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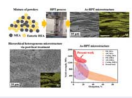
# Hierarchically heterogeneous microstructure and mechanical behavior of the multi-materials prepared by powder severe plastic deformation

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#### ABSTRACT

A heterostructuring strategy is an effective approach for abiding problems referred to as the strength-ductility trade-off. However, tailoring the heterostructure is an arduous problem. This work synthesized hierarchically heterogeneity by combining the eutectic high-entropy alloy (EHEA) and medium-entropy alloy powders through novel powder metallurgy-based severe plastic deformation (P-SPD). EHEA forms nano-scaled heterogeneity composed of FCC and B2, and the mixture of powder types forms micro-scale heterogeneity with hard and soft domains. This hierarchically heterogeneous microstructure leads to strong hetero-deformation-induced strengthening, achieving a yield strength of  $\sim$  1.5 GPa. The present P-SPD represents the feasibility of heterostructuring, aiding the development of HEAs.



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KEYWORDS High-entropy alloys; high-pressure torsion; multi-materials; heterostructures; mechanical properties

#### IMPACT STATEMENT

A hierarchically heterogeneous microstructure comprised of the eutectic high-entropy alloy and medium-entropy alloy was synthesized by the P-SPD procedures and achieved supreme mechanical properties attributed by hetero-deformation-induced strengthening.

# Introduction

Achieving a combination of high strength and ductility has been pursued by numerous metallurgists. Diverse strategies such as alloy design, severe plastic deformation (SPD), and heat treatment have been attempted. In considering the developing materials, the homogeneous structures have been deemed optimized microstructures to avoid early fracture initiation at the domain boundaries in the heterogeneous materials. However, the advantages of heterostructuring to achieve high strength and reasonable ductility are emerging recently [1–4]. Several studies reported synergetic strengthening to overcome the strength-ductility trade-off in heterostructures, such as harmonic, sandwich, gradient, and layered structures [5–7]. However, tailoring heterostructures in the microscale as intended is still problematic.

Powder metallurgy (PM) is a facile manufacturing method combining various powders, such as metals, alloys, ceramics, and polymers [8]. The heterostructure on a microscale can be manufactured by combining

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different types of powders. However, PM-processed parts usually lack ductility under tension because of pores, contaminations, and undesired phases formed during mechanical alloying and high-temperature sintering [9,10]. A novel PM-based fabrication route densifies powders through the high-pressure torsion (HPT) process at ambient temperature, leading to the utmost densification [11,12]. Furthermore, the HPT process is a well-known SPD technique, fabricating nano/ultrafinegrained materials [13–16].

In this work, heterogeneous multi-materials with medium-entropy alloy (MEA) and eutectic high-entropy alloy (EHEA) are fabricated through the abovementioned PM-based technique. As a single material, the achievable mechanical properties of MEA are limited. And, EHEA powder is too hard to be pre-compacted in powder metallurgy. Therefore, the mixture of MEA and EHEA can lead to a complementary relationship. EHEA acts as a reinforcement material in MEA and MEA acts as a binder material in EHEA. Consequently, hierarchically heterogeneous microstructure and exceptional mechanical properties of MEA-EHEA multi-materials were systematically investigated.

