Tribo-Corrosion Response of Additively Manufactured High-Entropy Alloy

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Tribo-corrosion response of additively manufactured high-entropy alloy

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High-entropy alloys (HEAs) with multiple principal elements represent a paradigm shift in structural alloy design and show excellent surface degradation resistance in corrosive environment. Here, the tribo-corrosion response of laser-engineered net-shaped CoCrFeMnNi HEA was evaluated in 3.5 wt% NaCl solution at room temperature. The additively manufactured (AM-ed) CoCrFeMnNi showed five times lower wear rate, regenerative passivation, and nobler corrosion potential during tribo-corrosion test compared to its arc-melted counterpart. A significant anisotropy was seen in the tribo-corrosion response with 45° to the build direction showing better performance compared to tests along the build direction and perpendicular to it. The open circuit potential curves were characterized by a sharp drop to more negative values as wear began, followed by continuous change for the active tribo-corrosion duration and finally a jump to nobler value at the end of the test indicating excellent surface re-passivation for the AM-ed alloy. The superior tribo-corrosion resistance of AM-ed CoCrFeMnNi was attributed to the refined microstructure and highly protective surface passivation layer promoted by the sub-grain cellular structure formed during additive manufacturing. These results highlight the potential of utilizing additive manufacturing of HEAs for use in extreme environments that require a combination of tribo-corrosion resistance, mechanical durability, extended service life, and net shaping with low dimensional tolerance.

**INTRODUCTION**

High-entropy alloys (HEAs), also known as complex concentrated alloys, represent a paradigm shift in alloy design strategy that demonstrate excellent mechanical properties and surface degradation resistance. There remains a tremendous interest in advanced manufacturing and processing of different HEA systems due to the tunability of properties for a wide range of structural and functional applications. Laser-engineered net shaping (LENS) has proven to be a versatile manufacturing route for fabrication of complex geometries with enhanced microstructural features that are difficult to obtain by conventional processing. The desirable attributes of LENS over other additive manufacturing (AM) techniques include rapid deposition time, microstructural refinement, and the capability for obtaining functionally graded and layered materials for tailored applications. The inherent localized heating and rapid cooling during AM tends to suppress elemental segregation and phase separation typically seen in multi-component alloys synthesized using conventional routes. There are several reports on AM of HEAs that include laser powder bed fusion (also called selective laser melting) as well as LENS. However, there are very few reports and limited understanding of the degradation mechanisms in additively manufactured (AM-ed) HEAs in terms of their wear, corrosion, and tribo-corrosion behavior.

Tribo-corrosion involves the synergistic combination of wear and corrosion that leads to accelerated material degradation. Tribo-corrosion is commonly seen in many applications including bio-implants, hydraulic systems, nuclear power plants, chemical reactors, marine industries, and deep-sea oil drilling rigs and pumps. HEAs offer a wide range of tunability of their surface properties due to the multi-principal alloying approach, which makes them appealing for tribo-corrosion applications. Prior studies on AM of HEAs have shown desirable mechanical properties, fine cellular microstructure, and excellent wear and corrosion behavior. However, there are no reports so far on the tribo-corrosion behavior of AM-ed HEAs and the associated structure–property–performance relationships.

Here, tribo-corrosion behavior of AM-ed (via LENS) CoCrFeMnNi HEA was studied and compared with its counterpart synthesized by conventional vacuum arc melt casting. The equimolar CoCrFeMnNi alloy was chosen because it is one of the most widely studied HEAs, with numerous reports on microstructural characterization, mechanical behavior, corrosion, and wear. There are no reports on tribo-corrosion degradation of this HEA. In addition, there is limited understanding of the effect of AM microstructure on wear-corrosion synergy. Here, we evaluated the effect of AM build orientation on tribo-corrosion response. The correlation between coefficient of friction (COF) and open circuit potential (OCP) was studied by our experimental setup to explain the tribo-corrosion degradation behavior of AM-ed versus as-cast CoCrFeMnNi HEA.

