Hierarchical microstructure strengthening in a single crystal high entropy superalloy

Yung-Ta Chen1,2, Yao-Jen Chang1,3, Hideyuki Murakami2,4, Taisuke Sasaki5, Kazuhiro Hono5, Chen-Wei Li6, Koji Kakehi6, Jien-Wei Yeh1,3 & An-Chou Yeh1,3*

A hierarchical microstructure strengthened high entropy superalloy (HESA) with superior cost specific yield strength from room temperature up to 1,023 K is presented. By phase transformation pathway through metastability, HESA possesses a hierarchical microstructure containing a dispersion of nano size disordered FCC particles inside ordered L12 precipitates that are within the FCC matrix. The average tensile yield strength of HESA from room temperature to 1,023 K could be 120 MPa higher than that of advanced single crystal superalloy, while HESA could still exhibit an elongation greater than 20%. Furthermore, the cost specific yield strength of HESA can be 8 times that of some superalloys. A template for lighter, stronger, cheaper, and more ductile high temperature alloy is proposed.

The development of high-entropy alloys (HEAs) has broken through the frame of conventional alloys by exploring the vast composition space of multi-principle elements1–6, and their extraordinary mechanical properties have been a subject of interest, for examples, single-phase CoCrFeMnNi HEA showed high tensile strength of 1,280 MPa with elongation up to 71% at cryogenic temperature7; the compressive strength could reach 2,240 MPa at 298 K for Al0.5CoCrFe0.5NiTi0.5 HEA8 and 1,520 MPa at 873 K for Al0.5CrNbTi2V0.5 HEA9 due to the presence of intermetallic phases, such as σ, B2 and Laves. Most of the high temperature mechanical properties data of HEAs in literatures are from compression tests3,4,10. Although some studies have reported tensile tested data of HEAs, there are only a few at elevated temperatures3,4,10–17. The tensile strength of HEA could be degraded severely at elevated temperatures4,17,18, for example at 1,023 K, the tensile yield strength of CoCrFeMnNi was lower than 100 MPa17. In conventional precipitation strengthened superalloys, precipitation of coherent Ni3(Al, Ti) L12 structured phase in FCC matrix can provide effective strengthening19. Recent studies have shown that coherent L12 phase could also be an effective strengthener in HEAs11,20–22, for example, L12 precipitation in Al7(Fe, Co, Ni)86Ti7 resulted a combination of high tensile yield strength (1,028 MPa) with large elongation (47.8%) at room temperature20. However, the elevated temperature tensile strength of HEA could be limited by insufficient fractions and relatively low solvus of strengthening phases in HEAs11. One of the highest reported tensile yield strength of HEAs at 1,023 K was 473 MPa for cast-type Al10Co25Cr8Fe15Ni36Ti6, which contained a FCC matrix with 46% L12 and 5% B2 phases by volume fractions9, although its yield strength could surpass those of solid solution type superalloys such as 800H and Inconel617, it was weaker than advanced precipitation strengthened cast-type superalloys23. Alloy design for higher L12 phase fraction and solvus temperature are required to further improve the high temperature tensile strength of HEAs. However, the high entropy composition scope3,4 of 5.0 at.% ≤ x ≤ 35.0 at.% could jeopardize the thermal stability of ordered phase such as L12 phase24–28, and Ni-rich HEA with Ni content beyond 35 at.% has provided a window of opportunity to design thermally stable L12 precipitation in HEA while retaining the compositional configurational entropy |ΔSconf.| > 1.5 R, where R is the universal gas constant3,4; this class of HEA has been named High Entropy Superalloys (HESA)25–28. Recently, the

1Department of Materials Science and Engineering, National Tsing Hua University, 101, Sec. 2, Kuang-Fu Road, Hsinchu 30013, Taiwan, ROC. 2Research Center for Structural Materials, National Institute for Materials Science, 1-2-1 Sengen, Tsukuba 305-0047, Japan. 3High Entropy Materials Center, National Tsing Hua University, 101, Sec. 2, Kuang-Fu Road, Hsinchu 30013, Taiwan, ROC. 4Department of Nanoscience and Nanoengineering, Waseda University, 3-4-1 Okubo, Shinjuku, Tokyo 169-8555, Japan. 5Research Center for Magnetic and Spintronic Materials, National Institute for Materials Science, 1-2-1 Sengen, Tsukuba 305-0047, Japan. 6Department of Mechanical Engineering, Tokyo Metropolitan University, 1-1 Minami-osawa, Hachioji-shi, Tokyo 192-0397, Japan. *email: yehac@mx.nthu.edu.tw
concept of HESA has been adopted by Zhang et al., Ni\textsubscript{L1\textsubscript{2}}(Fe, Co, Cr),\textsubscript{HE}(Al, Ti),\textsubscript{HE}H\textsubscript{6} based alloys were studied in as-cast condition; the tensile strength of these HESAs could reach 960 MPa at 1,023 K.

