Effect of printing parameters on sintered WC-Co components by binder jetting

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ABSTRACT
Hardmetals are materials employed to produce cutting and forming tools as well as wear resistant parts. Standard powder metallurgy suffers limitations in the manufacturing of shapes required by application-oriented design. Therefore, there is interest for the implementation of additive manufacturing, in particular low temperature techniques as binder jetting because they allow to preserve microstructures and peculiar properties. In our work, the powder was tungsten carbide with 12 wt.% cobalt (WC-Co). Shaping by binder jetting was followed by curing treatment to promote the binder polymerization, vacuum sintering and sinter-HIP to achieve near-full density. The powder was analysed in terms of size and shape, to determine its suitability for the procedure. Then, the effects of different combinations of printing parameters, layer thickness (50–100 μm) and binder saturation (60/75/90%), on the density of the green bodies were analysed. Finally, the relative density of the sintered components was measured and the pores shape and size were studied by SEM imaging, to assess possible consequences of the deposition procedure. Vickers hardness along the samples cross-section was measured and correlated to the printing conditions employed.

1. Introduction
The need of components with specific and complex geometries from hardmetals and cemented carbides has gathered the attention of the academy and the industry on the adoption of additive manufacturing
(AM) techniques applied to this class of materials. Indeed, the freedom of design granted by this kind of processes could allow for the commercialisation of customised products, with short production times, reduced costs and waste, and limited or no need for additional operations to adjust the size and shape by post-processing, which is challenging due to the high hardness and brittleness of cemented carbides (Aramian, Razavi, Sadeghian, & Berto, 2020; Gao et al., 2015; Lehmann & Zaeh, 2019; Lin, Xu, Shan, Chung, & Park, 2015; Seharing, Azman, & Abdullah, 2020).

At first, many efforts have been spent on employing this type of materials with more consolidated AM techniques, as selective laser sintering (SLS) and laser-powder bed fusion (L-PBF) (Campanelli, Contuzzi, Posa, & Angelastro, 2019; Chen et al., 2019; Grigoriev, Tarasova, Gusarov, Khmyrov, & Egorov, 2019; Ku, Pittari, Kilczewski, & Kudzal, 2019), laser engineered net-shaping (LENS) (Davoren, Sacks, & Theron, 2020) and direct metal laser sintering (DMLS) (Gu et al., 2012; Wang et al., 2002); however, the low control on the heating and cooling rates involved leads to a series of issues (e.g. thermal cracks, distortions, residual porosities, undesired carbides formation…) that prevent the quick adoption of these kind of processes, at least with current technologies and materials composition.

In the last years, indirect AM techniques, as fused deposition modeling (FDM) and binder jetting (BJ), have been extensively explored because they offer the advantages of application-oriented design combined to the possibility of controlling the thermal treatments required for the densification of the powders or granules employed. This allows to achieve a much tighter control on the microstructural evolution and retain a fine and homogeneous size distribution of equiaxed grains, with extremely limited amount of undesired phases and porosity (Cramer et al., 2020; Eguiluz, Ferr, Yus, & Ferrari, 2021; Enneti & Prough, 2019b; Lengauer et al., 2019; Mariani, Beltrami, et al., 2021; Mostafaei, De Vecchis, Kimes, Elhassid, & Chmielus, 2021; Mudanyi et al., 2021).

Although first attempts have displayed promising results, further research is needed to assess the effects of printing parameters on the final microstructural and mechanical properties. Currently, two types of WC-based powders have been employed in BJ, namely granulated (Carreño-Morelli, Alveen, Moseley, Rodríguez-Arbaizar, & Cardoso, 2020; Enneti et al., 2018; Enneti & Prough, 2019a, 2019b; Lehmann & Zaeh, 2019; Mostafaei, De Vecchis, et al., 2021) and fully dense feedstocks (Cramer, Nandwana, Lowden, & Elliott, 2019; Cramer, Nandwana, et al., 2019; Cramer et al., 2020; Cui et al., 2021; Mariani, Beltrami, et al., 2021; Mudanyi et al., 2021). The latter group comprehends works were BJ was
employed to produce preform, which were subsequently infiltrated with binder metal, as in Cramer et al. and Mudanyi et al., and one previous study of our group that features plasma spheroidised particles. Few studies have been realised on the effects of printing parameters, in particular:

- Enneti and Prough (2019a) observed the relevancy of layer thickness and binder saturation, rather than binder set time, on the properties of green samples (sintering was not performed);
- Mostafaei, De Vecchis, et al. (2021) described the effect of multiple parameters on green and densified components, obtained from powders produced by spray drying.

In our work, we explore the possibility of producing accurate and performing WC-Co components by binder jetting from a pre-densified feedstock. Specifically, an investigation on different combinations of layer thickness and binder saturation levels is presented, and the direct correlation of these two parameters to the organic binder distribution within the printed component, which has been demonstrated to affect the geometrical accuracy, the amount of residual carbon after debinding and the presence of excess porosity, among other effects (Enneti & Prough, 2019a; Mostafaei, De Vecchis, et al., 2021; Nandwana, Kannan, & Siddel, 2020).

2. Materials and methods

2.1. Materials and powder characterisation

WC - 12 wt.%Co powder by Global Tungsten & Powders Corp. (US) was employed, as in previous studies (Enneti et al., 2018; Mariani, Beltrami, et al., 2021). The raw material is plasma spheroidized to ensure a high shape regularity and it features a monomodal size distribution with a reduced average diameter ($D_{50} = 1.5 \mu m$, from the numerical distribution). The commercial aqueous-based binder “AquaFuse” by ExOne Inc. (US) was employed for all the printing campaigns.

Static image granulometry (ASTM E2651-19) was employed to determine the corresponding diameter and circularity degree of the particles from their 2D section projection. Circularity is calculated as the ratio of the circumscribed perimeter on the real perimeter of the particle. These features were confirmed by field emission-scanning electron microscopy (FE-SEM) with a ZEISS SIGMA 500. In addition, material composition was assessed by EDX analysis on a sufficiently large number of particles.
The packing and flowability behaviour of the powder was evaluated with a FT4 Powder Rheometer 2 (UK) by measuring:

- the apparent density ($\rho_{app}$), corresponding to that of the as-poured material;
- the tapped density ($\rho_{tap}$), i.e. the density after compaction by tapping the as-poured powder.

These values allowed to measure the Hausner ratio $H$, according to Eq. 1:

$$H = \frac{\rho_{tap}}{\rho_{app}}$$

### 2.2. Printing procedure

Components were produced with an Innovent+ printer by ExOne Inc. (US). The machine parameters (listed in Table 1) affecting the deposition and spreading of the powder on the printing plate were optimised to assure the reliable formation of the layers throughout the whole printing process. The drying time was kept fixed at 12 seconds. Instead, five combinations of layer thickness (LT) and binder saturation (BS) were tested to assess their effect on the geometrical accuracy and density of the components at the green and sintered stages. In detail, LT was set at 50 $\mu$m or 100 $\mu$m and BS was 60%, 70% or 90%. BS of 90% immediately showed unsatisfying results, therefore it was tested only with 100 $\mu$m of LT. Samples nomenclature is given in Table 2.

Then curing was performed at 180°C for 6 hours in a static air oven to produce resistant green bodies, which underwent depowdering by manual brushing. Debinding was performed at 500°C for 4 hours in inert
atmosphere and sintering at 1400°C in vacuum. Finally, a sinter-HIP process was realised with a 35 bar pressure at 1400°C for 20 minutes.

2.3. Microstructural and mechanical characterization

The relative density of the samples was calculated at all the steps of post-processing to evaluate the effect of the thermal treatments on the microstructure and accuracy. After curing density was measured by geometrical method given the large network of open porosity and samples brittleness, while Archimede’s method was applied to sintered and sinter-HIPed specimens (ASTM B311-17). In both cases, the reference density considered for WC with 12 wt.% of Co was 14.29 g cm⁻³.

