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## Structure motif of chemical short-range order in a medium-entropy alloy

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### ABSTRACT

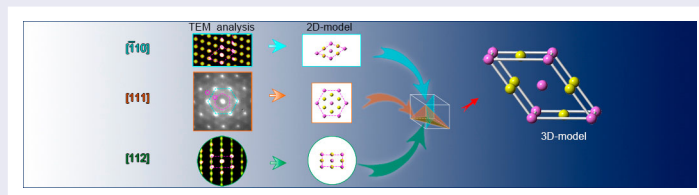
The chemical short-range orders (CSRO) are the sub-nanosized entities inherent in high-/medium-entropy alloys (H/MEA). To discern CSRO poses the challenge with pending issues as to under what zone axis in electron diffraction, signature scattering specific to CSRO will appear and what the structure motif of CSRO is. Here, we show extra diffuse scattering by CSRO from [111] and [112] directions in a VCoNi MEA and accordingly, construct the L<sub>1</sub>-type structure motif, and derive the spacial shape of flat cuboid of CSRO in virtue of diffractions under varying zone axis. Moreover, we demonstrate the methods to identify CSRO in lattice images.

### ARTICLE HISTORY

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### KEYWORDS

Chemical short-range order; electron diffraction; unit cell; medium-entropy alloy



### IMPACT STATEMENT

The structure motif is constructed as L<sub>1</sub>-type for chemical short-range order in a medium-entropy alloy in terms of electron diffractions under the multiple zone axis.

## 1. Introduction

High-/medium-entropy alloys (H/MEA) are the complex concentrated solid solution composed of multi-principal elements [1,2], showing an improved synergy between strength and ductility [3–7]. In H/MEA, constituent species would be much likely to make enthalpic interactions large enough locally such that the local chemical order (LCO) is induced in the basal solid solution [8–11]. The chemical short-range order (CSRO) is the incipient LCO in H/MEAs [9,10], akin to the counterparts in the conventional alloys [12–19]. By definition, CSRO refers to the preference/avoidance of some species in the alloy away from atomic fractions expected for random mixing, but the chemical order is short-ranged such that it prevails only in the first and the next couple of nearest-neighbor atomic shells around the atom at center.

In this sense, CSRO is usually at the sub-nanoscale in size, without lattice periodicity. CSRO is considered as an iconic entities inherent in H/MEA [20–29], playing a key role in dislocation behaviors [10,27], strain hardening [27], and macroscopic mechanical properties such as strength and ductility [11,21,22,25].

So far how to identify convincingly CSRO becomes undisputedly the top question. To demonstrate CSRO, there are two kinds of experimental techniques. One is indirect, such as the fitting of the X-ray or neutron data [30] as well as the three-dimensional atom probe tomography (3D-APT) [31]; while the other is the ‘seeing-is-believing’ strategy by the direct observation of CSRO using the state-of-the-art (e.g. high-resolution, HR) transmission electron microscope (TEM) [21,24,27]. For the latter, electron diffraction seems the most appropriate

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way capable of directly digging out the direct evidence of CSRO [14–17,24,27,32–34]. To be specific, in the selected-area electron diffraction pattern, CSRO may give off the extra superlattice scattering, usually in the form of diffuse disks with the diameters substantially larger than those of Bragg spots of base lattices [34]. Such diffraction information in reciprocal space demonstrates the presence of CSRO entities in real space, with the sizes at the (sub-)nanometer scale [34]. But for H/MEAs, the challenge emerges to identify CSRO in this way. On one hand, the basis to discern CSRO is the proprietary, CSRO-induced scattering, even much diffuse. Yet, these reflections show a much low intensity due to the moderate volume density of CSROs and their small sizes typically below one nanometer. The scattering signals are, therefore, much weak and highly diffuse such that it is hard to distinguish them from the noised background [24,27], even though an in-column  $\Omega$  energy filtered TEM is used to enhance the signal-to-noise ratio of diffraction contrast from CSRO [24]. On the other hand, an appropriate zone axis is necessary to display reflections by CSRO, as the diffraction intensity varies with crystallographic orientations [27,34]. On this account, it is better to check various zone axes for diffraction patterns. For example, scattering by CSRO is discernable under the [112] zone axis in an fcc base [27], yet disappear under the routinely used [110] one. Simply, two puzzling issues remain as to how likely scattering by CSRO appears under a specific zone axis and whether or not scattering under various zone axis is ascribed to the same CSRO.

