Coated Metal Powders for Laser Powder Bed Fusion (L-PBF) Processing: A Review

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Abstract: In the last years, functionalized powders are becoming of increasing interest in additive manufacturing (particularly in laser powder bed fusion processing, L-PBF), due to their improved flowability and enhanced processability, particularly in terms of laser absorbance. Functionalized powders may also provide higher final mechanical or physical properties in the manufactured parts, like an increased hardness, a higher tensile strength, and density levels close to theoretical. Coatings represent a possible interesting approach for powders' functionalizing. Different coating methods have been studied in the past years, either mechanical or non-mechanical. This work aims to present an overview of the currently obtained coated powders, analyzing in detail the processes adopted for their production, the processability of the coated systems, and the mechanical and physical properties of the final parts obtained by using L-PBF for the powders processing.

Keywords: coated metal powders; absorbance; L-PBF; mechanical and physical properties

1. Introduction

Manufacturing techniques based on powder metallurgy (PM) play a significant role in several different industrial fields including, among others, automotive [1,2], aerospace [3–5], and biomedical [6,7]. PM is also growing in the production of electro-magnetic components with properties difficult or impossible to be obtained with conventional forming techniques [8–11].

The necessity of obtaining more performing PM parts led to the development of functionalized base materials [12–14] and optimized processes. In this frame, the use of coated powders has been significantly increasing in PM, aiming to improve the powders’ processability or to modify the final part’s microstructural, mechanical, physical, and thermal properties. By coating the powders’ surface, it is possible to change melting temperatures [15], the flowability [15,16], and the absorbance [17–21] of the powder itself. Microstructural characteristics can also be modified by coating with a second phase, particularly nanoceramic particles such as ZrO2, Al2O3, B4C, SiC, TiB2, C-based structures [22–24]. These can induce grain refinement mechanisms, thus enhancing the mechanical strength [15,23]. Mechanical properties of the final parts, such as compressive strength [25,26], hardness [15,26,27], yield strength [15,28], and physical properties like density [19,29], are improvable by adopting coated powders.

Additive manufacturing (AM) represents a challenging topic for researchers, as demonstrated by the increasing number of industrial applications in several different sectors [30].
AM processes are classified after DIN EN ISO/ASTM 52900 standard [31]. They represent an alternative to subtractive and formative manufacturing [32]. In particular, laser powder bed fusion (L-PBF) is suitable for obtaining complex metallic objects owing to its geometric freedom when selectively sintering the powders [33–35]. L-PBF is a layer-by-layer process where a laser beam is applied to melt the metal powders selectively; the molten layer then solidifies at a high cooling rate [36–40]. Adopting a laser source implies using metal powders characterized by a proper laser absorptivity [41,42]. Poor absorbance causes many defects such as pores, low-quality surfaces, and un-melted powder in the 3D-printed parts [41,43–46].

Concerning the materials processable by L-PBF, beyond pure metals and alloys, also intermetallic-based compounds can be printed to obtain dense parts [47–49]. The possibility of integrating more parts into an assembly [37,38], obtaining high-density parts [50], and having low material wastage [45,49], are some of the further advantages provided by L-PBF. In order to guarantee uniform distribution of the powders in each layer, powders used for L-PBF should be characterized by good flowability and spherical morphology [36,51–53].

L-PBF printed parts can be post-treated with hot isostatic pressing (HIP) technique [54,55]. HIP removes residual stresses, promotes grain growth and recrystallization [56], improves mechanical properties, particularly creep and fatigue resistance [57,58]. In addition, HIP can reduce porosity generated during the previous process, as demonstrated with steels [59], Ti [60], and also Al [61] ingots obtained from casting.

Coated powders are adopted in the L-PBF process both for improving their processability (especially for increasing the laser absorptivity and powders’ flowability) and/or the properties of the final parts. For increasing laser absorptivity, at the emission wavelength of the laser commonly adopted in L-PBF (1064 or 1070 nm), an increase of powders’ surface roughness can be provided [62,63]. Another solution might be coating powders with a material characterized by a lower reflectance [17–19,21,63–65]. Poor flowability of powders can be attributed to the influence of van der Waals (vdW) forces. According to Rumpf [66], the effect of vdW forces can be reduced by adequately tailoring the surface roughness of the powders by coating them with smaller particles [67].

This work presents an overview of the most up-to-date coated systems (powder + coating), including the description of the applied coating methods, in order to analyze and understand not only the potentialities in improving materials’ processability through L-PBF (such as improving their absorbance) but also some of the mechanical properties of the final objects.

2. Base Metal Powders

Steels powders are widely used in an extensive range of industrial fields such as aerospace [68], automotive [69], oil and gas industries [70], biomedical engineering [7,68,71], and other fields, owing to their high ductility and high strength [72], biocompatibility and, considering stainless steel powders, also to their high corrosion resistance [68,72]. Austenitic stainless steel powders show however low hardness and relatively poor wear resistance; a strategy for improving the hardness and the wear resistance is coating the metal powders using ceramic nanoparticles (e.g., TiB₂) acting as a grain refiner. Extensive dislocations are introduced too, leading to a finer microstructure and higher mechanical properties [70,72].