The carbon content in all samples was measured by IR spectroscopy with a LECO WC230 according to ISO 11873. Sintered and sinter-HIPed microstructural features, as internal porosity, grain size and phase composition, were analysed by optical microscopy and FE-SEM on polished surfaces. The images were employed for analysis and manipulation by ImageJ through the trainable WEKA segmentation plugin to collect additional information on pores distribution with a resolution corresponding to 1 μm (Arganda-Carreras et al., 2017).

3. Results and discussion

3.1. Powder characterization

Granulometry (Figure 1) revealed that most of the particles stand in the 700 nm–10 μm range, thus the particles size distribution is shifted toward reduced diameters with respect to those usually employed in binder jetting and other additive techniques, which can go up to 100 μm (Mostafaei, De Vecchis, et al., 2021). This should ensure an optimal packing, as well as sintered density and geometrical accuracy, since the inter-particles porosity normally stands in the same size range.

A distinction was observed between the coarser fraction of the particles (diameter above 700 nm) that features a relevant circularity, consistently higher than 0.7, and the finer fraction (diameter below 400 nm), which displays a minimal shape homogeneity. Indeed, the latter fraction comprehends detected particles with a circularity degree below 0.3, meaning that the shape is elongated. These should be fragments that have detached from the main spheroidized particles during powder handling. SEM image of the as-receive powder in Figure 2 confirms the observations made on particles size and shape from granulometry. Fragments are visible in some spots (e.g. upper side of Figure 2).
Figure 1. Circularity and size distribution of the individual particles elaborated from granulometric analysis.

Figure 2. SEM image of the as-received powder.

Powder rheometry revealed that the employed material features an excellent flowability, considering that the Hausner ratio for a liquid would be 1.0. The result in Table 3 is impressive for such size distribution, and it is likely due to three main factors:

- accentuated sphericity of the particles
- high particles true density
- high hardness of the material
These features prevent particles mechanical interlocking and allows frictionless motion under their own weight without the need of powder de-humidification or similar operations, which is convenient in terms of time and cost.

3.2. Green specimens characterization

The printing parameters revealed to be highly influential on the repeatability and the accuracy of the process. Indeed, average density values display a limited variation in between the five combinations considered, ranging from 47.3% for 50_60 to 50.9% for 100_90, all of them being close to the powder packing value obtained by rheometric measurements in Table 3; however, the values distributions exhibit much different behaviours, as shown in Figure 3. The increase in average density with larger BS can be ascribed to the weight contribution of the binder itself, whereas the values distribution are correlated to the dimensional accuracy.

BS is the most influential parameter, particularly when combined with a larger LT (Rishmawi, Salarian, & Vlasea, 2018). It must be considered that the same BS level does not correspond to the same deposited binder volume at each layer with variable LT, as the volume to be filled increases with this parameter. Indeed, BS is defined as follows (Eq. 2):

\[ BS = \frac{V_{db}}{V_{por}} \]  

(2)

where \( V_{db} \) is the volume of binder deposited and \( V_{por} \) is the volume of the powder bed porosity. The latter parameter can be calculated as (Eq. 3):

\[ V_{por} = (1 - PPR) \times V_{sl} = (1 - PPR) \times LT \times S_{pgs} \]  

(3)

where \( PPR \) is the powder packing ratio, \( V_{sl} \) is the single layer volume and \( S_{pgs} \) is the surface of the printed geometry section. Eq. 2 and Eq. 3 can be combined to calculate the volume deposited at each layer, as described in Eq. 4 and Eq. 5:

\[ V_{db} = BS \times V_{por} = BS \times (1 - PPR) \times LT \times S_{pgs} = k_{p-g} \times BS \times LT \]  

(4)

Table 3. Relative apparent density (%), tapped density (%) and Hausner ratio (H) of the as-received powder.

|          | \( \rho_{app} \)/% | \( \rho_{tap} \)/% | H               |
|----------|--------------------|--------------------|-----------------|
|          | 50.69 ± 0.01       | 51.46 ± 0.01       | 1.02 ± 0.01     |
where $k_{p-g}$ is a factor that depends on the powder packing ratio (material feature) and the printed geometry (system feature), which were constant in our case. The $V_{db}$ ratios of our printing conditions are listed in Table 4.

