A mechanically strong and ductile soft magnet with extremely low coercivity

Soft magnetic materials (SMMs) serve in electrical applications and sustainable energy supply, allowing magnetic flux variation in response to changes in applied magnetic field, at low energy loss. The electrification of transport, households and manufacturing leads to an increase in energy consumption owing to hysteresis losses. Therefore, minimizing coercivity, which scales these losses, is crucial. Yet meeting this target alone is not enough: SMMs in electrical engines must withstand severe mechanical loads; that is, the alloys need high strength and ductility. This is a fundamental design challenge, as most methods that enhance strength introduce stress fields that can pin magnetic domains, thus increasing coercivity and hysteresis losses. Here we introduce an approach to overcome this dilemma. We have designed a Fe–Co–Ni–Ta–Al multicomponent alloy (MCA) with ferromagnetic matrix and paramagnetic coherent nanoparticles (about 91 nm in size and around 55% volume fraction). They impede dislocation motion, enhancing strength and ductility. Their small size, low coherency stress and small magnetostatic energy create an interaction volume below the magnetic domain wall width, leading to minimal domain wall pinning, thus maintaining the soft magnetic properties. The alloy has a tensile strength of 1,336 MPa at 54% tensile elongation, extremely low coercivity of 78 A m⁻¹ (less than 1 Oe), moderate saturation magnetization of 100 A m² kg⁻¹ and high electrical resistivity of 103 μΩ cm.

Lowest possible coercivity and highest possible electrical resistivity are primary goals for SMMs, to reduce hysteresis-related and eddy-current-related energy losses, noise and the associated material damage. Also, new SMMs with higher strength and ductility are needed, to operate under mechanically demanding loading conditions for safety-critical parts in transport and energy. High strength and ductility also serve as measures for many other mechanical properties, such as high hardness and fracture toughness. This multi-property profile creates a fundamental dilemma. The mechanical strength of metallic materials is produced by lattice defects and their elastic interactions with linear lattice faults that carry inelastic deformation, referred to as dislocations. However, the defects also interact with the magnetic domain walls and pin them. The loss in domain wall motion increases coercivity, so that the materials lose their soft magnetic features. Therefore, current SMMs follow the design rule of avoiding lattice defects to minimize coercivity. On the other hand, increasing the mechanical strength of an alloy requires enhancing its internal stress level through defects such as dislocations, grain boundaries and precipitates. This means that the task of making soft magnets mechanically strong is a trade-off between two mutually exclusive design strategies, namely, mechanical strength versus unaffected domain wall motion.

The theory of the grain size dependence of coercivity shows its proportionality to the sixth power of the grain size for the case of nanocrystalline materials, a relation that can also be applied to particles. Current design of SMMs has thus focused on using small particles (less than 15 nm) and grain sizes (less than 100 nm). According to magnetic strain theory, the coercivity depends on the energy required to displace domain walls to overcome lattice barriers. Here we introduce particles into a multicomponent massive solid solution matrix and increase their size from the commonly used range of 5–15 nm to 90–100 nm. With that, the internal stress level and the overall elastic coherency misfit energy are reduced through the smaller specific surface area (total surface area per unit of volume) of the particles, caused by the coarsening. We then propose that the particle design must follow four main rules. First, minimal pinning of domain walls requires a well-tuned and well-controlled particle size distribution with optimum balance between the decrease in specific surface area and the increase in magnetostatic energy during particle coarsening. Second, the particle size must remain smaller than the domain wall width to prevent strong pinning, that is, strong resistance against spin rotation. Third, the chemical composition and crystal structure of the particles determine their saturation magnetization; therefore, antiferromagnetic elements are usually excluded. Fourth, strengthening of the alloys is determined by the interaction between dislocations and particles and by the friction forces exerted on dislocations in the massive solid solution matrix. Thus, intrinsically strong intermetallic particles with minimal lattice misfit are targeted. These require high forces for dislocation cutting (providing strength), but repeated cutting...
by ensuing dislocations emitted by the same source shear them with gradual ease along the remaining and gradually reducing particle cross sections (providing ductility).

These different mechanism considerations had to be translated into a corresponding compositional alloy design concept. This is mainly guided by the requirement for (1) a ferromagnetic matrix with (2) high solid solution contribution and components that trigger the formation of (3) strong and stable intermetallic phases with (4) small lattice misfit relative to the matrix. These considerations have led us to the non-equiatomic iron–nickel–cobalt–tantalum–aluminium (Fe32.6Ni27.7Co27.7Ta5.0Al7.0 (at.%)) MCA. We synthesized the material in a vacuum induction melting furnace, followed by conventional hot rolling and homogenization (details of the processing procedures and chemical compositions are provided in Methods). Through further iso-thermal heat treatments (1–100 h at 1,173 K), we prepared samples with different average particle sizes, ranging from 24 ± 15 nm to 255 ± 49 nm (edge length is used to characterize the topological particle size). The particles have L12 structure and complex composition, as presented in detail below.

