulse frequency was 1 Hz, and the overlap of 40% was used for all spot sizes with and without a protective ablative coating. Also one sample with an overlap of 80% was treated with a 1 mm spot size without an ablative coating. In order to avoid ablation of the surface of the samples, in some cases an ablative layer was used. For this purpose a sacrificial aluminium tape of 70 µm thickness from 3MTM was placed on top of the sample. In other cases, the ablative layer was not applied, and the results in these two conditions were compared.

2.3. Residual stress determination using the hole drilling method

Residual stresses have been measured with the hole drilling method (HDM) according to the ASTM standard E837 [16,47]. It is a widely used technique for the determination of in depth residual stress profiles in parts especially after surface treatments such as LSP, USP, SP etc [47–52]. The measuring device is the RESTAN-MTS 3000 from SINT Technology (Fig. 2a). A 1.8 mm diameter hole is drilled into the piece to be analyzed. Residual stresses relax at the hole location causing strains also to change. A strain gage rosette with three grids measures these strains (Fig. 2b). Residual stresses are given by the theory of Kirsch [53], adjusted with experimental coefficients for blind hole analysis. Residual stresses were measured at a total of 36 points from the surface up to 1 mm in depth. Since the goal was to achieve more precise results of residual stresses in the near surface region, a variable depth increment was applied. In the region from the surface up to the depth of 100 µm, measurements were made every 10 µm. From 0.1 mm up to 0.5 mm in depth, measurements were made every 25 µm, and from 0.5 mm up to 1 mm every 50 µm.

3. Results and discussion

3.1. PH1 stainless steel

Fig. 3 shows some of the most important parameters of a residual stress curve. These are the maximum amount of CRS – Max CRS, depth at which maximum CRS are observed – Depth of max CRS, depth at which change from CRS to TRS – Depth of CRS, and the value of CRS in the near surface region, at the depth of 20 µm – Surface RS.

Residual stress measurements of the PH1 samples made by SLM in the as – built (AB) state show a low amount of compressive residual stresses (CRS) in the near surface region up to the depth of 183 µm (Fig. 4). Compressive residual stress in the near surface region of SLM samples is a rare observation. However, it was also observed that welded joints of martensitic stainless steels have such compressive residual stress state in the near surface zone [54–56]. The reason for their occurrence is that martensitic stainless steels experience a phase transformation during the cooling period. During this phase change, the crystal lattice expands and the accumulated stresses due to the shrinkage are relaxed and inverted.
Chemical composition of PH1 and 316L stainless steel, wt.% [43].

<table>
<thead>
<tr>
<th></th>
<th>Cr</th>
<th>Ni</th>
<th>Cu</th>
<th>Mn</th>
<th>Si</th>
<th>Mo</th>
<th>Nb</th>
<th>C</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>PH1</td>
<td>14–15.5</td>
<td>3.5–5.5</td>
<td>2.5–4.5</td>
<td>max 1</td>
<td>max 1</td>
<td>max 0.5</td>
<td>0.15–0.45</td>
<td>max. 0.07</td>
<td>Balance</td>
</tr>
<tr>
<td>316L</td>
<td>17</td>
<td>12</td>
<td>/</td>
<td>/</td>
<td>2.3</td>
<td>2.5</td>
<td>/</td>
<td>0.03</td>
<td>Balance</td>
</tr>
</tbody>
</table>

Table 2
LSP processing parameters: spot size; E – laser energy; I₀ – power density; P – estimated pressure at the surface; frequency.

<table>
<thead>
<tr>
<th>Spot size (mm)</th>
<th>E (J)</th>
<th>I₀ (GW/cm²)</th>
<th>P (GPa)</th>
<th>Frequency (Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.4</td>
<td>7.2</td>
<td>4.7</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>1.6</td>
<td>7.2</td>
<td>4.7</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>7.2</td>
<td>4.7</td>
<td>1</td>
</tr>
</tbody>
</table>

Fig. 1. Effect of tensile and compressive stresses on the crack growth propagation.

Fig. 2. a) Hole drilling device RESTAN-MTS 3000 from SINT Technology, b) sample with attached strain gage rosette for HDM.

to a small negative value (i.e. to a compressive residual stress value) [44].