Powders of titanium and its alloys are used in many industrial fields like chemical industries [73,74], defense industrial sectors, aerospace [73,75,76], and biomedical applications [6,75,77]. This is due to their high specific strength, high biocompatibility, low density, and optimal corrosion resistance [6,73,75,76,78]. However, titanium and its alloys are also characterized by low hardness and relatively poor wear resistance responsible for the limited industrial areas of use, especially for those applications characterized by erosive and abrasive-based wear processes. The addition of ceramic particles can raise the working temperature and the mechanical properties [74,79].
Metal powders made of aluminum and its alloys find application in a wide range of fields: aerospace [5,80], automotive [81,82], and domestic industries [80] owing to their lightweight, high specific strength, high corrosion resistance, good thermal conductivity, and good mechanical properties [5,80–82]. Aluminum matrix composites (AMCs) have been adopted to increase wear resistance and hardness. Coating metal powders can produce AMCs with reinforcement particles such as Al₂O₃ and SiC [83–85]; hexaborides such as CeB₆ and LaB₆ are used as grain refiners in aluminum alloys both in powder metallurgy [86] and in casting and wrought metallurgy [87,88].

Pure copper is widely used for electronic applications [89] and heat transfer components [90] in different sectors like automotive and naval sector [91] due to its excellent solderability and low electrochemical migration [92], superior electrical (58.7 × 10⁶ S/m) and thermal conductivity (400 W/(m·K)) [43,89,90,92]. Nevertheless, pure copper is soft, so alloying it with Sn, Zn, Cr, and Ni is mandatory for structural applications; these enhance the mechanical properties of copper but reduce its conductivity [93].

**Powders Absorptance**

The powders processed by L-PBF must have a good absorbance at the wavelength of 1064–1070 nm. Pure copper and aluminum are characterized by a low laser absorbance at 1070 nm [41,42,94], as shown in Figure 1 below.

![Figure 1](image-url)  
*Figure 1. Absorbance vs. wavelength for stainless steel AISI 304, titanium, aluminum, and copper; green, blue, and red laser emission wavelengths are highlighted.*

Figure 1 reports the absorbance of stainless steel 304, titanium, aluminum, and copper as function of the wavelength. Ti-based powders show a relative absorbance of about 70% in the range of 1064–1080 nm (corresponding to the red laser), whereas stainless steels (Figure 1 reports the case of AISI 304) are generally characterized by values around 30–35%. Copper and aluminum have a significantly lower absorbance in the red laser emission spectrum (below 10%). Moreover, copper also shows a quite critical absorbance since the energy provided by the laser is quickly dissipated or reflected [18,21]. Different approaches have been investigated to solve this problem: the first consists in the adoption of a high power laser (800–1000 W) [95,96], the second in processing specifically designed alloys characterized by lower thermal conductivity [97,98], the third in substituting the red laser source with a green or blue source (wavelengths of 515 nm [43] and 450 nm [99] respectively). At these wavelengths, copper shifts its optical absorption mechanism from the intra-band to the inter-band electronic transition [100]. Another possibility is by coating the powder’s
3. Coating Processes

Generally, coating processes consist of depositing particles or ions on metal powders; this target is achievable by mechanical or non-mechanical methods detailed in the article. The non-mechanical methods consist of an extensive range of physical and thermal processes with specific variations. Several processes involving solution, chemical, and physical methods for coating powders have been developed and used to solve some of the critical issues of mechanical coating methods.

3.1. Mechanical Methods

3.1.1. Ball Milling

Ball milling is a largely diffused technique that allows dispersing particles on metal powders: it is a non-equilibrium process generally operating at low temperatures, with conceptually linear operations, and a low overall cost [101, 102]. The optimal control of the results obtained from ball milling is however nontrivial; plenty of variables such as ball size, rotation speed, duration, atmosphere, and process control agents (PCA) have significant effects on its outcomes. PCA acts on powder’s purity, grain size, and shape. The ball milling process can be both dry and wet depending on the necessity: in wet ball milling, a solution is added as PCA to reducing the maximum temperature reached during the process [101]. The general scheme of the ball milling process is reported in Figure 2; the metal powders and the coating particles are inserted into a mill containing the balls and the PCA if needed; the balls lead to a faster homogenization, promoting the adhesion of the coating particles to the powder.

![Figure 2. Schematic diagram of ball milling process for coating powders, the objects in figures are not in scale.](image)

Ball milling is widely adopted for coating metal powders to be used in L-PBF [22, 23, 26–29, 47, 103–113] and in hot isostatic pressing (HIP) processes [27, 114, 115]. According to the energy level, ball milling can be high-energy (HEBM) or low-energy (LEBM). The main differences between the two lie in ball-to-power ratio, mixing speed, and processing time, with all the parameters being higher for HEBM. In particular, the ball-to-powder ratio ranges from 5:1 to 30:1 in HEBM, whereas it is 1:1 in LEBM [105]. HEBM is often coupled to nanoceramic particles [107, 116] since they tend to agglomerate due to their high aspect ratio easily. The promotion of strong vdW forces may cause the formation of aggregates leading to microstructural inhomogeneities. HEBM deforms the powders to a
high degree; indeed, they do not maintain the original spherical morphology, causing a reduction of flowability and making powders less suitable for L-PBF [29,105,116].

On the other hand, after LEBM, powders retain their original spherical morphology [105,116]. Zhai et al. [105] report LEBM to effectively coat steel particles with a uniform layer of Y2O3 nanoparticles. Powders kept their spherical geometry after processing for 7 h, and LEBM determined an increase in surface roughness leading to a slight decrease in flowability, apparent and tap densities.

Attar et al. [26] underlined the importance of milling time. In their study, the influence of milling time on the distribution of TiB2 particles on the powder surface is evaluated: 1 h is not sufficient to obtain a uniform coating, 4 h caused an excessive flattering of powders, whereas 2 h provided both a uniform coating and a suitable geometry. Han et al. [23] focused on the milling time, proposing to interval the milling with pausing steps to reduce the temperature reached by the powders due to the impacts with the balls. They also investigated the introduction of stearic acid lubricant to facilitate the process.

