Structure and magnetism of electrospun porous high-entropy $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})_3O_4$ and **(Cr1/5Mn1/5Fe1/5Ni1/5Zn1/5)3O⁴ spinel oxide nanofibers**

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High-entropy oxide nanofibers, based on equimolar (Cr,Mn,Fe,Co,Ni), (Cr,Mn,Fe,Co,Zn) and (Cr,Mn,Fe,Ni,Zn) combinations, were prepared by electrospinning followed by calcination. The obtained hollow nanofibers exhibited porous structure consisting of interconnected nearly strain-free $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})_3O_4$ and (Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})₃O₄ single crystals with pure Fd-3m spinel structure. Oxidation state of the cations at the nanofiber surface was assessed by x-ray photoelectron spectroscopy and cation distributions were proposed satisfying electroneutrality and optimizing octahedral stabilization. The magnetic data are consistent with a distribution of cations that satisfies the energetic preferences for octahedral vs. tetrahedral sites and is random only within the octahedral and tetrahedral sublattices. The nanofibers are ferrimagnets with relatively low critical temperature more similar to cubic chromites and manganites than to ferrites. Replacing the magnetic cations Co or Ni with non-magnetic Zn lowers the critical temperature from 374 K (Cr,Mn,Fe,Co,Ni) to 233 and 105 K for (Cr,Mn,Fe,Ni,Zn) and (Cr,Mn,Fe,Co,Zn), respectively. The latter nanofibers additionally have a low temperature transition to a reentrant spin-glass-like state.

Introduction

High-entropy materials (HEMs) are a new class of materials consisting in multicomponent solid solutions. Multicomponents, present in near-equimolar proportions, are randomly distributed in the lattice. $1,2$ They are attracting enormous scientific attention due to their unique structure, enhanced properties, novel functionalities and potential for application in many different fields ranging from catalysis to energy storage and conversion.³⁻²⁰

The class of HEMs encompasses materials having bonding character spanning from metallic to ionic and a variety of structures (hexagonal, cubic, rock-salt, fluorite, perovskite, spinel …), featured by high-entropy mixing at one or two cation sublattice(s).²¹ HEMs comprise high-entropy alloys (HEAs)^{1,2} and high-entropy ceramics (HECs), $2^{1,22}$ in turn including sulfides, 2^{3} silicides,²⁴ nitrides,²⁵ carbides,²⁶ oxides^{10,27-29} and other compounds.³⁰

After the synthesis of the first single-phase multicomponent rock-salt oxide via solid state sintering from an equimolar mixture of binary oxides in 2015, ^{28,31} many physical or chemical routes have been followed in the last years to prepare HEMs in various forms (bulk, nanoparticles, nanosheets ...).^{22,32-38} Very recently, it has been demonstrated that high entropy oxides (HEOs) in the form of nanofibers (NFs) can be successfully prepared by the electrospinning technique. $39-43$ Zhao et al. 39 have reported the synthesis of electrospun high entropy $(La_{1/5}Sm_{1/5}Eu_{1/5}Gd_{1/5}Tm_{1/5})₂Zr₂O₇$ NFs with excellent thermal stability, whereas Xing et al.⁴⁰ have produced $(Y_{1/5}Yb_{1/5}Sm_{1/5}Eu_{1/5}Er_{1/5})_2O_3$ NFs with cubic bixbyite structure.

High entropy rock-salt (Mg_{1/5}Co_{1/5}Ni_{1/5}Cu_{1/5}Zn_{1/5})O NFs prepared via electrospinning have shown interesting performance as active anode materials in rechargeable lithium ion batteries.^{41,42} Electrospinning is a simple, versatile and costeffective technique, widely adopted for the production of porous high aspect ratio nanostructures and suitable for the manufacturing of NFs on an industrial scale due to its scalability.⁴⁴⁻⁴⁸ This paper deals with the preparation and thorough characterization of electrospun porous NFs of highentropy (Cr,Mn,Fe,Co,Ni), (Cr,Mn,Fe,Co,Zn) and (Cr,Mn,Fe,Ni,Zn) oxides having spinel structure. This structure comprises a slightly distorted face-centered cubic (fcc) lattice of O^{2-} anions, where cations occupy tetrahedral 8a sites and octahedral 16d sites, thus forming two cation sublattices (Fig. S1).⁴⁹ High entropy spinel oxides (HESOs) have been the focus of in-depth studies.⁵⁰⁻⁵⁷

Among them, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, the first HESO to be reported,⁵⁰ has received particular attention and has been evaluated for several applications, such as electrocatalysis^{53,54} and energy storage.52,55,58 The magnetic properties of HESOs have already been summarized.^{59,60} They are governed by the oxido-mediated superexchange interactions within and between the two cation sublattices. The magnetic properties of (Cr1/5Mn1/5Fe1/5Co1/5Ni1/5)3O4, in both bulk and nanostructured form, ⁶¹⁻⁶⁵ have been the subject of several studies, which have

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firmly established that $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$ is a ferrimagnet below ca. 400 K. The low temperature behavior has been attributed to antiferromagnetically-coupled regions rich in Cr or Mn^{62} or locally varying inter-site exchange interactions. 65 Mao et al.⁶¹ have reported the magnetization isotherms at room temperature (RT) of $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})$ ₃O₄ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ and shown that substitution of nonmagnetic Zn^{2+} for magnetic Co^{2+} and Ni^{2+} decreases the saturation and remanent magnetization, and coercive field.

Here, a detailed magnetic characterization of electrospun $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})_3O_4$ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ NFs is presented. These HESO-NFs display a rich and diverse magnetic behavior depending on composition. The present study confirms that (Cr1/5Mn1/5Fe1/5Co1/5Ni1/5)3O⁴ NFs have ferrimagnetic structure between 5 and ≈400 K but, at variance with previous interpretation, the structure is non-collinear, i.e., the sublattice magnetizations are not parallel. We prove that both $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})$ ₃O₄ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ NFs are paramagnetic at RT and upon cooling they first assume a non-collinear ferrimagnetic structure and then a conical/spiral or canted ferrimagnetic structure with spin-glass-like character. The qualitative similarity of the Zn-containing NFs is, however, contrasted by the largely different magnetic parameters and transition temperatures. Therefore, the magnetic structure of HESO-NFs can be easily tuned by changing just a single cation. This study paves the way to the obtainment of one-dimensional nanostructures with controlled and easily tailored magnetic properties, which is of great applicative interest as they can potentially be used as building blocks for next-generation electromagnetic devices,⁶⁶ in magnetic sensors, and flexible magnets.⁶⁷

Results and discussion

Morphology of the HESO-NFs.

