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Invariant nature of substituted element in metal-hexacyanoferrate

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The chemical substitution of a transition metal (*M*) is an effective method to improve the functionality of materials. In order to design the highly functional materials, we first have to know the local structure and electronic state around the substituted element. Here, we systematically investigated the local structure and electronic state of the host (*M*_h) and guest (*M*_g) transition metals in metalhexacyanoferrate (*M*-HCF), Na_x(*M*_h, *M*_g)[Fe(CN)₆]_y (1.40 < x < 1.60 and 0.85 < y < 0.90), by means of extended X-ray absorption fine structure (EXAFS) and X-ray absorption near-edge structure (XANES) analyses. The EXAFS and XANES analyses revealed that the local structure and electronic state around *M*_g are essentially the same as those in the pure compound, *i.e*, *M*_g-HCF. Such an invariant nature of *M*_g in *M*-HCF is in sharp contrast with that in layered oxide, in which the *M*_g valence changes so that local *M*_g-O distance (*d*_{*M*-0}^g) approaches the *M*_h-O distance (*d*_{*M*-0}^h).

The chemical substitution of transition metal (M) is an effective method to improve the functionality of materials, such as electrochemical performance in sodium-ion secondary batteries (SIBs) and critical temperature of magnetic phase transition. Actually, the rate and cycle properties of manganese hxacyanoferrate (Mn-HCF) in SIBs are significantly improved by partial substitution of Fe, Co, and Ni for Mn¹⁻³. In addition, the critical temperature for the spin-crossover transition of $[Fe_{1-x}Zn_x (ptz)_6](BF_4)_2$ decreases with the Zn concentration $(x)^4$. To thoroughly comprehend the partial substitution effect on material properties, we first have to know the local structure and electronic state around substituted guest element (M_{e}) . The extended X-ray absorption fine structure (EXAFS) and X-ray absorption near-edge structure (XANES) analyses around the K-edges are powerful tools to investigate the local structure and electronic state around $M_{\rm g}$ and $M_{\rm h}$ in the mixed compounds. Here, let us consider the network material, such as coordination polymer and transition metal oxides, in which ligand-bridged M chains form one-, two-, or three-dimensional network. For example, in metal hexacyanoferrate (M-HCF), Na_x*M*[Fe(CN)₆]_y, which is the most prototypical coordination polymer, cyano-bridged -*M*-NC-Fe-CN-*M*-NC-Fe-CN-M-NC- chains form three-dimensional (3D) jungle-gym type network. In such a network material, the substitution of M_g for the host element (M_h) , $-M_h - L - M_h - L - M_g - L - M_h - L - M_h - (L represents ligand)$, causes local distortion, *i.e.*, elongation or compression of the M_g -L distance (d_{M-L}^g) reflecting the difference in the ionic radii (r) between $M_{\rm g}$ and $M_{\rm h}$. Such a lattice distortion increases the Gibbs free energy of the system. To minimize the Gibbs free energy, the local distortion should relax via variation in the electronic state of M_g . Actually, in layered oxide system¹, the M_g valence changes so that local M_g -O distance (d_{M-O}^g) approaches the M_h -O distance (d_{M-O}^h) .

Among the network material, the *M*-HCF is the oldest compound that the human being has synthesized. The *M*-HCF consists of a 3D jungle-gym type -*M*-NC-Fe-CN-*M*-NC- Fe-CN-*M*-NC- network and Na⁺ and H₂O, which are accommodated in the network nanopores. We note that the *M*-HCFs have considerable Fe(CN)₆ vacancies, where H₂O molecules coordinated to *M* instead of CN. Most of the *M*-HCFs show the face-centered cubic $(Fm3(-)m; Z=4)^5$ or trigonal $(R3(-)m; Z=3)^{6,7}$ structures. The *M*-HCFs are attracting current interest of material scientists, because they are promising materials for the lithium-ion secondary batteries (LIBs)⁸⁻¹⁰, SIBs^{2,3,11-20}, electrochromism^{21,22}, and thermoelectrics²³. In SIBs, the rate and cycle properties of Mn-HCF are significantly improved by partial substitution of Fe, Co, and Ni for Mn^{2,3}. Thus, *M*-HCF is a suitable system for investigation of the local structure and electronic state around M_g and M_h .