The ball milling process is also used for mechanical alloying (MA) [117,118]. HEBM can lead to MA, resulting in a modification of powders’ size, microstructure, and morphology. It can synthesize both equilibrium and non-equilibrium alloy phases such as supersaturated solid solutions, nanostructures, metastable crystalline phases, and amorphous alloys. The high collision between powders and balls determinate a mass transfer accelerating the diffusion of elements. Nevertheless, MA is characterized by a large amount of uncertainty on the final microstructure [118]. MA has been adopted in order to cover metal powders [22,27,47,114], which underwent characterization by X-ray diffraction (XRD). Gu et al. [47] investigated Ti powders mixed with graphite and Al nano powders; after 10 h processing, C peaks disappeared, while TiC was detected. A similar trend is obtained with Al peaks, which gradually disappears during the processing and is not detectable after 20 h of HEBM. AlMangour et al. [27] coated 316L powders with TiB2 particles (2.5 vol.%). After milling for 2 h, γ-Fe peaks were only detected (TiB2 was not detected due to the small amount), after milling for 4–8 h XRD evidenced TiB2 peaks, also α-Fe phase was identified, and its peaks intensity increased with milling time, due to the plastic deformation causing γ-Fe to transform into α-Fe. Wang et al. [114] also observed the formation of α-Fe and the reduction of γ-Fe peaks intensity after coating pre-alloyed AISI 304 powders with Ti and Y2O3. After milling for 15 h, Ti and Y2O3 were not detectable in the material, while YO1.401 and Ti3N1.29 peaks were indexable.

3.1.2. Turbula

Turbula proved to be effective in the mechanical coating of metals powders in several studies [15,16,19,119].

The Turbula mechanically mixes powders and coating particles without using balls; as a result, the deformation on the starting metal powders is strongly reduced. This technique has been adopted by Jadhav et al. [19] to coat the copper powders and increase their optical absorption and flowability. By adding 0.1 wt.% carbon particles, the authors enhanced the optical absorption to 67%, starting from 29% for pure copper. Moreover, Karg et al. [16] used turbula to add SiOx nanoparticles on aluminum alloy powders. The authors increased the powder’s flowability leading to an increase in the relative density of the final parts after L-PBF. Hentschel et al. [15] noticed that one of the most influential parameters is milling time; they found that achieving a homogeneous distribution of nanoparticles needed at least 2 h. The flowability of powders is increased by introducing 0.2 wt.% of nanoparticles; a further increase in the concentration of nanoparticles causes a decrease in flowability but remains acceptable for concentrations lower than 0.5 wt.%. 
3.2. Non-Mechanical Methods

3.2.1. Powders Immersion

Coating particles are dispersed in a solution into which metal powders are immersed, with a possible variation being the drop by drop addition of coating particles to the powders-containing solution. A general scheme of the process is shown in Figure 3.

![Schematic diagram of coating process involving solution for coating powders](image)

**Figure 3.** Schematic diagram of coating process involving solution for coating powders, the objects in figures are not in scale.

One coating process through powder immersion involving the use of solutions is electroless plating [120]. This technique is preferred to the electrolytic plating process owing to its lower cost, high efficiency, applicability independently of the shape of the powder, and a dense and uniform coating deposition [121–124] whose thickness depends on the powder morphology. Electroless plating has been used for coating powders in different fields [17,23,122–127]. Li et al. [122] observed that the powder surface became coarse and irregular; some coating clusters were detected, whose formation can be related to the high deposition rate. Intermediate layers may be introduced to improve adhesion, as investigated by Xu et al. [123]. Sn was used as an intermediate layer between the Cu powder and the Ag coating then this system was heat treated to form an alloyed transition layer increasing bonding strength and oxidation resistance. The coated powders maintained an excellent dispersion, while their sphericity and surface smoothness decreased marginally. Jadhav et al. [17] deposited a thin layer of 62 ± 14 nm of metallic Sn on copper powders; the best results were obtained with a 0.28 wt.% Sn coating layer that led to an increase in powders flowability and an enhancement from 19% to 51% of powders optical absorption. The authors also underlined the importance of using starting powders with low sulfur content, given their critical behavior in L-PBF. Geng et al. [125] homogeneously deposited Ni onto Al powders. The process led to an increased laser absorbance owing to the Ni lower laser reflectivity and increased surface roughness. In addition, flowability was comparable to that of the starting powder.

Electrostatic self-assembly is another possible way of coating powders. The electrostatic attraction allows to assemble spontaneously metal powders and the coating particles during hetero-agglomeration [128,129]. Electrostatic self-assembly guarantees good flowability and the possibility of depositing structures such as carbon nanotubes (CNTs), preventing their destruction [62,63,115,128]. Zhou et al. [116] used CNTs and nano Al₂O₃ to coat metal powders. The two nanoparticles are characterized by opposite charges (negative the former and positive the latter), so a CNTs/Al₂O₃ hybrid was obtained by mixing them. This structure resulted negatively charged since Al₂O₃ did not cover all CNTs, leaving some uncovered surface that can bond to the metal powder. The hybrid solution was slowly added to the metal powder colloid and then mechanically stirred. The deriving coated powders are characterized by homogeneous dispersion of the hybrid coating on the metal powders, suitable particle size and distribution, good flowability, and enhanced surface roughness leading to an increased laser absorptivity. In a more recent study Zhou et al. [62] used CNTs for coating titanium alloy powders. The oxidized surface
of the metal powder is positively charged and, due to the electrostatic attraction, it reacts with CNTs: an increase in the surface roughness has been detected coupled to a consequent increase in laser absorptivity. Dong et al. [63] used graphene oxide sheets to coat Al alloy powders; the coated powders are characterized by a decrease in thermal conductivity and an increase in laser absorptivity.