Based on the varying cations in the transition metal (TM) combinations, the produced (Cr,Mn,Fe,Co,Ni), (Cr,Mn,Fe,Co,Zn) and (Cr,Mn,Fe,Ni,Zn) HESOs were coded as CoNi, CoZn and NiZn (Fig. S2e).

Fig. S3 displays the results of the scanning electron microscopy/energy-dispersive X-ray spectroscopy (SEM/EDX) analysis on the as-calcined samples. SEM micrographs demonstrate the formation of NFs for all compositions. Nonetheless, slight changes are noticed in their morphology (Fig. S3a‒c). In sample CoNi (Fig. S3a), where bundles of NFs alongside isolated NFs are observed, the fibers look relatively uniform in diameter along their axis. On the contrary, in samples CoZn (Fig. S3b) and NiZn (Fig. S3c), the diameter is not uniform along the NF axis; the variation is more marked in the case of the latter sample. The center of the NF diameter distributions Fig. S3d–f), as measured by the image analysis software of the microscope, decreases in the order CoNi > CoZn > NiZn. EDX analysis (Fig. S3g-i) proves the spatially uniform distribution of oxygen and TMs throughout all samples at the micro-scale.

Fig. 1 and S4-S6 summarize the results of high-resolution transmission electron microscopy/scanning transmission electron microscopy (HRTEM/STEM)/EDX analysis. The TEM images with lower magnification (Fig. S4a,c,e) substantially confirm the morphological picture emerged from the SEM analysis. Apart from the above-described slight differences, the morphology of the NFs exhibits common features to all the compositions. The fibers, with diameters in the 150-450 nm range, are micrometer long and featured by a coral-like structure (Fig. 1a-c), as frequently observed in electrospun oxides.⁶⁸−⁷⁰ They consist of interconnected grains with size ranging between a few tens of nm to the hundred nm range.

Regardless of their composition, the primary oxide grains have well defined polyhedral shape, with more or less rounded vertices (Fig. 1d-h and S4). The oxide grains are crystalline with spinel structure and randomly oriented, as proved by the diffraction rings in the selected-area electron diffraction (SAED) patterns (Fig. 1i and S6a,d) for isolated NFs (Fig. 1j and S6b,e). They exhibit a spinel structure. The STEM/EDX elemental maps (Fig. 1k and S6c,f) reveal that in all samples the spatial distribution of metals and oxygen is homogeneous even at the nanometer scale.

The compositional analysis by STEM/EDX highlights the achievement of nearly equiatomic combinations of TMs (Fig. 1a–c), namely $(Cr_{0.18}Mn_{0.21}Fe_{0.18}Co_{0.22}Ni_{0.21})$, (Cr0.19Mn0.22Fe0.18Co0.22Zn0.19) and (Cr0.20Mn0.16Fe0.19Ni0.22Zn0.23) for samples CoNi, CoZn and NiZn, respectively.

The homogeneity of the cation distribution is further confirmed by the projection analysis of the EDX maps (Fig. 2). The normalized profiles of all the metals and oxygen overlap to within the experimental noise, except for a small discrepancy of the Zn profile in the CoZn NF (Fig. 2b). Model profiles for solid or hollow NFs were fitted to the experimental oxygen profiles of selected NFs (Fig. 2d-f). Comparing results for each NF, the NFs were assigned to three morphological categories, namely, quasi-solid, irregularly hollow, and hollow NF. All morphologies were observed for each NF composition and irregularly hollow NFs seem predominant in all samples. In addition to categorizing the NP morphology, the fitting procedure allowed measuring the NF outer and inner diameter of the selected NFs. The range of both diameters are collected in Table S2 along with that of the calculated shell thickness, and cross-section sketches of the three NF types are shown in Fig. 2g-i.

The decreasing trend of the outer NF diameter CoNi > CoZn > NiZn resulting from projection analysis agrees with the NP diameter distribution from the analysis of SEM images. Electron diffraction patterns proved that the NF comprise primary particles with spinel structure. HRTEM images of samples CoNi, CoZn and NiZn display crystal lattice fringes in primary particles that are suitably oriented with respect to the electron beam. Geometrical phase analysis (GPA) of the crystal lattice fringes allowed monitoring the crystallinity of the primary particles and the crystallographic relationship with neighboring particles (Fig. 3). Fringes were observed for the {111}, {220}, and {311} planes. No difference among the three NF samples was found, so the following discussion applies to all NF types. All fringe-displaying particles are single crystals and

are free from significant strain, except for thin strained layers where

the particles are in contact (Fig. 3g‒h)**.**

Fig. 1 Results of the HRTEM/STEM/EDX analysis on samples (a,d,g,h,l) CoNi, (b,e,i-k,m) CoZn and (c,f,n) NiZn. (a-h) HRTEM images; (i) SAED pattern for (j) an isolated NF, (k) corresponding STEM/EDX elemental maps and (I-n) relative concentrations of the TMs.

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Fig. 2 Projection analysis of the EDX elemental maps. (a-c) Projected profiles of a quasi-solid NF (a, CoNi), irregularly hollow NF (b, CoZn), and hollow NF (c, NiZn). (d-f) Experimental oxygen profiles (circles) and best-fit models (line). (g-i) Sketch of the typical cross section of (g) CoNi, (h) CoZn and (i) NiZn NFs. Note the different length scales in panels (a-f).

Fig. 3 Geometrical phase analysis of the NFs. (a-c) Intensity-colorized maps, (d-f) lattice rotation maps (°), and (g-i) lattice strain maps (% cell size). All primary particles are single crystals. The particle in panels c, f, i has no crystallographic relationship with its numerous neighbors whereas the particles in panels b, e, h are equi-oriented. The particle in panels a, d, g displays significant strain along its long edges that are not in contact with other particles. The shown data refer to NFs (a,d,g) CoNi, (b,e,h) CoZn and (c,f,i) NiZn.

A single case in a CoNi NF was observed, where a 40x60 nm polyhedral, round-edged particle displays compressive strain along its free edges (Fig. 3g). It is interesting that in all cases no significant strain is observed in the particle interior. In general, equal orientation of neighboring particles is not observed by HRTEM, even when they are face-sharing. Only in a few cases, equi-oriented particles are detected (Fig. 3b,e,h). In conclusion, GPA of HRTEM fringes showed that (i) the characteristics of the primary particles are common to the three NF samples, (ii) the primary particles are single crystals, and (iii) neighboring particles usually have different lattice orientation, the opposite is rarely detected in HRTEM images.

Crystalline phase of the oxide.

