Characterization of AlCoCrFeNi High Entropy Alloy Gas Atomized Powder

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Received: March 9, 2021; Revised: July 9, 2021; Accepted: October 18, 2021

AlCoCrFeNi alloys are widely studied due to the combination of high strength in a wide temperature range, good corrosion and wear resistance, but they typically present low ductility. Due to this limited ductility, traditional bulk fabrication processes that involve forming are difficult, and casting processes are also non-ideal due to pronounced segregation and coarse microstructure. Therefore, both the selective laser melting and the powder metallurgy are more promising routes for this alloy, both of these use powder as starting material. This work aimed to produce powder of Al20Co20Cr20Fe20Ni20 alloy by gas atomization and to perform its detailed characterization. The results show that the powder is mainly spherical, which resulted in good flowability. Most of the powder (62%) has a size smaller than 75µm, and microscopy analyses confirmed that a B2 dendritic microstructure was formed, with A2 cuboidal precipitates.

Keywords: High entropy alloys, gas atomization, powder metallurgy.

1. Introduction

High entropy alloys (HEAs), also named complex concentrated alloys (CCAs), or multi-principal elements (MPEAs) and multi-component alloys, have been developed over the last decade1-3. In contrast to conventional alloys that contain one or two main solvent elements, these alloys are composed of multiple alloying elements whose concentrations typically range from 5 to 35 atomic percent4-5. Among HEAs systems, AlCoCrFeNi is one of the most popular, owing to their relatively low-cost elements, coupled with the wide range of possible mechanical properties such as high yield strength at room and elevated temperatures, good wear behavior, and excellent corrosion resistance6.

Depending on their composition and processing history, their microstructure can be composed of FCC (A1) and/or BCC (A2) solid solutions together with their ordered versions (L12 and B2, respectively) and in some cases also sigma phase7. The microstructure of equiatomic AlCoCrFeNi is mostly composed of a B2/A2 mixture after solidification under several different reported cooling rates8-9.

Despite the aforementioned properties, the AlCoCrFeNi equiatomic HEA alloys exhibit poor ductility, which does not allow their processing by traditional thermomechanical processes (i.e., hot forging and/or hot rolling). The most used casting processes, such as vacuum arc melting and drop-casting, apply cooling rates of around 100 K/s or below, leading to undesirable segregation9.

In this context, some conventional consolidation methods of Powder Metallurgy, such as powder extrusion, spark plasma sintering (SPS) and hot isostatic pressing (HIP), are favorable alternatives to obtain bulk HEA material10. Indeed, there are multiple literature reports of HEA powders produced by mechanical alloying (MA) followed by conventional consolidation processes, resulting in better properties than as-cast materials. Nevertheless, during the MA process, the powders may be contaminated by the milling medium and the environment, affecting the mechanical properties11.

Hence, powders produced either by gas atomization (GA)12 or by plasma spheroidization13 if on the correct particle size and morphology, are perhaps even more suitable routes to produce such powders. The gas atomization is more common comparing these two methods and presents higher powder productivity. It is a rapid solidification process that involves the disintegration of a stream of liquid metal into droplets of different sizes through a jet of inert gas14-16. Many variables control the production of powders via GA, such as overheating temperature, gas composition, gas pressure, and GMR ratio, i.e., the ratio of gas mass flow rate (G) to metal mass flow rate (M), but the last two are the most important processing parameters17. Typically, increasing gas pressure reduces the median particle size (D50) due to increased gas velocity for a given nozzle. In other words, the particles’ size reduces as the relative velocity between the droplets and gas rises17. A high GMR ratio results in a high velocity of the atomized droplets, which in turn generates
high heat losses by convection to the atomization gas and also very high cooling rates (10⁶ - 10⁶ K/s), yielding a refined microstructure, metastable phases, and little segregation. This is an important advantage since some HEAs might require very long homogenization times, such as 20-60 hours at high temperatures.

The available studies on equiatomic AlCoCrFeNi powders are focused on: i) the effect of heat treatment on the microstructural evolution and the mechanical properties of AlCoCrFeNi powder; ii) analysis of the microstructural evolution and mechanical and corrosion behavior of AlCoCrFeNi alloy produced by spark plasma sintering, and iii) characterization of AlCoCrFeNi atomized and annealed powders for plasma spray coatings.

However, a detailed investigation of physical characteristics and microstructural evolution of AlCoCrFeNi powder under strictly controlled processing parameters has not been reported so far. Moreover, the phase formation in these rapidly solidified powders is also not well clarified yet either, and careful and in-depth microscopy analyses are necessary to evaluate the formation of B2, L12, A1, and/or A2 phases under these specific solidification conditions.

Thus, this paper’s noteworthy contribution was producing AlCoCrFeNi powders by gas atomization, ensuring their appropriate characteristics and detailed microstructural evolution for conventional consolidation processes.

