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# Surface functionalization of discrete metal-chalcogenide supertetrahedral clusters and the photocatalytic application†

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Atomically precise metal-chalcogenide supertetrahedral clusters (MCSCs) are supposed to be more attractive for functionalization than conventional metal-sulfide quantum dots owing to their potential ability to establish precise structure–composition–property relationships. However, the accurate surface functionalization of such cluster-based species remains difficult. In this paper, we present a facile method for synthesizing discrete MCSCs decorated with different functional groups *via* a one-step solvothermal reaction, which was demonstrated to have better solvent dispersibility compared with ligand-free ones. In addition, the composites were also prepared by combining ligand-free clusters (or ligand-partially protected or amino-modified ones) with two-dimensional MXene nanosheets. The composites derived from amino-modified clusters exhibited optimal performance of photocatalytic hydrogen evolution. Furthermore, the hydrogen bonding interactions between modified amino groups and MXene nanosheets were verified by <sup>1</sup>H-NMR spectroscopy. This work provides a facile approach for the surface functionalization of MCSCs, and facilitates the expansion of the functionality of atomically precise nano-species.

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## Introduction

Discrete metal-chalcogenide supertetrahedral clusters (MCSCs), known as ultra-small “quantum dots”, represent regularly fragmented units of zinc blende type materials.<sup>1–6</sup> Since their structures can be accurately identified at the atomic level, they have been widely studied in the fields of fluorescence,<sup>7–13</sup> photocatalysis,<sup>14–20</sup> and electrocatalysis<sup>21–25</sup> for establishing an accurate structure–composition–property relationship. However, such clusters usually exhibit poor dispersibility and conductivity during practical applications, and the weak binding forces between clusters and conductive substrates

limit their further development.<sup>26–32</sup> Therefore, rational modification of clusters for improving the interactions between clusters and conductive substrates would be helpful for their applications in catalysis.

Owing to the advancement of organic template-guided synthesis, most of the discrete T<sub>n</sub> and P<sub>n</sub> clusters (two main types of MCSC structures, where *n* represents the number of metal layers in the basic cluster units, Scheme S1†) reported in recent years are predominantly ligand-free types.<sup>29</sup> The high-degree negative charges of these clusters are usually balanced by protonated organic amines (*i.e.*, counterions), which results in poor dispersibility and instability. Alternatively, a group of C<sub>n</sub> clusters (fully coated with organic ligands, Scheme S1†) has superior dispersibility. However, their monomeric composition and poor structural tunability make them a suboptimal choice for subsequent functional applications. Therefore, by combining the advantages of the above two types of structures, a series of ligand-partially protected T<sub>n</sub> and P<sub>n</sub> clusters has been developed (Scheme S2†). The few ligands at the cluster surface help to improve their contact with the solution molecules and reduce their negative charge density, thereby improving structural stability and dispersibility. This feature is well-suited for functional expansion, and makes them a promising choice.

Since MCSCs generally exhibit poor electrical conductivity, combining them with a suitable conductive substrate is desirable for promoting the separation of the photogenerated electrons and holes, which may effectively enhance photo-

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† Electronic supplementary information (ESI) available: Supplementary tables and figures, PXRD patterns, EDS and <sup>1</sup>H-NMR analysis. CCDC 2254140–2254148 for ISC-24, ISC-25, ISC-25-4'-NH<sub>2</sub>, ISC-25-4'-CH<sub>3</sub>, ISC-25-4'-OH, ISC-25-4'-F, ISC-25-4'-OCH<sub>3</sub>, ISC-26-2'-NH<sub>2</sub> and ISC-26-4'-Cl. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3qi00983a>

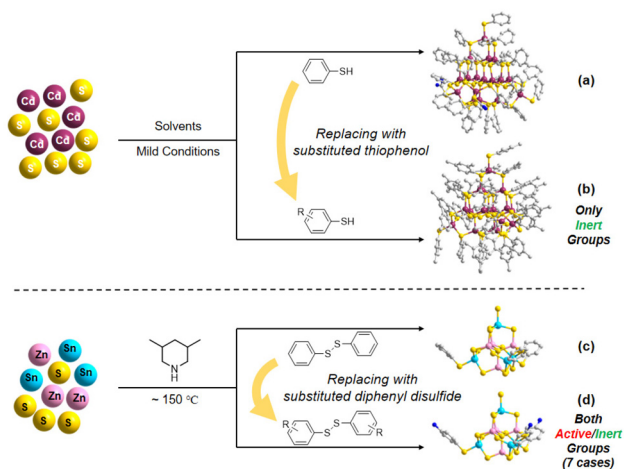
catalytic performance. In recent decades, MXenes, a layered transition metal carbide/carbonitride/nitride family, have attracted attention in many scientific fields, such as energy storage and conversion,<sup>33</sup> sensing,<sup>34</sup> and photocatalysis.<sup>35–38</sup> In the domain of photocatalytic hydrogen evolution (PHE),  $\text{Ti}_3\text{C}_2\text{T}_x$  is one of the most studied MXenes for the following reasons: (1) the favorable hydrophilic ability supports its chemical reactions within aqueous solutions;<sup>39</sup> (2) the excellent electrical conductivity promotes the separation and transfer of the photoinduced charge carriers from semiconductors;<sup>40</sup> and (3) the appropriate work function allows it to form a Schottky heterojunction with most semiconductors. Therefore, the ultra-thin  $\text{Ti}_3\text{C}_2\text{T}_x$  substrate is regarded as a suitable and promising substrate, in which abundant surface O/F sites can also serve as potential hydrogen bonding receptors for anchoring discrete MCSCs.