Fig. 4 shows the XRD patterns of the electrospun HESO-NFs. In all samples, the diffraction peaks correspond to the reflections of fcc spinel structure, belonging to Fd-3m space group (JCPDS no. 22-1084)^{51,52,56,61,71-76} No additional peaks attributable to secondary phases are detected, indicating that a single-phase is formed for any composition, namely $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})_3O_4$ $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})_3O_4$ and (Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})₃O₄ for samples CoNi, CoZn and NiZn, respectively. This finding is in agreement with the outcomes provided by the SAED patterns. The average size of HESO crystallites (i.e., coherence length) is calculated from the full width at half maximum (FWHM) of the (331) peak, through the Scherrer's equation, $d = k\lambda/\beta cos\theta$, where k (0.9) is the shape factor, λ is the wavelength of x-ray radiation, β is the FWHM in radians and θ is the Bragg's angle.⁷⁷

Fig. 4 XRD patterns of the HESO-NFs.

The obtained values increase in the order CoZn (83.7 nm) < CoNi (97.0 nm) < NiZn (98.5 nm), respectively.

The lattice constants, as inferred from Rietveld analysis (Fig. S7), increase in the order CoNi (8.3215(8) Å) < NiZn (8.3256(6)

Å) < CoZn (8.3712(8) Å). The variation is in the same order as the average ionic radius of divalent cations (Co²⁺, Ni²⁺ in sample CoNi, Co²⁺, Zn²⁺ in sample CoZn, and Ni²⁺, Zn²⁺ in sample NiZn).78,79

Surface composition of HESO-NFs and oxidation states of cations.

Fig. S8 shows the x-ray photoelectron spectroscopy (XPS) survey spectra of the investigated HESO-NFs. The high-resolution x-ray photoelectron (HR-XPS) spectra of the core levels are displayed in Fig. 5. The resulting surface composition is reported in Table S3. In all samples, the analysis reveals that Cr, Mn, Fe and O, as well as adventitious carbon, are present on the surface. Ni and Co, Co and Zn, Ni and Zn are also detected in samples CoNi, CoZn and NiZn, respectively.

The HR-XPS spectra of Cr2*p* core level (Fig. 5a,g,m) are featured by two couples of spin-orbit components centered at binding energies (BEs) of ca. 576.0 and 578.3 eV (2*p3/2*) and 585.6 and 587.8 eV (2*p1/2*) in all samples. This indicates the presence of two different Cr-based species on the surface of the investigated materials: (i) the dominant component (see Table S4), with its two spin-orbit features at the lower BE values, is attributed to Cr(III) species present into mixed TM spinel oxides, in accordance with the literature; $80,81$ and (ii) the second component points to the possible presence of chromium in its high oxidation state (i.e., VI).⁸² The co-presence of Cr⁶⁺ surface species has been reported also for $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$ HEO nanoparticles synthesized via a surfactant-assisted hydrothermal method.⁷⁴ It has been attributed to the oxidation of the sample surface, taking place during the 900°C heat treatment in air.

In the Mn2*p* region (Fig. 5b,h,n), HR-XPS spectra reveal the presence of one type of manganese species with the characteristic two peaks as result of the spin-orbit splitting. The Mn2*p3/2* component is peaking at ca. 641.8 eV, which is closer to the BE value typical of Mn(III) rather than other oxidation states, as observed in similar materials.⁸⁰ In samples CoNi and NiZn, the Mn2*p3/2* peak overlaps with a Ni LMM Auger line (Figure S8).⁸³

The HR-XPS spectra of Fe2*p* core level (Figure 5c,i,o) two peaks are detected located at BEs of ca. 710 and 724 eV (for the 2*p3/2* and 2*p1/2* features, respectively). Interestingly, the BE values of Fe2*p3/2* peaks decrease in the order CoZn (710.8 eV) > CoNi (710.4 eV) > NiZn (709.8 eV); the BEs of Fe2*p1/2* peaks behave analogously. All these values are comprised between the BEs of Fe(III) in $Fe₂O₃$ and that of Fe(II) in FeO, which are typically observed at ca. 711.2 and 709.5 eV, respectively.⁸⁰ Thus, on the surface of all investigated HESO-NFs, iron is expected to be found as a mixture of both III and II oxidation states, with the former mostly present in sample CoZn, and the latter in the sample NiZn. The co-presence of $Fe³⁺$ and $Fe²⁺$ species in spinel HEOs has been reported also by other authors.74

Fig. 5 High-resolution XPS spectra for samples (a-f) CoNi, (g-l) CoZn and (m‒r) NiZn and deconvolution in the different spectral regions: Cr2*p* (a,g,m); Mn2*p* (b,h,n); Fe2*p* (c,i,o); Co2*p* (d,j); Ni2*p* (e,p); Zn2*p* (k,q); and O1*s* (f,l,r).

In CoNi, CoZn and NiZn samples, the Fe2*p3/2* peak overlaps with a Ni LMM and/or Co LMM Auger lines (Figure S8).⁸³

In samples CoNi and CoZn, the HR-XPS spectra of Co2*p* core level (Fig. 5d and j, respectively) are featured by two spin-orbit components centered at BEs of ca. 780.2 eV (2*p3/2*) and 795.6 eV (2*p1/2*). Typically, Co(II) and Co(III) are difficult to be distinguished, due to a similar BE value at ca. 780.6 eV.80,83 Nevertheless, the appearance of strong satellite peaks, as observed in the investigated materials, indicates that cobalt atoms are present mainly as Co(II).80,84,85 The Co2*p3/2* peak overlaps with a Fe LMM Auger line (Fig. S8). 83

In the Ni2p region, HR-XPS spectra of samples CoNi and NiZn (Fig. 5e and p, respectively) reveal the presence of one type of nickel species with the characteristic two peaks as result of the spin-orbit splitting. The Ni2*p3/2* component is peaking at ca. 854.5 eV, which is similar to the BE value typical of Ni(II) in NiO and in mixed oxides of nickel and other metals.⁸⁰ In addition, the presence of satellite peaks corroborates the hypothesis that nickel atoms are present in the Ni(II) oxidation state. 80,84,85 The Ni2*p3/2* peak overlaps with a Mn LMM Auger line (Fig. S8).⁸³

In the Zn2p region of HR-XPS spectra of samples CoZn and NiZn (Figures 5k and q, respectively), two strong peaks centered at ca. 1020.8 and 1043.9 eV appear. They are assigned to the Zn2p3/2 and Zn2p1/2 spin-orbit components, respectively. The observed BEs are characteristic of ZnO,91,92 indicating that: (i) zinc is present as Zn(II); and (ii) apparently, zinc hydroxide is not present on the surface of these samples.