**Microstructure analysis**

Figure 1 shows the structural characterization of the MCA with medium particle size (M-MCA, in which ‘M’ stands for medium particle size) after annealing at 1,173 K for 5 h. The M-MCA shows an average grain size of 85.3 ± 25.6 μm (excluding annealing twin boundaries) according to electron backscatter diffraction (EBSD) analysis shown in Fig. 1a. The electron channelling contrast imaging (ECCI) analysis shows that the L12 particles have a high number density (7.2 ± 0.2) × 10^20 m⁻³ and a large volume fraction (55 ± 1%) in homogeneous distribution within the face-centred cubic (fcc) matrix (Fig. 1b). The lattice misfit (0.48%) between the fcc matrix and the L12 particles has been calculated using their lattice parameters acquired from the X-ray diffraction (XRD) patterns (Fig. 1c) by Rietveld simulation. Such a small lattice mismatch reduces the driving forces for further coarsening and the uniform dispersion prevents plastic localization at high strength 16. The central beam dark-field (DF) transmission electron microscopy (TEM) analysis shows that the average size of the L12 particles is 90.8 ± 35.8 nm (Fig. 1d). The corresponding selected-area electron diffraction (see inset of Fig. 1d) and high-resolution (HR)-TEM (Extended Data Fig. 1d) confirm the high coherency between the particles and the matrix.

The elemental partitioning between the L12 precipitates and fcc solid solution matrix is characterized by atom probe tomography (APT). Figure 1e shows the 3D distribution of the volume investigated by APT, highlighted by a set of isosurfaces delineating regions containing more than 25 at.% Fe. Figure 1f shows the 1D compositional profiles computed along the cylinder region in e (marked by the black arrow), showing the compositional changes across several interfaces. Error bars refer to the standard deviations of the counting statistics in each bin of the profiles.
Apart from these intragranular nanoparticles, we also observed two types of grain boundary variants: (1) coarse grain boundary particles (160.2 ± 55.3 nm) with the same crystal structure and composition as those inside the grains (Extended Data Fig. 1a–c) and (2) incoherent particles with a minor fraction (less than 0.3%) at the triple points of the grains with different structure (cubic Fd-3m, Fig. 1c) and composition (Ta_{40}Co_{26}Fe_{20}Ni_{11}Al_{3} (at.%), Extended Data Fig. 1e). These two types of particle are promoted by the high diffusion rate along the grain boundaries.

**Mechanical properties**

The current strategy of tuning the particle size also allows overcoming the strength–ductility trade-off, which is notable for advanced alloys with gigapascal-level strength. Figure 2a shows the tensile stress–strain curves of the M-MCA at room temperature, together with the average values for ultimate tensile strength ($\sigma_{UTS}$) and elongation at fracture ($\varepsilon_f$). The yield strength ($\sigma_y$) is 904 ± 11 MPa, with an ultimate tensile strength ($\sigma_{UTS}$) of 1,336 ± 21 MPa and an elongation at fracture ($\varepsilon_f$) up to 53.6 ± 1.5%, averaged from four tests. Accordingly, the M-MCA has a high $\sigma_{UTS} \times \varepsilon_f$ value of 71.6 GPa%. To show the improvement in strength and ductility achieved by the well-controlled particle size distribution, the mechanical response of a material variant with identical chemical composition, that is, Fe_{32}Co_{28}Ni_{28}Ta_{5}Al_{7} (at.%), but smaller particle size (S-MCA, annealed for 1 h, producing an average particle size of 24 nm), larger particle size (L-MCA, annealed for 100 h, average particle size of 255 nm) and the particle-free Fe_{35}Co_{30}Ni_{30}Ta_{5} (at.%) alloy^{11} are also presented in Fig. 2a.

Compared with the single-phase Fe_{35}Co_{30}Ni_{30}Ta_{5} (at.%) alloy with a relatively low value of $\sigma_y$ of 501 MPa (ref. 11), the notable increase in yield strength of the M-MCA can be attributed to the precipitation strengthening of the L1_{2} particles with a high volume fraction (55%). Especially, such an improvement in the strength of the M-MCA is achieved at no expense of ductility, which is fundamentally different from the case of the S-MCA, in which a pronounced loss in ductility is observed with increasing strength. The good ductility is correlated with the high work-hardening capability, as shown in Fig. 2b. Further increase of particle size at longer annealing time (100 h) in the L-MCA leads to the decrease of ductility (53%) and $\sigma_{UTS}$ (14%) as compared with those of the M-MCA. This is related to the mechanical weakness, strain localization and embrittlement in the particle-free zone adjacent to the grain boundary caused by solute depletion (Extended Data Fig. 2a). This interfacial weakening has been confirmed by the associated fracture
morphologies, with typical ductile fracture with fine dimples in the M-MCA material (see Fig. 2b inset) and intergranular fracture in the L-MCA material (Extended Data Fig. 2b).

To unravel the mechanisms responsible for the marked improvement in the strength–ductility combination of the M-MCA, we studied its deformation substructures using EBSD kernel average misorientation (KAM) analysis and ECCI at different hardening stages (Fig. 2c). In principle, high strength requires to impede dislocation movement, whereas good ductility needs mobility of dislocations and new production of dislocations17. At the early deformation stage, the M-MCA deforms by planar dislocation glide on {111} planes (Extended Data Fig. 3a–c), as commonly observed in fcc alloys 18. The dislocations extend through the grains. Corresponding pile-up configurations at the grain boundaries are shown by the higher KAM values (for example, at $\varepsilon_T = 5\%$; see KAM maps in Fig. 2c). The relatively large grain size (85.3 $\mu$m) of the current M-MCA enables higher mobility of dislocations compared with most of the previously reported strong MCAs that had smaller grain size19–21 (around 10 $\mu$m). Further straining refines the crystallographically aligned microbands and facilitates the shearing of L12 particles (for example, at $\varepsilon_T = 15\%$; see ECC images in Fig. 2c).