This work aims to improve the interactions between a cluster and a conductive substrate by manipulating the cluster surface environment, and eventually, improve the performance of photocatalytic hydrogen evolution (PHE). On the basis of the synthetic route of *Pn*-type clusters that are partially protected by ligands,<sup>32</sup> we changed the types of diphenyl disulfides to modify the cluster surface with different functional groups (Scheme 1). By taking advantage of the large number of O/F sites on the surface of MXene, which can easily form hydrogen bonds with amino-modified clusters, the PHE performance of amino-modified MCSCs (and ligand-free clusters) combined with MXene nanosheets was investigated.

## Experimental

### Materials

All chemicals were purchased from commercial sources and used without further purification. Deionized water was used in all synthetic procedures.



**Scheme 1** Schematics of the surface functionalization of discrete MCSCs: (a and b) using the modified thiophenol and (c and d) using modified diphenyl disulfide.

### Synthetic procedures

**Synthesis of ISC-24,  $10(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{17}]^{10-}\cdot 4\text{H}_2\text{O}$ .** The synthetic procedure for ISC-24 followed our previous work,<sup>32</sup> except that the manganese source was replaced by a zinc source. A mixture of zinc acetate tetrahydrate ( $\text{Zn}(\text{OAc})_2\cdot 4\text{H}_2\text{O}$ , 120 mg), tin dichloride ( $\text{SnCl}_2$ , 60 mg), sulfur powder (S, 120 mg), 3,5-dimethylpiperidine (3,5-DMP, 3.0 mL), and deionized water ( $\text{H}_2\text{O}$ , 1.0 mL) was added to a 25 mL Teflon-lined stainless steel autoclave and stirred for 30 min, then was heated to 125 °C for 7 days. After cooling down to room temperature, transparent colorless block crystals were obtained with a yield of 19.4% based on zinc acetate dihydrate.

**Synthesis of ISC-25,  $7(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{14}(\text{SPh})_3]^{7-}\cdot \text{H}_2\text{O}$ .** The synthetic procedure for ISC-25 followed our previous work,<sup>32</sup> except that the manganese source was replaced by a zinc source. Zinc acetate tetrahydrate ( $\text{Zn}(\text{OAc})_2\cdot 4\text{H}_2\text{O}$ , 120 mg), tin dichloride ( $\text{SnCl}_2$ , 60 mg), sulfur powder (S, 120 mg), diphenyl disulfide ( $\text{Ph}_2\text{S}_2$ , 160 mg), 3,5-dimethylpiperidine (3,5-DMP, 3.0 mL), and deionized water ( $\text{H}_2\text{O}$ , 1.0 mL) were mixed in a 25 mL Teflon-lined stainless steel autoclave and stirred for 30 min, then heated to 125 °C for 14 days. After cooling down to room temperature, transparent colorless rhombohedral crystals were obtained with a yield of 16.8% based on zinc acetate dihydrate.

**Synthesis of ISC-25-4'-NH<sub>2</sub>,  $7(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{14}(\text{SPhNH}_2)_3]^{7-}\cdot \text{H}_2\text{O}$ .** A mixture of zinc acetate tetrahydrate ( $\text{Zn}(\text{OAc})_2\cdot 4\text{H}_2\text{O}$ , 120 mg), tin dichloride ( $\text{SnCl}_2$ , 60 mg), sulfur powder (S, 120 mg), di-(4,4'-amino)-diphenyl disulfide ( $\text{S}_2(\text{NH}_2\text{Ph})_2$ , 160 mg), and deionized water ( $\text{H}_2\text{O}$ , 1.0 mL) was added to a 25 mL Teflon-lined stainless steel autoclave and stirred for 30 min, then heated to 125 °C for 14 days. After cooling down to room temperature, oblique octahedral or hexagonal transparent crystals were obtained with a yield of 12.6% based on zinc acetate dihydrate.

**Synthesis of other ISC-25 and ISC-26 series samples with different decorated functional groups.** Similar to the synthetic processes described above, the ligand of  $\text{Ph}_2\text{S}_2$  was replaced by substituted  $\text{Ph}_2\text{S}_2$  (e.g.  $(\text{ClPh})_2\text{S}_2$ , the categories are summarized in Table S2†) in the same molar ratio with the other components unchanged.