Finally, for all analyzed samples, a strong peak is observed in the O1*s* region (Fig. 5f,l,r). It can be decomposed in four different features (the relative amounts are reported in Table S4). In the HR-XPS spectra of O1*s* core level, (i) the dominant contribution, centered at ca. 529.5 eV, is assigned to lattice O^{2-} ions (OL) belonging to mixed TM oxides; $88-91$ (ii) the peak present at ca. 531.0 eV is attributed to native defects of O2 vacancies (OV); $88-91$ and (iii) the peaks centered at high BEs (i.e., at ca. 532.1 and 533.4 eV) are associated with adsorbed or chemisorbed oxygen species, such as O_2 or $H_2O.^{92-94}$ The fraction of OV increases in the order CoNi > NiZn ≈ CoZn (see Table S4 in SI), indicating that the chemical composition of the HEM can control the concentration of oxygen vacancies into the material.

Phase purity of the oxide and inversion degree of the spinel phase.

In order to get a deeper insight into the phase purity and crystallinity of the produced HESO-NFs, micro-Raman analysis was carried out. Spectra from several random positions on each specimen were recorded and compared to each other to evaluate the spatial homogeneity of the samples in terms of crystalline phase (Fig. S9). The lack of significant differences in the relative intensities and shifts in the positions of the bands confirms the formation of a single oxide phase in each sample, without appreciable local deviations from the average chemical composition.⁷¹

Fig. 6 displays the averaged spectra with the apparent peak positions. According to the factor group analysis, the predicted normal vibration modes of the *Fd-3m* spinels $(A_{1g} + E_g + F_{1g} +$ $3F_{2g} + 2A_{2u} + 2E_u + 4F_{1u} + 2F_{2u}$ include 39 optical modes and 3 acoustic modes.^{49,95-99} The triply degenerate F_{1u} modes are infrared-active, whereas the *A*1g, doubly degenerate *E*^g and triply degenerate F_{2g} modes are Raman-active.^{49,95,97,99} The remaining modes are inactive.^{49,96} All Raman-active modes involve the motions of oxygen along the cubic space diagonals.^{100,101} The A_{1g} mode is due to the vibration along the direction joining an oxygen atom to the tetrahedral M^{2+} cation and usually appears between 600 and 720 $cm^{-1}.$ ^{95,99,102-105} The $E_{\rm g}$ mode, detected between 250 and 360 cm⁻¹,¹⁰³⁻¹⁰⁵ is associated to symmetric bending of O2− anion with respect to octahedral M³⁺ cation.¹⁰³ The F_{2g} (1) mode, appearing between 160 and 220 cm⁻¹,¹⁰³⁻¹⁰⁵ originates from translation motion of M3+‒O2−, while higher frequency *F*2g(2) and *F*2g(3) modes, between 440 and 590 cm^{-1} , $95,102-104$ arise from the asymmetric stretching and asymmetric bending of oxygen, respectively.¹⁰⁴ Both positions of the Raman bands and their relative intensities strongly vary within the family of spinels.^{95,98,104-108} Just to cite

some cases, the most intense A_{1g} mode is detected at 630 cm⁻¹ in ZnFe₂O₄ and 690 cm⁻¹ in NiFe₂O₄;¹⁰⁴ the $E_{\rm g}$ mode, located at 320 cm $^{-1}$ in NiFe₂O₄, downshifts at 250 cm $^{-1}$ in ZnFe₂O₄,¹⁰⁴ whereas the *F*2g(1), *F*2g(2) and *F*2g(3) modes appear respectively at 180, 470 and 570 cm^{-1} in NiFe₂O₄ and at 160, 350 and 450 cm‒¹ in ZnFe2O4. ¹⁰⁴ In CoFe2O4, the *F*2g(1), *E*g, *F*2g(2), *F*2g(3) and A_{1g} modes are located at 162, 302, 463, 613 and 678 cm⁻¹, respectively.¹⁰⁵ In magnetite (FeFe₂O₄), the $F_{2g}(1)$ is not detected, while the *E*g, *F*2g(2), *F*2g(3) and *A*1g modes appear at 312 , 474, 548 and 677 cm $^{-1}$, respectively. 95

The same occurs in FeCr₂O₄, where the E_{g} , $F_{2g}(2)$, $F_{2g}(3)$ and A_{1g} modes are detected at 436, 493, 591 and 674 cm⁻¹, respectively.⁹⁵ In ZnCr2O4, the *F*2g(1), *E*g, *F*2g(2), *F*2g(3) and *A*1g modes appear at 182, 452, 511, 605 and 688 cm^{-1} , respectively, 95 whereas in ZnMn₂O₄, they are detected at 300, 321, 382, 475 and 678 cm^{-1} , respectively.¹⁰⁶ As a general behavior, the band positions decrease with the increase of the radii of both octahedral and tetrahedral cations, 95 e.g. in aluminate spinels, the position of the *E*^g mode downshifts with

Fig. 6 Averaged micro-Raman spectra of the HESO-NFs (the apparent peak positions, in cm⁻¹, are also reported).

increasing radius of tetrahedral cation (from 411 cm⁻¹ in ZnAl $_2$ O $_4$ to 395 cm $^{-1}$ in MnAl $_2$ O $_4)$; 95 the same occurs in chromites for the position of the $F_{2g}(2)$ mode (which moves from 541 cm⁻ ¹ in MgCr₂O₄ to 492 cm⁻¹ in FeCr₂O₄).⁹⁵ As a general behavior, the band positions decrease with the increase of the radii of both octahedral and tetrahedral cations.⁹⁵ The partial substitution of either divalent or trivalent cations with different divalent and trivalent cations, respectively, reflects in changes (in the lattice parameters and) in the spectral profile, with shifts of the Raman peaks and variations of their relative intensities.^{98,107} Other factors, such as the nanoparticle size¹⁰⁸ and the presence of defect-induced lattice distortions, 103 are also influential.

