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Fabrication and Characterization of Al_xCrCuFeNi₂ High-Entropy Alloys Coatings by Laser Metal Deposition

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Abstract

High-entropy alloys (HEAs) are becoming new hot spots in the metallic materials community, which are defined to contain equiatomic or close-to-equiatomic compositions. HEAs can possess many interesting mechanical properties, and in particular, they have the great potential to be used as coating materials requiring high hardness and wear resistance. In this study, the feasibility of fabrication $Al_xCrCuFeNi_2$ (x=0,0.75) HEAs was investigated via laser metal deposition from elemental powders. The microstructure, phase structure, and hardness were studied by an optical microscope, scanning electron microscopy with energy dispersive spectroscopy (SEM/EDS), electron backscatter diffraction (EBSD) and Vickers hardness tester. The bonding between the $Al_xCrCuFeNi_2$ (x = 0,0.75) HEAs and AISI 304 stainless steel were good combinations. The $Al_{0.75}CrCrFeNi_2$ alloy consisted of columnar dendritic microstructure with Al/Ni enrichment in the dendritic regions. The phase structure of the $Al_xCrCuFeNi_2$ (x = 0,0.75) HEAs were face center cubic structure as identified by EBSD. Vickers hardness results indicate that the average hardness of CrCuFeNi_2 HEA was 175 HV. With the addition of aluminium, the Vickers hardness of $Al_{0.75}CrCuFeNi_2$ HEA increased to 285 HV.

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Keywords: High-entropy alloys; additive manufacturing; laser metal deposition; microstructure; elemental powder

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1. Introduction

Conventional metallurgical theory suggests that the multiple alloying elements in an alloy may result in the formation of complex compounds. Recently this paradigm has been broken by high-entropy alloys (HEAs) developed by Yeh et al.[1] HEAs are composed of five or more principle elements in equimolar or near-equimolar ratios. The high mixing entropy of multi-principle elements induces the formation of solid-solution structure, e.g., face center cubic (FCC) or body center cubic (BCC) or FCC combined with BCC [1–6]. The discovery of HEAs has brought a new alloy design concept and generated researchers' interest in the past decade. An AlCrFeCoNi HEA was prepared by vacuum arc melting and exhibited excellent compressive strength 2004.23 MPa [2]. Another study of AlCrFeCuNi_x ($0.6 \le x \le 1.4$) HEAs was prepared by casting reported by Jinhong et al., which found the hardness of as-cast HEAs decreased as x increased from 1.0 to 1.4 [3]. Dong et al. investigated the AlCrFeNiMo_x (x = 0,0.2,0.5,0.8 and 1.0 in molar ratios) HEAs produced by vacuum melting [7]. Their work showed AlCrFeNiMo_{0.2} HEA possessed good fracture strength of 3222 MPa and plastic strain of 0.287, which implies its potential application in industrial areas. These HEAs were fabricated by casting or vacuum melting. Unlike the previous studies, this work will implement the laser metal deposition (LMD) method to fabricate the Al_xCrCuFeNi₂ (x = 0, 0.75 in molar ratios) HEAs.

As an advanced additive manufacturing technology, LMD can accomplish layer-by-layer fabrication of near netshaped components by introducing a powder stream through a high energy laser beam [4,5,8–10]. A melt pool is formed by rastering the laser beam, and the powders are injected into the melt pool to deposit each layer during the LMD process. Layer by layer composition changes, the introduction of a dissimilar metal interlayer and control over the melt zone size can be accommodated [4,9–13]. A FeCoNiCrCu HEA coating was synthesized, and its microhardness reached 375 HV_{0.5}, which was about 50% higher than that of the same alloy prepared by arc melting [4]. With the additional of titanium content, Al₂CrFeNiCoCuTi_x (x = 0, 0.5, 1.0, 1.5 and 2.0 in molar ratios) HEAs showed good corrosion and wear resistance on Q235 steel substrate [10]. Few research has been devoted to the fabrication of AlCrCuFeNi₂ HEAs by LMD.

In this paper, the feasibility of fabrication $Al_xCrCuFeNi_2$ (x = 0, 0.75 in molar ratios) HEA coatings on AISI 304 stainless steel (SS) was performed by laser metal deposition technology using elemental powders. The metallurgical bonding, microstructure, and Vickers hardness were investigated.