Here, we systematically investigated the local structure and electronic state around M_g and M_h in Na_x(M_h , M_g) [Fe(CN)₆]_y (1.40 < x < 1.60 and 0.85 < y < 0.90) by means of the EXAFS and XANES analyses. The analyses revealed

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Compound	$d_{M-N}^{pure}(\text{\AA})$	$d_{M-N}^{h}(\text{\AA})$	$d_{M-N}^{\mathrm{g}}(\mathrm{\AA})$	$d_{M-C}^{pure}(\text{\AA})$	$d_{M-C}^{h}(\text{\AA})$	$d_{M-C}^{\mathrm{g}}(\mathrm{\AA})$
Mn-HCF	2.217(5)			3.382(6)		
MnCo-HCF		2.210(7)	2.105(28)		3.377(7)	3.284(24)
MnNi-HCF		2.209(5)	2.100(8)		3.378(6)	3.265(11)
Co-HCF	2.100(8)			3.273(9)		
CoMn-HCF		2.103(8)	2.202(6)		3.276(8)	3.368(8)
CoNi-HCF		2.097(9)	2.088(8)		3.272(10)	3.253(11)
Ni-HCF	2.085(11)			3.237(11)		
NiMn-HCF		2.089(9)	2.197(6)		3.243(10)	3.359(8)
NiCo-HCF		2.085(11)	2.081(30)		3.238(11)	3.251(43)

Table 1. Interatomic distances, d_{M-N} and d_{M-C} in pure (M-HCF: Na_xM[Fe(CN)₆]_y) and mixed $(M_hM_g$ -HCF: Na_x (M_h, M_g) [Fe(CN)₆]_y) compounds. Note that M-HCF and M_hM_g -HCF have Fe(CN)₆ vacancies (0.85 < y < 0.90). The actual chemical compositions are shown in Table 2. Superscript, pure, in d_{M-N} and d_{M-C} means pure compound. Superscripts, h and g, in d_{M-N} and d_{M-C} mean host and guest elements, respectively.

Compound	composition	structure	a (Å)	$a_{\rm H}({ m \AA})$	<i>c</i> _Н (Å)
Mn-HCF	Na _{1.52} Mn[Fe(CN) ₆] _{0.88} 2.0H ₂ O	trigonal	(10.5448)	7.5253(8)	17.9307(20)
MnCo-HCF	Na _{1.56} Mn _{0.80} Co _{0.20} [Fe(CN) ₆] _{0.89} 1.7H ₂ O	trigonal	(10.5165)	7.5237(6)	17.7945(14)
MnNi-HCF	Na _{1.64} Mn _{0.75} Ni _{0.25} [Fe(CN) ₆] _{0.91} 4.2H ₂ O	trigonal	(10.4464)	7.4440(7)	17.8161(16)
Co-HCF	Na _{1.40} Co[Fe(CN) ₆] _{0.85} 3.9H ₂ O	trigonal	(10.3620)	7.4240(6)	17.4818(13)
CoMn-HCF	Na _{1.40} Co _{0.75} Mn _{0.25} [Fe(CN) ₆] _{0.85} 4.5H ₂ O	trigonal	(10.4047)	7.4544(5)	17.5545(11)
CoNi-HCF	$Na_{1.44}Co_{0.75}Ni_{0.25}[Fe(CN)_6]_{0.86}4.6H_2O$	trigonal	(10.3458)	7.4133(7)	17.4502(16)
Ni-HCF	Na _{1.52} Ni[Fe(CN) ₆] _{0.88} 5.8H ₂ O	cubic	10.2887(21)		
NiMn-HCF	Na _{1.60} Ni _{0.78} Mn _{0.22} [Fe(CN) ₆] _{0.90} 4.7H ₂ O	cubic	10.3455(24)		
NiCo-HCF	Na _{1.48} Ni _{0.82} Co _{0.18} [Fe(CN) ₆] _{0.87} 5.5H ₂ O	cubic	10.2982(21)		