In the present case, by comparing the spectra of the three samples (Fig. 6), no marked differences in apparent positions and relative intensities of the peaks are noticed below 400 cm⁻ 1 , which hints to a larger involvement of the trivalent cations^{99,101,103} (Cr³⁺, Mn³⁺, Fe³⁺, present in all samples) in the weaker $F_{2g}(1)$ and E_g modes. On the contrary, above 400 cm⁻¹, the spectral profiles are clearly different, pointing to a greater sensitivity of the stronger *F*2g(2), *F*2g(3) and *A*1g modes to the change in composition of the HEO-NFs (i.e. to the substitution of one of the divalent cations). In Ni-containing CoNi and NiZn NFs, the *F*2g(2) mode (apparently peaking between 504 and 517 cm^{-1}) is better resolved than in CoZn NFs, where it appears as a shoulder (at ~500 cm⁻¹), while the most intense A_{1g} mode, lying in the frequency range (620–700 cm^{-1}) peculiar to ferrites and chromites, $102,104$ apparently peaks at 658 cm⁻¹ in NiZn NFs and at lower frequencies (633 and 620 cm^{-1}) in CoZn and CoNi NFs. In the latter sample, a shoulder further appears on the higher frequency side of the *A*1g mode, suggesting the occurrence of inversion,71,95,109 in agreement with what reported for (Cr1/5Mn1/5Fe1/5Co1/5Ni1/5)3O⁴ nanoparticles, prepared via sol-

To ensure electroneutrality of the 2-3 spinel M_3O_4 phase, 1/3 of the cations must have charge +2 and 2/3 must have charge +3. Considering the relative stability of M^{2+} and M^{3+} cations in the spinel oxide lattice it is expected that chromium and zinc are present as Cr^{3+} and Zn^{2+} , respectively. The other cations (Mn, Fe, Co, Ni) can occur as both M^{2+} and M^{3+} . The preference for M^{3+} decreases as $Mn > Fe > Co > Ni$.

gel method⁴³ and reverse co-precipitation approach.⁷⁶

In spinel oxides, cations occupy tetrahedral 8a sites and octahedral 16d sites (Fig. S1). The energy difference between the 8a and 16d site for middle-late 3d ions ranges from 0 ($d⁵$ and d^{10} ions) to 158 kJ/mol ($d^3 Cr^{3+}$).^{110,111} This explains why ferrites $MFe₂O₄$ are often inverse (Fe³⁺ is d⁵), while chromites $MCr₂O₄$ are always normal.¹¹² Clearly, TEM is not able to resolve the 8a and 16d sites, so one may wonder whether entropic effects are strong enough to overcome the cation site preferences and achieve uniform cation distribution over all sites, or uniform distribution only occurs within 8a and 16d sites separately, leaving room to satisfy the cation site preferences. Using the crystal-field-theory based values for the octahedral stabilization energy¹¹⁰ and adapting the formula for the configurational entropy 113 of a crystal with sublattice to the spinel structure as

$$
S_{\text{config}}^{\text{FU}} = -R\left(\sum_{i} x_{i}^{(8a)} \ln x_{i}^{(8a)} + 2 \sum_{i} x_{i}^{(16d)} \ln x_{i}^{(16d)}\right) \tag{1}
$$

where the configurational entropy is calculated for a formula unit (FU) M_3O_4 , $x_i^{(8a)}$ and $x_i^{(16d)}$ are the fractions of metal *i* occupying the 8a and 16 sites, respectively, we can estimate the Helmoltz free energy FFU corresponding to a given distribution of cations. The distributions are strongly favored that have cations (*i*) partitioned between 8a and 16d sites to optimize octahedral stabilization, and (*ii*) randomly distributed within the 8a and 16d sites, under the electroneutrality constraint. At 1173 K (our highest calcination temperature), the cation distributions detailed in Table 1 have FFU lower than that of the completely random distribution by 37 to 45 kJ/mol (the details of the calculations can be found in the SI). The occurrence of intrasiterandom cation distributions is consistent with the presence of Mn as Mn³⁺, Fe as a mixture of Fe²⁺ and Fe³⁺, and of Co mostly as $Co²⁺$, evidenced by the XPS data. It also agrees with the XAS data **for** the contract of the

 $(\text{Co}_{1/5}\text{Mg}_{1/5}\text{Mn}_{1/5}\text{Ni}_{1/5}\text{Zn}_{1/5})(\text{Al}_{1/5}\text{Co}_{1/5}\text{Cr}_{1/5}\text{Fe}_{1/5}\text{Mn}_{1/5})_2\text{O}_4,^{114}$ and the XANES data for $(Ti_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_{3}O_4$, 115 which indicate that the cation site preferences are satisfied. Very recently, Sarkar et al.⁶⁵ demonstrated that micrometer-sized (Cr1/5Mn1/5Fe1/5Co1/5Ni1/5)3O⁴ HESO has intrasite-random cation distribution with $(Co_{3/5}Fe_{2/5})$ in 8a sites and (Cr3/5Mn3/5Fe1/5Ni3/5) in 16d sites using X-ray, Mössbauer, and neutron techniques. They also expect that Cr and Fe are present as trivalent cations, Ni as divalent cation, and Mn and Co as a mixture of M^{2+} and M^{3+} (M^{2+}/M^{3+} = 10:90 for Mn, 60:40 for Co). The important role of the excess octahedral stabilization was recognized, as we independently did. The data are close to those in Table 1. So, we are confident that the cation distributions we propose for $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})$ ₃O₄ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ are close to the true ones.

In order to infer additional information, Gaussian bands were utilized to reproduce the micro-Raman spectra (Fig. S10). Besides the $F_{2g}(1)$, E_g , $F_{2g}(2)$, $F_{2g}(3)$ and A_{1g} modes, additional bands were found to contribute to the Raman intensity, confirming that inversion occurs, to different extent, in all samples.71,76,95,109

Table 1. Cation distributions that satisfy electroneutrality and optimize octahedral stabilization.

a Inversion degree, defined as the fraction of trivalent cations in the 8a sublattice.

Magnetic properties.

General considerations. It is clear from the preceding discussion that the interpretation of the magnetic data will benefit from reference to chromites, manganites, and ferrites with spinel structure.

The primary particles comprising the HESO-NFs have diameter from a few tens of nm to the hundred nm range. From XRD data, the average crystallite diameter is ≈100 nm for CoNi and NiZn and ≈85 nm for CoZn. Magnetic lengths, such as the single-domain, coherent rotation, and superparamagnetic (SPM) blocking diameters,¹¹⁶ are a useful (though approximate) starting framework to develop the interpretation of magnetic data. We could find in the literature data for $Fe₃O₄$, CoFe₂O₄, and CoCr₂O₄ (Table S7). The NF primary particles have diameter similar to the single-domain diameter of $Fe₃O₄$ and much smaller than that of CoFe_2O_4 and CoCr_2O_4 , so they reasonably comprise a single domain. Magnetization inversion by coherent rotation diameter is expected if the NF behave as chromites but partially incoherent inversion (e.g., curling) might occur if the NFs behave similarly to ferrites. Finally, the NF primary particles are much larger than the SPM blocking diameter at 400 K of all

three spinel oxides and, therefore, superparamagnetism is not expected to be important.