In this work, the HESA (HESA-3\textsuperscript{25}) of interest is shown in Table 1; it is a cast-type alloy with a density of 7.96 g/cm\textsuperscript{3}, comparing to advanced cast superalloys such as CMSX-4\textsuperscript{36}, the raw materials cost of this HESA was 84% cheaper due to the absence of Re element content. This HESA in directionally solidified form\textsuperscript{27} had been prepared by the standard process\textsuperscript{24}. This work demonstrates that phase transformation pathway through metastability can induce an interesting hierarchical microstructure, which can further increase the elevated temperature tensile strength of HESA. The aim of this article is to present a microstructure template for developing future advanced high temperature alloys with improved cost-performance.

Results

Hierarchical microstructure of HESA. Single crystal samples of HESA, with the composition shown in Table 1, were heat treated by two different heat treatment steps, i.e. HT-1 and HT-2; Fig. 1 shows the heat treated microstructures. HT-1 consisted a single ramp from room temperature to 1,500 K in 20 h followed by air cooling, and the microstructure contained cuboidal precipitates and nano-particles, Fig. 1a, superlattice diffraction pattern observed on the [001] zone axis indicates that these precipitates possessed L\textsubscript{1\textsubscript{2}} structure; the TEM dark field image taken from the L\textsubscript{1\textsubscript{2}} (001) superlattice spot is shown in Fig. 1b. The precipitation of L\textsubscript{1\textsubscript{2}} phase (24 nm in size) and secondary L\textsubscript{1\textsubscript{2}} particles (1 nm in size) occurred during the air cooling process in HT-1, since the solvus of L\textsubscript{1\textsubscript{2}} phase in this HESA was 1,472 K\textsuperscript{22}. After an additional heat treatment at 1,023 K for 20 h followed by water quenching, the HT-2 process caused the L\textsubscript{1\textsubscript{2}} phase to grow from 143 to 153 nm in average and secondary L\textsubscript{1\textsubscript{2}} particles to coarsen into an average of 14 nm as shown in Fig. 1c; Furthermore, there were nano size particles appeared inside the cuboidal L\textsubscript{1\textsubscript{2}} precipitates. Figure 1d shows the selected area diffraction pattern from the L\textsubscript{1\textsubscript{2}} phase on the [001] zone axis together with the dark field TEM image taken from the L\textsubscript{1\textsubscript{2}} (001) superlattice spot. Since L\textsubscript{1\textsubscript{2}} precipitate has ordered FCC structure with the lattice constant close to the disordered FCC phases, one way to identify L\textsubscript{1\textsubscript{2}} phase was by the L\textsubscript{1\textsubscript{2}} superlattice spots, which could be used in dark field image to enhance the contrast between L\textsubscript{1\textsubscript{2}} phases (brighter contrast) and the FCC matrix and nano-particles (darker contrast), Fig. 1d. Since the diffraction patterns exhibited only FCC and L\textsubscript{1\textsubscript{2}} diffraction spots, so these nano-particles were very likely to possess disordered FCC structure.