The maximum amount of the observed CRS is −230 MPa at the depth of 94 μm which presents a 20% of its UTS of the material which is 1150 MPa. The value of the surface RS is −41 MPa (4% of the UTS). Beyond 183 μm, the residual stress state changes towards tensile state. Above the depth of 183 μm where the stresses are neutral, we can observe a sudden increase in TRS up to the depth of around 340 μm where the TRS have a value of 435 MPa (38% of the UTS). Beyond this point we can see a slower increase in the amount of tensile stresses throughout the whole depth of the measurement.

Other samples made by SLM were treated with Laser shock peening. Table 3, gives an overview of results of residual stress measurements done on samples in the as – built and LSP treated state and a graphical representation is given in Fig. 4.
Fig. 3. Residual stress curve with the most important parameters indicated: Max CRS – maximum amount of CRS; Depth of max CRS – depth at which maximum CRS are observed; Depth of CRS – depth at which residual stresses change from CRS to TRS; Surface RS – value of CRS in the near surface region, at the depth of 20 μm.

Table 3
Results of RS measurements: Max CRS/normalized by UTS; depth of max CRS; depth of CRS; Surface RS/normalized by UTS. Measurements are made in the as-built state (AB), or with LSP treatments of 1, 2 and 5 mm, 40 and 80% overlap, and with and without an ablative coating (C/NC).

<table>
<thead>
<tr>
<th>LSP treatment</th>
<th>Max CRS [MPa]/percentage of the UTS [%]</th>
<th>Depth of max CRS [μm]</th>
<th>Depth of CRS [μm]</th>
<th>Surface RS [MPa]/percentage of the UTS [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 mm 40% NC</td>
<td>–421/37</td>
<td>142</td>
<td>406</td>
<td>–117/10</td>
</tr>
<tr>
<td>1 mm 40% C</td>
<td>–530/46</td>
<td>115</td>
<td>403</td>
<td>–187/16</td>
</tr>
<tr>
<td>2 mm 40% NC</td>
<td>–401/35</td>
<td>136</td>
<td>495</td>
<td>–103/9</td>
</tr>
<tr>
<td>2 mm 40% C</td>
<td>–494/43</td>
<td>120</td>
<td>647</td>
<td>–165/14</td>
</tr>
<tr>
<td>5 mm 40% NC</td>
<td>–325/28</td>
<td>139</td>
<td>620</td>
<td>–69/6</td>
</tr>
<tr>
<td>5 mm 40% C</td>
<td>–412/36</td>
<td>113</td>
<td>788</td>
<td>–163/14</td>
</tr>
<tr>
<td>1 mm 80% NC</td>
<td>–798/69</td>
<td>247</td>
<td>&gt;1 mm (~266 MPa at 1 mm)</td>
<td>–169/15</td>
</tr>
<tr>
<td>AB state</td>
<td>–230/20</td>
<td>94</td>
<td>184</td>
<td>–42/4</td>
</tr>
</tbody>
</table>

3.1.1. Maximum value of CRS
3.1.1.1. LSP treatment without an ablative coating. For an LSP treatment without an ablative coating (Fig. 5), maximum CRS are –421 MPa for 1 mm spot size (37% of the UTS), –401 MPa (35%) for 2 mm and –325 MPa (28%) for 5 mm, and they are observed at similar depths (136–142 μm). These values make a significant improvement of 17%, 15% and 8% of the UTS respectively, compared to the as-built state which has CRS of –230 MPa (20%) at a depth...
of 94 µm. It can be observed that smaller spot size leads to larger maximum values of residual stresses. This effect was observed in literature [57] and it is presumed that since LSP with smaller spots is more often applied for a same surface area compared to LSP with larger spots, the effect is more pronounced. It is somehow equivalent to a higher number of shots, which is discussed in section 3.1.4.

3.1.2. LSP treatment with an ablative coating. Similar trends are observed in treatments with an ablative coating: −530 MPa for 1 mm (46% of the UTS), −494 MPa (43%) for 2 mm and −412 MPa (36%) for 5 mm. In this case, this maximum values are higher compared to the treatment without the ablative coating and they are observed closer to the surface (113–120 µm). These values present an even larger improvement compared to the as – built state of 26%, 23% and 16% of the UTS respectively which is shown in Table 4. Also an increase in CRS for LSP treatments with an ablative coating compared to those without an ablative coating can be observed. This is explained by the protective role of the ablative coating, since the plasma is created on its surface and not on the surface of the SLM sample. This prevents ablation and local melting of the sample surface and creation of TRS due to this melting. When the ablative coating is not applied, these TRS are decreasing the overall effect of the LSP process in both the max value of CRS and their depth.