A suspension characterized by a pH ranging between the two isoelectric points (IEPs) of two different materials leads to a dielectrophoretic deposition, representing an alternative way of coating powders [130,131]. Different studies have adopted coating ferritic steel powders with Y$_2$O$_3$ nanoparticles [25,131–133]. Before the deposition step, the suspension containing Y$_2$O$_3$ nanoparticles has been irradiated to de-agglomerate the nanoparticles. The pH of the suspension is then set to a value able to guarantee that particles and powders are oppositely charged. Finally, the steel powders are added to the suspension, and the pH controls the adsorption. Thus, the dielectrophoretic deposition is scalable and economically feasible [132].

Other solution-based methods for coating powders are reported in literature [21,56,134–136]. Ma et al. [136] adopted an electro-codeposition process where mechanical mixing and sonication were used to distribute Al$_2$O$_3$ nanoparticles and maintain dispersion. Metal powders coated by nanoparticles dissipate less laser-transferred heat, thus generating an increase in the melt pool depth. The study also reports a reduced heat-affected zone. Garmendia et al. [135] coated an aluminum alloy powder with a 1 wt.% copper formate–methanol solution; powders were then heat-treated in vacuum, and the copper precursor was reduced to metal copper. This coating process did not particularly affect the size, flowability, and morphology of the powders. Smith et al. [56] used acoustic mixing to coat NiCoCr powders with Y$_2$O$_3$ nanoparticles. The acoustic mixing uses a wave that attained a resonance among the container, the powders, and the vibrating spring system [137], providing homogenization of the powder in one hour. Zhang et al. [134] coated aluminum alloy powders with styrene to process them through direct light processing (DLP), a specific AM process requiring the dispersion of powders in a monomer and its photoinitiator [138,139]. The necessity of coating aluminum alloy powders before dispersing them into the monomer and its photoinitiator is related to the agglomeration of fine powders (size < 30 µm), having a high surface-to-volume ratio [140], and to the high refractive index difference between the powders and the photosensitive resin; the latter causing a low light penetration depth. Styrene monomer and its initiator are added to the metal powders. They are stirred and heated to 75 °C; after 6 h, the powders are coated and ready to be mixed with the monomer and its photoinitiator for printing. The coating has a uniform thickness (200–400 nm), and the shape of the powder is not modified by the coating process.

3.2.2. Chemical Vapor Deposition and Physical Vapor Deposition

Babul et al. [141] produced graphene-coated Cu powders using a chemical vapor deposition (CVD) technique, including several steps. The first consisted of fluidizing the metal powder in the coating chamber. For this purpose, vibrations were used, coupled to hydrocarbons-containing gases flowing in the chamber. Next, hydrocarbon decomposition was induced by enhancing the temperature, generating the carbon source to produce graphene. Lastly, graphene was nucleated, growing on the surface of the metal powder.

Fluidized bed CVD (FBCVD) has recently been adopted for coating metal powders [142]. It can form homogeneous coatings due to its capability of suspending each particle in the reactor. Particles are surrounded by the flowing gas containing the reactive gaseous precursor. Liu et al. [48] used C$_2$H$_2$ as the gaseous precursor of CNTs and Ar as the carrier gas. The decomposition temperature has been set to 550 °C; at this temperature, C$_2$H$_2$ decomposed, nucleating on Ti-6Al-4V powder surface. With a continuous flux of gases, the CNTs started to grow on metal powders leading to the formation of a homogeneous coating. FBCVD does not impact on powders geometry and guarantees uniform coating and the maintenance of the original spherical geometry of the powders. In the
paper authors noticed that the degree of sphericity corresponding to the ratio (equivalent surface area diameter)/(equivalent circumscribed circle diameter) varies from 91.6 to 90.2. Such values lead to a good flowability of the particles and consequentially to acceptable processability [48,63,143–145]. FBCVD also allows depositing particles characterized by strong vdW interactions, otherwise very difficult to be uniformly deposited [143]. A scheme of FBCVD is shown in Figure 4.

![Figure 4](image-url)

**Figure 4.** Schematic diagram of FBCVD process for coating powders, the objects in figures are not in scale.

Li et al. [64] underline the importance of adopting a catalyst to synthetize CNTs. For this reason, they evaluated how to introduce catalysts on the surface of the powder before the FBCVD processing. Fe and Ni impurities in Ti-6Al-4V and other alloys have a catalytic effect: a chemical etching was performed to activate these species [48,64,143]. However, catalytic impurities cannot be introduced in Al powders; a 0.1 wt.% catalyst was therefore inserted through electro-less plating [64,144]. A catalyst on the powders’ surface leads to a better interfacial bonding strength and an excellent interfacial relationship between the powders and the FBCVD coating.

Pannitz et al. [146] coated steel powders with few-layer graphene (FLG) or silicon carbide (SiC) adopting a top spray configuration of the fluidized bed coating process; in particular, they used the scheme of the process reported in Figure 5. Moreover, Lüddecke et al. [67] applied this method for coating steel and aluminum powders with FLG or SiC or iron oxide black (IOB).

The deposition consists in spraying the solution containing the coating particles on metal powders; then, a drying step is provided. The coatings deposited via this technique are homogeneous [67,146]. After the coating step, no significant variation in powders’ size is noticeable. As a general result, the laser absorption increases for all the coatings investigated in these studies (few-layer graphene (FLG), silicon carbide (SiC), iron oxide black (IOB)). In particular, FLG resulted in being the most efficient in reducing the metal reflectance; SiC particles can enhance the powder’s flowability, while IOB is not as efficient due to its high water content.
Figure 5. Schematic diagram of fluidized bed process for coating powders, the objects constituting figures are not in scale.