The quantification of the evolution of the average microband spacing shows a microband refinement process during straining (Extended Data Fig. 3d). The gradually reduced microband spacing causes higher passing stress and, thus, enhanced strain hardening. This has been proposed to explain the good strength–ductility combinations in high-manganese steels22 and MCAs23. Hence, the observed dynamic microband refinement and particle shearing are the prevalent deformation mechanisms in the current MCAs. No Orowan looping was observed, even when increasing the average particle size up to 255 nm for the L-MCA (Extended Data Fig. 2c), as the average particle spacing remains far below the critical value (3,094 nm, see Methods) for the activation of dislocation bowing around particles, a mechanism referred to as Orowan effect. Furthermore, the stress required for shearing particles in the M-MCA with a medium particle size (91 nm) with high volume fraction (55%) is 2.2 times larger than that for the S-MCA with a smaller particle size (24 nm) with low volume fraction (43 ± 1%) (see Extended Data Table 1). Therefore, the high critical shear stress required for cutting the L12 particles and the dynamic microband refinement during plastic deformation lead to the strong strain-hardening capacity of the M-MCA.
**Fig. 4 | Mechanical and magnetic features combined in the new Fe₃₂Co₂₈Ni₂₈Ta₅Al₇ (at.%) M-MCA material.** a, Ashby map compiling room-temperature ultimate tensile strength (σₓᵧ) × elongation at fracture (εₓᵧ) and intrinsic coercivity compared with other SMMs, such as Fe–Ni (refs. 24,25), Fe–Si (refs. 26,27), Fe–Co (refs. 28,29), Fe (ref. 30), amorphous alloys 42–44 and established MCAs 30–32. b, Ashby map showing σₓᵧ × εₓᵧ values versus average grain size compared with data for other strong and ductile MCAs 26–29,47–50.

**Magnetic properties**

Figure 3a,b shows the magnetic properties of the MCAs. All alloys show typical soft ferromagnetic behaviour. The M-MCA shows an excellent combination of extremely low coercivity ($H_c$) of 78 ± 3 A m⁻¹ (less than 1 Oe) and moderate saturation magnetization ($M_s$) of 100.2 ± 0.2 A m⁻¹ kg⁻¹. We identified a higher $M_s$ for the alloy variant with larger average particle size (see inset in Fig. 3a). The reason for this is the change in intrinsic magnetic behaviour, as indicated by the higher Curie temperature ($T_c$), shown by the thermomagnetic curves (Extended Data Fig. 4a). Two distinct changes of the slope, indicating two ferromagnetic phases, are observed in the S-MCA. By contrast, only one sharp drop is observed in the M-MCA and L-MCA materials, indicating the presence of only one ferromagnetic phase.

This is further confirmed by measuring the magnetic behaviour of the MCAs at elevated temperatures (Extended Data Fig. 4b). Considering that both the fcc and L₁₂ phases contain high concentrations of ferromagnetic elements, we investigated their individual magnetic response through casting both phases as separate bulk samples with their respective nominal compositions acquired previously from APT analysis (see Methods for details). The results show that the L₁₂ bulk phase is paramagnetic, whereas the fcc matrix is ferromagnetic in the M-MCA (Extended Data Fig. 4c). Owing to different partitioning, the magnetic behaviour of the L₁₂ phase varies from ferromagnetic in the S-MCA material variant to paramagnetic in the M-MCA and L-MCA materials. The mechanism behind this transition is the change in the intrinsic spin alignment, which is related to the change in chemical composition (Extended Data Fig. 5) and ordering during annealing. The overall increase in saturation magnetization of the MCAs as a function of particle coarsening is attributed to the change in fcc matrix composition owing to elemental partitioning (Extended Data Fig. 5), that is, specifically to the resultant higher concentration of (Fe+Co) in the fcc matrix. This effect enhances the total average magnetic moment per formula unit and leads to a higher $M_s$.

To gain further insight into the mechanism behind the magnetic response of the M-MCA, we investigated the domain structure using magneto-optical Kerr effect (MOKE) microscopy (Fig. 3c) under different applied magnetic field strengths. Starting from the AC demagnetized state to an applied field of 40 kA m⁻¹, the nucleation of magnetic domains is uniformly distributed within the grain. Further increasing the applied field (155 kA m⁻¹) leads to domain wall movement and growth of energetically favourable domains. The domains grow unaffected inside the grains but get impeded at grain and twin boundaries (Extended Data Fig. 6). Figure 3d summarizes the statistically averaged particle size distribution with respect to the coercivity of all the MCA samples at different annealing states. The data are acquired by developing an automated processing protocol, as shown in Extended Data Fig. 7. The coercivity first decreases from 763 A m⁻¹ (S-MCA, average particle size 24 nm) to 78 A m⁻¹ (M-MCA, average particle size 91 nm) and then increases to 1,745 A m⁻¹ (L-MCA, average particle size 255 nm). Both the average particle size and the grain size increase monotonously with increasing annealing time (Extended Data Table 1). Because the grain size of the MCA materials is above the critical single-domain size, its coercivity decreases with grain coarsening, following the model for the grain size dependence of the coercivity as $H_c = 1/D$ (in which $D$ is the grain size)\(^\text{13}\).