2. Experimental

2.1. LMD processing

Gas-atomized elemental powders of aluminium (Al), chromium (Cr), copper (Cu), iron (Fe) and nickel (Ni) purchased from Atlantic Equipment Engineers Inc. was used as precursor materials. The particle size of the elemental powders provided by Atlantic Equipment Inc. is as tabulated in Table 1. The elemental powders were weighted in the required ratios and then mixed by a Turbula mixer (Glen Mills Inc., Clifton, NJ, USA) for 30 mins to obtain homogeneous blends. Elemental compositions (atomic %) of the as-blended HEAs are given in Table 2.

Materials	US Standard Mesh				
Al	-100				
Cr	-100				
Cu	-100				
Fe	-100				
Ni	-100/+325				

Table 1. Particle size distribution of the elemental powders.

Table 2. Nominal compositions (atomic %) of HEAs.

Alloys	Al	Cr	Cu	Fe	Ni
CrCuFeNi ₂	0	20	20	20	40
Al _{0.75} CrCuFeNi ₂	13	17	17	17	36

The schematic of the LMD system is shown in Figure 1. The 1 kW continuous-wave YAG fiber laser (IPG, Photonics, Oxford, MA, USA) was used as a heat source with a beam diameter of 2 mm. The metallic powders were fed through a vibration X2 powder feed system (Powder Motion Labs, MO, USA). The powders were introduced into the melt pool by an alumina tube. Argon gas was used as a carrier gas to deliver the powder mixtures to the melt pool. The movement during the laser deposition was achieved through a computer numerical control (CNC) table.



Fig. 1. Schematic of the laser metal deposition (LMD) system.

Commercially procured AISI 304 SS bar stock (dimension: 2 inch \times 2 inch \times 0.25 inch) was used as the substrate and cleaned with acetone to clean the surface. A preheating scan was performed by running the laser across the substrate. The thin wall structure was built, and the laser power of the initial three layers was conducted at 700 W and 8.5% (3.36 g/min) powder feed rate. The remaining of the deposition was carried out at 600 W and 8.5% (3.36 g/min) powder feed rate with 1 mm layer thickness.

2.2. Characterization

For microstructural characterization, the deposits were transverse cross-sectioned and prepared with standard metallographic methods. The samples were polished with 320-1200 grit SiC grinding paper, and the final mechanical polish was 0.05 µm silica suspension. The specimens were given the electrolytic etching in the nitric acid solution (70 mL nitric acid and 30 mL distilled water).

A Hirox optical microscope investigated the $Al_xCrCuFeNi_2$ HEAs morphology. The scanning electron microscopy (SEM), elemental analysis using energy dispersive spectroscopy (EDS), and electron backscatter diffraction (EBSD) studies of the specimens were carried out in a Helios Nanolab 600 SEM coupled with an EDS and an EBSD detector. The obtained EBSD data was processed and analyzed using Aztec software. The hardness was obtained with a Struers Duramin hardness tester (Struers Inc., Cleveland, OH, USA) using a 9.81 N force and 10 s load duration.

3. Results and discussion

3.1. Microstructure

Figure 2 shows the optical images of the deposited HEAs. The deposit was shown in the top area in Figure 2a while the bottom part was AISI 304 SS substrate. An explicit interface was seen between the deposit and the AISI 304 SS substrate. The columnar dendrite microstructure was observed from Figure 2b. Similarly, a good metallurgical bonding existed between $Al_{0.75}CrCuFeNi_2$ HEA and the AISI 304 SS substrate. The dendritic continued in $Al_{0.75}CrCuFeNi_2$ alloy. The growth direction of these columnar was identified to be along with the deposition direction, which could be correlated with the solidification direction during the LMD process.



Fig. 2. Optical images of (a) the interface between CrCuFeNi₂ HEA and AISI 304 SS substrate, (b) microstructure of CrCuFeNi₂ HEA, (c) the interface between Al_{0.75}CrCuFeNi₂ HEA and AISI 304 SS substrate and (d) microstructure of Al_{0.75}CrCuFeNi₂ HEA.