### **Materials and methods**

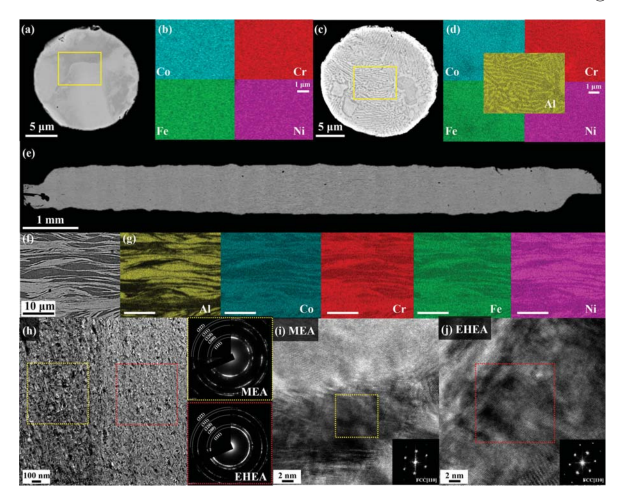
The CoCrFeNi MEA and AlCoCrFeNi<sub>2.1</sub> EHEA powders were fabricated using gas atomization. The produced powders were mixed in proportions targeting the 50 at% of MEA and EHEA. The powders were pre-compacted at a pressure of 40 MPa to manufacture a disk-shape sample with a diameter of 10 mm and a thickness of 1.2 mm. The pre-compacted disks were subjected to the HPT process with a pressure of 5 GPa for 4 turns at a rate of 1 rpm. Then, the HPT-processed samples with a thickness of 0.8 mm were annealed at 700 and 800 °C for 15 and 60 min under an Ar atmosphere. The HPT-processed and post-HPT annealed samples were denoted as As-HPT, A715, A760, A815, and A860, respectively, according to the annealing temperature and time.

For microstructural analysis, X-ray diffraction (XRD), scanning transmission electron microscope (STEM) and scanning electron microscopy (SEM) with backscatter electron (BSE), energy-dispersive spectroscopy (EDS), and electron backscatter diffraction (EBSD) detectors were utilized. Furthermore, the synchrotron XRD characterization was conducted at the 8D beamline of the Pohang Accelerator Lab. Tensile tests and loadingunloading-reloading (LUR) tests were performed on the dog-bone-shaped samples with a gauge length of 1.5 mm, a thickness of 0.8 mm, and a width of 0.7 mm. The quasistatic strain rate of  $10^{-3}$  s<sup>-1</sup> was used for the mechanical tests. A precise tensile strain was taken by a digital image correlation (DIC) method.

#### **Results and discussion**

In Figure 1(a-d), the BSE micrographs and EDS maps are obtained in the spherical-shaped gas-atomized MEA and EHEA powders. There is no elemental segregation in the MEA powder (Figure 1(b)), while the phase separation between the Al-rich phase and Al-lean phase is observed in the EHEA powder (Figure 1(d)). In AlCoCrFeNi<sub>2.1</sub> EHEA, the phase separation from liquid to Al-lean face-centered cubic (FCC) and Al-rich B2 structures is reported at below  $\sim$  1320 °C [17]. Figure 1(e) exhibits an overall cross-section of the as-HPT sample with a defectfree structure. In detail, the magnified microstructure of the as-HPT sample displays two domains elongated along the HPT rotation direction (Figure 1(f)). The corresponding EDS maps in Figure 1(g) exhibit the Al-lean MEA and Al-rich EHEA domains. Due to severe friction stress during the HPT process, the interfaces between MEA and EHEA powders are strongly bonded [11]. The EDS line profile across the domain interface is shown in Fig. S1. According to the distribution of Al, the diffusion layer formed during the HPT process is less than 1 μm. Notably, the EHEA domain in the as-HPT sample presents a segregation-free structure, unlike the starting EHEA powder. To distinguish nano-scaled elemental segregation, STEM analysis was performed. Figure 1(h) exhibits a sound interface between the MEA and EHEA domains without bonding defects. The SAED rings of both domains are obtained from the selected regions (yellow region: MEA and red region: EHEA). The SAED rings of the MEA domain indicate that the FCC structure remains after the HPT process. Similarly, SAED rings of the FCC phase are detected in the EHEA domain. The fast Fourier transform (FFT) patterns from MEA and EHEA also exhibit FCC structures (Figure 1(i, j)). In other words, the B2 phase observed in starting EHEA powder (Figure 1(c, d)) disappears after the HPT processing, attributable to the dissolution of B2 phase under SPD [18-20]. From the diameter of SAED rings, the lattice parameters of the FCC phase in the MEA and EHEA are calculated as 3.55 and 3.59 Å, respectively. The dissolution of the B2 phase could be inferred by the decreased diameter of SAED rings in EHEA compared to MEA. The penetration of Al elements in the FCC matrix by the dissolution of B2 phase leads the larger lattice parameters, shrinking the SAED rings. In addition, the MEA and EHEA domains are refined as a nanocrystalline microstructure by the HPT process. The magnified HR-TEM images of the MEA and EHEA domains exhibit ultrafine-grained structures in Fig. S2. The grain sizes of the MEA and EHEA are  $\sim$  29.4 and  $\sim$  25.3 nm, respectively.