The electron diffractions under multiple zone axes would help to establish the structure motif of CSRO, i.e. the three-dimensional (3D) unit cell of chemical occupation of species [30]. The structure motif inevitably exists in CSRO even without the definite lattice periodicity due to small sizes. The structure motif will facilitate an in-depth understanding of the CSRO formation and particularly, the relationship between CSRO and host phase, and the evolution of CSRO later into the medium-/long-range order.

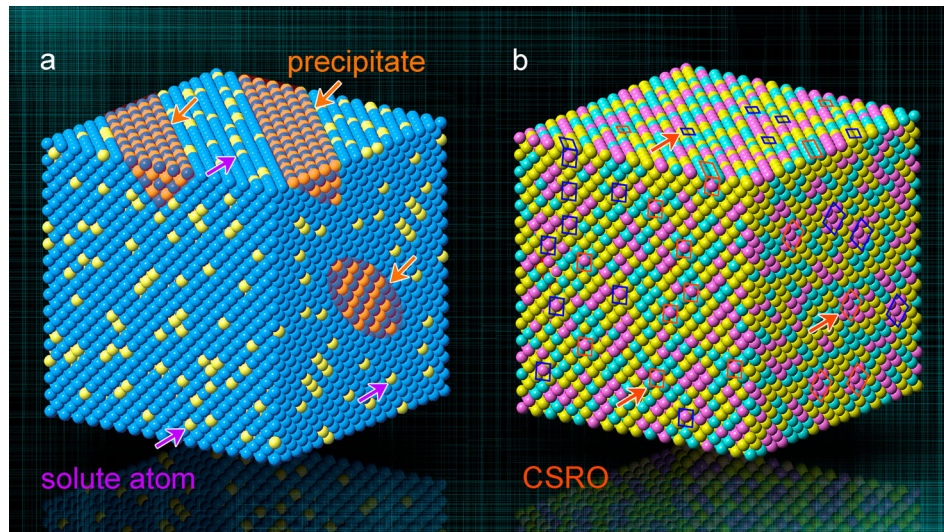
Here, an equiatomic ternary VCoNi MEA was selected as the model material [27,35,36]. Following the processing conditions in the early investigation [27], the microstructure was obtained where the CSRO is formed in a face-center-cubic (fcc) base after recrystallization annealing at 880°C for 3 min. We will look for more zone axes having electron diffractions by CSRO, in addition to the [112] zone axis reported already [27]. On this basis, we will build up the structure motif of CSRO with the help of mutual relationship and lattice correspondence between real and reciprocal spaces. Finally, the three-dimensional spacial shape of CSRO is derived.