Physical vapor deposition (PVD) techniques have successfully been used to deposit coatings on metal powders, to be further processed through AM, HIP, and press and sinter [20,147–150]. An homogeneous coating of stainless steel on copper powder and coating of copper on stainless steel powder were obtained by Matos et al. [147]. They used a high-frequency vibration system to keep metal powders moving during the deposition coating, adopting a laboratory-made dc magnetron sputtering system; this process was performed in a vacuum chamber. The coating layer is nanocrystalline regardless of the particle shape, and coated powders show a reduction in interparticle friction. Fernandes et al. [148,150] adopted this technique to obtain a homogeneous coating on WC powders, providing the complete coverage of powders’ surface and low coating contents (~1 wt.%). The coating layer led to an increase in the average powder size from 9.1 ± 0.5 \( \mu \text{m} \) to 10.7 ± 0.3 \( \mu \text{m} \) [148]. The increase of the average powder size is also related to a high nanoporosity of the coating layer (around 20%), implying an increase in surface roughness and consequentially in the specific surface area. The coating is permeable to air; however, an increase in coating thickness leads to a decrease in permeability and provides a better oxidation resistance. Simões et al. [149] used PVD for coating powders for HIPping. They noticed an agglomeration of particles during the deposition caused by the non-spherical geometry of the particles, the finer particles, and the broad particle size distribution.

PVD-coated Cu powders were used by Lassègue et al. [20] and Tiberto et al. [46] for L-PBF processing. Lassègue et al. [20] used CrZr particles to coat Cu powders; the authors obtained a coating non-homogeneous in thickness (from 32 to 445 nm). Despite the dishomogeneities and the poor adhesion of the coating film on the Cu powders, a considerable increase of optical absorbance is reported, without major impacts on the processability of the coated powder. Tiberto et al. analyzed powders coated with titanium and reported an increase in laser absorbance of titanium coating film, characterized by a thickness of approx. 86–95 nm [46].

Jadhav et al. [18,151,152] developed a specific coating method for high reflective metals powders in L-PBF manufacturing, consisting of pre-alloying the metal powder with an alloying element that can react with carbon and nitrogen generating nitrides, carbides, or carbonitrides. In addition, the alloying element must have a low or zero miscibility at room temperature with the highly reflective metal. The alloyed metal powder can then be treated in two different ways: (a) Coated using CVD, PVD, or other methods with a coating containing C or N and then treated in a furnace; (b) directly heat-treated in nitrogen or carbon or nitrogen-carbon atmosphere. In both cases, the heat treatment temperature is set at 600–850 °C for at least 1 h. During the treatment, chemical reactions occur between
the alloy element and C or N contained in the coating or the atmosphere, leading to carbides, nitrides, or carbonitrides. Powders after the heat treatment are characterized by the structure shown in Figure 6.

The powder can be ideally divided into three parts: (1) the surface, corresponding to the diffusion layer having a thickness around 459 nm, composed of carbides, nitrides, or carbonitrides of the alloying element and this last in its metallic form; (2) a depleted layer characterized by the highly reflective metal that has been depleted of the alloying element passed in the diffusion layer; (3) the core, whose composition is that of the pre-alloyed powder. This process leads to a considerable increase in absorbance. Nitrogen is finally released from the powders in the L-PBF processing not to affect the chemical composition of the final part.

Based on the aforementioned, in Table 1 a summarizing scheme is provided.

Table 1. A schematic summary of the coating methods’ characteristics.

| Coating Method     | Feasibility of the Process | Dispersion of the Coating Particles | Cost of the Process               |
|--------------------|----------------------------|-------------------------------------|-----------------------------------|
| HEBM               | Easy                       | Good                                | Low                               |
| LEBM               | Easy                       | Good                                | Low                               |
| Powders immersion  | Complex                    | Excellent                            | Depends on the specific adopted method |
| Vapor based        | Complex                    | Excellent                            | High                              |

4. Properties of L-PBF Processed Parts

AlMangour et al. [104] obtained a relative density of 99.99% using 316L powders with 5% TiB\textsubscript{2}; they also detected increased compressive mechanical yield strength ($\sigma_{0.2} = 827.5 \pm 17.0$ MPa @ TiB\textsubscript{2} = 5 vol.%; $\sigma_{0.2} = 980.9 \pm 10.9$ MPa @ TiB\textsubscript{2} = 10 vol.%) and ductility. Parts produced with coated powders through ball milling and parts deriving from the powders direct mixing were analyzed [27]. As for the hardness, no relevant difference was detected after adding 2.5 vol.% TiB\textsubscript{2} while the parts obtained from ball-milled powders showed a higher hardness when the TiB\textsubscript{2} content is 15%. The effect of HIP after L-PBF was also evaluated, which resulted in the elimination of significant cracks and pores. A long HIP cycle however caused a decrease in hardness and wear resistance, due to the transformation of equiaxed grains into nanoparticles agglomeration. Pannitz et al. [146] processed stainless steel 1.4404 grade powders coated with SiC (4 vol.%), SiC (1 vol.%), and FLG (0.75 vol.%). Relative densities of 93.9%, 99.9%, and 99.9% respectively
were obtained. The addition of few-layer graphene (FLG) led to an improvement in laser absorbance and faster heat dissipation into the solidified layer during the printing process. Doñate-Buendia et al. [25] registered an increase of 29% in compressive strength using a ferritic stainless steel powder PM2000 (74.90 wt.% Fe, 20.40 wt.% Cr, 3.94 wt.% Al, and 0.58 wt.% Ti, Si, and Cu traces) nanocoated with Y$_2$O$_3$. Wilms et al. [131] obtained a final porosity of 0.5% processing a ferritic stainless steel powder (Nanoval—21.03 wt.% Cr, 4.67 wt.% Al and 0.47 wt.% Ti) homogenously decorated with nano-scaled Y$_2$O$_3$ powder particles. No cracks and no increment in hardness have been detected, while an increase in compressive strength was reached. AlMangour et al. [29] detected an improved density by introducing finer coating particles onto 316L; they also noticed an increase in hardness when increasing the TiC content.