Phases in HESA after HT-1 and HT-2 states have also been revealed by atom probe tomography reconstructions as shown in Fig. 2. To distinguish between FCC matrix and L\textsubscript{1\textsubscript{2}} precipitates, Cr and Ti atoms have been depicted as these elements strongly partitioned to the FCC matrix and the L\textsubscript{1\textsubscript{2}} precipitates, respectively. Ti 4.8 at.% iso-surface is set to highlight the FCC matrix—L\textsubscript{1\textsubscript{2}} precipitate interface for clear visualization. In Fig. 2a, HT-1 state shows an FCC matrix with L\textsubscript{1\textsubscript{2}} precipitates and secondary L\textsubscript{1\textsubscript{2}} particles in the FCC matrix channel. The chemical composition of each phase was measured and listed in Table 1. In addition, partitioning coefficient which is defined as the concentration of element in FCC matrix divided by the concentration of the same element in the L\textsubscript{1\textsubscript{2}} phase, can be calculated. In the HT-1 state, the partitioning coefficients of Al (0.39), Ti (0.14), Nb (0.23) and Ni (0.53) are below unity and that indicates a preferential partitioning behavior toward the L\textsubscript{1\textsubscript{2}} phase, the partitioning coefficients of Co (1.83), Cr (8), Fe (4.27), Mo (3) and W (1.5) are greater than unity, suggesting that these elements preferred to partition to the FCC matrix; the elemental partitioning behaviors were similar to those reported in superalloys\textsuperscript{19,31}. In Table 1, the magnitude of the configuration entropy (\(\Delta S_{\text{conf}}\)) for the FCC matrix is calculated to be 1.63 R, which agrees with the high-entropy composition definition. So, in the HT-1 condition, HESA possessed a High Entropy (HE) FCC matrix and Medium Entropy (ME) precipitates (L\textsubscript{1\textsubscript{2}}).
By contrast, the APT reconstructions in Fig. 2b shows an interesting hierarchical microstructure, in addition to the FCC matrix and the L1₂ phases, there were nano-particles dispersion within the L1₂ phase. These nano-particles were only present in the HT-2 state, not in the previous HT-1 state. Compositions of each phase in HT-2 state are summarized in Table 1. The calculated partitioning coefficients of Al, Ti, Nb, and Ni are 0.27, 0.09, 0.08 and 0.41, respectively; Co, Cr, Fe, Mo, and W are 2.24, 11.33, 5.56, 3.2, and 1.67, respectively. Partitioning coefficients in HT-1 and HT-2 states are similar. The nano-particles in the L1₂ phase of HT-2 state possessed similar compositions as that of the FCC matrix, Table 1. According to the TEM analysis on structure (Fig. 1d) and APT composition measurement (Table 1), these nano-particles could be determined indirectly as disordered FCC particles. The magnitude of configuration entropy (ΔSconf) for the FCC matrix and the FCC particles are 1.61 R and 1.55 R, respectively, and both satisfy the high-entropy composition definition. Therefore, microstructure in HT-2 state contained High Entropy FCC matrix with a dispersion of Medium Entropy L1₂ phase that also possessed nano-sized High Entropy FCC particles. Elemental partitioning behaviors and the concentration profiles evolution between the FCC and L1₂ phases from HT-1 to HT-2 are shown in the Supplementary Fig. S1a,b.

The composition analysis of each phase in Table 1 have been confirmed by the lever rule analysis 26,27,31, which is based on the principle of mass conservation. Plots of CL₁₂–CFCC versus Cₙ–CFCC from the HT-1 (Supplementary Fig. S2a) and HT-2 (Supplementary Fig. S2b) conditions can be obtained, where CL₁₂, CFCC and Cₙ represent the chemical compositions of ME-L1₂ phase, HE-FCC matrix and bulk alloy, respectively. Line fittings could be conducted to verify the measured compositions, since the slopes could represent the mole fraction of the L₁₂ phase. The linear fitting slopes in Supplementary Fig. S2a,b correspond to 68.4% and 70.6% of L₁₂ precipitates in mole fractions, respectively; these values match well with the volume fractions of phases determined in Fig. 1.

**Thermodynamic simulation and phase evolution.** To elucidate the possible underlying mechanisms for the formation of the HE-FCC particles in the HT-2 condition, CALPHAD-based simulation, ThermoCalc 32 (TCHEA3 databases) was used to predict phase diagrams of HESA and ME-L₁₂ phase in both HT-1 and HT-2 conditions, Fig. 3a,b. FCC and L₁₂ are the main phases predicted by the phase diagram in Fig. 3a; although minor B₂ and Mu phases were also predicted, they were not found in the microstructure experimentally. Inter-
estingly, CALPHAD phase fraction predictions (Fig. 3b) indicate that both ME-L12 phase in HT-1 and HT-2 conditions could decompose into FCC and L12 equilibrium phases, these results suggest the tendency to form FCC-structured particle inside the L12 phase. However, the microstructure of the ME-L12 phase in HT-1 state did not contain FCC particles. So, it can be deduced that the ME-L12 phase in HT-1 condition was in a supersaturated state, Figs. 1a and 2a. On the other hand, the HT-2 heat treatment could promote the formation of FCC particles within ME-L12 phase, Figs. 1c and 2b. To further clarify the supersaturation and decomposition of ME-L12 phase, ThermoCalc was applied to determine the equilibrium phase compositions at 1,023 K, and the calculated partition coefficients are Al (0.19), Ti (0.02), Nb (0.41), Co (2.16), Cr (22.78), Fe (9.13), Mo (11.5) and W (11.0). Detailed equilibrium phase compositions are given in Supplementary Table S1. By comparing equilibrium elemental partitioning coefficients with those of the HT-1 and HT-2 states in Table 1, the HT-2 condition appeared to be closer to the equilibrium state than that of the HT-1 condition. The phase fractions calculated from the ME-L12 phase composition of HT-1 state indicate that the L12 phase could decompose into 7.7 mol% FCC phase at 1,023 K (Fig. 3b). By contrast, FCC phase fraction at 1,023 K calculated from the HT-2 composition was 6.2 mol% (Fig. 3b); the actual fraction of HE-FCC particles measured experimentally was 4.3 vol. % in the L12 phase, indicating that the HT-2 state was still not at the equilibrium state. Therefore, the formation of FCC particles in HESA was driven by the supersaturation of FCC phase formers inside the L12 phase. The microstructural evolutions of HESA from HT-1 to HT-2 and prolonged ageing are illustrated in Fig. 3c. The higher energy level represents the HT-1 state, which was relatively unstable and would evolve into a lower energy state by the decomposition of supersaturated L12 phase. With additional heat treatment at 1,023 K for 20 h, phase decomposition occurred and lead to the formation of metastable FCC particles in the L12 phase at a lower energy level as the HT-2 state. With further ageing at 1,023 K for 200 h, the metastable FCC particles would mostly be eliminated by diffusing FCC formers into the surrounding FCC matrix, eventually, the whole system would reach the equilibrium state.