Table 2. Chemical composition and lattice constants of pure (*M*-HCF) and mixed (M_hM_g -HCF) compounds. a_H and c_H are the lattice constants in the hexagonal setting. The lattice constants in the brackets are approximate values ($=2^{1/3}/3^{1/6}a_H^{-2/3}c_H^{-1/3}$) in the pseudo-cubic setting.

that the local structure and electronic state of M_g are essentially the same as that in the pure material, *i.e.*, M_g -HCF Such an invariant nature of M_g in *M*-HCF is in sharp contrast with that in layered oxide, in which the M_g valence changes so that d_{M-O}^{g} approaches d_{M-O}^{h} . We will discuss the origin for the difference in the substitution effect between *M*-HCF and the layered oxide.

EXAFS analysis

We performed careful EXAFS analysis on three pure (*M*-HCF) and six mixed (M_hM_g -HCF) compounds. Details of synthesis and characterization are described in the Method section. The x-ray diffraction (XRD) patterns for the nine compounds are shown in Fig. S1. In the EXAFS analyses, we included the contributions from the first-(N) and second- (C) nearest neighbor elements. In order to include the Fe(CN)₆ vacancy effect, the coordination numbers (N_N) of N are treated as an adjustable parameter with restriction of $N_N + N_O = 6$ and $d_{M-O} = d_{M-N}$, where N_o is the coordination number of O. With use of the EXAFS equation, least-squares fittings are performed for the FT[$\chi(k)k^3$]-R plots (Fig. S2). Thus obtained interatomic distances are listed in Table 1.

Interatomic distance between M and N

Figure 1(a) shows correlation diagram of the *M*-N distances (d_{M-N}) between the mixed compounds (d_{M-N}^{h}) around M_h and pure compounds (d_{M-N}^{pure}) . We found that d_{M-N}^{h} in the mixed compound is the same as d_{M-N}^{pure} in the pure compound, indicating that partial substitution of M_g for M_h has no effect on d_{M-N}^{h} . Figure 1(b) shows correlation diagram of d_{M-N} between the mixed compound (d_{M-N}^{g}) around M_g and d_{M-N}^{pure} . Importantly, d_{M-N}^{g} in the mixed compound is the same as d_{M-N}^{pure} .

As-mentioned above, $d_{M\cdot N}^{g} (d_{M\cdot N}^{h})^{h}$ is the same as that in M_{g} -HCF (M_{h} -HCF). Such an invariant nature causes significant difference between $d_{M\cdot N}^{h}$ and $d_{M\cdot N}^{g}$. In Fig. 1(c), we compared $d_{M\cdot N}^{h}$ and $d_{M\cdot N}^{g}$ in the mixed compounds. In CoMn-, NiMn-, MnCo-, and MnNi-HCFs, we observed significant difference between $d_{M\cdot N}^{h}$ and $d_{M\cdot N}^{g}$. The difference ($|d_{M\cdot N}^{h} - d_{M\cdot N}^{g}| = 0.1$ Å) corresponds 5% of $d_{M\cdot N}$. If such a local distortion around M_{g} influences the $d_{M\cdot N}^{h}$ value at the nearest-neighbor M_{h} site, the partial substitution causes significant distribution of $d_{M\cdot N}^{h}$. Such a distribution of $d_{M\cdot N}^{h}$ should increase the Debye–Waller factor ($\sigma^{2}_{M\cdot N}^{h}$) around M_{h} . Figure 2 shows correlation diagram of $\sigma^{2}_{M\cdot N}$ between the mixed compounds ($\sigma^{2}_{M\cdot N}^{h}$) around M_{h} and pure compounds ($\sigma^{2}_{M\cdot N}^{h}$) in the mixed compound is the same as $\sigma^{2}_{M\cdot N}^{pure}$ in the pure compound. This indicates that the partial substitution of M_{g} has no effect even at the nearest-neighbor M_{h} site.