3.1.3. Surface residual stresses
If we look at the values of the surface RS, it can be observed that the smaller spot sizes lead to larger CRS. The same tendency was observed in sections 3.1.1 and 3.1.2 in the case of the maximum amount and the depth of CRS. For samples treated without an ablative coating residual stresses are −117 MPa (10%) for 1 mm spot size, −103 MPa (9%) for 2 mm and −69 MPa (6%) for 5 mm. This makes an increase of 6%, 5% and 2% of the UTS respectively, compared to the as – built state which has CRS of −42 MPa (4%).

For samples treated with an ablative coating, measured surface RS are −187 (16%), −165 (14%) and −163 MPa (14%) for 1, 2 and 5 mm spot size respectively. This presents an increase compared to the AB state of 12%, 11% and 10% of the UTS. In the case of all 3 spot sizes, we can again see a significant influence of the ablative coating on the amount of CRS in the near surface region. For a 1 mm spot size we can observe an increase from −117 MPa (10%) to −187 MPa (16%), for 2 mm from −103 MPa (9%) up to −165 MPa (14%) and for the 5 mm from −69 MPa (6%) up to −163 MPa (14%) in the uncoated/coated state respectively.

3.1.4. Overlap rate
One sample was treated with a higher overlap rate of 80% and without an ablative coating. For this sample the smallest spot size of 1 mm was used, as this value might be preferential for the LSP treatment of small and complicated parts made by SLM. These

### Table 4
Percentage of increase of maximum and surface CRS compared to the as built state and normalized by UTS.

<table>
<thead>
<tr>
<th>LSP treatment</th>
<th>Max CRS compared to the AB state, and normalized by UTS; [%] of UTS</th>
<th>Surface RS compared to the AB state, and normalized by UTS; [%] of UTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 mm 40% NC</td>
<td>17 6</td>
<td>26 12</td>
</tr>
<tr>
<td>2 mm 40% NC</td>
<td>15 5</td>
<td>15 5</td>
</tr>
<tr>
<td>5 mm 40% NC</td>
<td>23 2</td>
<td>16 2</td>
</tr>
<tr>
<td>1 mm 80% NC</td>
<td>49 11</td>
<td>49 11</td>
</tr>
</tbody>
</table>

Fig. 5. Residual stress curves measured for samples in the AB and LSP treated states without an ablative coating.
measurements showed a high increase in all the important features of the residual stress profile. Maximum residual stresses were $-798 \text{ MPa} (69\%)$, observed at a depth of 247 $\mu\text{m}$, compared to $-421 \text{ MPa} (37\%)$ at 142 $\mu\text{m}$ for 1 mm 40% overlap LSP treatment done also without a coating. These values presented the largest increase compared to the AB state with 49% increase of the UTS in the amount of CRS and a 163% increase in depth of the maximum CRS. Surface RS were $-169 \text{ MPa} (15\%)$, which was an increase of 11% of the UTS in the AB state. Depth of the CRS, was above 1 mm which was the range of the measurement. At the depth of 1 mm they were still compressive with a value of $-266 \text{ MPa} (23\%)$.

With an increase by a factor of 2 in the overlap rate from 40% to 80%, we have an increase by a factor of 9 of the total number of shots on the treated surface. This large increase in the total number of shots leads to a significant increase in all measured RS parameters, i.e., maximum CRS, depth of max CRS, depth of CRS and surface RS, shown in Fig. 5.

### 3.2. 316L stainless steel

Since the PH1 SLM samples in the as built state showed unusual CRS in the near surface region, 316L SLM samples were used to show the potential of the LSP treatment on other materials and the ability to fully transform TRS to CRS. 316L was chosen because of its wide spread application, but also because it is an austenitic stainless steel that has a TRS state in the near surface region that is more common for SLM parts. The characteristic values in the AB and LSP treated states are shown in Table 5 and the curves of both samples made of 316L and PH1 with same treatments in Fig. 7.