However, the magnitude of the decrease in coercivity owing to grain coarsening according to the model is negligible compared with the experimentally observed values: the difference according to the model calculation between the S-MCA and M-MCA material variants is 2 A m⁻¹, but the experimentally observed difference is 775 A m⁻¹. Accordingly, the energy required for the irreversible displacement of domain walls within the grain is the determining effect for the extremely low coercivity.

Therefore, the notable decrease in coercivity at the early particle coarsening stage ($≤$91 nm) is attributed to the gradual reduction of the overall coherency stresses between matrix and particles, owing to their average size increase. More specifically, the values of the specific surface area × integrated lattice misfit decrease from 1.09 × 10⁶ m⁻¹ in the S-MCA material to 4.08 × 10⁶ m⁻¹ in the M-MCA material. The dislocation density in the matrix decreases from 1.50 × 10¹⁴ m⁻² in the S-MCA material to 9.32 × 10¹³ m⁻² in the M-MCA material when increasing the annealing time from 1 h to 5 h (see Methods). The reduction in dislocation density lowers the associated elastic distortion fields that can pin the domain walls. Although the elastic distortion and dislocation density decrease with particle coarsening, the coercivity of the L-MCA material increases. Two main mechanisms are proposed to explain this. First, the average particle size and the associated strain field in the L-MCA becomes larger than the domain wall width ($\delta_w$), leading to domain wall pinning. The L-MCA has a $\delta_w$ of 117 nm and a much larger average particle size of 253 nm, whereas the average particle size of the M-MCA is 91 nm, which is below its $\delta_w$ of 112 nm (see Methods). Second, the increased magnetostatic energy ($E_s$) associated with the paramagnetic particles causes a stronger individual pinning effect of each particle.
on the domain wall motion. More specifically, the $H_c$ of the L-MCA is estimated to be 23 times larger than that of the M-MCA (see Methods). When considering only particle size, it should be noted that, for the M-MCA, the coherent particles distributed along the grain boundaries with an average size (160 nm) above the $\delta_\text{m}$ (112 nm) are expected to have a stronger pinning effect on the domain walls than those in the grain interiors (91 nm). However, these coarser particles occupy only a small fraction (1.2 ± 0.2%) in M-MCA, hence, with a negligible effect on domain wall motion.

These considerations show that the nanoscale size distribution of the particles must be carefully controlled to minimize their pinning effect on domain wall movement, which determines the coercivity of the alloy. This is achieved here by an optimal balance between the release of structural defects (for example, interfacial elastic distortion, dislocation density) just down to a level required to maintain high mechanical strength and the increase of the pinning effect from the magnetostatic energy, while keeping the particle size below the domain wall width during particle coarsening.

**Overview of several property profiles**

To highlight the good combination of mechanical and magnetic properties of the M-MCA with optimal particle size, we compare it with existing SMMs in an Ashby plot showing the $\sigma_{\text{UTS}} \times \nu I$ values against the $H_c$ (Fig. 4a). This comparison shows that the $\sigma_{\text{UTS}} \times \nu I$ value of the new M-MCA material outperforms all other SMMs. Notably, the $H_c$ of the new material is lower than that of all Fe–Ni (refs. 13, 14) alloys and other MCAs 26–37, comparable with that of Fe–Si (ref. 30), Fe–Co (refs. 38, 40) alloys and pure Fe (ref. 41). Amorphous and nanocrystalline soft magnetic alloys 42–45 can show ultralow $H_c$ (less than 10 A m$^{-1}$) and high mechanical strength, yet their limited ductility, damage tolerance and workability prohibit their application in cases in which the load path changes or high stresses, forming or machining are applied. A radar plot comparing the various soft magnetic and mechanical properties of the current M-MCA with several typical commercial SMMs is shown in Extended Data Fig. 8. Although the saturation induction ($B_s$) of the current M-MCA is not comparable with that of typical commercial SMMs (Extended Data Fig. 9), it has higher electrical resistivity ($\rho$) (103 ± 0.8 $\mu$Ω cm, see Extended Data Fig. 10), a feature that makes it suited for applications with AC conditions. The high $\rho$ of the M-MCA is expected to be derived from the high resistance to electron movement from the larger lattice distortion 46. Figure 4b compares the $\sigma_{\text{UTS}} \times \nu I$ values versus the grain size of the M-MCA material with recently reported strong and ductile MCAs 26–37, 21, 47–50. The analysis shows that the current alloy reaches high values of $\sigma_{\text{UTS}} \times \nu I$ even without the substantial contribution from grain boundaries, confirming the notable strengthening effect provided by the nanoparticles and the massive solid solution.

**Conclusions**

In summary, we have developed a material that unifies so far mutually exclusive properties, namely, high mechanical strength (1,336 MPa), high tensile ductility (54%), low coercivity (78 A m$^{-1}$), moderate saturation magnetization (100 A m$^{-2}$) and high resistivity (103 $\mu$Ω cm). We realized this in a new class of bulk SMMs through a nanoparticle dispersion with well-controlled size (91 nm), magnetic properties, coherency strain, strength and interface energy. The design strategy is opposite to that generally applied in conventional SMM design. Instead of using the smallest microstructure features (particle size <15 nm) to avoid magnetic wall pinning as in conventional SMMs, we have chosen a relatively coarse particle dispersion with tuned particle/matrix interfacial coherency stresses and paramagnetic properties to minimize magnetic pinning of domain walls on the one hand (soft magnetism) and maximize interaction strength with dislocations on the other (strength and ductility).