**Preparation of  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets.**  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets (here T represents the oxygen terminus generated by HF etching) were synthesized according to the reported method (Schemes S2 and S3†).<sup>41</sup>  $\text{Ti}_3\text{AlC}_2$  powders (>98 wt% purity; particle size <75 μm, i.e., 200 mesh) were immersed in HF (10 wt%) solution at 300 K by stirring for 24 h. Then the resulting suspension was washed several times with deionized water and centrifuged to remove the remaining impurities and HF until the pH value of the solution reached 5–6. The wet sediment was washed twice with ethanol and transferred to a wide-necked jar to air-dry for 3–4 days. The finally obtained sample was named  $\text{Ti}_3\text{C}_2\text{T}_x$ . The multilayered  $\text{Ti}_3\text{C}_2\text{T}_x$  was then stirred with dimethyl sulfoxide (DMSO) at 310 K for 24 h. The colloidal suspension was centrifuged to obtain the intercalated

powders. The powders were then washed several times with deionized water to remove the residual DMSO. Next, the residue was dispersed in deionized water in a weight ratio of  $\text{Ti}_3\text{C}_2\text{T}_x$  to water of 1:100. Afterward, ultrasonication treatment was carried out for 4 h to exfoliate the  $\text{Ti}_3\text{C}_2\text{T}_x$  powder into 2-D sheets under argon. The suspension was centrifuged at 3500 rpm for 10 min to remove the unexfoliated MXenes. Finally, the  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets were obtained by filtration and dried in a vacuum for further experiments.

**Preparation of P1-CNPs (CNPs: cluster-derived nanoparticles).** P1-CNPs were prepared in the mixed solution of formamide and water (1:1 in volume). The corresponding crystals of 10 mg of ISC-24, ISC-25, and ISC-25-4'-NH<sub>2</sub> were ground into powder and dispersed in 10 mL of 1:1 formamide/water. After 30 min of ultrasonication treatment, the formamide/water solution of P1-CNPs with a concentration of 1 mg mL<sup>-1</sup> was prepared by mechanical stirring for 4 h.

**Preparation of CMn samples.** CMn samples were prepared by mixing 1 mg mL<sup>-1</sup> of the corresponding P1-CNP formamide aqueous solution with the one containing the ultra-thin  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets according to the mass percentage (CM0 is P1-CNPs without  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets).

## Results and discussion

### Synthetic procedure and crystal structure description

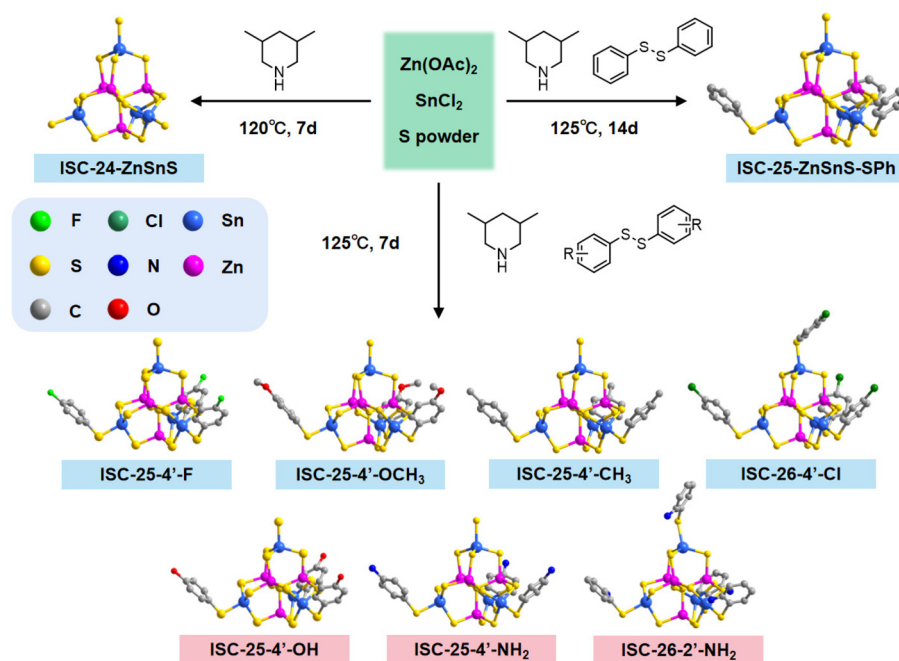
The title compounds constructed from supertetrahedral P1-ZnSnS clusters were obtained by a facile one-step solvothermal reaction (see the Experimental section and the ESI† for

details). To directly realize the functionalization of the partially protected P1-ZnSnS, various diphenyl disulfides decorated with different functional groups were selected, and the synthetic routine and molecular structure of the relevant compounds are shown in Fig. 1.

The obtained clusters mentioned in this work are all composed of the  $[\text{Zn}_4\text{Sn}_4\text{S}_{17}]^{10-}$  core, so the functionalized samples are represented in a simplified form (e.g., ISC-25-ZnSnS-SPH is simplified to ISC-25, and the functionalized ISC-25 are represented as the site of the substituent on their benzene ligand, e.g., ISC-25-4'-NH<sub>2</sub>). The derivatives of ISC-26 are also showcased in a similar pattern. The structural difference between ISC-25 and ISC-26 is the number of protecting ligands (three-corner protected in ISC-25 and four-corner protected in ISC-26, Fig. 1).