In the AB state, we observed TRS in the whole range from the surface up to the 1 mm depth. The maximum value in the near surface region was 342 MPa at about 131 $\mu\text{m}$ which represents 45% of the UTS of the material (760 MPa).

LSP treatment with a 1 mm spot size, no coating and 40% and 80% overlap was done. For the sample treated with a 40% overlap, CRS were observed up to the depth of 416 $\mu\text{m}$. The maximum value of CRS was $266 \text{ MPa} (35\%)$ at 128 $\mu\text{m}$, and surface RS were $-103 \text{ MPa} (14\%)$. For the sample treated with an 80% overlap, CRS were observed up to the depth of 804 $\mu\text{m}$. The maximum value of CRS was $-729 \text{ MPa} (96\%)$ at 94 $\mu\text{m}$ and in the surface region $-418 \text{ MPa} (55\%)$ at a 20 $\mu\text{m}$ depth. The maximum value of CRS represents 96% of the UTS of the material which indicates material strain hardening due to the high number of LSP shots on the surface in the 80% overlap LSP condition.

Tensile residual stresses are very efficiently converted to compressive residual stresses for 316L samples, and the asymptotic profile is qualitatively similar to that of the PH1 samples.

### 4. Conclusions

In this paper, the capability of the LSP treatment to alter the residual stresses of SLM parts was demonstrated. It was shown that in the case of the martensitic PH1 stainless steel, the maximum amount, depth of the profile and amount in the near surface region of the beneficial CRS can be drastically increased. Also, for the austenitic 316L stainless steel, the highly tensile state of the AB sample was changed to a beneficial CRS state. Various LSP processing parameters were used, with and without an ablative coating. It can be concluded that:

- Laser shock peening can be used to effectively and easily change the residual stress profile. Even with a single pass, the RS changes from tensile to compressive in the case of 316L, or in the case of PH1, the values of CRS are drastically increased and observed to larger depths.
- Smaller spot size leads to larger amount of residual stresses both in the near surface region and in the depth. These trends were observed both with and without an ablative coating.
- Larger spot sizes lead to increased depths of CRS.
- Ablative coating leads to larger and deeper CRS. This is true for the entire profile, in particular in the near surface region (at 20 $\mu\text{m}$) and at the maximum value of CRS.

<table>
<thead>
<tr>
<th>LSP treatment</th>
<th>Max RS [MPa]/percentage of the UTS [%]</th>
<th>Depth of max RS [(\mu\text{m})]</th>
<th>Depth of CRS [(\mu\text{m})]</th>
<th>Surface RS [MPa]/percentage of the UTS [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>316L – AB</td>
<td>342/45</td>
<td>131</td>
<td>/</td>
<td>105/14</td>
</tr>
<tr>
<td>316L – 1 mm 40% NC</td>
<td>$-266/35$</td>
<td>128</td>
<td>416</td>
<td>$-103/14$</td>
</tr>
<tr>
<td>316L – 1 mm 80% NC</td>
<td>$-730/96$</td>
<td>94</td>
<td>804</td>
<td>$-418/55$</td>
</tr>
</tbody>
</table>
- When using an ablative coating the CRS increase is more pronounced for spot sizes of 2 and 5 mm
- Higher overlap rates (80%) lead to higher CRS and a deeper CRS profile. Although these LSP parameters give significantly better results, the associated higher number of impacts leads to longer processing time.

Further investigations will be done on the effect of the LSP treatment on the microstructure and the fatigue life of SLM parts. Also, the possibility of combining LSP treatment with SLM into a single machine will be addressed. The goal of using LSP during the building phase of SLM as a “3D LSP” method would be to tailor residual stresses throughout the part, especially focusing on critical points and in the near surface zone. This would possibly lead to increased depth and volume of CRS in the near surface region, which is known to have beneficial effects on fatigue life. Another goal would be to reduce the accumulation of TRS during the building phase, to avoid process failure for certain materials which are hard or even impossible to process by SLM (e.g. Ni-based superalloys).

References

Fig. 7. Residual stress curves measured for both PH1 and 316L samples in the AB and LSP treated states without an ablative coating. Spot size was 1 mm and overlaps 40% and 80%.