The infinite composition space of MCAs allows realizing materials with good combinations of soft magnetic and mechanical properties. The new alloy design approach allows for tailoring SMMs for magnetic parts exposed to severe mechanical loads, be it from manufacturing and/or during service, for which conventional SMMs are mechanically too soft or too brittle. Future efforts on developing advanced magnetic MCAs could target variants with further improved soft magnetic properties (for example, higher magnetic saturation) while preserving their outstanding mechanical properties, at lower alloy costs, and use high-throughput experiments combined with computational techniques, for example, machine learning, to accelerate the discovery of new alloy variants.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-022-04935-3.

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Methods

Materials preparation
The bulk MCA ingot with a predetermined nominal composition of Fe_{32.6}Co_{28.7}Ni_{27.7}Ta_{4.0}Al_{7.0} (at.%) was first cast in a vacuum induction furnace using pure metallic ingredients (purity higher than 99.8 wt.%) under high-purity argon (Ar) atmosphere. The as-cast ingot with dimension 40 mm × 60 mm × 20 mm (length × width × thickness) was then hot rolled at 1,473 K to an engineering thickness reduction of 50% (final thickness 10 mm). After hot rolling, the alloy sheets were then homogenized at 1,473 K for 10 min in Ar atmosphere, followed by water quenching. To obtain a wide size distribution of the particles, further isothermal heat treatments were conducted at 1,173 K, lasting from 1 h up to 100 h (1 h, 2 h, 5 h, 20 h, 50 h and 100 h) in Ar atmosphere and followed by water quenching. The exact chemical composition of the MCA measured by wet-chemical analysis is Fe_{32.6}Co_{28.7}Ni_{27.7}Ta_{4.0}Al_{7.0} (at.%), which is close to the predesigned composition. In addition, the bulk ingots (50 g) with compositions identical to that of the fcc (FeCoNi40Ta3) matrix phase and the L12 particle (Ni40Co26Ta3 (at.%) in the M-MCA derived from the APT analysis were also cast, respectively, by arc melting under Ar atmosphere. The ingots were remelted six times to achieve chemical homogeneity.

Analytical methods
XRD measurements were carried out in an X-ray system (Diffractometer D8 Advance A25-XI) with Cu Kα radiation (λ = 1.78897 Å, 35 kV and 40 mA). EBSD characterizations were conducted in a ZEISS Crossbeam focused ion beam scanning electron microscope at 15 kV. ECCI characterizations were performed using a ZEISS MERLIN high-resolution field-emission electron microscope at 30 kV. TEM analysis including selected-area electron diffraction was conducted in a JEOL JEM-2100 at 200 kV. Scanning transmission electron microscopy (STEM) images were collected using a probe-corrected Titan Themis 60-300 (Thermo Fisher Scientific) microscope. To modify the Z-contrast characteristics of the imaging mode, high-angle annular dark-field (HAADF) micrographs with a convergence angle of 23.8 mrad were acquired at 300 kV. The resulting collection angle ranges from 73 mrad to 200 mrad. Further energy-dispersive X-ray spectroscopy (EDS) analysis was conducted using Thermo Fisher Scientific’s Super-X windowless EDS detector at an acceleration voltage of 300 kV. APT experiments were performed in a local electrode atom probe (LEAP 5000 XR) from Cameca Instruments Inc. and analysed with commercial AP Suite software (v6.1). A pulse frequency of 125 kHz, a pulse energy of 40 pJ and a temperature of 60 K was used. The detection rate was kept at a frequency of 1 ion per 100 pulses.

Mechanical response measurements
Room-temperature uniaxial tensile tests were performed using flat tensile specimens at an initial strain rate of 1 × 10^{-3} s^{-1}. The tensile specimens were machined along the rolling direction from the alloy sheets by electrical discharge machining. The specimens with a total length of 60 mm, a gauge length of 30 mm, a gauge width of 5 mm and a thickness of 2 mm were used to investigate the bulk tensile properties. Further, smaller tensile specimens with a total length of 20 mm, a gauge length of 10 mm, a gauge width of 2 mm and a thickness of 1 mm were used to measure the local strain evolution by the digital image correlation method. At least four specimens for each condition were tested to confirm reproducibility. Further, to clarify the relation between global strain—stress behaviour and microstructure evolution, we also conducted interrupted tensile tests on different global true strains (that is, 5%, 15% and 25%), and the microstructures in the middle part of the deformed regions were then characterized accordingly.

Magnetic response measurements
The magnetic response was evaluated using the Quantum Design Magnetic Property Measurement System (MPMS) equipped with a standard Vibrating Sample Magnetometry (VSM) option. Cuboid specimens of dimensions 3 mm × 3 mm × 1 mm (length × width × thickness) were used for the measurements. The hysteresis loops M(H) were performed in an external magnetic field of ±800 kA m^{-1} at a constant magnetic-field-sweeping rate of 1 kA m^{-1} at 10 K, 300 K, 500 K and 800 K, respectively. The temperature dependence of magnetization M(T) analysis was carried out under an applied field of 40 kA m^{-1} from 10 K to 1,000 K with a temperature-sweeping rate of 10 K min^{-1}.

The magnetic domain patterns were characterized by a MOKE ZEISS microscope (Axio Imager.D2m). The domain wall movement was captured under an applied magnetic field of ±155 kA m^{-1}. Before the measurement, a background image was collected as a reference in the AC demagnetized state. The images acquired at different applied fields were enhanced by subtracting the background image using KerrLab software.