## 2. Results and discussion

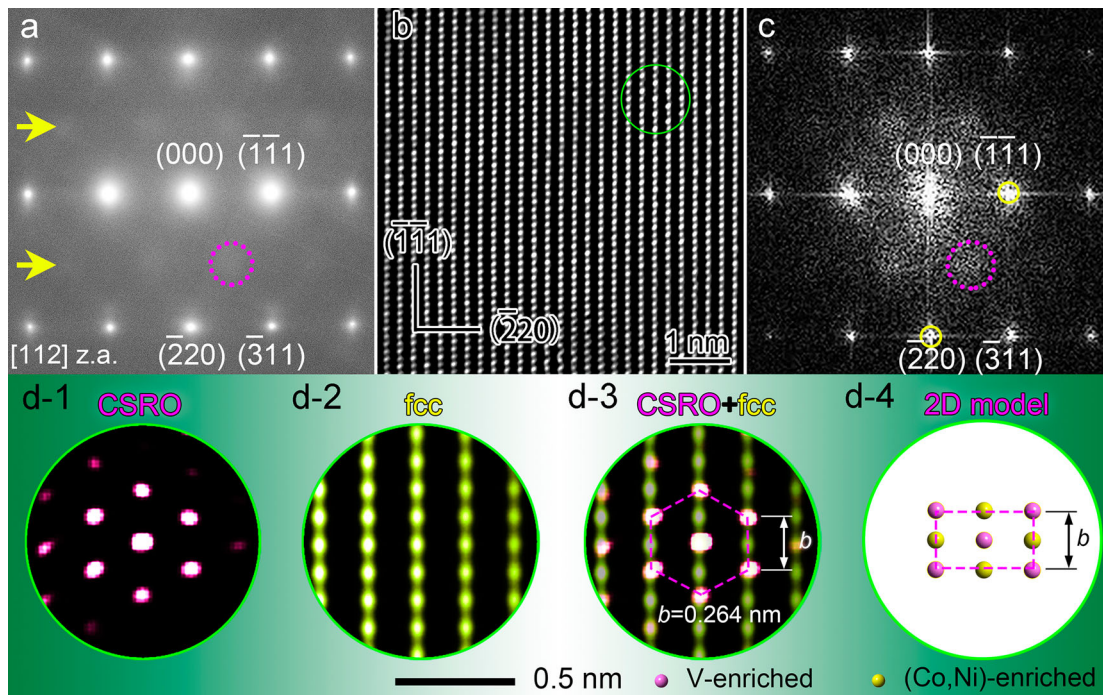
Solute atoms, labeled by solid spheres in yellow, Figure 1(a), in conventional alloys usually show the typical configurations of two kinds, i.e. those in a solution of solvent atoms and those in the form of a precipitate. Thereinto, atoms are randomly mixed in the former (purple arrows), while appear periodic in the latter (orange arrows) with solute concentration often much higher than that in the base. In contrast, the configuration of atoms in the H/MEA is different. Simply taking a ternary MEA as an example, Figure 1(b), three principal elements (labeled by solid spheres, respectively, in yellow, pink, and turquoise), all are random and disordered in a base solution of single phase. Lots of CSROs dispersedly distribute (marked by blue and red frames). As a result, CSRO shows two features by comparison to a precipitate. Firstly, atoms in CSRO still occupy the lattice sites of disordered base, but with at least one element forming the short-range order. Second, CSRO is of the short ordered period, usually less than 1 nm.

In the following, the electron diffraction patterns under several zone axes (z.a.) are demonstrated to find out the z.a. as many as possible with reflections specific to CSRO. Figure 2(a) is the selected-area electron diffraction (SAED) pattern under the [112] z.a. The extra superlattice scattering appears in the form of diffuse disks, lining up at the  $\frac{1}{2}\{\bar{3}11\}$  positions of fcc lattice as indicated by two yellow arrows and one is marked by pink circle. These diffuse disks in reciprocal space are the conclusive evidence for CSRO entities of small sizes in real space [27,28,34]. These scattering is the corollary to the rule-of-shape-factor between real and reciprocal space, i.e. ‘small becomes large’ and vice versa [16,32,33].

The HRTEM image of fcc base is obtained with no sign of CSRO, Figure 2(b). Next, a simple approach is proposed to find out where CSRO is. The first step is to obtain the fast Fourier transform (FFT) pattern, Figure 2(c), showing diffuse scattering by CSRO (one is circled in pink) and sharp reflections of fcc base (yellow circle). The second is to obtain the inverse FFT (IFFT) images for CSRO (white spots, Figure 2(d-1)) and fcc base (green spots, Figure 2(d-2)) by using their respective reflection. These two IFFT images come from the small area (green circle) in Figure 2(b). The final step to take is to merge above two IFFT images, Figure 2(d-3), where one CSRO (pink hexagon by white spots) is clearly visible on the fcc lattice, with spacing ( $\lambda$ ) of (113) plane in CSRO twice that in fcc base. Based on the measurements by the atomic-resolution EDS mapping [27], CSRO consists of a (Co,Ni)-enriched plane sandwiched by two V-enriched planes, as illustrated in Figure 2(d-3). Resultantly, the two-dimensional CSRO model is established



**Figure 1. Schematic distribution of atoms.** (a) Conventional alloy internally with solute atoms (yellow atoms) and precipitates (orange area). (b) Ternary medium-entropy alloy with CSROs labeled by blue and red frame.