High temperature tensile properties. Tensile tests were conducted at 298 K, 723 K, 923 K, 1,023 K and 1,173 K on HESA in HT-1 and HT-2 states, the whole tensile stress–strain curves are presented in Supplementary Fig. S3. Figure 4a shows the tensile yield strength versus temperature plot, which includes HESA, some advanced superalloys and some conventional HEAs. The single-phase CoCrFeMnNi HEA possessed a yield strength of 362 MPa at 298 K and 156 MPa at 1,023 K[30]. Al_{10}Co_{25}Cr_{8}Fe_{15}Ni_{36}Ti_{6} showed a yield strength of 627 MPa at 298 K and 473 MPa at 1,023 K[30]. By contrast, HESA in HT-1 state could achieve a yield strength of 880 MPa at 298 K and 954 MPa at 1,023 K, both were higher than those of CoCrFeMnNi and other HEAs, approaching the yield strength level of advanced superalloy such as CMSX-4 (888 MPa at 298 K and 913 MPa at 1,023 K)[30]. For fair comparison in this work, the data of CMSX-4 were selected from those of as-cast single crystals treated by the standard heat treatments without hot isostatic pressing (solutioned heat treatment at 1,549 K/2 h + 1,560 K/2 h + 1,577 K/3 h + 1,588 K/2 h + 1,594 K/2 h + 1,597 K/2 h (air cooling), and two step ageing treatments at 1,353 K/4 h (air cooling) + 1,144 K/20 h (air cooling)). The standard heat treatments of CMSX-4 varied in literatures, so the reported yield strength at 1,023 K could vary as well[30,31,37,38].
consistency, the yield strength data of CMSX-4 in Fig. 4 were from the same literature\(^33\). Impressively, HESA in the HT-2 state exhibited even higher yield strength (993 MPa at 298 K and 1,023 MPa at 1,023 K) comparing to those of the HT-1 state. The yield strength of the HT-2 state has surpassed several advanced superalloys from room temperature to elevated temperatures. At 1,023 K, the yield strength of HESA in HT-2 state was 110 MPa higher than that of CMSX-4\(^33\). However, when the testing temperature was raised to 1,173 K, yield strength of the HT-1 and HT-2 states became similar.

Figure 4b,c summarize the specific yield strength, cost-specific yield strength and elongation at 1,023 K for HESA and several advanced superalloys\(^24,27,38,39\). Specific yield strength is the yield strength divided by density, and cost specific yield strength is the specific yield strength divided by the raw material cost. In Fig. 4b, HESA shows an excellent high temperature specific yield strength, which is 11% higher than that of CMSX-4. The CMSX-4 data shown in Fig. 4b,c was taken from previous work by Matsubara et al.\(^24\), which provided data of the specific tensile yield strength and elongation at 1,023 K. Figure 4c also includes CMSX-8\(^39\), which is an improved cost-performance version of single crystal superalloy and possesses 1.6 times the cost specific yield strength of CMSX-4. By contrast, HESA presents a remarkable advancement with 8 times the cost specific yield strength of CMSX-4. In addition, at 1,023 K, HESA exhibited a tensile elongation greater than 20%, which was three times
greater than that of CMSX-4, Fig. 4b,c. These results suggest HESA as a cheaper, lighter, stronger and more ductile alloy as comparing to advanced superalloys.