Attar et al. [26] reached a relative density of 99.50% by coating titanium powders with TiB$_2$; such coating increases hardness due to the formation of titanium monoboride from the reaction between TiB$_2$ and the Ti matrix. Kun et al. [110] used TiC nanoparticles, resulting in an increased tensile strength (914 MPa) with a ductility comparable to the uncoated Ti powders. Zhou et al. [62] determined that CNTs react with the Ti matrix generating TiC and leading to an increase of hardness compared to Ti and the titanium matrix composites produced by traditional processes; the final microstructure is finer with a uniform distribution of TiC particles. Liu et al. [48] produced Ti-6Al-4V samples starting from powders coated with CNTs; this material is characterized by a higher ultimate tensile strength (1255 MPa instead of 1078 MPa), yield strength (1162 MPa instead of 964 MPa), and lower elongation (3.2% from 4.2%) compared to the material produced from uncoated Ti-6Al-4V powders.

Geng et al. [144] noticed that CNTs react with Al during printing; this led to the formation of Al$_4$C$_3$ distributed inside the grains and has a strengthening effect. Based on the results presented by the authors, compared to uncoated aluminum powders, both the ultimate strength and the yield strength increased, the first from 82 MPa to 129 MPa while the second varied from 66 to 99 MPa. No difference was detected in the tensile elongation. Geng et al. [125] evaluated the variation of mechanical properties at increased fractions of Ni deposited on the surface of pure Al powders. The authors found that the optimal fraction of Ni is 5 wt.%, since it increases the ultimate tensile strength from 124 ± 2.5 MPa to 182 ± 0.5 MPa and the yield strength from 95 ± 2 MPa to 146 ± 1 MPa without reducing its elongation. A chemical reaction occurred between the coating (Ni) and the matrix, generating the second-phase Al$_2$Ni. In another study, Geng et al. [126] observed that also Co coating reacts with Al matrix leading to the formation of Al$_4$Co$_2$; this second phase and the Al matrix have a coherent interfacial relationship. The authors also measured the surface roughness of the printed objects, which was lower than the pure Al parts: surface roughness decreased as Co content was raised (the maximum content of Co introduced is 1.1 wt.%). Zhou et al. [128] observed the precipitation of Al$_4$C$_3$ nanorods when printing aluminum powders coated with graphene oxide (GO). Such precipitates are monocrystalline and increase the mechanical properties owing to their intimate contact with the matrix. Jue et al. [22] obtained a relative density of 97.30% and a hardness equal to 175 HV0.1 by coating Al powders with Al$_2$O$_3$. Gu et al. [107] reached a hardness of 188.3 HV0.1 and tensile strength of 486 MPa while the elongation remained constant after coating Al powders with TiC. Mair et al. [28], in their study about 2024 Al alloy decorated with CaB$_6$, were able to print parts characterized by an equiaxed microstructure, with a relative density of 99.5%, hardness equal to 132 ± 4 HV5, a tensile strength of 391 ± 22 MPa, a yield strength of 348 ± 16 MPa, and elongation equal to 12.6 ± 0.6%. Dong et al. [63] noticed that GO sheets applied onto an AlSi10Mg powder partially reacts with the matrix leading to the formation of Al$_4$C$_3$. They also found two different types of porosity; the smaller ones represented by gas porosities, while the bigger and irregular ones to be attributed to the fact that the GO sheets retain their shape during melting. Nevertheless, an increase in hardness has been detected.
By processing a CuCr powder with surface modified with N, Jadhav et al. [18] obtained samples characterized by a thermal and electrical conductivity lower than the un-modified material, but with a higher elongation at break and yield and tensile strengths. Solution annealing and age hardening caused an improvement of all the aforementioned properties, a part from elongation at break. In a previous study, on the evaluation of the addition of 0.1 wt.% carbon nanoparticles on gas atomized Cu powder with purity of 99.7, Jadhav et al. [19] produced parts with a relative density of 98% and a tensile strength of $125 \pm 11$ MPa. Using a CrZr-coated Cu powder, Lassègue et al. [20] obtained samples with a relative density of 94.3% instead of 87.3% for the uncoated powders. Jadhav et al. [17] evidenced the importance of having sulfur-free copper powders as starting material in order to reduce the occurrence of defects such as porosity and solidification cracks. By using sulfur-free powders, authors obtained a tensile strength of $256 \pm 14$ MPa, a yield strength of $203 \pm 4$ MPa, and hardness of $90 \pm 3$ HV0.3, with electrical and thermal conductivity respectively of $80 \pm 1\%$ international annealed copper standard (IACS) and $334 \pm 4$ (W/(m·K)). Lindström et al. [21] underlined the importance of incrementing the laser absorbance to reach higher final densities in the printed parts.

Smith et al. [56] performed a HIP cycle on the AM-produced parts, highlighting that ceramic nanoparticles have a pinning effect leading to the suppression of grain growth and recrystallization.