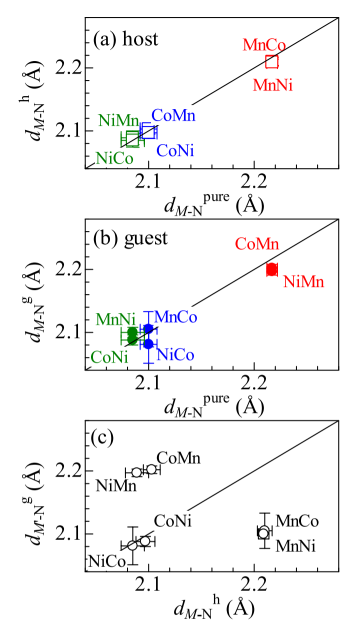


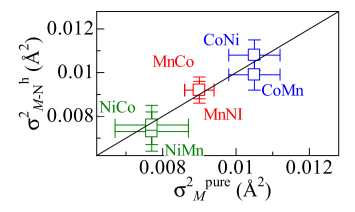
Figure 1. (a) Correlation diagram of the M-N distances (d_{M-N}) between the mixed compounds (d_{M-N}^{h}) around M_h and pure compounds (d_{M-N}^{pure}) . (b) Correlation diagram of d_{M-N} between the mixed compounds (d_{M-N}^{g}) around M_g and d_{M-N}^{pure} . (c) Comparison between d_{M-N}^{g} and d_{M-N}^{h} in the mixed compounds. $M_h M_g$ represents $Na_x(M_h, M_g)$ [Fe(CN)₆]_y. Red, blue and green colors represent the interatomic distances around Mn, Co, and Ni, respectively.

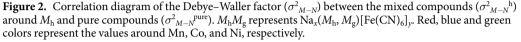
Interatomic distance between M and C

Figure 3(a) shows correlation diagram of the M - C distances (d_{M-C}) between the mixed compounds $(d_{M-C})^{\text{pure}}$ around M_{h} and pure compounds $(d_{M-C})^{\text{pure}}$. We found that d_{M-C}^{h} in the mixed compound is the same as d_{M-C}^{pure} , indicating that partial substitution of M_{g} for M_{h} has no effect on d_{M-C}^{h} . Figure 3(b) is the correlation diagram of d_{M-C} between the mixed compounds (d_{M-C}^{g}) around M_{g} and d_{M-C}^{h} . Figure 3(b) is the correlation diagram of d_{M-C} between the mixed compounds (d_{M-C}^{g}) around M_{g} and d_{M-C}^{pure} . d_{M-C}^{g} in the mixed compound is the same as d_{M-C}^{pure} . Here, recall that d_{M-N}^{g} in the mixed compound is the same as d_{M-C}^{pure} . Here, recall that d_{M-N}^{g} in the mixed compound is the same as d_{M-C}^{pure} . These observations indicate that the ligand (CN) environment around M_{g} in the mixed compound is the same as that in the pure compound, *i.e.*, M_{g} -HCF. In Fig. 3(c), we compared d_{M-C}^{h} and d_{M-C}^{g} in the mixed compounds. In CoMn-, NiMn-, MnCo-, and MnNi-HCFs, we observed significant difference ($|d_{M-C}^{\text{h}} - d_{M-C}^{\text{g}}| = 0.1 \text{ Å}$) between d_{M-C}^{h} and d_{M-C}^{g} .

Electronic state

Now, let us proceed to the electronic state of M_g and M_h . Figure 4 shows the XANES spectra of the pure (*M*-HCF) and mixed (M_hM_g -HCF) compounds around the (a) Mn K-edge, (b) Co K-edge, and (c) Ni K-edge. The thick black curve represents the spectra of pure compounds. The thin solid and broken curves correspond to the spectra around M_h and M_g in the mixed compounds, respectively.