Physical response measurements
The electrical resistivity response was evaluated using the Quantum Design Physical Property Measurement System (PPMS) equipped with an Electrical Transport Option (ETO) option. Cuboid specimens of dimensions 6 mm × 2 mm × 1 mm (length × width × thickness) were used for the measurements. The resistivity ρ values are calculated by:

$$\rho = \frac{RA}{l}$$

in which R is the reported resistance, A is the cross-sectional area through which the current is passed and l is the voltage lead separation. The resistance value of each measurement is obtained by averaging those from 100 times of current passing. At least three specimens for each condition were tested.

Thermodynamic calculations
The equilibrium compositions of the fcc matrix and L12 particles in the FeCoNi40Ta3 (at.%) alloy at 1,173 K were calculated using the Thermo-Calc software (v.2022a) equipped with the High Entropy Alloys database TCHEA v.4.2. The calculated equilibrium compositions for the fcc and L12 phases in the FeCoNi40Ta3 (at.%) are Fe3 segregation of the particles.

Estimation of particle size (edge length)
The size distribution is statistically analysed by applying a batch image-processing protocol with several 2D-projected ECC images of all the MCA samples at different annealed states (Extended Data Fig. 7). The average particle size (edge length) of the L12 particles is estimated by:

$$d = \sqrt{\frac{\sum S}{I}}$$

in which d is the average particle size, Si is related to the area of each particle acquired from the 2D-projected ECC images by the batch image-processing protocol and I is the total particle number. The particle size of the M-MCA is also characterized by DF-TEM (Fig. 1d) and bright-field TEM (Extended Data Fig. 1). The TEM results fit well with the value acquired by ECC images.

Estimation of interfacial coherency stress
The coherency stress at the L12–fcc interface is determined by integrating the lattice misfit across the interface as:

$$\delta = S_{L12/fcc} \sum \delta_s$$

in which S_{L12/fcc} is the L12–fcc interface area related to the average particle size (d), the volume fraction of the L12 particles (f) and the overall volume (V) as follows:
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\[ S_{L1/fcc} = \frac{\sqrt{3}}{2a^2} \cdot 6d^2 = \frac{6\sqrt{3}f}{d} \]

\( \delta \) is the varying lattice misfit as a function of distance \( x \) from the L12–fcc interface determined by the following equation:

\[ \delta_x = 2 \times \left( \sum \frac{a_{L1}^x - a_{fcc}^x}{a_{L1}^x + a_{fcc}^x} \right) \]

\( a_{L1}^x \) and \( a_{fcc}^x \) are the lattice parameters of the L12 and fcc phases at the interface region, respectively. Such values were calculated using the L12–fcc interfacial chemical compositions acquired from the APT datasets with Vegard’s relation49:

\[ d_{L1} = d_{0L1} + \sum f_{L1}t_{L1}x_{L1} \]

\[ d_{fcc} = d_{0fcc} + \sum f_{fcc}x_{fcc} \]

in which \( d_{L1}^x \) and \( d_{fcc}^x \) are the average lattice parameters for the L12 particles and fcc matrix, respectively, derived from the Rietveld simulation based on the XRD measurements, as shown in Extended Data Table 1. \( f_{L1}^x \) and \( f_{fcc}^x \) are the Vegard coefficients for the L12 and fcc phases, respectively, obtained from the ordered Ni3Al phase and the disordered fcc phase in the Ni-base superalloys50, as shown in Extended Data Table 2. Note that the above-calculated lattice misfit \( \delta \) represents the theoretical unconstrained state. This can be related to the constrained misfit (\( \varepsilon \)) by elasticity theory as below51:

\[ \varepsilon = 3 \frac{\delta}{2a} \]

The estimated interfacial constrained misfit value is 1.09 × 10^6, 4.08 × 10^5 and 1.96 × 10^5 for the S-MCA, M-MCA and L-MCA, respectively. On the basis of the experimental observation (Fig. 2c and Extended Data Table 1), particle shearing is the primary deformation mechanism in the investigated MCAs. The strengthening contribution of particle shearing (\( \Delta \tau \)) is estimated according to56:

\[ \Delta \tau_{\text{shearing}} = k \sqrt{\frac{F}{b \cdot 2 \lambda}} \]

by using the relation \( F = d^{1/2} \) and introducing constant \( k \). The effect of particle strengthening of the M-MCA is then estimated to be two times larger than that of the S-MCA (\( \Delta \tau_{S-MCA}/\Delta \tau_{M-MCA} = k_{S-MCA}/k_{M-MCA} \)).

When considering the volume fraction of the particles to be constant, the mean spacing of the particles increases with increasing particle size. As a result, the force required for shearing particles increases until the Orowan mechanism is activated, that is, dislocations bowing the particles becomes easier than shearing. The critical mean spacing of the particles is determined by56:

\[ \Delta = \frac{F}{b \cdot 2 \lambda} = \Delta \tau_{\text{Orowan}} = \frac{Gb}{2\lambda} \]

\( G = 84 \text{ GPa} \) is the adopted shear modulus57. Consequently, the critical mean spacing of the particles is calculated as 3.094 nm. However, in the current MCAs, the volume fraction of the L12 phase is not constant even after annealing at 1,173 K for 100 h. This is because the alloys have not yet reached the thermodynamical equilibrium state, as indicated by both thermodynamical calculations and APT analysis (Extended Data Fig. 5).