Therefore, the amount of binder deposited at each layer for the five samples varies even when BS is the same. Ideally, the binder should be sufficient to equally fill the layer porosity and join the particles with the precedent layer; however, the wetting phase is deposited on the top surface of the layer regardless of its thickness, thus oversaturation may not be expected overall, but it might occur locally. This mechanism could lead to wetting out-of-bounds particles, due to capillarity in all directions, also sideward, thus reducing the printed shape accuracy, rather than

$$V_{db} \cdot k_{p-g}^{-1} = BS \cdot LT$$

Figure 3. Relative density distributions of the green samples obtained from the combination of LT = 50/100 µm and BS = 60/75/90%.

| LT/µm | BS  | $V_{db} \cdot k_{p-g}^{-1}$ |
|-------|-----|-----------------|
| 50    | 0.60| 30              |
| 50    | 0.75| 37.5            |
| 100   | 0.60| 60              |
| 100   | 0.75| 75              |
| 100   | 0.90| 90              |
completely infiltrate along the Z-axis (Chun et al., 2020; Mariani, Beltrami, et al., 2021). The risk of incomplete infiltration, thus of insufficient wetting at the layer-layer interfaces, with larger LT and same BS seems to be confirmed by the reduction in green strength measured by Enneti and Prough (2019a), under similar conditions, and by other studies on different materials (Barthel, Janas, & Wieland, 2021; Jimenez et al., 2019; Lecis et al., 2021; Mao, Li, Li, Cai, & Wei, 2021).

Consequently, the printed volume varies with respect to the desired geometry and this mechanism is accentuated when the overall liquid amount is higher, which occurs with an increase of either BS or LT or both. The limited control on the green samples geometry, when a larger volume of binder is employed, is responsible for the lower repeatability of the 100_75 and 100_90 printing combinations, as also observed by Mostafaei, De Vecchis, et al. (2021).

### 3.3. Sintered and sinter-HIPed specimens characterization

The sintering process allowed to achieve above 95% relative density values regardless of the printing conditions. As shown in Table 5, the densification degree was differentiated and varies inversely with the average relative density of the corresponding green specimens. In particular, the samples produced with BS = 60% experienced a densification of almost 50% and featured the largest density values. The amount and the distribution of the binder within the printed structure affects the characteristics of the samples: as already described in previous works, localised over-saturation may generate inhomogeneous and enlarged porosity after debinding, which hinders or reduces pores filling from solid-state or liquid-state diffusive mechanisms.

Sintering also has the beneficial effect of producing more homogeneous samples. This should be due to the overall reduction of the porosity, which is mostly of spherical shape, as the process achieve final stage sintering (Rishmawi et al., 2018). Pore size distributions and circularity (Figure 5) obtained from 2D transverse sections (examples are shown in Figure 4) confirm that meaningful differences in between samples are not detected. Therefore, the effect of irregular macro-porosities generated by improper binder distribution is mitigated. Samples with BS of 60%

| Sample | 50_60 | 50_75 | 100_60 | 100_75 | 100_90 |
|--------|-------|-------|--------|--------|--------|
| Avg. relative density/% | 96.5±0.1 | 95.8±0.2 | 97.4±0.2 | 96.0±0.3 | 96.4±0.3 |
| Densification degree/% | 49.2±0.6 | 47.6±0.5 | 49.6±1.3 | 45.2±2.9 | 45.4±4.4 |
feature slightly larger average pore size, which could be due to the more progressed sintering process leading partially to porosity coalescence.