In the Mn K-edge spectra [Fig. 4(a)], the main peak is attributable to the Mn1s-4p transition, whose position is a crude measures of the Mn valence. The blue- (red-) shift of the main peak suggests increase (decrease) in the Mn valence. In Mn-HCF (thick black curve), the main peak is observed at 6548.0 eV. We observed no detectable peak-shift in MnCo- and MnNi-HCFs, indicating that the Mn valence is robust against the partial substitution. Importantly, the peak positions of the substituted Mn in CoMn- and NiMn-HCFs are the same as that of Mn-HCF. Figure 4(d) shows magnified Mn K-edge spectra in the pre-edge region, which is dominated by the Mn1s-3d transition. We observed no detectable peak shift in MnCo- and MnNi-HCFs. These observations indicate that electronic states of the substituted Mn in MnCo- and MnNi-HCFs are the same as that in Mn-HCF. This is consistent with the fact that the ligand (CN⁻) environments around substituted Mn in CoMn- and NiMn-HCFs are the same as that in Mn-HCF [Figs 1(b) and 3(b)].

Similarly to the case of the Mn K-edge spectra, we observed no detectable peak-shift in the Co K- and Ni-K-edge spectra. In Fig. 4(b), the peak positions of substituted Co in MnCo- and NiCo-HCFs are the same as that in Co-HCF. In Fig. 4(c), the peak positions of substituted Ni in MnNi- and CoNi-HCFs are the same as that in Ni-HCF. We observed no detectable peak shift in the respective pre-edge region [Fig. 4(e) and (f)]. Thus, the electronic states of the substituted Co and Ni are the same as those in Co- and Ni-HCFs, respectively. This is consistent with the fact that the ligand (CN⁻) environments around substituted Co and Ni are the same as those Co- and Ni-HCFs, respectively [see Fig. 1(b) and Fig. 3(b)].

Discussion

Now, let us discuss the difference in the local structures between the M-HCF and layered oxides. In M-HCF, the local structure around M_{ρ} is essentially the same as that in the pure compound. In layered oxide, however, the $M_{\rm g}$ valence changes so that $d_{M-0}{}^{\rm g}$ approaches $d_{M-0}{}^{\rm h}$. Such a change in $d_{M-0}{}^{\rm g}$ relaxes the local distortion and lowers the Gibbs free energy of the system. Why is local structure around M_{g} invariant in M-HCF and is not in layered oxide? We ascribed the invariant nature in M- HCF to (1) strong ionicity of $[Fe(CN)_6]^{4-}$ and (2) the structural flexibility in the jungle-gym type network. First of all, $[Fe(CN)_6]^{4-}$ is a hard anion with a stable closed electronic configuration. In fact, the profiles of the XANES spectra around Fe K-edge are the same for pure Mn-, Co-, and Ni-HCFs [Fig. 3S(a) and (c)]. Then, it is difficult for M_g to exchange charge with $[Fe(CN)_6]^{4-}$. Consistently, the XANES spectra around Fe K-edge hardly change in the mixed compounds [Fig. 3S(b) and (d)]. By means of the EXAFS analyses, we further investigated $d_{\text{Fe-C}}$ (and $d_{\text{Fe-N}}$) in the nine compounds (Table S2). We found that $d_{\text{Fe-C}}$ (and $d_{\text{Fe-N}}$) is the same for the nine compounds within the experimental errors. By contract, in transition metal oxides, the hybridization between the O 2p and M 3d orbitals effectively causes the M-dependent charge exchange between O and M. Such a hybridization can modify the M_{σ} valence to minimize the Gibbs free energy. Secondary, the jungle-gym type network of M-HCF is flexible against the partial substitution. This is because the network is fairly sparse and has considerable $Fe(CN)_6$ vacancies. Actually, the density (~1.9 g/cm₃) of *M*-HCF is much smaller than that (=5.0 g/cm³) of layered oxides. With such a structure, local elongation (compression) of d_{M-N} ^g in the *M*-NC-Fe-CN- M_g -V-*M*-NC-(V represents the vacancy) chain can be compensated by off-axial rotation of the $Fe(CN)_6$ unit (axial displacement of M_g). By contract, in layered oxides, the Na, M, and O ions form respective triangular lattices. The atomic sheets stacks along the *c*-axis in the order of Na, O, M, O, and Na. In such a pseudo-close-packed structure, the local distortion induced by M_g is hard to be compensated.