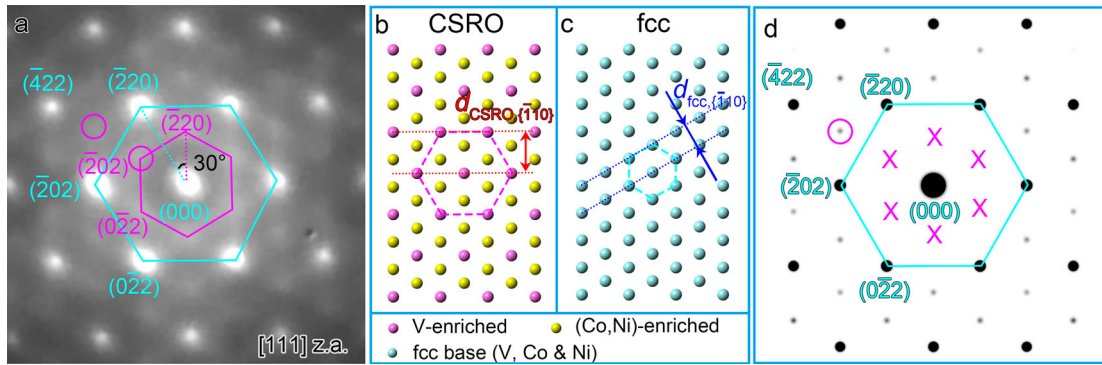


**Figure 2. Electron diffraction of CSRO under [112] zone axis.** (a) SAED pattern. Yellow arrows and dash circle in pink: extra diffuse scattering by CSROs at  $\frac{1}{2}\{\bar{3}11\}$  positions of fcc base. (b) Lattice image of fcc base under [112] z.a. (c) FFT pattern corresponding to (b). Pink and yellow circles: diffuse scattering by the CSRO and sharp diffraction spot by base fcc. (d-1 & d-2) Inverse FFT patterns of CSRO and fcc base, respectively, in the area circled in green in (b). (d-3) Merged inverse FFT pattern of (d-1) and (d-2). Dash hexagon: CSRO.  $b$ : lattice parameter of unit cell in the CSRO, which is measured to be 0.264 nm. (d-4) Two-dimensional model of the CSRO. Solid spheres in pink and yellow: V-enriched and (Co,Ni)-enriched atom column. Note that the pink column is not entirely V atoms, also with (Co,Ni) atoms to certain contents.

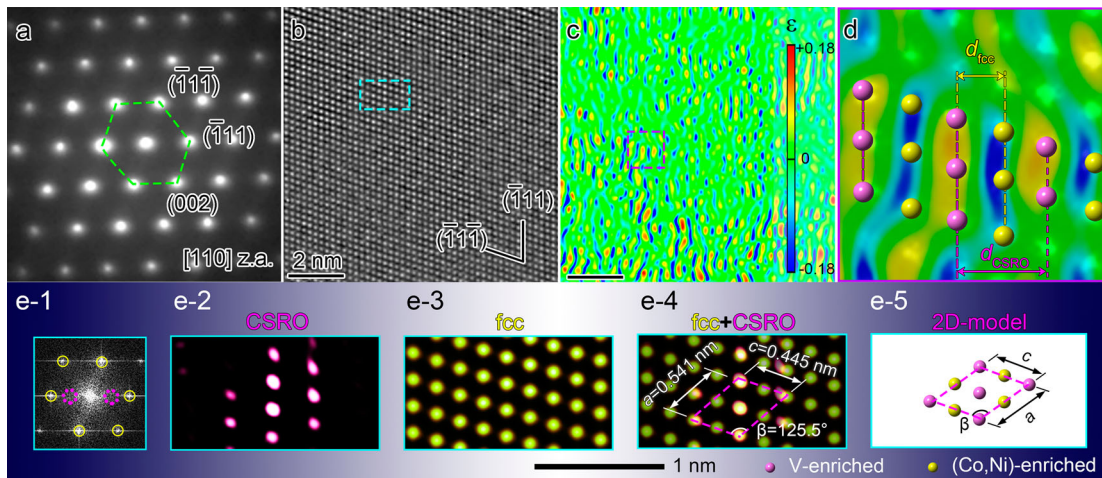
in the [112] direction, Figure 2(d-4). V atoms (spheres in pink) occupy the vertices of unit cell, while both Co and Ni atoms (in yellow) locate in-between [27].