Tables 2–5 provide a summary of the L-PBF process parameters applied on different coated powders, including the relative density and mechanical properties measured on printed parts.
Table 2. A summary table reporting L-PBF process parameters, relative density and mechanical properties of printed parts using Fe-based alloys.

| Metal Powder | Coating | Coating Technique | Ref. | Laser Power (W) | Scanning Speed (mm/s) | Hatching Distance (µm) | Layer Thickness (µm) | Relative Density | Mechanical Properties |
|--------------|---------|-------------------|------|----------------|-----------------------|------------------------|---------------------|---------------------|-----------------------|
| TiB₂         | Ball milling | [104] | 100 | 83.33 | - | 50 | 99.99% | \( \sigma_{0.2}^2 = 980.9 \pm 10.9 \text{ MPa} \) |
| TiC          | Ball milling | [106] | 100 | - | 120 | - | 98.22% \( @ \eta^3 = 300 \text{ J/mm}^3 \) |
| \( \text{Y}_2\text{O}_3 \) | Ball milling | [105] | 250 | 1200 | - | 40 | 99.6% \( @ 0.3 \text{ wt.\% } \text{Y}_2\text{O}_3 \) |
| TiB₂         | Ball milling | [27] | 100 | 83.33 | 120 | 50 | 91.5% | \( \text{Max compressive yield strength } @ \eta^3 = 67 \text{ J/mm}^3 \) |
| steels       | TiC      | Ball milling | [29] | 100 | 250 | 50 | 50 | \~96\% \( @ 2.5 \text{ vol.\% nm-TiC} \) |
| SiC          | Ball milling | [119] | 100 | 330 | 40 | 50 | - | \~99.9\% \( @ 1 \text{ vol.\% SiC} \) and 0.75 vol.\% FLG |
| FLG/SiC      | Fluidized bed | [146] | 130 | 700 | 80 | 30 | - | \( \text{Hardness = 403 HV0.2 @ } 15 \text{ vol.\% nm-TiC} \) |
| \( \text{Y}_2\text{O}_3 \) Solution | [25] | 160 | 800 | 60 | 30 | - | \( \sigma_{c,p0.2}^5 = 538 \pm 17 @ T_{\text{amb}} \) |
| \( \text{Y}_2\text{O}_3 \) Solution | [131] | 160 | 800 | - | 30 | 99.2% | \( \sigma_{c,p0.2}^5 = 620 \pm 30 @ 600 \text{ °C} \) |

1 Solution: referred to processes described in Section 3.2.1. 2 \( \sigma_{0.2} \): 0.2% offset yield strengths. 3 \( \eta \): volumetric energy densities. 4 COF: coefficient of friction. 5 \( \sigma_{c,px} \): compressive strength at plastic deformation of \( x \). 6 TS: tensile strength.
Table 3. A summary table reporting L-PBF process parameters, relative density and mechanical properties of printed parts using Ti-based alloys.

| Metal Powder | Coating Technique | Ref. | Laser Power (W) | Scanning Speed (mm/s) | Hatching Distance (µm) | Layer Thickness (µm) | Relative Density | Mechanical Properties |
|--------------|------------------|------|-----------------|-----------------------|-------------------------|---------------------|-------------------|----------------------|
| TiB₂         | Ball milling     | [108] | 300            | 800                   | 100                     | 30                  | 92.18% @ 1 wt.% TiB₂ | Nanohardness = 9.96 ± 0.50 GPa @ 1 wt.% TiB₂ |
|              |                  |      |                |                       |                         |                     | 91.33% @ 2 wt.% TiB₂ | Nanohardness = 10.57 ± 0.53 GPa @ 2 wt.% TiB₂ |
|              |                  |      |                |                       |                         |                     | 85.16% @ 3 wt.% TiB₂ | Nanohardness = 9.98 ± 0.49 GPa @ 3 wt.% TiB₂ |
| TiB₂         | Ball milling     | [26]  | 180            | 118–154               | 100                     | 100                 | >99.5%           | Maximum strain = 17.8 ± 3.2% |
| Ti and its alloys |        |      |                |                       |                         |                     |                   |                      |
| TiC          | Ball milling     | [109] | 90             | 300                   | 50                      | 50                  | 98.3%             | Hardness = 577 HV0.2 |
|              |                  |      |                |                       |                         |                     |                   |                      |
| Al           | Ball milling     | [47]  | 800            | 100                   | 150                     | 150                 | Near fully dense | -                    |
| TiC          | Ball milling     | [110] | 100            | 200                   | 70                      | 50                  | 98.2%             | TS ⁵ = 914 MPa      |
| CNTs         | Solution ¹       | [62]  | 20.6           | 10                    | 100                     | 25                  | -                 | El ⁶ = 18.3%        |
| CNTs         | FBCVD            | [48]  | 135            | 600                   | 50                      | 30                  | 99.9%             | Hardness = 581 ± 28 HV @ 3 wt.% CNTs |
|              |                  |      |                |                       |                         |                     |                   | UCS ⁴ = 1162 MPa    |
|              |                  |      |                |                       |                         |                     |                   | El ⁶ = 3.2%         |
|              |                  |      |                |                       |                         |                     |                   | UTS ⁷ = 1255 MPa    |
|              |                  |      |                |                       |                         |                     |                   | CS ⁸ = 23.8%        |

¹ Solution: referred to processes described in Section 3.2.1. ² COF: coefficient of friction. ³ YS: yield strength. ⁴ UCS: ultimate compressive strength. ⁵ TS: tensile strength. ⁶ El: elongation. ⁷ UTS: ultimate tensile strength. ⁸ CS: compressive strain.
Table 4. A summary table reporting L-PBF process parameters, relative density and mechanical properties of printed parts using Al-based alloys.