Summary

By means of the systematic EXAFS and XANES analyses, we found that the local structure and electronic state of M_g in *M*-HCF are essentially the same as those in the pure material, *i.e.*, M_g -HCF. Such an invariant nature observed in *M*-HCF is in sharp contrast with the layered oxide systems. We ascribed the invariant nature to (1) strong iconicity of [Fe(CN)₆]₄₋ and (2) the structural flexibility in the jungle-gym type network. Our observation indicates that *M*-HCF is a novel platform of transition metals, in which magnetic, electronic, local structural properties of them are the same as those in the pure compound. With these characteristics, we can easily design and/or calculate the macroscopic magnetic, electric, structural properties of mixed *M*-HCF from the atomic level.

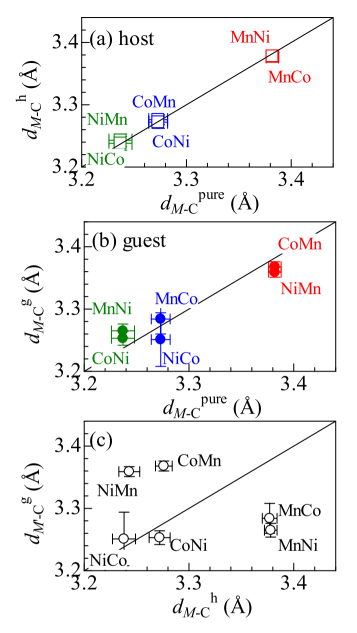


Figure 3. (a) Correlation diagram of the M-C distances (d_{M-C}) between the mixed compounds $(d_{M-C})^h$ around M_h and pure compounds. (b) Correlation diagram of d_{M-C} between the mixed compounds $(d_{M-C})^g$ around M_g and pure compounds. (c) Comparison between d_{M-C}^g and d_{M-C}^h in the mixed compounds. M_hM_g represents $Na_x(M_h, M_g)$ [Fe(CN)₆]_y. Red, blue and green colors represent the interatomic distances around Mn, Co, and Ni, respectively.

Method

Sample preparation and chemical composition. We prepared pure (*M*-HCF; M = Mn, Co, and Ni) and mixed (M_hM_g -HCF; M_h and $M_g = Mn$, Co, and Ni) compounds by participation method from aqueous solutions in air at 40 °C. In the synthesis of pure compounds, an aqueous solution (40 mM MCl_2 : M = Mn, Co, and Ni) was slowly dropped to an aqueous solution (40 mM Na₄[Fe^{II}(CN)₆] and 4M NaCl). In the synthesis of mixed compounds, an aqueous solution (32 mM M_hCl_2 , 8 mM M_gCl_2 , and 4 M NaCl). In the synthesis of mixed slowly dropped to an aqueous solution (40 mM Na₄[Fe^{II}(CN)₆] and 4 M NaCl). In both the cases, the latter solution was slowly dropped to an aqueous solution (40 mM Na₄[Fe^{II}(CN)₆] and 4 M NaCl). In both the cases, the latter solution was stirred at 250 rpm with a magnetic stirrer during the instillation. The dropping rate (= 100 ml/hour) was controlled with use of a tube pump. After the instillation, the solutions were kept for 12 hours. Then, the precipitates were gathered with a 0.1 µm filter, washed well with distilled water, and dried in air. The colors of the obtained powders were white (Mn-HCF), light yellow (MnNi-HCF), light green (Ni-, MnCo-, and CoMn-HCF), light blue (NiMn-HCF), and dark green (Co-, CoNi-, and NiCo-HCF).

The chemical compositions of the pure compounds were determined so as minimize the trial function:

$$\mathbf{F}(y, z) = \sum_{i=\mathrm{Na}, M, \mathrm{Fe}} (\mathrm{wt}\%^i_{\mathrm{obs}} - \mathrm{wt}\%^i_{\mathrm{cal}})^2,$$