TEM analysis on tensile-tested samples are shown in Fig. 5; HT-1 samples tested under 723 K, 923 K and 1,173 K are shown in Fig. 5a,c, respectively; there were dislocations accumulated at the HE-FCC matrix and ME-L12 precipitates interfaces, and ME-L12 precipitates were sheared by pairs of superdislocations that appeared to be straight and parallel. TEM analysis on HT-2 samples tested under the same conditions are shown in Fig. 5d–f. At 723 K (Fig. 5d) and 923 K (Fig. 5e), pairs of superdislocations in ME-L12 precipitates appeared to be wavy, this indicates that HE-FCC particles could act as obstacles for superdislocations within ME-L12 precipitates. At 1,173 K (Fig. 5f), the HE-FCC particles disappeared, and the yield strength of HESA in the HT-1 and HT-2 states became similar at 1,173 K, Fig. 4a.

Discussion

There have been several important HEA studies reporting breakthroughs in strength-ductility synergy at cryogenic and room temperatures7,20–22,40; however, none of them have addressed the high temperature mechanical properties of these HEAs. Therefore, this article uniquely addresses the issue of high temperature tensile strength of HEAs, especially the HESA in the HT-2 condition has performed exceptionally well comparing reported HEAs and advanced superalloys, Fig. 4. The dispersion of HE-FCC particles in ME-L12 precipitates can pin superdislocations, Fig. 5, attributing to an average increase of 113 MPa in yield strength comparing to that of the HT-1 condition. The distinctiveness of the hierarchical microstructure achieved in this work and the strengthening contribution associated with HE-FCC particles are discussed in this section.

Microstructure observations indicate that minor secondary L12 could be identified in HT-1 and HT-2 conditions, Figs. 1 and 2. In traditional cast superalloys, secondary L12 particles could form during the cooling process...
after solution heat treatment\textsuperscript{19,30,41–43}. And, HT-1 sample also possessed some secondary L\textsubscript{12} phase possibly during the air cooling process. Secondary L\textsubscript{12} particles appeared to coarsen after HT-2 process, Figs. 1 and 2. Although this microstructure evolution suggested the metastability of the FCC matrix, the amount of secondary L\textsubscript{12} particles identified was very minor in both HT-1 and HT-2 states in this work. Previous studies on superalloy have speculated that high fractions of secondary L\textsubscript{12} phase may decrease the net width of FCC channel and improve mechanical strength\textsuperscript{41}; in this work, secondary L\textsubscript{12} fractions were too little to affect mechanical properties. In the future, metastable FCC matrix channel should be explored further by heat treatment design to investigate if additional hierarchical microstructure strengthening can be achieved.

To the best of authors’ knowledge, only few literatures have reported the dispersion of FCC particles in L\textsubscript{12} matrix\textsuperscript{44–46}. Vogel et al.\textsuperscript{44} studied FCC particles in Ni\textsubscript{73}Al\textsubscript{13}Ti\textsubscript{9} L\textsubscript{12} matrix alloy with the partitioning coefficients of Al (0.6), Ti (0.42) and Ni (1.14); Ni showed preferential partitioning toward FCC phase, and high Ni content in the L\textsubscript{12} phase was deduced to be responsible for the FCC particle formation. Meher et al.\textsuperscript{45} designed a Re and Ru bearing Ni-based superalloy, and FCC particle dispersion in L\textsubscript{12} phase was found within the dendritic microstructure; the supersaturation of Co, Re and Ru in the L\textsubscript{12} phase was suggested to be the driving force for the formation of FCC particles. So, the supersaturation of FCC phase partitioning elements in the L\textsubscript{12} phase is the cause for the formation of the FCC particles as shown in Fig. 3. The supersaturation achieved in the L\textsubscript{12} phase of HT-1 sample promoted the formation of FCC particles and reduced the degree of supersaturation in the L\textsubscript{12} phase of the HT-2 state. The ME-L\textsubscript{12} phase in HESA possessed the formula (Ni, Co, Fe, Cr)(Al, Ti, Cr, Nb), which contained more FCC phase partitioning elements and a higher entropy value of 1.38 R than that of the L\textsubscript{12} phase in CMSX-4 (1.1 R)\textsuperscript{46}. High contents of Co and Fe are reported to narrow the L\textsubscript{12} phase field and affect phase boundary gradient\textsuperscript{47,48} between FCC and L\textsubscript{12} phases, hence high content of Co and Fe in HESA promotes the supersaturation in the L\textsubscript{12} phase during air cooling process of HT-1. By contrast, very minor fractions of FCC particles (< 0.3 vol%) was reported in the L\textsubscript{12} phase of CMSX-4\textsuperscript{46}, in this work HESA possessed much more nano FCC particles (4.3 vol%) in the ME-L\textsubscript{12} phase. So, the high entropy composition of HESA might have contributed to high degree of supersaturation in ME-L\textsubscript{12} phase and promoted the formation of relatively high fraction of HE-FCC particles. Moreover, alloying with Co and Fe were found to improve the intrinsic ductility of L\textsubscript{12}-Ni\textsubscript{3}Al phase\textsuperscript{49,50}, which could lead to ductile L\textsubscript{12} precipitate strengthened HEAs\textsuperscript{20}. Therefore, ME-L\textsubscript{12}-(Ni, Co, Fe, Cr)(Al, Ti, Cr, Nb) phase may contribute to the outstanding tensile elongation of HESA in additional to the ductile FCC phase as comparing to that of CMSX-4, Fig. 4b,c.