**Estimation of magnetostatic energy**

The magnetostatic energy (\( E_s \)) determines the coercive force that interacts between the paramagnetic particles (for M-MCA and L-MCA) and domain wall movement according to the formula58:

\[ E_s = \frac{1}{2} \mu_0 M_s^3 d^3 \]

in which \( \mu_0 = 4\pi \times 10^{-7} \text{ H m}^{-1} \) is the permeability of vacuum, \( d \) is the average particle size and \( M_s \) is the saturation magnetization of the fcc matrix. For the M-MCA and L-MCA, in which the L12 phase is paramagnetic (Extended Data Fig. 4), the \( M_s \) of the fcc matrix is considered as the overall \( M_s \) of the alloy. The values of \( E_s \) markedly increase with increasing particle size, that is, it varies from 1.57 × 10^14 (M-MCA) to 3.65 × 10^13 (L-MCA). The notable increase in magnetostatic energy results in a strong magnetic pinning effect.

**Estimation of domain wall width**

Strong pinning arises and results in the deterioration of coercivity when the microstructure defects have a comparable dimension to the domain wall thickness (\( \delta_w \)). As a result, the estimation of the \( \delta_w \) helps to understand the extremely low coercivity in the current work is given by59:

\[ \delta_w = \pi (A_{eff}/K_I)^{1/2} \]

in which \( A_{eff} = k_b T_s / 2a_0 \) is the exchange stiffness, \( k_b = 1.380649 \times 10^{-23} \text{ J K}^{-1} \) is Boltzmann’s constant and \( T_s \) and \( a_0 \) are the Curie temperature and lattice parameter of the fcc matrix, respectively (Extended Data Fig. 4d and Extended Data Table 1). \( K_I \) is the first magnetocrystalline anisotropy constant. The value of \( K_I \) (M-MCA) is taken from the Co–Fe system50,51 based on the composition of the fcc matrix (Fig. 1f) as 10.4 kJ m^3 (Al and Ta are non-ferromagnetic elements that do not show any magnetic moment, the chemical composition of the fcc phase Fe_{58}Co_{30}Ni_{12}Al_{1.34}Ta_{0.65} (at.%), in the M-MCA is thus considered as Co_{58}(Fe+Ni)_{30} (at.%)). The domain wall thickness of the M-MCA is therefore estimated to be 112 nm. Similarly, the domain wall thicknesses of the S-MCA and L-MCA are calculated as 103 nm and 117 nm, respectively.
Data availability

Data and codes are available from the authors.

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Author contributions

L. H., Z. L. and D. R. designed the research project; L. H., N. J. P. and F. M. characterized the alloys; L. H., I. R. S. F., Y. W. and B. G. analysed the data; L. H., Z. L., O. G. and D. R. conceptualized the paper. all authors contributed to the discussion of the results.

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Competing interests

The authors declare no competing interests.

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Extended Data Fig. 1 | STEM-HAADF micrographs and corresponding STEM-EDS analysis showing the structure and chemical composition of the particles with different morphologies. a, Overview of a STEM lamella showing the variation in particle morphology near the grain boundary. b, Enlarged view (red frame in a) showing particles inside the grains and along the GBs as well as their respective EDS elemental maps. c, Enlarged view (orange frame in a) illustrating a heterogeneous non-coherent particle at the triple junction of GBs and its EDS elemental maps. d, Atomically resolved HAADF-STEM micrographs and fast Fourier transforms of representative matrix and particle regions, indicating that the particles (ordered, bright) are coherent with the matrix (disordered, dark). The micrographs were obtained along the [011] zone axis. e, Comparison of the average chemical composition of different particles and the matrix determined by the EDS mapping. GB, grain boundary.
Extended Data Fig. 2 | Deformation mechanisms of the investigated MCA.

a, ECC images of the S-MCA and L-MCA. The formation of the particle-free zone can be identified in the L-MCA.
b, Fracture surface morphologies of the L-MCA. The enlarged view shows the presence of intergranular fracture at the grain boundary.
c, ECCI analysis showing the microbands and the shearing of the particles in the S-MCA and L-MCA.
Extended Data Fig. 3 | Evolution of the microband refinement. a–c, ECCI analysis showing the evolution of microbands with increasing local strain ($\varepsilon_{loc}$) at 10%, 30% and 80%, respectively. d, Quantification of the decreases in average microband spacing with increasing local strain. The local strain has been estimated by the digital image correlation method.
Extended Data Fig. 4 | Magnetic behaviour of the MCAs in a wide temperature range. 

**a**, $M(T)$ curves measured in the temperature range 10–1,000 K under a magnetic field of 40 kA m$^{-1}$. **b**, $M(H)$ loops of the MCAs measured at 800 K, confirming that the second phase in the S-MCA is ferromagnetic, with a $T_c$ higher than 800 K. The measuring temperature of 800 K was selected on the basis of the $M(T)$ results as marked by a black dashed line in **a**. **c**, $M(H)$ loops of the individual cast fcc and L1$_2$ phases in the M-MCA measured at 300 K. **d**, Temperature dependence of the $M_s$ of the investigated MCAs.

| Alloy  | $M_s (\text{Am}^2/\text{kg})$ @ 10, 300, 500, 800 | $T_c$ (K) |
|--------|--------------------------------|-----------|
| S-MCA  | 121.6, 97.2, 76.0, 9.3 | $T_{c1} = 667.8, T_{c2} > 800$ |
| M-MCA  | 123.6, 99.8, 77.1 | / 693.8 |
| L-MCA  | 125.3, 102.2, 78.6 | / 704.5 |
Extended Data Fig. 5 | Atom probe characterization showing the tomography and composition of the L1₂ particles and fcc matrix in the MCAs. a, APT datasets showing the reconstructed S-MCA and L-MCA containing L1₂ particles. b, Average chemical compositions of the fcc and L1₂ phases acquired from the APT subvolumes.
Extended Data Fig. 6 | MOKE microscopy observation showing the pinning of the magnetic domain wall during the magnetization process in the M-MCA.