The extra superlattice scattering by CSRO is further observed under the [111] z.a., Figure 3(a). These diffuse reflections are at the positions of  $\frac{1}{3}\{\bar{4}22\}$

of fcc base, one for each is circled in pink. The diffraction patterns are indexed by pink and turquoise lines, respectively, for CSRO and fcc base. Two (110) planes in CSRO and fcc base form an angle of  $30^\circ$ , i.e.  $(110)_{\text{CSRO}}$  and  $(110)_{\text{fcc}}$ . The distance (d) from the center of fcc base (dash line in turquoise) to  $(\bar{2}20)$  spot is



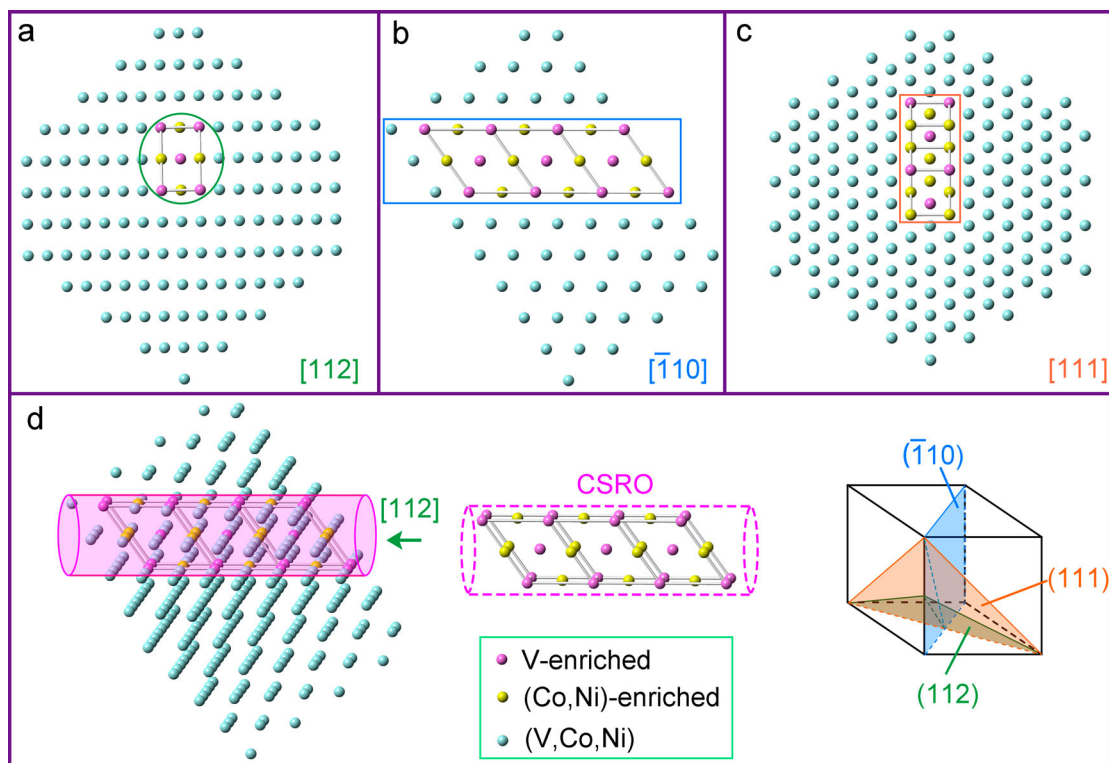
**Figure 3. Electron diffraction pattern of CSRO under [111] zone axis.** (a) Diffraction pattern. Note two positions for superlattice scattering at both  $1/3\{422\}$  and  $2/3\{422\}$  of fcc base; one for each is circled in pink. (b) and (c) 2D model for CSRO and fcc base. Spacing ( $\lambda$ ) of (110) planes are marked. (d) Simulated diffraction pattern. Pink circle: scattering by CSRO. Pink cross: extinction of CSRO.