Magnetic data and analysis. The high field magnetization M_{50k} (H=50 kOe) and remanence M_{rem} were measured between 5 and 300 K (Fig. 7). All samples have significant high-field and remanent magnetization, at least at low temperature, and are thus able to sustain a magnetically ordered structure. M_{50k} and Mrem at 5 K are in the 17–40 emu/g and 10–30 emu/g range, respectively, that correspond to molar magnetization per formula unit (FU) in the range 0.7–1.7 μ_B and 0.4–1.3 μ_B , respectively (μ_B is the Bohr magneton). Since the cations, except Zn^{2+} , have molar magnetization in the 3.2-5.9 μ_B range, the measured M_{50k} and M_{rem} demonstrate that the NFs have an ordered magnetic structure of ferrimagnetic (FIM) nature, excluding ferromagnetic (FM) and (canted) antiferromagnetic (AFM) structures, which would display much larger and smaller M_{50k} and M_{rem} , respectively. The squareness M_{50k}/M_{rem} is large at 5 K, it extrapolates to ≈0.75 at 0 K, a value typical of Stoner-Wohlfarth particles with cubic anisotropy. Incoherent magnetization inversion and interparticle interactions seem thus to be relatively unimportant even at low temperature.¹¹⁷

The magnetization for a completely collinear FIM structure, such as that found in the mixed ferrites $MFe₂O₄$, can be calculated by a simple Monte Carlo approach based on the cation distributions in Table 1 (This approach is detailed in the ESI). The calculated values are 2.5, 7.0, 6.3 μ_B per FU for CoNi, CoZn and NiZn NFs, respectively. Thus, the observed magnetization is not consistent with a collinear, antiparallel alignment of the ionic magnetic moments (sublattice magnetizations), like in the typical Néel FIM structure. It is however consistent with a FIM structure where the sublattice magnetizations are not collinear.

 M_{50k} and M_{rem} decrease with increasing temperature. M_{rem} of CoNi at 300 K (1.35 emu/g) is 5% of the remanence at 5 K. The remanence of Zn-containing (CoZn and NiZn) NFs vanishes well below RT. Therefore, CoZn and NiZn NFs are paramagnetic (PM) at 300 K and CoNi NFs probably becomes PM slightly above RT. We now turn our attention to the transition temperature T_c between the FIM and PM phases. Considering that the primary particles have size ranging from a few tens of nm to the hundred nm range, we expect a distribution of transition temperatures. The temperature, at which M_{rem} vanishes, marks the T_c of the primary particles with highest T_c , presumably the largest particles. Due to noise and artifacts, it is not possible to accurately measure where M_{rem} vanishes. The best estimate is that $T_c \approx 150$ and 200 K for CoZn and NiZn NFs, respectively. This ordering of the $T_c = (CoNi > NiZn > CoZn)$ can also be appreciated in the M_{50k} plot (Fig. 7). M_{50k} of CoZn changes concavity at ≈150 K, suggesting that at higher temperatures it follows the Curie-Weiss-Néel law for ferrimagnets, 112

$$
\frac{1}{\chi} = \frac{T}{c} + \frac{1}{\chi_0} - \frac{\sigma}{T - \theta_p} \tag{2}
$$

where $χ = M/H$ is the susceptibility, C is the Curie constant, $χ_0$ and σ are related to the intra- and inter-sublattice molecular fields, and θ_p is the paramagnetic Curie point, which is slightly higher than the true transition temperature T_c.¹¹⁸ As can be seen in Fig. 7d, 1/χ_{50k} of CoZn

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closely follows the Curie-Weiss-Néel law for T > 200 K, so we can identify the PM and FIM Curie points $T_c \approx \theta_p = 137$ K, in agreement with T_c from M_{rem} data. Interestingly, the asymptotic temperature θ_N $= -C/\chi_0$ is close to zero, indicating the presence of frustrated FM and AFM exchange interactions that compete and partly compensate.**¹¹⁹** The Curie constant corresponds to an effective moment μ_{eff} = $(3k_B C/N_A)$ = 5.0 μ_B per FU that is just 66% of that calculated by a Monte Carlo approach. This discrepancy is probably due to the lack of data in the high temperature range where the T/C term of Equation 2 is dominant.

Confirmation of the presence of a FIM-PM transition and details about the FIM phase are provided by the magnetization isotherms M(H) measured at different temperature (Fig. 8). Since the magnetization is not always saturated at 50 kOe, we analyzed how M approaches saturation¹¹⁷ to more accurately estimate the true M_{sat}.

Fig. 7 Temperature dependence of the high field magnetization M_{50k} (a, H = 50 kOe), remanence M_{rem} (b), squareness M_{50k}/M_{rem} (c), and inverse high-field susceptibility $1/\chi_{50k}$ (d) of the NFs. The dashed line in panel d is the best-fit Curie-Weiss-Néel curve for CoZn NFs.

The magnetic parameters are collected in Table 2 (see Materials and methods section).

The behavior of the NFs is markedly different from each other. The CoNi NFs have open loops that do not change shape up to 300 K. At low T, the coercivity H_c is moderate (0.83 kOe). On increasing temperature, M_{sat} , M_{rem} , H_c , and the coercivity of the remanence H_{cr} decrease slowly due to the approach to the critical FIM-PM T_c . This interpretation is supported by the large size of the primary particles and the dependence of coercivity on T. Indeed, neither H_c nor H_{cr} follow Kneller's law¹²⁰ typical of thermally activated magnetization inversion process. They instead follow a power law (H_c / H_{c,0}) \approx (M_{sat} / $M_{sat,0}$ ^{4.5}, where the 0 subscript indicates the value at 5 K, which, provided that H_c is proportional to the anisotropy field 2 K/Msat, is consistent with Zener's law¹²¹ typical of bulk materials. Further supporting evidence will be provided below.

The magnetization isotherms of CoZn and NiZn NFs have similar behavior though at shifted temperature mirroring the different T_c . From 5 to 150 K, they are open loops with almost constant M_{sat} and dramatically decreasing H_c (and H_{cr}). The main difference is that CoZn NFs have much larger H_c at 5 K (6.37 kOe) than NiZn NFs have (1.17 kOe). It is noteworthy that CoNi NFs have H_c lower than that of CoZn NFs, despite the presence of Co in both NFs. This difference can be attributed to the fact that in CoZn NFs a larger fraction of Co is present in octahedral 16d sites (see Table 1). At 150 K, the magnetization of NiZn NFs still is non-linear, while that of CoZn NFs is almost linear. At 300 K, both NFs are definitely in a linear regime: at H = 50 kOe, the paramagnetic magnetization $M_p = \chi_p$ H is 83% and 99% of the total M_{50k} magnetization for NiZn and CoZn NFs, respectively. M_p is truly related to primary particles that have entered the PM regime, as shown by the small magnitude of its slope χ_{p} , which corresponds to a magnetization of a few μ_{B} per FU. In the SPM regime, the initial susceptibility would be several orders of magnitude larger and strong non-linearity would be observed already at H < 1 kOe.