GBs and TBs act as the pinning sites for domain wall movement. The applied magnetic field is horizontal to the viewing plane. GB, grain boundary; TB, twin boundary.
Extended Data Fig. 7 | Process for analysing the size distribution of the L1\(_2\) particles by batch image-processing protocol. First, an edge detection algorithm was applied to the raw images to isolate particles from the matrix. Second, a contour detection algorithm was used to search for closed contours. Third, the particle shape was approximately evaluated and converted into a square shape. Finally, the size distribution of closed contours (particles) was computed. The algorithm was constructed using a Python\(^\text{\textregistered}\). For each sample, the particle size distribution was derived from more than 20 ECC images of different sample regions.
Extended Data Fig. 8 | Comparison of the mechanical and functional property spectra of the new MCAs and existing commercial SMMs. The representative commercial SMMs include silicon steel (Fe–4 wt.%Si (ref. 38)), a Fe–Ni alloy (Fe–78.5 wt.%Ni (ref. 38)) and a Fe–Co-based alloy (Fe–49 wt.%Co–2 wt.%V (ref. 63)).
Extended Data Fig. 9 | Ashby map showing $\sigma_{UTS} \times T_e$ as a function of saturation induction ($B_s$). The investigated M-MCA is compared with a wide class of conventional SMMs, including Fe–Ni (refs. 24,25), Fe–Si (ref. 38), Fe–Co (refs. 39,40), Fe (ref. 41), amorphous alloys 42–45 and other MCAs 26–37. The $B_s$ is calculated using the equation $B_s = 4\pi M_s \rho_m / 10,000$, in which $\rho_m = 8.6$ g cm$^{-3}$ is the mass density of the M-MCA.
Extended Data Fig. 10 | Electrical resistivity of the Fe$_{32}$Co$_{28}$Ni$_{28}$Ta$_{5}$Al$_{7}$ (at.%) MCAs. The investigated MCAs are compared with a wide class of conventional SMMs, including pure iron, Fe–Ni (refs. 24,25) alloys, Fe–Si (ref. 38) alloys, Fe–Co (refs. 39,40) alloys and other established MCAs 26,28,29,64–66. The inset shows the evolution of electrical resistivity with increasing annealing time of the current MCAs.
Extended Data Table 1 | Structural parameters of the present MCAs

| Alloys                  | S-MCA   | M-MCA   | L-MCA   |
|-------------------------|---------|---------|---------|
| $a_0^{\text{fcc}}$ (Å)  | 3.6062± | 3.6050± | 3.6030± |
|                        | 0.0001  | 0.0001  | 0.0001  |
| $a_0^{\text{Ll2}}$ (Å) | 3.6223± | 3.6225± | 3.6228± |
|                        | 0.0001  | 0.0001  | 0.0001  |
| Lattice misfit, $\delta_0$ (%) | 0.445  | 0.484  | 0.548  |
| Volume fraction of the particle (%) | 43.2 ± 0.5 | 55.2 ± 0.8 | 60.5 ± 1.0 |
| Average particle size (nm) | 23.9 ± 12.3 | 90.8 ± 35.8 | 255.5 ± 50.1 |
| Average grain size (µm) | 72.6 ± 16.9 | 85.3 ± 25.6 | 150.2 ± 50.4 |
| Mean spacing of the particles (nm) | 192.8 ± 74 | 65.5 ± 40.6 | 30.3 ± 17.3 |
| Specific surface area (m$^2$/m$^3$) | $1.08 \times 10^8$ | $3.65 \times 10^7$ | $1.42 \times 10^7$ |
| Specific surface area $\times$ integrated lattice misfit (%-m$^2$/m$^3$) | $1.09 \times 10^6$ | $4.08 \times 10^5$ | $1.96 \times 10^5$ |
| Microstrain, $\varepsilon_s$ | 0.00084± | 0.00085± | 0.00087± |
|                        | 0.00002 | 0.00002 | 0.00002 |
| Crystallite size, $D$ (nm) | 76 ± 4 | 124 ± 9 | 220 ± 40 |
| Dislocation density (m$^{-2}$) | $1.50 \times 10^{14}$ | $9.32 \times 10^{13}$ | $5.38 \times 10^{13}$ |
### Extended Data Table 2

Vegard coefficients in the disordered fcc phase and the ordered Ni$_3$Al phase taken from the Ni-base superalloys

| Element (at%) | Fe  | Ni  | Co  | Ta  | Al  |
|--------------|-----|-----|-----|-----|-----|
| $\Gamma_i^Y$ ($\Gamma_i^{fcc}$) | 0.123 | 0   | 0.02 | 0.725 | 0.179 |
| $\Gamma_i^{Ni_3Al}$ ($\Gamma_i^{L1_2}$) | 0.01 | 0   | 0   | 0.51 | 0   |