**Figure 4. Electron diffraction of CSRO with [110] zone axis.** (a) Nano-beam diffraction pattern. (b) Lattice image with [110] z.a. (c) GPA strain map imaged by  $(\bar{1}11)$  reflection. Scale bar: tensile (positive) and compressive (negative) strain. (d) Close-up view of GPA map in the purple square in (c). (e-1) FFT pattern of fcc base in (b). (e-2) and (e-3) Inverse FFT images for CSRO and fcc. (e-4) Merged inverse FFT image, showing the CSROs on fcc base.  $a$ ,  $c$  and  $\beta$ : unit cell parameters. (e-5) 2D unit cell model. Solid spheres in pink and yellow: atom column of V- and (Co,Ni)-enriched.

$\sqrt{3}$  times of that (dash line in pink) for CSRO. In other words,  $d_{CSRO, \{110\}} = \sqrt{3}d_{fcc, \{110\}}$ , see the graphic details as shown in Figure 3(b,c). Accordingly, the 2D model of CSRO is established when viewing along the [111] z.a., Figure 3(b). Thereinto, the hexagon (dash line in pink) is the periodic, minimum unit cell of CSRO, in contrast to that (turquoise line) in fcc base, Figure 3(c). Further, the simulated diffraction pattern is obtained, Figure 3(d), where the  $\{202\}$  reflections (labeled by Xs in pink) are forbidden for CSRO. This is in contrast to the scattering by CSRO at the positions of  $\frac{1}{3}\{422\}$  fcc base, even faint but clearly visible, Figure 3(a). The reason is that the model in Figure 3(d) is derived under the assumption of the long-range, disordered arrangement of atoms in a precipitate. However, CSRO is chemically ordered, even short-ranged. As such, scattering by CSRO does not satisfy the strict extinction law and makes the diffraction deviate from the theoretical extinction distance.

Finally, the [110] z.a. is checked. At least seemingly, this is the most suitable z.a. to identify CSRO [24], but unfortunately, no convincing scattering by CSRO is available so far. The same is true in the present VCoNi MEA, Figure 4(a). Here, another method is demonstrated to find out CSRO based on the atomic-scale geometric phase analysis (GPA). The first step is to obtain the HRTEM lattice image under the [110] z.a., Figure 4(b). The second is to obtain the GPA strain map, Figure 4(c), with the locally periodic distribution of strain contrasts. This local strain is derived based on the displacement field [37]. This method provides strain distribution induced by CSRO. In a simplified way, here, just considering the difference of atomic radii in V (radius of 1.35 Å), Co (1.26 Å), and Ni (1.24 Å), the column of larger V atoms will cause positive strains, as schematically shown in Figure 4(d), provided that these columns belong to CSRO which projects to the (111) plane. In



**Figure 5. 3D unit cell of the CSRO.** (a–c) 2D models with the [112],  $[\bar{1}10]$ , and [111] zone axes, respectively. Pink and yellow spheres: V and (Co,Ni)-enriched atom positions; turquoise spheres: randomized (V,Co&Ni) atom positions. (d) 3D unit cell of the CSRO with a monoclinic lattice (Left panel). Middle panel: 3D unit cell which is embedded in fcc base. Note the [112] direction which is parallel to the major axis of the CSRO. Right panel: three typical crystallographic orientations in fcc lattice.