**RESULTS AND DISCUSSION**

**Microstructural characterization**

Near-full density was obtained for the AM-ed CoCrFeMnNi alloy under optimized processing conditions. The corresponding microstructural characterization are summarized in Fig. 1. Scanning electron microscopy (SEM) image in Figure 1a shows some spherical micro-scale pores (indicated by arrows) of size less than 10 μm. These pores likely originated from the powder feedstock and evaporation of constituent elements as well as gas entrapment from the melt pool turbulence. Figure 1b shows...
Fig. 1  Microstructural analysis. a SEM image of laser-engineered net-shaped CoCrFeMnNi alloy showing near-full density microstructure with some micropores indicated by arrows. b EBSD IPF map along build direction showing columnar grains. c SEM image of as-cast CoCrFeMnNi alloy with columnar grains. d Optical microscopy (OM) image of the longitudinal section of the LENS sample. e Magnified image of the square box area in (d) showing melt pool, columnar grains, and sub-grain cells. f SEM image showing transverse section of the sub-grain cellular structures. g SEM image showing the longitudinal section of the sub-grain cellular structures. h High-magnification IPF image showing local misorientation within primary grains and the inset showing misorientation profile along the arrow. i Corresponding kernel average misorientation (KAM) map indicating dislocation arrays along the cell boundaries. j X-ray diffraction patterns showing single-phase FCC structure for AM-ed and as-cast CoCrFeMnNi.
the inverse pole figure (IPF) map of the AM-ed sample along the build direction (BD) with columnar grains of average size \( \sim 36.4 \pm 23.4 \mu m \). The AM melt pool, semi-circular in shape, had a diameter-to-depth ratio of 5:2 and overlap rate of 60% with the adjacent pool as shown in Fig. 1c. Elongated columnar grains growing towards the melt pool center were formed near the melt pool boundary due to large thermal gradient \( (G) \) and varying solidification rate \( (R) \) as shown in the zoomed in section in Fig. 1d. The large thermal gradient and solidification rate leads to a solidification mode dependent on the degree of constitutional undercooling between the melt pool center and boundary. This results in the microstructure ranging from equiaxed at the center to columnar at the boundary. Similar microstructural features of the melt pool have been previously reported. Sub-grain cell structures with an average size of \( \sim 2-3 \mu m \) were observed within the melt pool (Fig. 1e). The occurrence of sub-grain cellular structures is typical of the rapid solidification conditions of the LENS process. In the longitudinal section of the sub-grain cells (Fig. 1f), secondary columnar grains were seen, indicating side instabilities of the solid–liquid interface perpendicular to the primary growth direction with a transition of sub-grains from cellular to dendritic growth. High-magnification electron backscattered diffraction (EBSD) analysis was performed to investigate fine sub-grain cells within the primary grains. The IPF and kernel average misorientation (KAM) maps are shown in Fig. 1g, h, respectively. The IPF map clearly shows local misorientations in the grain interior across the sub-grain cells. The average local misorientation between sub-grain cell structures along the arrow shown in the IPF image was \( \sim 1.25^\circ \) as shown in the misorientation plot (inset of Fig. 1g), suggesting a build up of strain in the AM-ed sample. Furthermore, many fine cell-like features in the size range of 2–3 \( \mu m \) were observed in the KAM map (Fig. 1h), which may be attributed to arrays of geometrically necessary dislocations at the cell boundaries. Microstructure of the as-cast CoCrFeMnNi alloy is shown in Fig. 1i with characteristic parallel columnar grains. The phases present in both AM-ed and as-cast CoCrFeMnNi were identified using XRD as shown in Fig. 1j. Both AM-ed and as-cast CoCrFeMnNi showed single-phase face-centered cubic (FCC) microstructure as previously reported.

![Fig. 2](image)

**Fig. 2**  **Effect of build orientation on tribo-corrosion behavior of AM-ed CoCrFeMnNi.**  
**a** Schematic representation of the three orientations studied.  
**b** Coefficient of friction.  
**c** Open circuit potential.  
**d** Wear volume loss.  
**e** Wear rate for the three build orientations.
Effect of build orientation on tribo-corrosion of AM-ed CoCrFeMnNi

Anisotropy resulting from AM build orientation\textsuperscript{21}, grain refinement, and grain-size distribution directly affect the mechanical properties, wear behavior, and consequently tribo-corrosion response of alloys. The three orientations considered in the present study were: (i) along the build direction (BD), (ii) perpendicular to BD (90° to BD), and (iii) 45° to BD, as marked in Fig. 2a. The results for COF and OCP are shown in Fig. 2b, c, respectively. The OCP curves in Fig. 2c were characterized by a sharp drop to more negative values as wear began, followed by continuous change in OCP for the active tribo-corrosion duration, and finally a jump to nobler value at the end of the test indicating surface re-passivation. The sharp drops in OCP (Fig. 2c) corresponded exactly to the spikes in COF as reported previously for Ti6Al4V\textsuperscript{45,46}. As OCP moved towards nobler values for the three orientations, both COF and wear rate decreased. The wear volume loss and wear rate for the three orientations are shown in Fig. 2d, e, respectively. The wear volume loss and wear rate along 45° to BD was found to be lowest at $9.28 \times 10^{-4}$ mm$^3$ and $5.9 \times 10^{-6}$ mm$^3$/Nm, respectively. This may be attributed to the orientation of most grains at 45° to the BD, thereby resulting in lower grain-boundary density along that direction. Thus, 45° to BD showed the best tribo-corrosion resistance with lowest overall wear rate among the three directions studied, and was used for the remaining tests for comparison with the as-cast alloy.