Interestingly, ISC-26-2'-NH<sub>2</sub> was obtained using 2,2'-dinitrodiphenyl disulfide, instead of 2,2'-diaminodiphenyl disulfide. This may be attributed to excess sulfur powder in the reaction system, which reduces the nitro group to the amino group. A similar situation also occurs in ISC-25-4'-NH<sub>2</sub>, while ISC-25-4'-NH<sub>2</sub> can also be obtained by directly using 4,4'-diaminodiphenyl disulfide. This process is also shown in detail in Table S2.†

A series of functionalized crystalline ISC-25 and ISC-26 samples was obtained in the transparent rhombohedral form, but ISC-26-4'-Cl appeared in the form of a transparent decagonal octahedron. Single crystal X-ray structural analysis revealed that the ISC-24 sample crystallized in  $P\bar{4}3n$  (No. 218), all ISC-25 series samples and ISC-26-2'-NH<sub>2</sub> crystallized in the  $R\bar{3}$  space group (No. 148), and the ISC-26-4'-Cl sample crystallized in the  $C2/c$  space group (No. 15). These functionalized clusters are



**Fig. 1** Synthetic processes and molecular structures of ISC-24, ISC-25, and ISC-26 with different functional groups. The products with functional groups have been distinguished by different colored backgrounds in comment boxes (light blue: inert functional groups, e.g., -CH<sub>3</sub>, -OCH<sub>3</sub>, -Cl, -F; pink: active functional groups: e.g., -OH, -NH<sub>2</sub>).

basically isomorphic to the structures previously reported.<sup>32</sup> Combined with the results of C, H, and N elemental analysis of **ISC-24**, **ISC-25**, and **ISC-25-4'-NH<sub>2</sub>** (Table S3<sup>†</sup>) and their SCXRD data (Table S1<sup>†</sup>), the formulas of the above three compounds were determined as follows: **ISC-24**:  $10(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{17}]^{10-}\cdot\text{H}_2\text{O}$ ; **ISC-25**:  $7(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{14}(\text{SPh})_3]^{7-}\cdot\text{H}_2\text{O}$  and **ISC-25-4'-NH<sub>2</sub>**:  $7(3,5\text{-DMPH}^+)[\text{Zn}_4\text{Sn}_4\text{S}_{14}(\text{SPhNH}_2)_3]^{7-}\cdot\text{H}_2\text{O}$ . Compared to the P1 cluster reported previously, the cluster reported here retained the pristine atomic arrangement of the divalent and tetravalent metal ions, *i.e.*, each P1-ZnSnS contained one ( $\text{M}_4^{\text{II}}\text{S}$ )<sup>6+</sup> and four ( $\text{M}^{\text{IV}}\text{S}_4$ )<sup>4-</sup> units to meet Pauling's electrostatic valence rule.

### Morphology of Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets and dispersed P1-CNPs

Layered Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets were prepared by exfoliating bulk Ti<sub>3</sub>AlC<sub>2</sub> powders in HF aqueous solution. The corresponding structure information for Ti<sub>3</sub>AlC<sub>2</sub> and the preparation process are shown in Schemes S3 and S4.<sup>†</sup> After immersing them in HF solution for 24 h, the Ti<sub>3</sub>AlC<sub>2</sub> powders (200 mesh) were successfully expanded and converted to the layered Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> structure (Fig. 2a). The emergence of layers was caused by the fact that the Al layers in Ti<sub>3</sub>AlC<sub>2</sub> nanoparticles were etched away, and the interlayer space was increased by released H<sub>2</sub>. After intercalation by dimethyl sulfoxide (DMSO), the layered structures were further delaminated into 2D Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets by ultrasonication (Fig. 2b). The AFM image in Fig. 2c shows that the thickness of Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets is about 4 nm. In addition, PXRD characterization was performed on the Ti<sub>3</sub>AlC<sub>2</sub> MAX phase and Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets to distinguish the outcome of Al etching. As shown in Fig. 2d, there are a number of characteristic peaks at  $2\theta = 9.95^\circ, 19.50^\circ, 34.38^\circ, 37.49^\circ, 39.21^\circ, 42.13^\circ, 48.72^\circ,$  and  $56.66^\circ$ , which correspond to the (002), (004), (101), (103), (104), (105), (107), and (109) planes of the Ti<sub>3</sub>AlC<sub>2</sub> MAX phase. This result is in agreement with the one in the published literature.<sup>42,43</sup> The degree of the

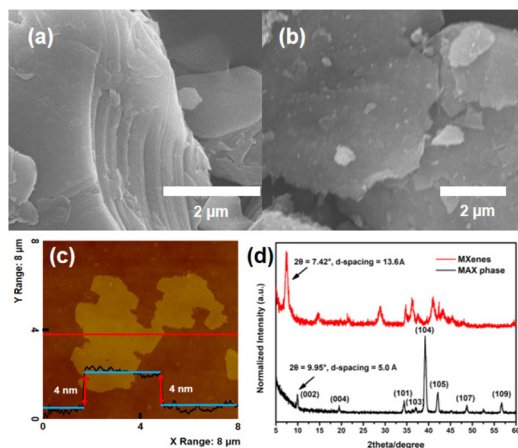
(002) plane in PXRD is generally used to calculate the inter-layer spacing of the layered material by applying Bragg's law, and the *d*-spacing of the Ti<sub>3</sub>AlC<sub>2</sub> MAX phase is calculated to be approximately 5.0 Å. Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> MXene nanosheets show a (002) peak at  $2\theta = 7.42^\circ$ , referring to a *d*-spacing of 13.6 Å (Fig. 2d). It is clearly demonstrated that the Al layer in the Ti<sub>3</sub>AlC<sub>2</sub> MAX phase has been well etched, which allows water molecules to further exfoliate the Al layer for preparing single-layered Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> MXene nanosheets.<sup>45</sup>