XRD spectra of all samples shown in Figure 2(a) are consistent with the HR-TEM results. Initial MEA

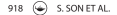


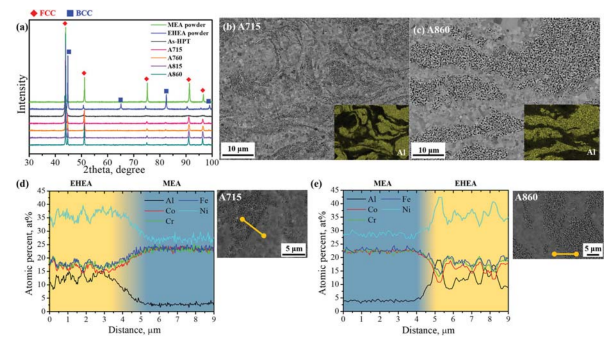
**Figure 1.** (a, c) SEM-BSE micrographs and (b, d) EDS maps of the gas-atomized (a, b) MEA powder and (c, d) EHEA powder, (e) the crosssection SEM image of the As-HPT sample, (f) SEM-BSE image of the As-HPT sample with (g) corresponding EDS maps, (h) bright field TEM micrographs of the MEA/EHEA interface with SAED ring patterns from MEA and EHEA regions, and HR-TEM micrographs of MEA and EHEA is shown in (i) and (j) accompanied by FFT patterns, respectively.

powders contain FCC single-phase, and EHEA powders consist of FCC and B2 dual-phase (Figure 2(a)). However, the peaks related to the B2 phase are not detected in the spectrum of the as-HPT sample, owing form the dissolution of B2 phase under severe HPT-induced strain. Subsequent annealing on the HPT-processed samples is conducted to tailor the microstructure to reach a good combination of strength and ductility. The phase decomposition to FCC and B2 phase occurs in all annealed samples according to the B2 peaks detected at 2theta of  $\sim$  82°. After HPT processing, the dissolution of B2 makes a supersaturated FCC phase in the EHEA which is thermodynamically unstable [21]. Also, the diffusion process is generally accelerated in severely deformed alloys due to lattice defects [22]. Therefore, the post-HPT annealing, even for a short time, leads to the decomposition of the supersaturated FCC matrix in the EHEA domains. To verify the recrystallization behavior, EBSD and convolutional multiple whole profile (CMWP) analyses were

performed on the A715 sample (Fig. S3). Even with the lowest annealing temperature and shortest annealing time, the A715 sample exhibitsed a low average KAM value of  $\sim 0.32^{\circ}$  and dislocation density of  $8.77 \times 10^{13}$  m<sup>-2</sup>, as determined from CMWP fitting, indicating a fully recrystallized microstructure [23].

The distribution of the constituent phases is displayed in Figure 2(b, c). The MEA and EHEA domains are distinguished by the enveloped Al element maps. And, EDS line profiles of A715 and A860 samples are shown in Figure 2(d, e). The fluctuations of elemental distribution are observed in the EHEA domain because of the phase separation to FCC and B2 phases. Between EHEA and MEA domains, the diffusion layers are extended over 1  $\mu$ m after annealing. Interestingly, the presence of Al over 3 at% is observed in MEA domains, which indicates some EHEA domains dissolved in MEA domains rather than forming separated EHEA domains. The compositions of the domains were summarized in Table S1. The





**Figure 2.** (a) X-ray diffraction patterns of the powders, as-HPT sample, and annealed samples at different temperatures and times, SEM-BSE micrographs of the (b) A715 and (c) A860 samples, and (d, e) EDS line profiles obtained from the yellow lines indicated in the enveloped BSE images.