| Metal Powder | Coating  | Coating Technique | Ref. | Laser Power (W) | Scanning Speed (mm/s) | Hatching Distance (µm) | Layer Thickness (µm) | Relative Density | Mechanical Properties |
|--------------|----------|-------------------|------|-----------------|-----------------------|------------------------|---------------------|-------------------|------------------------|
| CNTs         | Solution | 1 [144]           | 350  | 1250            | 60                    | 30                     | 99.1%               |                   | YS = 99 MPa          |
| Ni           | Solution | 1 [125]           | 400  | 1000            | 60                    | 30                     | Near fully dense @ 0.5 wt.% Ni |                   | El 6 ~ cost          |
| Co           | Solution | 1 [126]           | 400  | 1000            | 60                    | 30                     | Near fully dense @ 0.5 wt.% Co |                   | UTS = 129 MPa         |
| SiO₂         | Turbula   | [16]              | 100  | 250             | 50                    | 30                     | 99.98% @ atm Ar     |                   |                       |
| Al and its alloys |          |                   |      |                 |                       |                        |                      |                   |                       |
| GO           | Solution | 1 [128]           | 20,6 | 10              | 100                   | 25                     | 97.36%              |                   |                       |
| Al₂O₃        | Ball milling | [22]     | 130  | 550             | -                     | 70                     | 97.3%               |                   | Wear resistance = 4.75 × 10⁻⁵ mm³/(N·m) |
| TiC          | Ball milling | [107]   | 120  | 200             | 50                    | 50                     | -                   |                   | Hardness = 188.3 HV0.1 |
| CaB₆         | Ball milling | [28]    | 200  | 1000            | 100                   | 30                     | >99.5%              |                   | TS = 486 MPa          |
| GO           | Solution | 1 [63]           | 95   | 200             | 45–105                | 30                     | 2.35 g/cm³          |                   | El 6 = 10.9%          |
| Cu           | Solution | 1 [135]          | 200  | -               | 80                    | 30                     | 99.1 ± 0.2%         |                   | YS = 132 ± 4 HV      |

1 Solution: referred to processes described in Section 3.2.1. 2 η: volumetric energy densities. 3 COF: coefficient of friction. 4 YS: yield strength. 5 TS: tensile strength. 6 El: elongation. 7 UTS: ultimate tensile strength. 8 E: Young’s modulus.
Table 5. A summary table reporting L-PBF process parameters, relative density and mechanical properties of printed parts using Cu-based alloys.

| Metal Powder | Coating | Coating Technique | Ref. | Laser Power (W) | Scanning Speed (mm/s) | Hatching Distance (µm) | Layer Thickness (µm) | Relative Density | Mechanical Properties |
|--------------|---------|-------------------|------|----------------|-----------------------|------------------------|---------------------|----------------------|------------------------|
| Nitrides,    | Thermochemical | [151] | 500 | 800 | 90 | 30 | 98.6% | YS $^2 = 183 \pm 7$ MPa | TS $^3 = 254 \pm 5$ MPa |
| carbides     |         |       |     |     |    |    |       | El $^4 = 39 \pm 2\%$ | K $^6 = 24 \pm 0.2$ IACS% | Thermal conductivity $= 112 \pm 1$ W/(m·K) |
| C            | Turbula | [19]  | 725 | 400 | 120 | 30 | 98% | Hardness $= 0.637 \pm 0.02$ GPa | TS $^3 = 125 \pm 11$ MPa |
|              |         |       |     |     |    |    |       | E $^5 = 105 \pm 2$ GPa | Proof (yield) strength $= 64 \pm 7$ MPa |
| Cu and its   | Mixing + heat treatment | [152] | 500 | 700 | 90 | 30 | 96.1% @ 400 W and 200 m/s | | |
| alloys       |         |       |     |     |    |    |       | | |
| CrZr         | PVD     | [20]  | 270 | 300 | 90 | -  | 94.3% | | |
| Sn           | Solution $^1$ | [17]  | 500 | 600 | 105 | 30 | 99.6% | YS $^2 = 203 \pm 4$ MPa | TS $^3 = 256 \pm 14$ MPa |
| Ni/Sn        | Solution $^1$ | [21]  | 200 | 100 | 100 | 20 | -     | YS $^2 = 203 \pm 4$ MPa | TS $^3 = 256 \pm 14$ MPa |

$^1$ Solution: referred to processes described in Section 3.2.1. $^2$ YS: yield strength. $^3$ TS: tensile strength. $^4$ El: elongation. $^5$ E: Young’s modulus. $^6$ K: electrical conductivity.
5. Conclusions

Powders coating processes are of increasing interest; the possible applications of coated metal powders are wide and involve many aspects such as processability, mechanical, and physical properties. Different ways of coating are possible. The so-called mechanical coating techniques were among the first adopted. However, when high energy is involved, a severe plastic deformation occurs, modifying powders’ shape and impairing the powders’ flowability. On the other hand, coating processes involving chemical solutions do not affect the geometry and size distribution of the powders, usually resulting in homogeneous and continuous coatings. These processes are characterized by different difficulties such as the limited quantity of powder that can be treated together, the costs and the complexity of the process itself; for these reasons it is mandatory to evaluate new techniques not currently in the market in order to obtain constant and homogenous coating which can provide higher processability.

Coated powders can improve L-PBF processability, in terms of higher flowability and absorbance, especially for those metals characterized by a high laser reflectivity. The surface modification of powder can pursue a higher laser absorbance. Two main factors are involved: the surface roughness, which is usually increased by coatings, and the choice of coating material with a high absorbance coefficient. Several parameters influence flowability as well. A higher surface roughness can negatively affect it, whereas ceramic nanoparticles can increase it, due to the intrinsic low van der Waals forces.

Mechanical properties of the final parts can also be improved by processing suitably coated powders. Hardness can increase by using ceramic coating particles or by the grain refining effect. Using metal powders homogeneously coated by nanoparticles may result in an improvement of tensile strength and hardness without sacrificing ductility.

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