2. Materials and Methods

AlCoCrFeNi ingots were prepared from elements with purity above 99.5% using an induction furnace Inductotherm 50-30R. After, these ingots were atomized in a PSI – HERMIGA under a gas atomization pressure of 60 bar, nozzle diameter of 2.5 mm, gas-metal ratio of 5.9 m³/Kg, and an atomization temperature of 1487°C, which was around 100°C above the alloys liquidus temperature. These parameters were established in order to obtain spherical and fine powders. When the atomization process was completed, the powders were cooled down to room temperature in an argon atmosphere. In order to characterize the resulting powders, they were sieved into seven particle size ranges: greater than 180, 106-180, 75-106, 53-75, 32-53, 20-32µm, and below 20µm. Also, the flowability behavior was measured using Hall Funnel according to ASTM B213 Standard. The average value and standard deviation were calculated based on seven measurements. The true powder density was obtained with Accupyc II 1330 gas displacement pycnometry system, using the process described at ASTM B 293 standard. To ensure measurement accuracy, five measurements were carried out.

Microstructural characterization was carried out by X-ray diffraction (XRD) using Rigaku Geiger-Flex X-ray diffractometer with Cu-Kα radiation (0.15406 nm). Scanning and transmission electron microscopy (SEM and TEM) analysis were also carried out using, respectively, a Philip XL-30 FEG SEM, operating at 30kV or below, and an FEI Talos F200X TEM, operating at 200kV. Both microscopes are coupled with energy-dispersive X-ray spectroscopy (EDX) system for chemical analysis.

The TEM samples were prepared by argon ion milling in a Gatan Precision Ion Polishing System (PIPS). Only splats of powders with a size greater than 180 microns were analyzed because it was not possible to prepare good-quality samples of powder with smaller particle sizes.

To investigate the microstructure of atomized AlCoCrFeNi powders, the cross-sections were prepared using standard metallographic procedure finishing with alumina 1µm polishing. The polished samples were etched with aqua regia (1:3 vol mixture of nitric acid and hydrochloric acid) for 3 minutes.

3. Results and Discussion

The powder’s chemical composition by EDX/SEM is presented in Table 1, showing that the elements’ uniform distribution is close to its nominal equiatomic composition. Figure 1 displays the particle size distribution by both mass and accumulated mass percentages. About 62% of the particles had a diameter smaller than 75µm, and the mean mass diameter ($D_3,2$) is 63µm.

SEM micrographs of the atomized powders in the range of 20 - 180µm are shown in Figure 2. Distinct morphologies are seen among different granulometric ranges. Smaller powders present more spherical morphology due to the higher cooling rate, which is inversely proportional to the powder size, in agreement with the observation of Mazzer et al.

The spherical shape leads to better flowability and packing density. On the other hand, larger powder sizes usually have more irregular morphologies and more satellite particles, which may compromise the Hall Flow rate. The flowability of powders in the range 20-75 µm showed a good flow behavior (19.3 ± 0.3 s/50g) compared to commonly used alloys Ti-6Al-4V, IN718, and AlSi10Mg: 32.4, 28.3, and 62.0 s/50g, respectively. The density of the atomized powder was calculated by gas displacement pycnometry using the AccuPyc II 1330 machine.
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was 7.11 ± 0.01 g/cm³, which is below the theoretical value (8.0 g/cm³)²².

Although the XRD pattern result (Figures 3a, b) exhibit pronounced peaks of the A2 phase, there are (100) and (111) weak reflections of the B2 phase in accordance with the literature¹⁹,²³. For example, Cheng et al.²³ stated that their powders (Al₂₀.₆Co₂₀.₁Cr₂₀.₄Fe₁₉.₄Ni₁₉.₄) consisted of a major amount of A2 phase and also B2 phase in a minor amount with the characteristic reflection (100).

It is known that A2 and B2 phases formed together may have similar lattice parameter, especially on this alloy, in which both these phases are formed by a multi-step decomposition from a high-temperature A2 phase into two nanometric phases: another A2 phase enriched Fe-Cr plus an Al-Co-Ni rich B2⁷,⁹. The B2 superlattice reflections are not very intense, given the relative proximity of this system’s elements in the periodic table, which yields a small structure factor for these extra reflections. Therefore, it is necessary to perform the further investigation by TEM to confirm the presence, morphology, and fraction of the A2 and B2 phases²⁴.

As shown by the dark field image (DF), the microstructure of the splat is composed of fine precipitates of A2 phase into B2 matrix (Figure 4a). The selected area electron diffraction pattern (SAEDP) indicated in Figure 4b taken from [0 0 1] zone axis confirmed the presence of the two phases⁷–⁹,²⁴.