3.2. EDS and EBSD analysis

The evolution in chemistry from the substrate to the CrCuFeNi₂ HEA was characterized by EDS line scan first. The quantitative results are plotted in Figure 3a. The results measured by EDS of the AISI 304 SS substrate (Cr: ~18-19 atomic %, Fe: ~70-71 atomic %, Ni: ~9-10 atomic % in Figure 3a) did not derive from the nominal AISI 304 SS chemical compositions. Mn (~ 1-2 atomic %) was detected by EDS in AISI 304 SS substrate but was not shown in Figure 3. The elemental compositions of Cu (~18-21 atomic %) and Ni (~35-38 atomic %) increased while that of Fe (~23-26 atomic %) reduced, and Cr (~20 atomic %) remained changed from the substrate to the CrCuFeNi₂ HEA deposit. A small amount of Cu (~1-2 atomic %) was detected in the substrate because the substrate was mixed with the HEA deposit. The distribution of the consisted compositions from the substrate to the Al_{0.75}CrCuFeNi₂ HEA was shown in Figure 3b. The constituents of the Al_{0.75}CrCuFeNi₂ HEA were determined by EDS (Al: ~9-10 atomic %, Cr: ~19 atomic %), Cu: ~17 atomic %, Fe: ~20 atomic %, Ni: ~ 32-34 atomic %). The difference between the as-blended (13 atomic %) and as-deposited aluminium (~9-10 atomic %) compositions could be attributed to the inconsistency of powder capture efficiency and evaporation due to its low melting point.



Fig. 3. Elemental composition evolution (a) the interface between AISI 304 SS substrate and CrCuFeNi₂ HEA, (b) the interface from AISI 304 SS substrate to Al_{0.75}CrCuFeNi₂ HEA.

EBSD and EDS measurements were conducted in the aim of differentiating structure and phase information of the $Al_xCrCuFeNi_2$ (x = 0, 0.75) alloys. Regions of interest and phase analysis of CrCuFeNi_2 and $Al_{0.75}CrCuFeNi_2$ alloys are shown in Figures 4 and 5 respectively. Figures 4 and 5 indicate an FCC structure in both HEA fabrications. The phase fractions and the corresponding lattice parameter identified by EBSD are listed in Table 3. The zero solution is the fraction of the selected area whose crystal structure could not be solved by the software.

Alloy	Phase Name	Space Group	Lattice Parameter (Å)	Fraction (%)
CrCuFeNi ₂	FCC	Fm-3m (225)	3.66	88.2
	BCC	Im-3m (229)	2.93	0.13
	Zero solution	-	-	11.67
Al _{0.75} CrCuFeNi ₂	FCC	Fm-3m (225)	3.66	99.3
	BCC	Im-3m (229)	2.93	0.09
	Zero solution	-	-	0.61

Table 3 Summary of the lattice parameter and phase fraction (%) of the Al₃CrCuFeNi₂ HEAs obtained from EBSD analysis.



Fig.4. EBSD phase map indicates predominating FCC phase in the CrCuFeNi₂ alloy. (a) The region of interest on CrCrFeNi₂ alloy (b) Phase map shows predominantly FCC phase (represented by blue color) within the region of interest.



Fig.5. EBSD phase map indicates predominating FCC phase in the $Al_{0.75}$ CrCuFeNi₂ alloy. (a) The region of interest on $Al_{0.75}$ CrCuFeNi₂ alloy, (b) Phase map shows predominantly FCC phase (represented by blue color) within the region of interest.

Figures 6 and 7 show the EDS element maps obtained from the Al_xCrCuFeNi₂ HEAs. The Fe-K α , Cr-K α , Ni-K α , Cu-K α and Al-K α signals were used to estimate the elemental compositions within the regions of interest in the deposits. EDS elemental compositions were gathered from the dendritic microstructures for the Al_xCrCuFeNi₂ (x = 0, 0.75) HEAs. The standardless measurements are listed in Table 4. The microstructure of CrCuFeNi₂ alloy exhibited a dendritic microstructure as reported previously. Based on the previous EBSD phase analysis, this dendritic phase was likely to be a single FCC phase. While it was observed a distinct contrast between the dendritic and interdendritic regions (as seen in Figure 6), this contrast could be attributed to the segregation of Cu (which tended to partition and segregate readily [14,15]) as in Figure 6e. Table 4 shows that Cu was enriched in the interdendritic regions. Figure 7 indicates that with the addition of aluminium, Al_{0.75}CrCuFeNi₂ alloy contained predominantly two phases. Associated with the results from Table 4, the dendritic phase was observed to be Al and Ni rich (Al+Ni: ~41 atomic %, Fe+Cr: ~18 atomic %), while the interdendritic microstructure was rich in Fe and Cr (Al+Ni: ~28 atomic %, Fe+Cr: ~52 atomic %). The Cu was deficient in the interdendritic regions.