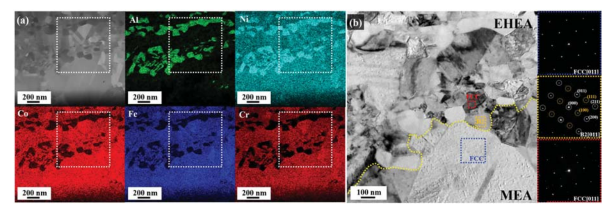


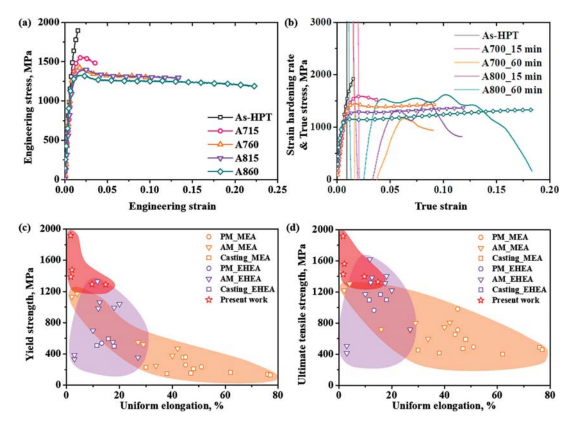
Figure 3. (a) TEM micrographs of the interface of the MEA and EHEA domains with the corresponding EDS maps and (b) HRTEM images with the FFT patterns of FCC and B2 phases in A715.

increased contents of Ni in MEA domains also support the dissolution of EHEA domains in MEA domains.

In Figure 3, TEM micrographs and corresponding EDS maps of the domain interfaces in the A715 sample show the decomposition of the EHEA to the Al-rich and Al-lean regions. From the FFT pattern obtained in the HR-TEM image, the Al-rich and Al-lean regions are confirmed to be the B2 and FCC phases.

Since the EHEA domain decomposes to a dual-phase structure during the post-HPT annealing, the grain growth is retarded in this domain because one phase can effectively resist the growth of the other phase, resulting in the finer grain size [22,24]. As a result, the FCC grains in the EHEA domain contain a finer grain size

of ~463 nm than the FCC grains in the MEA domain of ~778 nm in the A715 sample. Also, the grain size of the B2 phase (~393 nm) is similar to the FCC grain size in the EHEA. In the A860 sample, as the annealing temperature and time increase, the grain size increases in both MEA and EHEA domains. However, the difference in the grain size gets distinctly in the A860 sample. The FCC grain size in the MEA grows up to ~1.282 µm, while the grain size of FCC and B2 phases in the EHEA remains about ~400 nm. Consequently, a hierarchically heterogeneous structure is achieved, containing coarse FCC grains in the MEA domain, and fine FCC and B2 grains in the EHEA domain. Furthermore, the nanoindentation hardness of each phase is reported as 9.7 (B2),



**Figure 4.** (a) Engineering stress-strain curves of the as-HPT and annealed samples, (b) SHR curves. Comparison of (c) YS vs. uniform elongation and (d) UTS vs. uniform elongation of the samples in the present work and literature processed by the casting, conventional powder metallurgy (PM) routes, and additive manufacturing (AM) [12,38–52].

5.8 (FCC in the EHEA domain), and 4.6 GPa (MEA), respectively [25,26]. Because the B2 and FCC phases in the EHEA domain have higher hardness and finer grain size, the difference in mechanical properties between the MEA and EHEA domains become more pronounced. The present hierarchically heterogeneous structure consists of grain size heterogeneity in FCC grains, chemical composition heterogeneity stemming from different powders, and crystal structure heterogeneity originating from phase decomposition to FCC and B2 phases.