The nanoparticles generated from the cluster dispersion are denoted as **P1-CNPs** (CNPs: cluster-derived nanoparticles) in this work. Three examples of clusters were selected for comparison to test their dispersibility: **ISC-24** without ligand protection, **ISC-25** with a partially protected ligand, and **ISC-25-4'-NH<sub>2</sub>** with a partially protected ligand modified with amino (-NH<sub>2</sub>) groups. The corresponding **P1-CNP** solutions were prepared by adding the ground crystals of the above three compounds to the formamide/water solution (1:1), followed by stirring for 4 h and sonicating for 1 h. The particle sizes of **P1-CNPs** formed from **ISC-24** were found to be 25–55 nm (Fig. 3a, mean size ~40 nm), which far exceeded the size of individual clusters (~1.6 nm). This suggests that the dispersed particles existed in the aggregated form of multi-clusters. In contrast, the mean particle sizes of **P1-CNPs** formed by ligand-partially protected **ISC-25** and amino-modified **ISC-25-4'-NH<sub>2</sub>** were found to be 4–7 nm and 1–4 nm, respectively (Fig. 3b and c). The latter two were significantly more accessible to solution. This can be explained by the low negative charge of the clusters and the improved solubility.

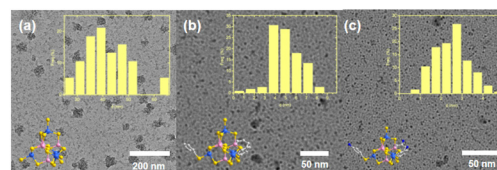
In addition, no effective lattice striations were observed in the HRTEM images of the **P1-CNPs** formed by the dispersed clusters. This suggests that **P1-CNPs** have adopted an amorphous state, which is probably due to the small size of the individual clusters (~1.6 nm).

### Fabrication and morphology of P1-CNPs/Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanocomposites (CM<sub>n</sub>)

The nanocomposites consisting of **P1-CNPs** and exfoliated Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> are denoted as **CM<sub>n</sub>** (*n* indicates the weight percentage of the added Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>, *n* = 0 indicates that no Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> was added). The fabrication process for the composite is shown in Scheme S4.<sup>†</sup> The results of PXRD characterization of **CM1** (composed of **ISC-25-4'-NH<sub>2</sub>** derived **CM0** and Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>) are shown in Fig. S11.<sup>†</sup>



**Fig. 2** SEM images of layered (a) and exfoliated Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets (b) obtained by HF treatment of Ti<sub>3</sub>AlC<sub>2</sub> powder; (c) AFM images of dispersed single-layered Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets; and (d) PXRD results of Ti<sub>3</sub>AlC<sub>2</sub> powder and exfoliated Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets.



**Fig. 3** TEM images of **P1-CNPs** formed by (a) **ISC-24**; (b) **ISC-25**, and (c) **ISC-25-4'-NH<sub>2</sub>**, and the insets show the size distribution and the molecular structures of the corresponding clusters.