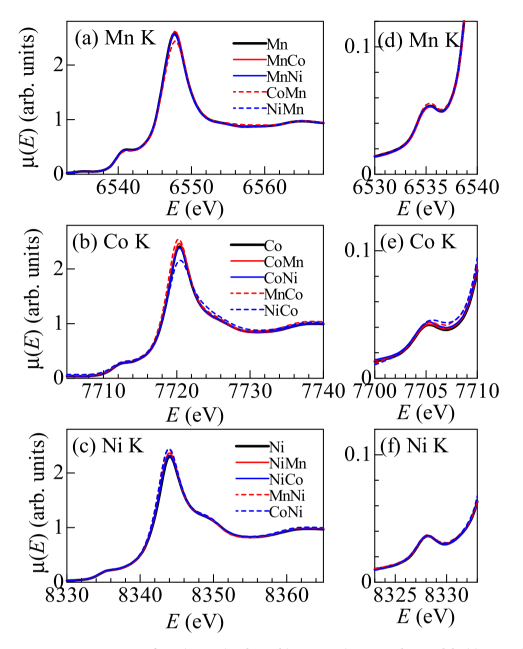


Figure 4. XANES spectra of pure (*M*-HCF) and mixed (M_hM_g -HCF) compounds around the (**a**) Mn K-edge, (**b**) Co K-edge, and (**c**) Ni K-edge. Thick black curves represent the spectra of pure compounds. Thin solid and broken curves correspond to the spectra around M_h and M_g in the mixed compounds, respectively. (**d**)–(**f**) Magnified spectra in the pre-edge region.

where wt%_{obs}^{*i*} and wt%_{cal}^{*i*} are the experimentally-obtained and calculated weight percent of the *i*-th elements, respectively. wt%_{obs}^{*i*} for the metal elements were determined by the inductively-coupled plasma (ICP) method. In the calculation, we assume the chemical formula: Na_{4y-2}*M*[Fe(CN)₆]_y*z*H₂O. The chemical compositions of the mixed compounds were determined so as minimize the trial function:

$$\mathbf{F}(x, y, z) = \sum_{i=\mathrm{Na}, M_{\mathrm{g}}, M_{\mathrm{h}}, \mathrm{Fe}} (\mathrm{wt}\%^{i}_{\mathrm{obs}} - \mathrm{wt}\%^{i}_{\mathrm{cal}})^{2}.$$

wt%_{obs}ⁱ for the metal elements were determined by the ICP method. In the calculation, we assume the chemical formula: $Na_{4y-2}M_{g1-x}M_{hx}[Fe(CN)_6]_yzH_2O$. Thus determined chemical compositions are listed in Table 2.

Crystal structure and lattice constants. Synchrotron-radiation XRD measurements were performed at the BL8A beamline of the Photon Factory, KEK. The wavelength (=0.689028 Å) of the X-ray was calibrated by the lattice constant of standard CeO₂ powders. The samples were finely ground and placed in 0.3 mm ϕ glass capillaries. The capillaries were sealed and mounted on the Debye-Scherrer camera. The XRD patterns (Fig. S1) were detected with an imaging plate. The exposure time was 5 minutes. The XRD patterns of Ni-, NiCo-, and

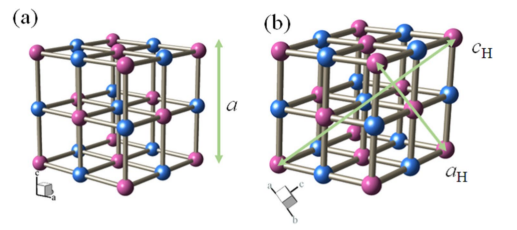


Figure 5. Schematic pictures of (a) fcc and (b) trigonal (hexagonal setting) structures. Red and blue spheres represents Fe and *M*, respectively. Bars represent CN. Na and H_2O are omitted for simplicity.

NiMn-HCFs can be indexed in the face-centered cubic (fcc) structure (Fm3(-)m; Z=4). The lattice constants, a, were refined by the Rietveld method (Rietan-FP²⁴). The XRD patterns of Co-, CoMn-, CoNi-, Mn-, MnCo-, and MnNi-HCFs can be indexed in the trigonal structure (R3(-)m; Z=3). The lattice constants, a_H and c_H , were refined in the hexagonal setting. The lattice constants, a, a_H , and c_H , are listed in Table 2. Figure 5 shows schematic pictures of the (a) fcc and (b) trigonal (hexagonal setting) structures. The two structures are essentially the same expect for the slight elongation along the (111) direction in the trigonal structure.