In a previous work on Ni\textsubscript{86.1}Al\textsubscript{8.5}Ti\textsubscript{5.4} alloy\textsuperscript{44,45,48,52}, the FCC particles could coarsen into directional plates in the L\textsubscript{12} matrix after prolonged ageing at 1,023 K. Meher et al.\textsuperscript{45} also found that ageing at 1,073 K could coarsen...
the FCC particles and decreased its fraction significantly. In this work, since the FCC particles in the L1_2 phase were in metastable states, all the FCC particle forming elements would diffuse across the L1_2 phase into the surrounding FCC matrix after prolonged ageing, so that the whole system could reach the equilibrium state, Fig. 3c. This could also explain the disappearance of HE-FCC particles in HESA at 1,173 K, Fig. 5f. Although the thermodynamic process is inevitable during thermal exposure, engineering alloys are rarely designed to be used in thermodynamic equilibrium state. In future work, it might be possible to prolong the rate of dissolution of FCC particles by further heavy elements addition, such as Mo, since Mo possesses low diffusivity and preferentially partition toward FCC phase. Addition of more heavy elements might be able to slow down the evolution of metastable hierarchical microstructure further and prolong its service life for high temperature application.

There is a very limited number of papers reporting benefit of the kind of hierarchical microstructure by experiments\(^{44, 45, 52–59}\). Vogel et al.\(^{48}\) showed a 50 HV hardness increase with the presence of FCC precipitates in Ni\(_{73}\)Al\(_{15}\)Ti\(_{12}\) L1_2 matrix alloy. In addition, Tian et al.\(^{53}\) reported a 78 MPa increase in compression strength at 973 K by the FCC precipitation in Ni\(_{73}\)Al\(_{15}\)Ti\(_{12}\) matrix alloy. Meher et al.\(^{45}\) showed the L1_2 phase with nano-sized FCC particles could possess enhanced coarsening resistance for the L1_2 phase. Smith et al.\(^{56}\) found that the FCC particles inside L1_2 phase might increase compression creep life from 20 to 90 h under 1,033 K/414 MPa. Although these studies demonstrated the strengthening contribution of the FCC precipitation in L1_2 matrix, these were mainly L1_2 matrix in bulk and could be brittle in tension. Notably, this article is the first to report tensile properties of this kind of hierarchical microstructure in tension, even at elevated temperatures.

To elucidate the outstanding tensile yield strength of HESA, each strengthening factor needs to be examined. The yield strength of HESA in the HT-2 state was 113 MPa higher than that of the HT-1 state. Since the phase compositions of both FCC matrixes and both L1_2 phases in the HT-1 and HT-2 states (Table 1) were very similar, their difference with respect to anti-phase boundary (APB) energies of L1_2 phase, stacking fault energies of the FCC matrix, and the lattice misfits should be minimal. This implies that the additional strengthening contributions from the intrinsic properties of HE-FCC matrix and ME-L1_2 phase could be excluded. So, higher strength in the HT-2 state might arise from higher ME-L1_2 phase volume fraction, larger ME-L1_2 phase size, and the presence of HE-FCC particles. According to the TEM analysis in Fig. 5b and e, paired dislocations resided within the ME-L1_2 precipitate. The pair-coupling model\(^{19, 27}\) could be applied to calculate contributing factors of L1_2 precipitate sizes and fractions on critical resolved shear stress (CRSS):

\[
\tau_c = \sqrt{\frac{3}{2}} \times \left( \frac{G_{bfcc}}{R} \right) \times \sqrt{\frac{F}{\pi^2}} \times \sqrt{\frac{2\pi RY_{APB}}{\gamma_{APB} \phi G_{bfcc}^2}} - 1
\]