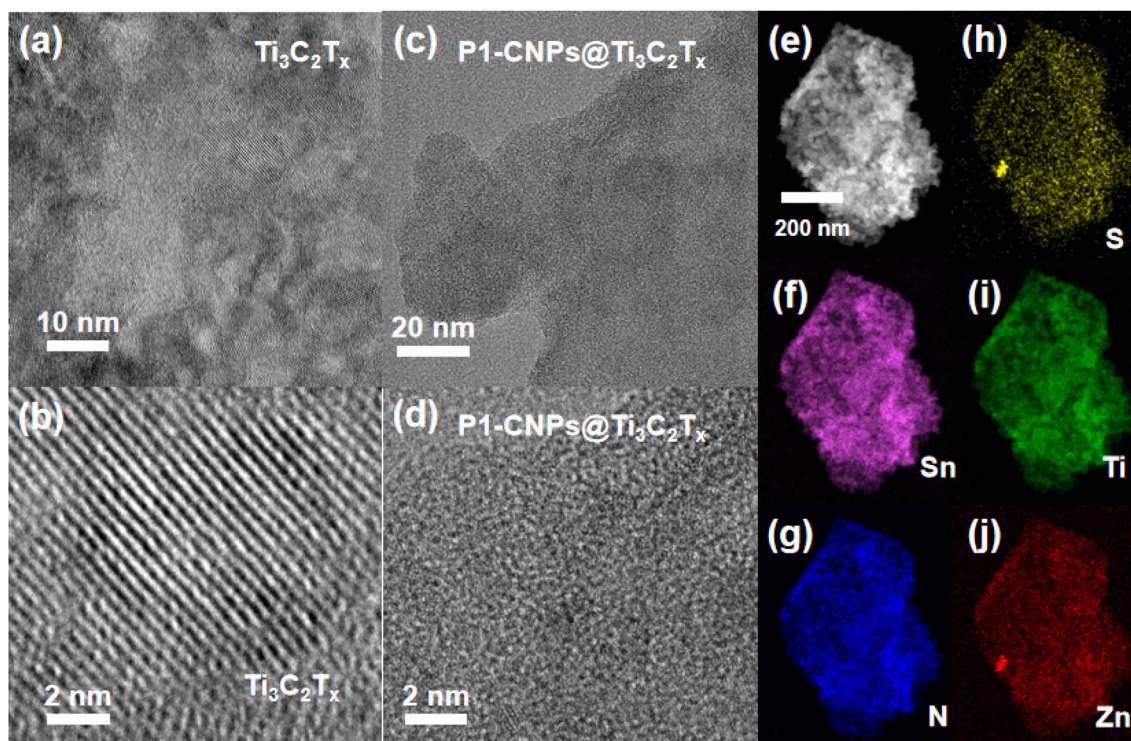
Due to the low loading amount and low crystallinity, only the diffraction peaks of MXene nanosheets were observed. In order to characterize the morphology and structure of **CMn**, we performed HRTEM tests on the corresponding samples. Firstly, the morphology of  $\text{Ti}_3\text{C}_2\text{T}_x$  was characterized, and clear lattice stripes were observed as shown in Fig. 4a and b. However, for **CM1**, the lattice stripes were not detected in the HRTEM images (Fig. 4c and d). This was also confirmed by the selected area electron diffraction (SAED) characterization. As shown in Fig. S3a,† single-crystal diffraction patterns of  $\text{Ti}_3\text{C}_2\text{T}_x$  were observed when selected electron diffraction was performed on the regions shown in Fig. 4a and b, while the **CM1** composites exhibited a polycrystalline form (Fig. S3b†), consistent with the states shown in Fig. 4c and d. In addition, we also performed HAADF-STEM characterization on **CM1** prepared with **ISC-25-4'-NH<sub>2</sub>** as the precursor, and performed EDS mapping tests (Fig. 4e–j). The constituents of Zn, Sn, S, and N in the clusters were uniformly distributed on the  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets, demonstrating the successful preparation of the composites.

### PHE performance of **CMn**

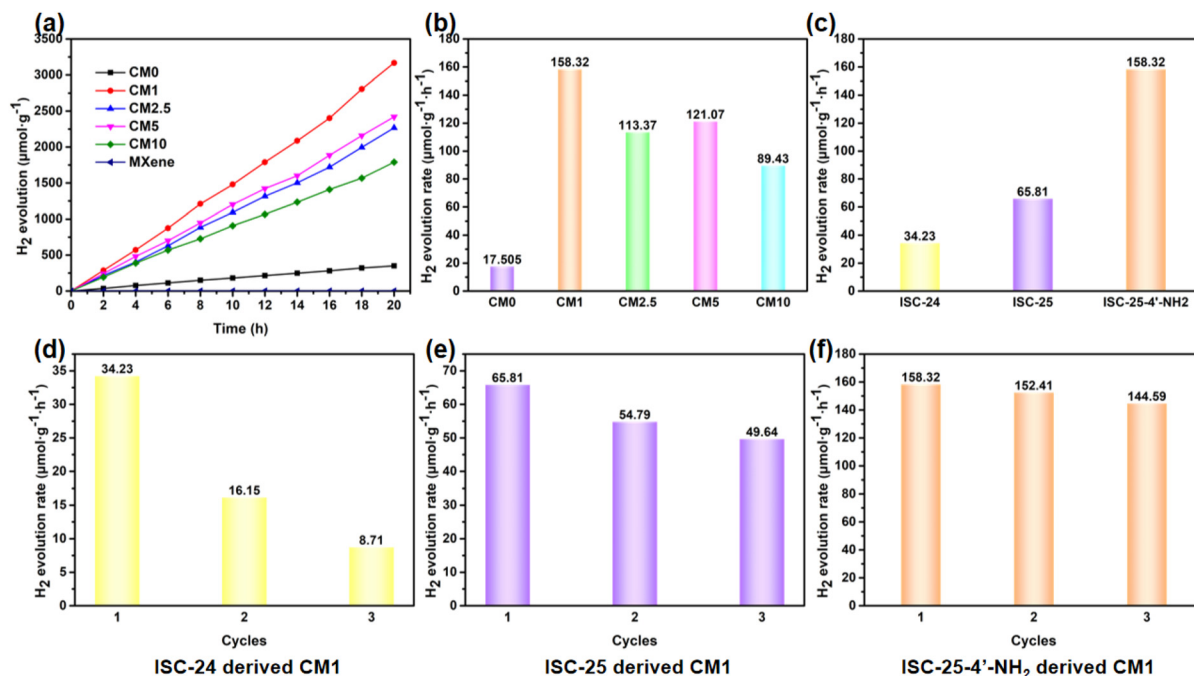
The photocatalytic performance of the above composites was tested by conducting PHE reactions under full-spectrum irradiation using the  $\text{Na}_2\text{S}/\text{Na}_2\text{SO}_3$  reagent as the sacrificial reagent. Firstly, **ISC-25-4'-NH<sub>2</sub>** derived **CM0** was selected for the PHE reaction, and the  $\text{H}_2$  production rate was calculated to be  $17.51 \mu\text{mol g}^{-1} \text{h}^{-1}$  according to the data shown in Fig. 5a.