The tensile stress-strain curves are displayed in Figure 4(a). The tensile strength of the as-HPT sample reaches ~1.92 GPa, representing the strong bonding between different powders achieved by cold-consolidation using P-SPD procedure. Due to the restricted dislocation movement in the as-HPT sample, this sample demonstrates a limited plastic deformation [12,27]. After the post-HPT annealing, the ductility is improved in the cost of strength. The yield strength (YS) values of the annealed samples decrease from ~1476 (A715), ~1388 (A760), ~1293 (A815), to ~1289 MPa (A860) due to the reduced dislocation density and grain growth, while total elongation continuously increases from ~4% (A715), ~10 (A760), ~14 (A815), to 22% (A860). Noticeably, all annealed samples exhibit remarkable

YS over 1.2 GPa, attributing to the nano-crystalline microstructure, utmost densification, high dislocation density, strong B2 phases, and hierarchical heterostructure achieved by the present fabrication technique.

To clarify the source of high YS, the expected YS is calculated using the rule of mixture (ROM) equation [28]. The A860 sample consists of MEA and EHEA domain with an atomic percent of 50%, which can be recalculated to 49.4 Vol% of MEA and 50.6 Vol% of EHEA using the atomic weight of constituent elements and density of alloys. First, the strength of the MEA domain is obtained from the following Hall-Petch equation [29]:

$$\sigma_{MEA} = \sigma_{0\_MEA} + K_{MEA} \cdot d^{-\frac{1}{2}},\tag{1}$$

where  $\sigma_{MEA}$ ,  $\sigma_{0\_MEA}$ ,  $K_{MEA}$ , and d are YS, friction stress (256 MPa), Hall-Petch coefficient (226 MPa  $\cdot \mu m^{\frac{1}{2}}$ ), and grain size of MEA domain, respectively [30]. By substituting the grain size of the A860 sample ( $\sim 1.3 \mu m$ ) into the equation, the YS of the MEA domain is calculated as  $\sim 454$  MPa.

Meanwhile, the Hall-Petch-type equation can be used in the EHEA [31]:

$$\sigma_{EHEA} = \sigma_{0\_EHEA} + K_{EHEA} \cdot \lambda^{-\frac{1}{2}}, \qquad (2)$$

where  $\sigma_{EHEA}$ ,  $\sigma_0$  <sub>EHEA</sub>,  $K_{EHEA}$ , and  $\lambda$  are YS, friction stress, Hall-Petch coefficient, and inter-lamellar spacing of the EHEA domain. The inter-lamellar spacing  $(\sim 0.86 \ \mu m)$  is calculated as the sum of grain sizes of FCC and B2 phases in the EHEA domain. The  $\sigma_0$  *EHEA* and  $K_{EHEA}$  are obtained as 163 MPa and 907 MPa  $\cdot \mu m^{\frac{1}{2}}$ , by plotting the data of YS and inter-lamellar spacing in Ref. [32]. The YS of the EHEA domain is calculated as  $\sim$  1142 MPa. Using ROM equation, the YS of MEA-EHEA multi-materials is predicted to be 802 MPa, which cannot represent the experimental YS. The difference between the calculated and experimental YS is assumed to be attributed to residual stress, dislocations, contamination of powders, and HDI strengthening. Furthermore, the large Al atoms added to MEA domains during the annealing process can induce additional solid solution strengthening [3]. The microstructural heterogeneity could induce massive plastic incompatibility between the domains. To maintain plastic compatibility, geometrically necessary dislocations (GNDs) could be generated and piled up at the domain boundaries, leading to heterodeformation induced (HDI) strengthening [33–35]. The HDI strengthening contributes to YS and additional strain hardening, resulting in a good combination of strength and ductility [3,33,34].