where \(\tau_c\) is the CRSS, \(\gamma_{APB}\) is the APB energy, \(F\) is the L1_2 phase volume fraction, \(R\) is the L1_2 phase radius, \(b_{fcc}\) is the Burgers vector in the FCC matrix, \(G\) is the shear modulus and \(\phi\) is a dimensionless constant accounts for the elastic repulsion between the paired dislocations. To assist the calculation, the values of \(\gamma_{APB}, b_{fcc}, G\) and \(\phi\) were taken from the previous work\(^{27}\) and JMatPro calculations\(^{60}\) to be 0.22 J/m^2, 0.257 nm, 81 GPa and 1, respectively. The HT-1 condition contained an average L1_2 phase size of 143 nm and the L1_2 phase volume fraction of 68.4%; the HT-2 condition possessed an average L1_2 phase size of 153 nm and the L1_2 phase volume fraction of 70.6%. The calculation result indicates that the difference in CRSS between these two states was less than 1%, which means that slight difference in the L1_2 phase sizes and fractions between HT-1 and HT-2 had very limited impact on yield strength. Furthermore, previous study has shown the limited effect of secondary L1_2 particles on tensile properties\(^{45}\). With only little amount of secondary L1_2 particles (< 0.5 vol%) found in this study for both HT-1 and HT-2 samples, its influence on strength can be minimal. Therefore, the origin of additional strengthening contribution should be associated with the presence of HE-FCC particles inside ME-L1_2 phase at HT-2 state.

Previous studies from Nemoto et al.\(^{55}\), Hirsch et al.\(^{59}\), Pretorius et al.\(^{61, 62}\), Liu et al.\(^{63, 64}\) and Ardell et al.\(^{65}\) have deduced the strengthening contribution for ordered matrix by disordered particles; dislocations inside the L1_2 phase could be attracted to the disordered FCC particles that had no APB penalty and possessed shorter magnitude of burgers vector. This can explain the wavy superdislocations observed by TEM in Fig. 5d,e; to make dislocations wavy is to make it longer, hence energy requirement for dislocation motion would be higher with the presence of HE-FCC particles. In this work, Hirsch’s model\(^{59, 65}\), was applied to evaluate the strengthening contribution in CRSS as:

\[
\tau_c = \frac{1.1455 K}{r_{b_{L1_2}}} \times \sqrt{\frac{3f}{4\pi}} \times \sqrt{\frac{f\gamma_{APB}}{K}} - 1
\]

where \(\tau_c\) is the CRSS, \(r\) is the average radius of FCC particles, \(f\) is the volume fraction of FCC particles in the L1_2 phase, \(b_{L1_2}\) is the Burgers vector in L1_2 phase, \(\gamma_{APB}\) is the APB energy and \(K\) is the repulsive force between the edge superdislocations which can be stated as:

\[
K = \frac{G_{bfcc}^2}{2\pi (1 - \nu)}
\]

And, \(G\) is the shear modulus, \(b_{bcc}\) is the Burgers vector in FCC phase, \(\nu\) is the Poisson's ratio. To derive CRSS, the \(\gamma_{APB}, b_{bcc}, R_{L1_2}, G\) and \(\nu\) were taken from the previous work\(^{27}\) and JMatPro calculations\(^{60}\) as 0.22 J/m^2, 0.257 nm, 0.514 nm, 0.313 and 81 GPa, respectively. Also, the average HE-FCC particle radius and volume fraction were measured as \(r = 4\) nm and \(f = 4.3\)%, respectively. The predicted CRSS is 78 MPa, then, the CRSS value should be multiplied by 0.706 to fit the L1_2 phase fraction in HESA. The estimated increase in yield strength should be two times higher than the CRSS values, since the maximum value of Schmid factor is 0.5. Hence the increase in yield strength due to nano HE-FCC particles is 110 MPa, which is almost the same as the actual yield strength.
increase determined by experiment, i.e. 113 MPa, from the HT-1 state to the HT-2 state. This reveals the profound strengthening contribution of nano HE-FCC particles inside ME-L1$_2$ phase. The calculated data for the increase in yield strength as a function of the average FCC particle radius (Supplementary Fig. S4) shows the 4 nm of particle radii in the HT-2 state was very close to the peak strength of 110.18 MPa with 3.6 nm particle radius.

Hierarchical HESA has shown outstanding high temperature tensile properties in this work. Creep resistance is also an important property for high temperature application. Although the disappearance of FCC particles at 1,023 K after 200 h, Fig. 3c, suggested that HESA would have to rely on remaining L1$_2$ precipitates for strengthening, our previous work$^{27}$ has shown that without nano FCC particles, HESA possessed similar creep resistance as that of CMSX-2. The future challenge would be to prolong the thermal stability of FCC particles inside L1$_2$ phase, this might be the future direction to develop more creep resistant HESA. Alloy design may utilize more slow diffusing species such as Mo to improve the stability of hierarchical microstructure, and CALPHAD method will need to be utilized for alloy design to avoid formation of detrimental topologically close-packed (TCP) phases$^{19,30}$.