Sinter-HIP revealed to be a fundamental step in the process since it increased the average relative density of all samples above 99%. The application of pressure combined with high temperature allowed to remove the internal closed porosity. This can be clearly observed in Figure 6, where the sinter-HIPed sample micrographs (Figure 6C and D) do not feature relevant macro-porosity as in the sintered samples (Figure 6A and B). At both stages, fine and coarse grains can be observed, which are due to the microstructure of the powder employed and have been studied in previous works (Enneti et al., 2018; Mariani, Beltrami, et al., 2021).

Figure 7B displays the hardness values obtained from micro-indentation on longitudinal section of sintered and sinter-HIPed samples. The main

![Figure 4. SEM micrographs of the sintered samples transverse sections. In order: (A) 50_60; (B) 50_75; (C) 100_60; (D) 100_75; (E) 100_90.](image-url)
difference stands in the presence of values below 1100 HV of some samples, which are due to residual porosity that may produce an enlargement of the indented track. Sinter-HIPed components feature an extremely limited porosity, thus the risk of pore-affected measurements is reduced, and lower values close to 1200 HV are likely generated by cobalt pools that are observed throughout the microstructure of all the samples. Effects of the presence of graphitized carbon in the microstructure could be excluded since relevant differences were not detected in between five printing parameters combinations and among sintered and sinter-HIPed specimens, as can be seen from Table 6. The risk could have arisen from the incomplete burnout of the polymer during debinding in inert atmosphere (Lecis et al., 2021), however the process was effective in removing the organic binder prior to densification to an extent sufficient at avoiding the risk of printing-dependent microstructural defects or anomalies.

Once again, distinctions in between the tested printing combinations were not detected at the microstructural level, coherently with the homogeneity observed in the microstructural features, while macroscopic geometrical inaccuracies can be easily observed with higher LT and BS, as can be seen in Figure 8, where the samples with the lowest and largest amount of binder at each layer are presented (50_60 and 100_90, respectively).

4. Conclusions

The work describes the effects of the pivotal parameters for the organic binder distribution within binder jetted specimens, i.e. the layer thickness (LT) and the binder saturation (BS), on the geometrical accuracy and microstructural properties. The tested conditions include two LT (50 and
Figure 6. SEM micrographs of sintered longitudinal (A) and transverse (B) polished section surfaces and of sinter-HIPed longitudinal (C) and transverse (D) polished section surfaces of the 50_60 specimen.

Figure 7. Vickers hardness values measured on sintered (gray) and sinter-HIPed (red) samples obtained from the combination of LT = 50/100 µm and BS = 60/75/90%.

Table 6. Average carbon content values (%) before and after sinter-HIP treatment.

| Process step | 50_60  | 50_75  | 100_60 | 100_75 | 100_90 |
|--------------|--------|--------|--------|--------|--------|
| Carbon content/wt.% | Sintering | 5.54±0.01 | 5.51±0.02 | 5.56±0.02 | 5.52±0.03 | 5.53±0.04 |
|                | Sinter-HIP | 5.54±0.03 | 5.52±0.01 | 5.57±0.12 | 5.55±0.06 | 5.54±0.01 |
100 μm) and three BS levels (60, 75 and 90%). The material employed is tungsten carbide with a 12 wt.% Co matrix powder, obtained from plasma spheroidization.

The analyses performed at the different stages of the process have underlined that:

- both LT and BS strongly affect the reliability of the shaping phase, given its dependence on the distribution of the binder within the powder bed, highlighting improved green parts quality for lower values of both parameters;
- after sintering and sinter-HIP, the differences at the microstructural level are minimised, thus the correlated mechanical properties are comparable for all the tested conditions, which is certainly advantageous in terms of flexibility of the production cycle.

In conclusion, the main effect of the printing parameters choice on the finished products is that on their level of geometrical accuracy. This is of pivotal importance for commercialisation because the huge advantage of producing this class of material from AM would be obtaining complex design and details without the need of surface material removal.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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