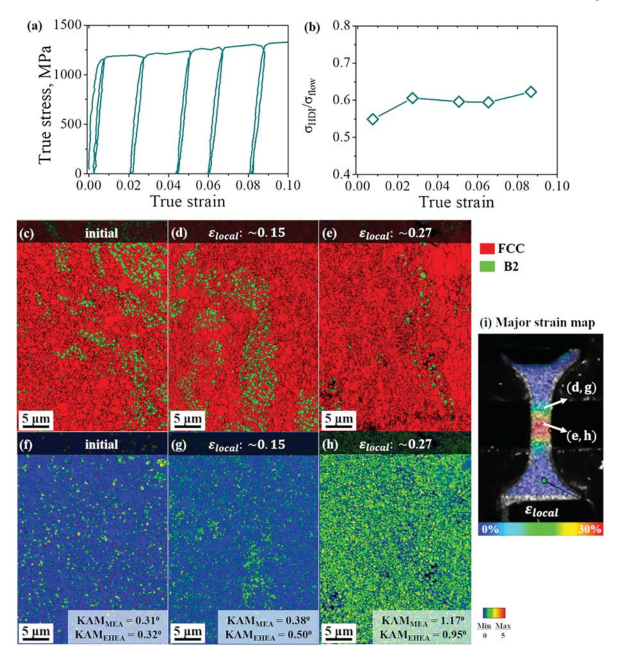
Meanwhile, it is known that post-elongation after necking can be affected by the sample dimension, while the uniform elongation before necking is independent of sample thickness and gauge length [36]. To exclude the sample size effects, the uniform elongation is determined from the intersection of true stress-strain curves and strain-hardening rate (SHR) curves in Figure 4(b). To obtain the uniform elongation and valuate the contribution of HDI strengthening, the strain-hardening behaviors of the samples are examined using the strainhardening rate (SHR) curves in Figure 4(b). The as-HPT sample shows a decreasing trend in SHR until fracture. This behavior is usually observed in SPD materials with high dislocation density [27]. Meanwhile, the SHR curves of the annealed samples show dramatic drops under subzero values after yielding, which was observed in ultrafine-grained materials due to the yield drop phenomenon [12]. Then, the SHR values of the A760, A815, and A860 samples return to positive and fluctuate in positive values until fracture. The different domain sizes for GND accumulation and the saturation of dislocation at different strain levels lead to the fluctuation of the SHR values. Among the intersection of the true stress-strain curves and the SHR curves, the highest strain is considered the onset of necking [37]. The uniform elongation increases from  $\sim 2\%$  (A715),  $\sim 2\%$ (A760),  $\sim 10\%$  (A815), to 15% (A860). With controllable annealing, an exceptional strength-ductility combination

of YS  $\sim$  1289 MPa and uniform elongation  $\sim$  15% is obtained in the A860 sample.

The YS vs. uniform elongation and the ultimate tensile strength (UTS) vs. uniform elongation of the samples in this study are compared to those of the MEAs and EHEAs from literature (Figure 4(c, d)) [12,38–52]. To exclude the size effect, the values of uniform elongation were used for comparison. The multi-materials fabricated in this study exhibit superior mechanical properties to all MEAs in the previous studies. Moreover, the strength of multi-materials is higher or similar to that of EHEAs fabricated by casting, powder metallurgy, and additive manufacturing. Considering the simple calculation from the ROM [28], the mechanical properties of this study should be positioned in the intermediate range of the MEAs and EHEAs. However, the mechanical properties of these multi-materials are placed above the those of single EHEAs and MEAs. The impressive mechanical properties achieved in this study are attributed to grain refinement, dislocation density, and heterostructures, which can be manufactured by the novel P-SPD technique. Furthermore, superior mechanical properties would be expected by controlling the ratio of powders and adding reinforcement particles to fabricate HEA/MEA-matrix composites.

To evaluate the contribution of HDI strengthening in the present hierarchically heterostructure, the ratios of HDI stress to flow stress are calculated at different true strains from the LUR test of the A860 sample (Figure 5(a,b)). The HDI stress and effective stress of the sample are summarized in Table S2. The effective stress of the A860 remains around 500 MPa, while the values of HDI stress steadily increase from  $\sim 637$  to  $\sim 802$  MPa. Also, the strain hardening of the A860 sample is predominantly attributed to HDI stress, which occupies  $\sim 60\%$  of flow stress.