This poor PHE performance could be attributed to the low separation efficiency of the photogenerated electron–hole pairs of **CM0**. Enhanced photocatalytic activities were obtained for the **ISC-25-4'-NH<sub>2</sub>** derived samples containing a certain percentage of  $\text{Ti}_3\text{C}_2\text{T}_x$  (1%, 2.5%, 5%, and 10%, *i.e.*, **CM1**, **CM2.5**, **CM5**, and **CM10**, respectively). The PHE rate from **CM1** was calculated to be  $158.32 \mu\text{mol g}^{-1} \text{h}^{-1}$  (Fig. 5a and b). This result indicates that the  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets play an important role in separating the photogenerated electron–hole pairs for **P1-CNPs**. The PHE activity of **CM2.5**, **CM5** and **CM10** decreased to 113.37, 121.07, and  $89.43 \mu\text{mol g}^{-1} \text{h}^{-1}$ , respectively. These results were caused by the aggregation of  $\text{Ti}_3\text{C}_2\text{T}_x$  and **P1-CNPs**, which was verified by its TEM images (Fig. S4†). Under such situation, the  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets partially mask the active sites of **P1-CNPs**, thereby leading to a reduced photocatalytic efficiency.

The photocatalytic stability of **ISC-24**, **ISC-25**, and **ISC-25-4'-NH<sub>2</sub>** derived **CM1** composites was also investigated (Fig. 5d–f). The activity of the **ISC-24** derived **CM1** decreased by 52.9% for the second round of photocatalysis, and further decreased by 25.4% to  $8.71 \mu\text{mol g}^{-1} \text{h}^{-1}$  in the third round of testing (Fig. 5d). This can be attributed to the poor stability of the **ISC-24** derived sample. For the **CM1** composites prepared with **ISC-25** and **ISC-25-4'-NH<sub>2</sub>** as precursors, the photocatalytic activity of the second and third rounds decreased by 83.2%, 75.4% and 96.3%, 91.3%, respectively, compared to the first round. The results show that the composites containing **P1-CNPs** with the amino-modified clusters have good stability.



**Fig. 4** HRTEM images of MXene nanosheets (a and b) and **ISC-25-4'-NH<sub>2</sub>** derived **CM1** (c and d). HAADF-STEM image of **P1-CNPs/Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>** nanocomposites (e), and EDS elemental mapping images of (f) Sn, (g) N, (h) S, (i) Ti, and (j) Zn, respectively.



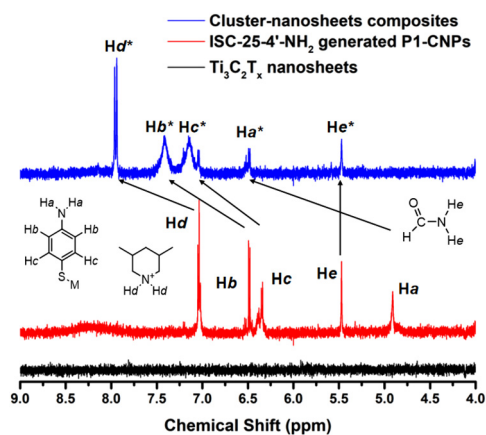
**Fig. 5** Investigation of photocatalytic properties of  $CM_n$ . (a and b) Comparison of PHE performance of  $CM_0$ ,  $CM_1$ ,  $CM_{2.5}$ ,  $CM_5$ , and  $CM_{10}$  nano-composites under irradiation using  $ISC-25-4'-NH_2$  derived  $CM_n$ . (c) Comparison of PHE performance of  $CM_1$  composite materials using  $ISC-24$ ,  $ISC-25$ , and  $ISC-25-4'-NH_2$  derived  $CM_n$ . (d–f) Test results of multiple cycles on  $CM_1$  composite materials using  $ISC-24$ ,  $ISC-25$ , and  $ISC-25-4'-NH_2$  derived  $CM_n$ .

### <sup>1</sup>H-NMR test for P1-CNPs and $CM_n$

Considering that the  $CM_1$  derived from  $ISC-25-4'-NH_2$  has good PHE performance and high stability, we assume that the amino groups in the modified P1 cluster play an important role. Therefore, the <sup>1</sup>H-NMR test was conducted to verify the above conjecture (Fig. 6). The formation of hydrogen bonds between the amino groups and the oxygen terminus of  $Ti_3C_2T_x$  nanosheets caused the chemical shift values of hydrogen atoms to be shifted towards the lower field due to this effect. Similar situations have been reported in the case of an S...H–N hydrogen bond.<sup>44</sup> In addition, protonated 3,5-dimethyl-piperidine, which acts as counterion during cluster dispersion, also interacts electrostatically with the oxygen terminus on the  $Ti_3C_2T_x$  nanosheets to form hydrogen bonds, as evidenced by the result of the  $H_d$  signal moving from 7.1 ppm to 8.0 ppm in Fig. 6. This interaction also provides a theoretical basis for the enhanced photocatalytic performance of  $ISC-24$  and  $ISC-25$  based  $CM_1$  (Fig. 5c).