Effect of normal load on tribo-corrosion response

To evaluate the tribo-corrosion behavior of both AM-ed and as-cast CoCrFeMnNi in a dynamic loading environment, 5 Hz reciprocating frequency was used with varying normal loads of 5 N, 10 N, and 15 N. The COF and corresponding OCP for AM CoCrFeMnNi are shown in Fig. 3a, b, respectively. At all three loads, a distinct drop in COF coincided exactly with increase in OCP between 50 and 450 s, which was attributed to the breakdown in protective passive layer of the AM-ed CoCrFeMnNi. However, the surface re-passivated with further progression of the test as supported by the reduction in COF and shift in OCP towards nobler value. The average COF was in the range of 0.1–0.2 for the three loads for AM CoCrFeMnNi (Fig. 3a). The COF and OCP recorded simultaneously for the as-cast CoCrFeMnNi are shown in Fig. 3c, d, respectively. At 5 N (red plot), COF decreased initially before rising continually to higher values while the OCP moved towards nobler values with progression of the tribo-corrosion test. At 10 N and 15 N loads, the COF curves (Fig. 3c) were relatively smooth while both the OCP curves (Fig. 3d) moved...
towards less noble values indicating the lack of passive layer regeneration. The average COF was in the range of 0.1–0.2 for the three loads while average OCP was in the range of −0.45 to −0.30 V (vs Ag/AgCl). The wear volume loss for both the AM-ed and as-cast CoCrFeMnNi, shown in Fig. 3e, increased monotonically with increase in load but at a different rate going from 5 N to 10 N than from 10 N to 15 N as contact stresses changed with increase in load. The overall wear volume loss for the as-cast CoCrFeMnNi was about six times higher compared to its AM-ed counterpart at all the studied loads. In Fig. 3f, the wear rate for AM CoCrFeMnNi (black dashed plot) showed a monotonic positive slope. While the as-cast CoCrFeMnNi (red dashed line) showed an increase in wear rate from 5 N to 10 N, but a reduced wear rate going from 10 N to 15 N. The overall wear rate for AM CoCrFeMnNi was about six times higher compared to its AM-ed counterpart at all the studied loads. This was attributed to the refined microstructure and sub-grain cellular structures in the case of AM-ed CoCrFeMnNi (shown in Fig. 1), providing greater deformation resistance and a more adherent surface passivation layer.

Wear morphology and mechanism

The surface morphologies of AM and as-cast CoCrFeMnNi HEA after reciprocating tribo-corrosion at different loads with a frequency of 5 Hz are shown in Fig. 4. Both alloys exhibited a change in wear mechanism as the load was varied from 5 N to 15 N. Reciprocating sliding wear morphology of AM-ed alloy at 5 N in Fig. 4a showed typical abrasive wear with mild groves and deformation ridges parallel to the sliding direction along with some degree of oxide pullout. This was ascribed to the surface oxide layer being sufficiently thick and regenerative to provide normal load shielding and passivation. For 10 N load shown in Fig. 4b, a smoother profile with mild oxidative wear was seen because of increased normal load causing a polishing effect with smaller groves and ridges along with dark spots representing oxide patches. Spot energy dispersive spectroscopy (EDS) analysis of the wear track at points S1 and S2 in Fig. 4b showed oxygen content of 3.90 at% and 48.26 at%, respectively, indicating the presence of oxide patches as summarized in Table 1. The change from mild abrasive wear to oxidative wear may be attributed to the increased normal load which tends to polish the surface as a hard-on-hard contact is formed with a lubricating oxide layer in between. For 15 N load (Fig. 4c), similar abrasive wear was observed with mild ridges and polishing effect. The stable and regenerative passive layer for AM CoCrFeMnNi contributed to its better tribo-corrosion behavior. For the as-cast alloy at 5 N and 5 Hz (Fig. 4d), abrasive wear was observed similar to AM CoCrFeMnNi with mild ridges parallel to the sliding direction and dark oxide patches. This was ascribed to the protective passive layer being insufficiently thick, less regenerative, and unable to shield and passive to the extent of the corresponding AM-ed alloy. At 10 N load (shown in Fig. 4e), oxidative wear was observed with large sections of oxide pullout from the combination of high load and corrosive environment that deteriorated the protective passive layer and exposed the underlying alloy. Spot EDS analysis of the wear track at points S3 and S4 in Fig. 4e showed oxygen content of 4.15 at% and 45.24 at%, respectively, again indicating the presence of oxide patches as summarized in Table 1. The larger pullout of oxide patches indicates relatively lower stability of the surface oxide layer for the as-cast alloy. At 15 N load (Fig. 4f), severe deformation ridges and deterioration of the surface oxide led to severe abrasive wear. Thus, the AM-ed alloy showed better tribo-corrosion behavior and stable regenerative oxide compared to its as-cast counterpart under varying loads, which supports the results of OCP and COF discussed in the previous sections.
The tribo-corrosion degradation mechanism for AM-ed CoCrFeMnNi HEA was distinctly different compared to its as-cast counterpart in 3.5 wt% NaCl as discussed in the previous sections. The refined microstructure of the AM-ed CoCrFeMnNi HEA led to strengthening of the alloy and increased wear resistance compared to the as-cast CoCrFeMnNi. Furthermore, the AM-ed alloy had a more protective passive layer which may be attributed to the sub-grain cellular structure promoting better anchoring of the surface oxide. In contrast, the as-cast CoCrFeMnNi HEA showed a less protective surface passivation layer resulting in extensive oxide pullout and severe abrasive wear.