Fig. 8 Magnetization isotherms M(H) of the (a,b) CoNi, (c,d) CoZn and (e,f) NiZn NFs. The panels (b,d,f) show the central part of the isotherms. Note the different horizontal scale of the panel (d).

Temperature dependent M(H) curves have thus strengthened the conclusion that our HESO-NFs have a low-T FIM phase and a high-

T PM phase with transition temperature T_c decreasing as CoNi > NiZn > CoZn. They have provided further evidence that SPM effects are not dominant, as anticipated on the basis of the size of the primary particles. The small χ_{p} , corresponding to a FU magnetization of a few μ_B , is consistent with a non-collinear FIM structure.

More detailed information about the magnetic structure of the NFs is obtained from the zero-field cooled (ZFC) and field cooled (FC) magnetization (M_{ZFC} , M_{FC}) measured with H_{cool} = H_{meas} = 100 Oe (Fig. 9). All (Z)FC curves differ from the typical (Z)FC curves of ferrite NPs.

For instance, the ZFC-FC bifurcation occurs at temperature much higher than the maximum M_{ZFC} . The NFs display (Z)FC curves with similar shape that have different temperature span in correspondence to the unequal T_c . Extending the temperature range up to 400 K allowed us to observe the transition of CoNi NFs to the PM phase. There clearly is a dispersion of the transition temperature due to the particulate nature of the NFs.

Therefore, to estimate T_c from M_{FC} data, the intercept of the maximum slope line was extrapolated and its intersection with

the M = 0 axis was interpreted as an average T_c . For CoNi NFs, the FIM-PM transition occurs at $T_c = 374$ K (Table 3). On decreasing T, M_{FC} increases almost linearly and then levels off at T \leq 20 K. No low T transition is observed. The M_{FC} of CoNi NFs is almost exactly proportional to M_{50k} (M_{FC} \approx 0.44 M_{50k}) showing that there is a common thermal process underlying the M increase at low T. This process is the thermal dealignment of the spins about the magnetization axis. Thermally activated magnetization inversion (superparamagnetism) can be excluded since it strongly depends on the applied field. The large difference between M_{ZFC} and M_{FC} may be related to several causes.

It can be due to the presence of sizeable magnetic anisotropy typical of spinel oxides. The power dependence of H_c on M_{sat} supports this hypothesis. Another possible explanation is the formation of a spin-glass-like phase below RT. This seems less likely, at least for CoNi, as M_{ZFC} and M_{FC} are already unequal at RT.

Table 2. Magnetic parameters of the NFs calculated from the magnetization isotherms M(H). Parameters may be absent when the NFs are mostly paramagnetic.

a M_{sat} is the saturation magnetization as obtained from the analysis of the approach to saturation.¹¹⁷

The dashed line is the best-fit Curie-Weiss-Néel curve for CoZn NFs.

Table 3. Critical temperatures of the NFs.^a

a Estimated from the intercept of the maximum slope line with the M = 0 axis (T_c) or M = M(5 K) line (T_s).

Fig. 9 Temperature dependence of the low-field (100 Oe) magnetization of the NFs. Left: ZFC and FC magnetization of the NFs. Right: inverse low-field susceptibility $1/\chi_{100}$ of CoZn NFs.

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The (Z)FC magnetization of Zn-containing NFs is close to zero at RT and rapidly increases when the transition to the FIM phase approaches. The PM phase gives way to a FIM phase at $T_c = 233$ (NiZn) and 105 K (CoZn), temperatures much lower than that of CoNi NFs. M_{FC} reaches a maximum at 50 K and then decreases and levels off at low T. This marks the transition to a phase with lower M at the (average) critical temperature $T_S = 11$ and 20 K for NiZn and CoZn NFs, respectively. The M decrease suggests the formation of a spin-glass-like phase. The shape of MZFC does not allow us to exclude SPM phenomena, but the thermal unblocking of the spin-glass-like phase and the thermal decrease of Msat (clearly observed in the CoNi NFs) seems sufficient to explain the temperature dependence of M_{ZFC} . Therefore, the (Z)FC magnetization of Zn-containing NFs gives evidence of the occurrence of another magnetic structure at very low T.

The low-field susceptibility χ_{100} = M_{FC}/H_{meas} sufficiently extends above T_c to be significantly modelled by the Curie-Weiss-Néel law (Eq. 2) only for the CoZn NFs. 1/χ₁₀₀ closely follows Eq. 2 for T > 150 K (Fig. 9). The asymptotic temperature $θ_N ≈ 0$ and the effective moment $μ_{eff} = 6.1 μ_B$ (80% of $μ_{eff}$ calculated by a Monte Carlo approach) are both in agreement with the results of the modelling of $1/\chi_{50k}$. The best-fit parameters provide $T_c \approx \theta_p = 150$ K, in agreement with M_{rem} data and the modeling of the high-field susceptibility. This T_c is not inconsistent with that in Table 3 since the former is the T_c of the largest particles while the latter represent an average over all particles.

Discussion of the magnetic properties and structure.

As anticipated, the magnetic results can be discussed with reference to the magnetic structure of ferrites, chromites, and manganites. The magnitude of M_{sat} and the temperature dependence of M(Z)FC and H_c are not consistent with the usual behavior of ferrites, which have Néel FIM phases with collinear sublattice magnetization and T_c above \approx 600 K. It is also interesting to note that when Zn replaces a magnetic cation (Co, Ni) in our NFs, the magnetization decreases, at variance with ferrites that show the well-known magnetization increase upon Zn substitution.¹²²

In spinel-structure chromites MCr_2O_4 and manganites MMn₂O₄, the interplay between strength and frustration of the superexchange interactions is such that they display a noncollinear FIM phase with M_{sat} and T_c much lower than those the of corresponding ferrites MFe₂O₄, and often a transition (T_s) to a low temperature reentrant spin-glass-like phase.¹¹⁹ The magnetic structure of the low-T phase in chromites typically is conical/spiral with a long-range ordered longitudinal component and a short-range (a few nm) ordered transverse component.¹²³⁻¹²⁵ The low-T phase of manganites has been less studied. The neutron diffraction data of $NIMn_2O_4$ at T = 1.5 K were interpreted in terms of a strongly canted magnetic structure where the sublattice magnetizations form an angle of 27°.