Figure 5(c-e) and Figure 5(f-h) represent the phase maps and KAM maps of the A860 sample in the different local strains to assess the evolution of GNDs in the present hierarchically heterostructure. The KAM evolutions in the MEA and EHEA domains at the different local strains are summarized in Fig. S4. The initial microstructures are obtained in the grip part of the tensile sample, which exhibits a low level of KAM values. The low KAM value means that the HPT-induced dislocations are recovered during annealing. Meanwhile, at the local strain of  $\sim 0.15$  (Figure 5(d,g)), the KAM values of the FCC phases (0.50°) in the EHEA domain increase first, owing to profuse phase boundaries between FCC and B2 phases. The hard B2 phase and soft FCC matrix in the EHEA develop GNDs to maintain strain compatibility [33,53]. Ultrafine grains of FCC and B2 phases in the EHEA domains offer profuse grain boundaries



**Figure 5.** (a) Loading-unloading-reloading curves of the A860, (b) ratio of HDI stress to flow stress taken from the LUR test, and (c-h) EBSD-KAM maps taken from (i) the major strain map of the deformed A860 sample at (c, f) an initial state and local strain of (d, g)  $\sim$  0.15 and (e, h) 0.27, respectively. The KAM maps are measured up to the third nearest neighbor with a maximum misorientation of 5°.

and phase boundaries. Furthermore, the B2 phases act as strong obstacles to dislocation movement [45]. Abundant obstacles like the B2 phase, phase boundaries, and grain boundaries attribute the steep increase of dislocation density in the FCC phases of EHEA domains. However, the dislocation accumulation shows a different aspect in the intensely deformed microstructures at the local strain of ~0.27 (Figure 5(e,h)). The KAM value of the MEA domain (1.17°) increased significantly, surpassing the KAM value of the FCC in the EHEA domain (0.95°). Because the EHEA domains act as hard domains, and the MEA domains act as soft domains in the microscale, the GND evolves in the MEA domains to compensate for the strain incompatibility, leading to an additional HDI strengthening.

Those KAM evolutions displayed in Figure 5(c-h) can help elucidate the trend of the ratio of HDI stress to flow stress in Figure 5(b). The microstructural inhomogeneity between the FCC and B2 phases in the EHEA is repeated at the nanoscale, leading to a rapid increase and subsequent saturation of dislocation density. The ratio of HDI stress to flow stress sharply increases from 0.55 ( $\epsilon_{true} = 0.01$ ) to 0.61 ( $\epsilon_{true} = 0.03$ ), which can be attributed to the increased accumulation of GNDs within the EHEA domain. On the other hand, the MEA and EHEA domains repeat at the in microscale, resulting in a relatively slower increase in the ratio of HDI stress to flow stress.

# Conclusion

While the importance and expediency of the heterostructuring are getting more and more attention, it is noteworthy that a hierarchical heterostructuring can be obtained by selecting appropriate materials and consolidating their powder mixtures. In this work, a sound bulk disk with hierarchically heterogeneous structures was fabricated by combining the ductile CoCrFeNi MEA and hard AlCoCrFeNi<sub>2.1</sub> EHEA powders with a dual-phase structure. HPT processing and adequate heat treatment led this multi-material to have a hierarchically heterogeneous structure, consisting of (1) nano-scaled phase separation within EHEA and (2) EHEA-MEA domain separation, leading to a superior combination of mechanical strength and ductility compared to EHEA and MEA of the previous studies. Furthermore, the ultrafine grains fabricated by SPD process effectively improved the strength of multi-materials. These materials could be considered in high-tech applications such as micro-gears. This study suggests an attractive solution to the fabrication of hierarchically heterogeneous structures, which have been considered troublesome. A complicated heterostructure with two or more hierarchies can be fabricated by including several powders of various physical properties, combining powders with heterostructures, or controlling the powder size and powder ratio. The suggested solution for the hierarchically heterogeneous structuring would be a breakthrough to achieve supremum properties through a PM-based technique.

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# **Disclosure statement**

No potential conflict of interest was reported by the author(s).

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#### Data and materials availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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