In summary, the tribo-corrosion response and mechanism of AM-ed CoCrFeMnNi HEA was evaluated in 3.5 wt% NaCl solution at room temperature and compared with the corresponding cast alloy. A significant anisotropy was seen in the tribo-corrosion response with 45° to the BD showing the best performance. Both AM-ed and as-cast CoCrFeMnNi exhibited abrasive wear mechanism with the AM-ed alloy showing six times lower wear volume loss and five times lower wear rate compared to its as-cast counterpart. Superior tribo-corrosion resistance of AM-ed CoCrFeMnNi was attributed to the refined microstructure and highly protective passivation layer promoted by the sub-grain cellular structure formed during AM.
METHODS
Laser-engineered net shaping
CoCrFeMnNi plates with dimensions of 29 mm × 10 mm × 10 mm were additively manufactured using laser-engineered net shaping (LENS) on an OPTOMEC LENS 450 equipped with a 400 W Nd:YAG laser. Pre-alloyed CoCrFeMnNi powder with average particle size of 36.8 µm was delivered from the powder feeder to the melt pool through four co-axial nozzles. A standard low-carbon steel plate with a thickness of 6.35 mm was used as the printing substrate. The standoff distance between the nozzles and workpiece was kept ~ 9 mm to maintain deposition efficiency and stability. The processing chamber was filled with high-purity argon with the oxygen content maintained below 10 ppm during printing. The processing parameters were optimized to obtain a near-fully dense printed plate as summarized in Table 2. Bi-directional scan strategy was used and the scan direction was rotated 90° for obtaining consecutive layers. In parallel, equiatomic CoCrFeMnNi alloy ingot was synthesized by vacuum-magnet arc melting using high-purity elements (>99.9%). The as-cast CoCrFeMnNi was flipped and re-melted five times for homogeneity.

Surface analysis and microstructure characterization
Surface morphology, microstructural characterization, and chemical composition of the AM-ed and as-cast CoCrFeMnNi HEAs were obtained using SEM (FEI Quanta-SEM 200™) and EDS. X-ray diffraction analysis using Rigaku Ultima III with 1.54 Å Cu-Kα radiation was performed to determine the crystal structure of the alloys. EBSD analysis was performed to determine the grain-size distribution and orientation. Surface topography and volume loss from tribo-corrosion was determined using non-contact white light interferometry (WLI, RTEC instruments, San Jose, CA, USA) and analyzed using Gwyddion software (version 2.55).

Tribo-corrosion test
Tribo-corrosion experiments were carried out on a reciprocating ball-on-flat tribometer (RTEC Instruments, San Jose, CA, USA) coupled with a potentiostat (Gamry Instruments, Ref-3000, Warminster, PA, USA) in accordance with ASTM G119. Figure 5a shows the tribo-corrosion module potentiostat (Gamry Instruments, Ref-3000, Warminster, PA, USA) in accordance with ASTM G119. Figure 5a shows the tribo-corrosion module.

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**AUTHOR CONTRIBUTIONS**


**COMPETING INTERESTS**

The authors declare no competing interests.

**ADDITIONAL INFORMATION**

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