Although TEM analysis was performed on the splats produced at a cooling rate of 10⁵ K/s slightly lower when compared to 10⁵–10⁶ K/s for 20-120 powders size range¹⁷,²⁵, it is still a high value. This allowed noting the presence of A2-rich precipitates and bands into the B2 matrix (Figure 4a). In addition, the A2/B2 phases can be seen in XRD pattern in all particle ranges (Figures 3a, b). Therefore, given the similarity of the final microstructure (Figure 4a) and its pattern diffraction (Figure 4b), which enabled us to consider the solid-state transformation path for such alloy which is described as follows. According to recent Monte Carlo simulations⁷ at high temperatures (above 800°C), the equiatomic AlCoCrFeNi alloy most likely presents only the A2 phase. At approximately 800°C, the alloy transforms into a partially B2 ordered phase. Upon further cooling, the partially ordered phase (B2) transforms into a mixture of

Figure 2. Secondary electron (SE) images showing the spherical and irregular morphology of the powders.
another disordered Cr-Fe enriched A2 and ordered AlCoNi enriched B2 phases by short atomic motions. This phenomenon can be explained by the dominant role of aluminum, which has a very negative bonding energy with the other elements in this alloy, explaining the first B2 ordering followed by decomposition into another a with nickel and cobalt-rich B2 phase. This configuration is energetically favored at lower temperatures since it promotes a larger number of pairs of Co/Ni and Al demonstrated by the very negative interaction energies calculated (-7.58 and -8.15 KJ/mol, respectively), in
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contrast to the lower interaction energies between other pairs such as Co-Cr, Co-Fe, Co-Ni, Cr-Fe, Cr-Ni and Fe-Ni (0.06; 0.72; -0.25; -0.10; -0.36 and -0.07 KJ/mol, respectively)\textsuperscript{7}. This is further evidenced by a higher Al-containing material, the Al\textsubscript{33}Co\textsubscript{16.7}Cr\textsubscript{16.7}Fe\textsubscript{16.7}Ni\textsubscript{16.7} alloy, which does not exhibit the A2 phase at all since the ordering temperature (Tc) would be higher than the liquidus temperature\textsuperscript{7}.

Based on this, it is clear that such decomposition’s kinetics are very fast, and increasing the cooling rate during solidification would not change the final morphology since this is a pure solid-state transformation. However, it is expected that smaller particles would also experience a higher cooling rate in the solid-state compared to bigger particles like splats. In this case, considering the experimental evidence displayed in Figure 3 where we clearly see the presence of the same phases for all powder size ranges, the A2/B2 domains are expected to be increasingly more refined for finer particle sizes, which frequently occurs in the similar systems\textsuperscript{25}.

In short, the order-disorder transition cannot be observed experimentally step by step. Therefore, we suggest this decomposition kinetics as the most plausible explanation for the resultant morphology.

It can be seen in Figure 5a the microstructure of the powder in the range of 20-75 µm with secondary dendrite arm spacings of about 1.8 µm, 3.9 µm, 5.4 µm, and 7.0 µm for the particle size of 20 µm, 32 µm, 53 µm, and 75 µm, respectively. This result highlights the close relationship between secondary dendritic arm spacings and the cooling rates/size powders as mentioned in the literature\textsuperscript{17}.

Although it is expected to observe more segregation with increasing powder size, a relatively homogeneous elemental distribution was found in the different powders sizes, with no evidence of chemical segregation within the EDX mapping detection limits (Figure 5b). It should be highlighted that the same alloy produced by arc melting, which involves much slower cooling rates, the chemical segregation was also small, as reported by Ferrari et al.\textsuperscript{24}.

In short, the gas atomization process was a promising route in producing spherical particles with the homogeneous chemical composition of the equiatomic AlCoCrFeNi alloy powder, and it will probably be to other alloys with high aluminum content.

4. Conclusions

In this work, AlCoCrFeNi powder was produced by gas atomization. The physical properties of the powder and its resulting microstructure were analysed. From the results, the following conclusions were drawn:

- AlCoCrFeNi powder was successfully produced through inert gas atomization, with 62% of the powder smaller than 75µm.
• Gas atomized powders below 75µm were mostly spherical and show a small number of satellite particles, with 63µm of mean diameter and high flowability.
• The XRD and TEM analyses indicated the formation of dendrites composed of an A2/B2 nanometric mixture and the secondary dendrite arm spacings were 1.8µm, 3.9µm, 5.4 µm and 7.0µm for particle sizes 20µm, 32µm, 53µm and 75µm, respectively.
• These results show that the powder obtained presents adequate properties for powder metallurgy processes.

5. Acknowledgments

The authors would like to thank the Brazilian research funding agencies – This study was financed in part by the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior- Brasil (CAPES)- Finance code 001, São Paulo Foundation Research – FAPESP (Grant 2017/27031-4) and National Council of Technological and Scientific Development – Brazil / CNPQ (Grant 167845/2018-7). Additionally, the Laboratory of Structural Characterization of Department of Materials Engineering at the Federal University of São Carlos (LCE/DEMA/UFSCar) for the microscopy facilities.

6. References

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