X-ray absorption spectroscopy. The X-ray absorption spectroscopy (XAS) measurements were conducted at BL-9C of the Photon Factory, KEK. The powder was finely ground, mixed with BN, and pressed into pellets with 5 mm in diameter. The XAS were recorded in a transmission mode with a Si(111) double-crystal monochromator at 300 K. The wavelengths of the monochromator were calibrated with the absorption edge of Fe, Mn, Co, and Ni foil. In X-ray absorption near edge structure (XANES) analyses, the background subtraction and normalization were performed using the ATHENA program²⁵.

EXAFS analysis. After the background subtraction and normalization of the XAS spectra, oscillatory components χ were extracted against the angular wavenumber k. k is defined by $k = \sqrt{2m(E - E_o)/\hbar}$ where m, E, and E_0 are the electron mass, energy of the incident X-ray, and absorption edge energy, respectively. Then, Fourier transformation of $\chi(k)k^3$ was performed in the k-range from 3.0 to 11.0 Å⁻¹ These data procedures were performed with use of ATHENA program²⁵.

In order to refine the structural parameters, least-squares fittings are performed for the $FT[\chi(k)k^3] - R$ plots with use of ARTEMIS program²⁵. In the plane wave and single-scattering approximation, $\chi(k)$ around the K-edge is expressed by the EXAFS equation as:

$$\chi(k) = -S_0^2 \sum_j \frac{N_j}{kR_j^2} F_j(k) e^{-2\sigma_j k^2} \sin\{2kR_j + \varphi_j(k)\},\$$

where S_0 , N_i , R_j , F_j , σ_j^2 , and φ_j are the passive electron reduction factor, degeneracy of path, path length, effective scattering amplitude, mean square displacement, and effective scattering phase shift of the *j*th atom, respectively. The least-squares fittings were performed in the *R* range from 1 Å to 3.2 Å. S_0^2 for each absorption edge were estimated from the pure compounds: $S_0^2(Fe) = 0.92$, $S_0^2(Mn) = 0.97$, $S_0^2(Co) = 1.09$, and $S_0^2(Ni) = 1.03$. These S_0^2 values were use in the analyses of the mixed compounds. Practically speaking, S_0^2 lies within a general range between 0.7 and 1.1^{26} . In this sense, the other structural parameters obtained the EXAFS analysis are valid, even though $S_0^2 > 1$ is physically meaningless. In the analyses around the Mn, Co, and Ni K-edges, we included the contributions from the first- (N) and second-(C) nearest neighbor elements. Because of the linear $_{M-N}$ -C-Fe coordination, single (*M*-C-*M*), double (*M*-C-N-*M*), and triple (M-N-C-N-*M*) scattering paths were taken into account in the analyses of second nearest neighbor elements (C). In order to include the Fe(CN)₆ vacancy effect, N_N are treated as an adjustable parameter with restriction of $N_N + N_O = 6$ and $d_{M-O} = d_{M-N}$. Figure 2S shows prototypical examples of the fitting. The obtained structural parameters are listed in Table S1. In the analyses around the Fe K-edge, we included the contributions from the first- (C) and second- (N) nearest neighbor elements. Because of the linear Fe-C-N-*M* coordination, single (Fe-N-Fe), double (Fe-N-C-Fe), and triple (Fe-C-N-C-Fe) scattering paths were taken into account in the analyses of second nearest neighbor elements (N). We fixed N_C and N_N at 6. The obtained structural parameters are listed in Table S2.

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Author Contributions

H.N. performed sample preparation and characterization, XAS measurements, and careful EXAFS analyses. W.K. supported the X-ray diffraction measurements at PF 8 A. T.S. provided advice on the synthesis of *M*-HCF. H.N. (KEK) collaborated in the EXAFS measurements at PF 9 C as beamline staff. Y.M. planned the research, refined the lattice constants, and wrote the manuscript.

Additional Information

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