the vicinity, negative strain is caused by the columns of smaller Co/Ni atoms in CSRO. Note that the spacing of  $(\bar{1}11)$  planes in CSRO is twice that in fcc base, Figure 4(d). It is therefore supposed that extra scattering by CSRO should be at the positions of  $\frac{1}{2}\{\bar{1}11\}$ , see dash circles in pink, Figure 4(e-1), i.e. the FFT pattern from Figure 4(b). Indeed, scattering by CSRO (two pink circles) is visible, even much weak in intensity. The last is to obtain the IFFT images for both CSRO and fcc base, Figure 4(e-2 and e-3), corresponding to the area (dash parallelogram in turquoise) in Figure 4(b). Indeed, there exists a CSRO, Figure 4(e-2). By merging these two IFFT images, this CSRO is clearly seen to lay on the fcc base, Figure 4(e-4). Based on it, the 2-D model is established with the [110] z.a., Figure 4(e-5). Three parameters,  $a$ ,  $c$  and  $\beta$ , are obtained in Figure 4(e-4), both as lattice parameters of unit cell.

Three 2D models of CSRO are summarized, see Figure 5(a–c), according to diffraction patterns from three independent [112], [111], and  $[\bar{1}10]$  z.a. The principle, similar to the engineering drawing, was applied to build up the 3D model. To be specific, by analyzing three 2D models and respective TEM projection in a way similar to the three orthographic views of a component [38], complete information is obtained of both

lattice structure and order occupation of three elements. As a result, the structure motif of CSRO, i.e. 3D unit cell, is reconstructed, left panel in Figure 5(d). To well recognize CSRO from fcc base, V and (Co,Ni) atoms in CSRO are colored, respectively, in pink and yellow; while randomly distributed V, Co, and Ni atoms are turquoise in the fcc base. Based on it, the space group is determined to be P2 for CSRO, which is a monoclinic lattice structure, with  $\beta = 125.5^\circ$ , and lattice parameters of  $a = 0.541$  nm,  $b = 0.264$  nm, and  $c = 0.445$  nm, which are obtained from the measured real space of atoms in HRTEM images, Figure 2(d) and 4(e). The stoichiometry is  $V_2Co_xNi_{(2-x)}$  according to the 3D unit cell. Meanwhile, the symmetry is much lower than that in fcc base (Fm-3m). After the atomic occupation in CSRO is rearranged as a result of the chemical order, interestingly, this CSRO unit cell is much similarly to that of  $L1_1$  [39], with  $a = 0.747$  nm and  $\gamma = 89.9^\circ$ .

Based on the determined structure motif of CSRO, the 3D spacial shape of CSRO is a flat cuboid, outlined in pink, left panel in Figure 5(d). It follows that to see CSRO depends on not only this 3D spacial shape of CSRO but also the orientation relationship of CSRO with fcc base. First, more CSROs will be seen if viewed from the [112] z.a. which is parallel to the major axis of CSRO. However,

this does not necessarily mean that it is true under all  $< 112 > z.a.$  For example, both  $[121]$  and  $[211]$  z.a. form an angle of  $33.6^\circ$  with the major axis of CSRO. This leads to a decrease in the proportion of CSRO inside the spacial column for diffraction imaging in TEM thin film. The diffraction intensity by CSRO will become much weak such that it is hard to distinguish them from the noised background [24,27]. Second, the  $[110]$  z.a. is parallel to the minor axis of CSRO, right panel in Figure 5(d). Accordingly, the diffraction intensity by CSRO is much weak by comparison with that under the  $[112]$  z.a. This is the reason that it is so difficult to observe extra scattering along the  $[110]$  z.a. Generally, CSRO may be seen or not under all  $[110]$  z.a., depending on the orientation relationship between CSRO and zone axis. The same is true for the  $[111]$  z.a. Simply to say, whether or not diffuse diffraction by CSRO appear depends on the spacial shape of CSRO and the orientation relationship between CSRO and zone axis, along with the thickness of TEM thin film for CSRO characterization.

### 3. Conclusion

It is a nontrivial challenge to find out CSRO in the microstructure of H/MEA. With regard to electron diffraction for seeking out scattering signals by CSRO, a step forward in this letter is the  $[111]$  zone axis showing diffuse scattering specific to CSRO. More importantly, the structure motif of CSRO, i.e. 3D unit cell of species occupation, is constructed as the ordered  $L1_1$ -type intermetallic compound, based on the crystallographic orientation relationship between the CSRO and host fcc base by virtue of multiple electron diffractions.

### Disclosure statement

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

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