In summary, this work has introduced a novel hierarchical microstructure in HESA, which has the potential to surpass advanced superalloys in terms of tensile properties and cost-performance. HESA can be 8% lighter and 84% cheaper than that of commercial single crystal superalloy CMSX-4; at 1,023 K, the cost specific yield strength of HESA is 8 times that of CMSX-4 while its tensile strain can reach 20%, which is 3 times that of CMSX-4. HESA is strengthened by an interesting hierarchical microstructure consisted of FCC matrix, L1$_2$ precipitates inside FCC matrix, and FCC particles inside L1$_2$ particles. Superdislocations inside L1$_2$ precipitates appeared to be pinned by FCC precipitates, and this could explain why HESA could be stronger than those containing only FCC matrix + L1$_2$ precipitates. The high entropy composition of HESA is an important factor for achieving this hierarchical microstructure by phase transformation pathway through metastability. A template for lighter, stronger, cheaper, and more ductile high temperature alloy is proposed.

Methods
The HESA of interest is Ni$_{43}$Co$_{15}$Al$_{10.5}$Fe$_{7.4}$Cr$_7$Ti$_{3.8}$Nb$_{1.2}$Mo$_{0.9}$W$_{6.8}$ (at.%)$^{27}$. The configurational entropy ($\Delta S_{\text{conf}}$) of the nominal composition is 1.58 R, which under the HEA definition ($\Delta S_{\text{conf}} \geq 1.5$ R). HESA samples were fabricated into single crystal bars supplied by the Superalloys and High Temperature Materials Group in National Institute for Materials Science (NIMS), Japan. These single crystal bars were cast by Bridgman process. Laue X-ray method was used to make sure the <100> orientation of each single crystal bar. Then, the as-cast single crystal bars were solution heat treated by a single ramp process from room temperature to 1,500 K in 20 h and air cooled to room temperature (this heat treatment process is termed HT-1). After HT-1 process, an additional heat treatment was performed at 1,023 K for 20 h followed by water quench (the entire heat treatment process is termed HT-2). To further examine the microstructure evolution of HESA, a prolonged ageing after HT-1 was done at 1,023 K up to 200 h followed by water quench.

Scanning Electron Microscope (SEM: JEOL 7200F) and Transmission Electron Microscope (TEM: JEOL JEM-F200, 200 kV) were utilized for microstructure characterization. The specimens for SEM observation were polished and electro-etched in the etchant containing 20% H$_3$PO$_4$ + 80% H$_2$O at 2.5 V. Measurement of the phase fraction and size were based on the SEM images. The size and fraction measurements were performed by Nano-Measurer image analysis software and ImageJ software, respectively; at least three SEM images and over one hundred precipitates were measured. As for TEM analysis, specimens were prepared by grinding and twinjet polishing in the solution containing 10% HClO$_4$ + 90% C$_2$H$_5$OH at 30 V and 243 K. Atom Probe Tomography (APT) analysis was carried out by using laser assisted Local Electron Atom Probe (LEAP 5000 XS). APT specimens, along [100] direction, were prepared by a focus ion beam (FIB: Helios 650) system with the standard lift-out procedure. The APT data were collected under a laser mode with laser wavelength 355 nm, laser pulse energy 25 pJ, laser pulse rate 250 kHz, detection rate 1% and specimen temperature 30 K. And, the collected data were reconstructed and analyzed using CAMECA IVAS 3.8.2 software. Measurement of the FCC nano-particle size and fraction were based on the APT data.

High temperature tensile tests were conducted by Shimadzu testing machine. Flat tensile specimens were machined along the < 100 > direction with the gauge length, width and thickness to be 19.6 mm, 2.8 mm and 3 mm, respectively. The specimens were polished and tested at 298 K, 723 K, 923 K, 1,023 K and 1,173 K with a constant strain rate of 10$^{-3}$ s$^{-1}$.

Data availability
All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. The research data of this study is available from the corresponding author A.C.Y. upon reasonable request.

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**Author contributions**

Y.T.C., Y.J.C., H.M., J.W.Y. and A.C.Y. conceived this research. Y.T.C., T.S. and K.H. contributed to the APT measurements and analyses. Y.T.C., C.W.L and K.K. performed the high temperature tensile tests and data analysis. Y.T.C., Y.J.C., H.M., J.W.Y. and A.C.Y. contributed to the APT measurements of the CRSS and TEM observations of dislocation processes in Ni69Co9Al18Ti4. Y.T.C., Y.J.C., H.M., J.W.Y. and A.C.Y. conducted the TEM analysis. Y.T.C. and A.C.Y. wrote the manuscript. Y.T.C., A.C.Y., H.M., K.H. and J.W.Y. reviewed the manuscript. All authors participated in the discussion and finalized this paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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**Correspondence** and requests for materials should be addressed to A.-C.Y.

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