Fig.6. EDS elemental maps of CrCuFeNi₂ alloy, (a) region of interest, (b) element map of Fe, (c) element map of Cr, (d) element map of Ni and (e) element map of Cu.



Fig.7. EDS elemental maps of $Al_{0.75}$ CrCuFeNi₂ alloy, (a) region of interest, (b) element map of Fe, (c) element map of Cr, (d) element map of Ni, (e) element map of Cu and (f) element map of Al.

Area	Al	Cr	Cu	Fe	Ni
Nominal	0	20	20	20	40
Dendritic	0	18.5	18.7	27.1	35.7
Interdendritic	0	16.9	25.9	21.4	35.8
Nominal	13	17	17	17	36
Dendritic	15	7.3	41.4	10.3	26
Interdendritic	8	20	24	28	20
	Area Nominal Dendritic Interdendritic Nominal Dendritic Interdendritic	AreaAlNominal0Dendritic0Interdendritic0Nominal13Dendritic15Interdendritic8	AreaAlCrNominal020Dendritic018.5Interdendritic016.9Nominal1317Dendritic157.3Interdendritic820	Area Al Cr Cu Nominal 0 20 20 Dendritic 0 18.5 18.7 Interdendritic 0 16.9 25.9 Nominal 13 17 17 Dendritic 15 7.3 41.4 Interdendritic 8 20 24	AreaAlCrCuFeNominal0202020Dendritic018.518.727.1Interdendritic016.925.921.4Nominal13171717Dendritic157.341.410.3Interdendritic8202428

Table 4 Elemental compositions of the elements at different regions in atomic % for the CrCuFeNi₂ and $Al_{0.75}CrCrFeNi_2$ alloys.

From the mixing enthalpy of atom-pair as listed in Table 5, it clearly shows that the mixing of enthalpy of Al and Ni is higher (-22 kJ/mol) than of other atom-pair. It indicates that Al and Ni atoms tend to form atom pairs and segregate. Similar results have been reported previously, with this microstructure being attributed to the spinodal decomposition [2,6,16–18].

Table 5 Mixing enthalpy of different atom-pair in the CrCuFeNi2 and Al_{0.75}CrCrFeNi2 alloys [19].

ΔHmix (kJ/mol)	Cu	Cr	Al	Ni
Fe	13	-1	-11	-2
Cu	-	12	-1	4
Cr	-	-	-10	-7
Al	-	-	-	-22

3.3. Vickers hardness

Figure 8 gives the Vickers hardness profiles of the $Al_xCrCuFeNi_2$ (x = 0, 0.75) alloys deposits on the AISI 304 SS substrates. The Vickers hardness of CrCuFeNi_2 alloy was around 175 HV, which could be attributed to the solid solution strengthening. Table 6 gives the Vickers hardness of various alloys, including AISI 304 SS, Inconel 625 and 7075-T6 aluminium [20,21], and $Al_{0.75}CrCuFeNi_2$ alloy has the highest average hardness of 285 HV. With the addition of aluminium, the average Vickers hardness of $Al_{0.75}CrCuFeNi_2$ HEA reached 285 HV, because the second phase strengthening blocked the dislocation [6,18]. The high hardness of $Al_{0.75}CrCuFeNi_2$ HEA coating is expected to correlate with good performance in strength and wear resistance [10,13].



Figure 8 Vickers hardness profiles of the $Al_xCrCuFeNi_2$ (x = 0,0.75) alloys.

Allov Hardness (HV) Reference CrCuFeNi₂ 175 This work Al_{0.75}CrCuFeNi₂ 285 This work AISI 304 SS 160 This work Inconel 625 156 [21] 7075-T6 aluminium 118 [20]

Table 6 Vickers hardness of various alloys.

4. Conclusions

 $Al_xCrCuFeNi_2$ (x = 0,0.75 in molar ratios) HEAs were coated on AISI 304 stainless steel substrate via laser metal deposition technology. The metallurgical bonding, microstructure, and the Vickers hardness were investigated and discussed. The good metallurgical bonding was observed between the HEA coatings and the substrate. The $Al_xCrCuFeNi_2$ (x = 0,0.75) HEAs coating exhibited columnar dendritic microstructure and FCC structure identified by EBSD. CrCuFeNi_2 HEA was found to have an average hardness of 175 HV, while $Al_{0.75}CrCuFeNi_2$ HEA has a hardness of 285 HV.

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