The magnetic behavior of Zn-containing CoZn and NiZn NFs closely resembles that of chromite (and manganites) under many respects. For instance, the dramatic decrease of coercivity at low T indicates the transition between the spin-glass-like and the non-collinear FIM phase.¹²⁶ Therefore, the magnetic behavior of CoZn and NiZn NFs is consistent with the presence of an intermediate non-collinear FIM phase and a low-T reentrant spin-glass-like phase. The CoNi NFs share some magnetic characteristic with chromites, e.g., the low Msat, but under other respects, e.g., the relatively high T_c and the slow decrease of M_{sat} , M_{rem} , and H_c with increasing T, they behave like a ferrite. This behavior can be understood with reference to the parameter u ,¹²⁶ which quantifies the magnetic frustration:

$$
u = (4 J_{BB} S_B) / (3 J_{AB} S_A)
$$
 (3)

where J_{AB} is the superexchange integral between the nearest-neighbor spins at the 8a and 16d sites, J_{BB} is that between the nearest-neighbor spins at two 16d sites, and SA and *S*^B are the magnitudes of the spins at the 8a and 16d sites. For small u (< 8/9), Néel collinear FIM structure occurs, but at larger u the increasing frustration gives rise to non-collinear FIM structures with decreasing order range.¹²³ In CoNi the full network of superexchange interactions counteracts the magnetic frustration and the effects of the latter are weaker than those in MCr₂O₄ and MMn₂O₄ spinel oxides. The replacement of a magnetic cation by Zn^{2+} in CoZn and NiZn NFs significantly reduces the average values of both J_{AB} and S_{A} , leading to a lower T_c , and it increases u and the magnetic frustration, leading to lower collinearity and the occurrence of a reentrant spin-glass-like-phase at low T.

The above argument about magnetic structure is based on the close similarities of the chemical composition, crystal structure, and magnetic behavior between our HESO NFs and the spinel oxides chromites, manganites, and ferrites. Of course, we do not claim that we have proved the HESO NF magnetic structure, which would require additional experiments, such as magnetic neutron scattering, that are outside the scope of this paper. It however seems more likely that, for compounds of identical crystal structure and similar chemical composition, similar magnetic behavior stems from similar magnetic structure than that different magnetic structures accidentally give rise to similar magnetic behavior. Caution is however in order as HE compounds might display properties far from those of the corresponding low-entropy compounds.

Finally, we compare our results with preceding magnetic characterization of HESOs. The magnetic properties of HESOs have already been summarized.^{59,60} Spinel oxides with composition (Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})₃O₄, equal to that of CoNi NFs, have attracted some attention. The literature magnetic data (Table 4) are in general agreement with those recorded for our CoNi NFs but they were rarely discussed in detail. Magnetization isotherms were first measured on NP aggregates obtained by solution combustion followed by calcination. 64,127 The M(H) curves at RT were slightly open loops similar to that of CoNi NFs. M(H) at 300 K was also measured on aggregates of ≈100 nm particles synthesized by a sol-gel procedure, and in this case a closed M(H) curve typical of SPM behavior was recorded.⁶³ A more in-depth investigation of bulk $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})$ ₃O₄ obtained by sintering at 1273 K was later reported.⁶⁵ The results from Mössbauer and

magnetometric data, including the FIM-PM T_c , are very similar to those of CoNi NFs. A broad investigation of high-entropy spinel oxides M_3O_4 , ferrites MFe₂O₄, and chromites MCr₂O₄ has **Table 4.** Magnetic parameters of HESOs from the literature.

been reported upon, which includes $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5}$ ₃O₄, spinel oxide.⁶² The HEOs were synthesized by ball milling, compression, and sintering at 1523

a At 20 kOe. **b** At 30 kOe. c At 5 kOe. ^d At 90 kOe. **e** From magnetization data. f From Mössbauer data. ^g At 40 kOe.

K. The M(H) isotherms (5–300 K) of $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$ are consistent with the present data for CoNi NFs, but the M_{ZFC} and M_{FC} curves are rather different. The latter have a shape loosely similar to the curves in Fig. 9, but they are quantitatively very different, e.g., M_{FC} (5 K, H_{cool} = H_{meas} = 100 Oe) for the bulk HESO (6.6 emu/g) unexpectedly is much lower than that of CoNi NFs (18.2 emu/g). The magnetic data was explained in terms of the occurrence of Cr or Mn rich regions with AFM structure. We think that the behavior of $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$ HESO is better explained on the basis of the presence of Cr and Mn, reasonably as Cr^{3+} and Mn³⁺ in the 16d sites. Indeed, in the same paper the ZFC-FC curves of high-entropy chromites $MCr₂O₄$ are displayed, which closely resemble those of the CoZn and NiZn NFs and confirm that frustration effects are induced by Cr and increased by the replacement of magnetic cations by non-magnetic ones. Finally, micrometer-sized $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$ particles were recently studied.⁶⁵ The 300 K isotherm is reversible whereas at 5 K a small coercivity (0.23 kOe) was observed. These data agree with those here measured for CoNi NFs, the coercivity difference is explained by the larger size of the particles, which are multidomain (see Table S7). The (Z)FC magnetization is similar to the present one except for an inflection at ≈75 K, which was attributed to the depinning of domain walls.

HESOs with composition equal to CoZn and NiZn NFs have been much less studied.⁶⁴ The RT M(H) curves of aggregates of 24-nm NPs, obtained by solution combustion followed by calcination, were slightly open loops ($H_c < 0.1$ kOe) in stark contrast with CoZn and NiZn NFs, which are essentially PM at RT. We can only speculate that this striking difference is due the different synthetic route and morphology of the oxides.

Conclusions

Electrospun $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})$ ₃O₄ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ NFs, exhibit porous structure consisting of interconnected nearly strain-free oxide single crystals with pure Fd-3m spinel structure. The in-depth investigation of their magnetic properties shows that $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Ni_{1/5})_3O_4$, $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Co_{1/5}Zn_{1/5})$ ₃O₄ and $(Cr_{1/5}Mn_{1/5}Fe_{1/5}Ni_{1/5}Zn_{1/5})$ ₃O₄ NFs have a diverse and rich magnetic behavior, grounded into the random cation distribution within (but not across) sublattices. In particular, the Zn-containing NFs display complex magnetic structures at low temperature. The substitution of a single cation (20% of the cationic content) changes the FIM-PM T_C by about 130 K. More insight into this fascinating magnetism could be obtained by investigating the details of the cation distribution, as recently carried out for (Cr0.2Mn0.2Fe0.2Co0.2Ni0.2)3O4.⁶⁵ Furthermore, the suggested magnetic structure of our HESO NFs should be proved by direct techniques such as neutron diffraction. Of course, the randomness inherent in HEMs makes such detailed investigations a difficult endeavor.

Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare

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