### Optical and electrochemical properties of $CM_n$

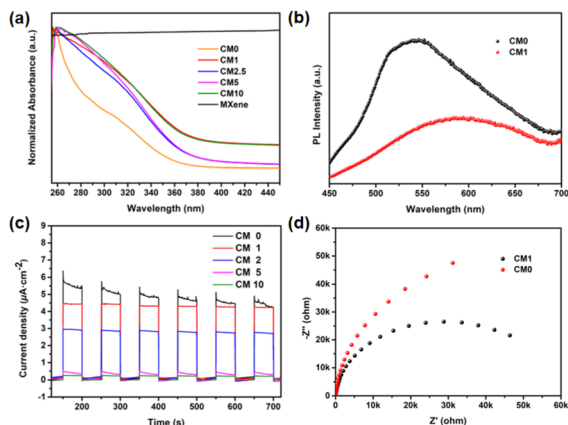
To further understand the enhanced photocatalytic performance of  $CM_n$  composites compared to P1-CNPs, a series of optical and electrochemical properties were characterized. Firstly, UV-vis absorption spectra were obtained to characterize the light absorption capability of the samples. As shown in Fig. 7a, the light absorption of  $CM_1$  was significantly increased compared to  $CM_0$  in the whole range of 260–460 nm due to the dispersed  $Ti_3C_2T_x$  nanosheets, and similar phenomena



**Fig. 6** <sup>1</sup>H-NMR spectra of  $Ti_3C_2T_x$  nanosheets, P1-CNPs formed by  $ISC-25-4'-NH_2$ , and their composite materials with a chemical shift value of 4.0–9.0 ppm (see Fig. S5–S7† for original data).

were observed for  $CM_{2.5}$ ,  $CM_5$  and  $CM_{10}$ . Subsequently, steady-state photoluminescence (PL) spectra were recorded for the  $ISC-25-4'-NH_2$  derived  $CM_0$  and  $CM_1$ , as shown in Fig. 7b. The emission intensity of  $CM_1$  was significantly weakened compared to that of  $CM_0$ , suggesting that the electron–hole recombination in  $CM_1$  composites is well-suppressed because of the effective charge transfer between P1-CNPs and  $Ti_3C_2T_x$ .

In addition, the photoelectric response and electrochemical impedance spectra (EIS) were recorded to detect the separation of photogenerated electron–hole pairs and the interfacial



**Fig. 7** (a) UV-vis absorption spectra of MXene nanosheets and **CM<sub>n</sub>**; (b) PL spectra of ISC-25-4'-NH<sub>2</sub> derived **P1-CNPs** and **CM1** (Ex = 380 nm); (c and d) photoelectric response curves (c) and electrochemical impedance spectra (d) of ISC-25-4'-NH<sub>2</sub> derived **CM<sub>n</sub>**.

charge transfer. As shown in Fig. 7c, **CM0** showed the highest transient photocurrent densities, but were less stable compared to the prepared **CM<sub>n</sub>** decorated electrodes. According to this situation, **CM0** should have a higher PHE efficiency. However, considering the feature of sharp reduction of the photocurrent (Fig. 7c), we also compared the EIS curves of **CM0** and **CM1**, as shown in Fig. 7d. The fitted EIS spectra of **CM1** showed a smaller semicircle and a lower interfacial resistance, revealing the more efficient electron-hole pair separation and transfer efficiency of the **CM1** heterostructure. We therefore attribute the low photocatalytic efficiency of **CM0** to poor stability and weak conductivity. **CM1** exhibited the highest photocurrent density in the **CM<sub>n</sub>** series, corresponding to its highest PHE performance and stability, while **CM2.5**, **CM5**, and **CM10** exhibited lower photocurrent densities due to the formation of a sandwich-like structure, as shown in Fig. S4,† where the multilayered Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets may cover the active sites of **P1-CNPs**.

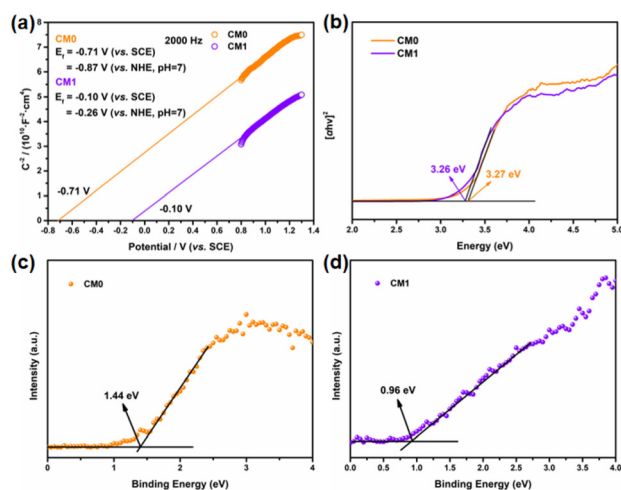
X-ray photoelectron spectroscopy (XPS) tests were also carried out on ISC-25-4'-NH<sub>2</sub> derived **P1-CNPs**, Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets, and **CM1** composites to further verify the formation of composites between the two substances, and the results are shown in Fig. S8–S10.† The Zn, Sn and S signals of ISC-25-4'-NH<sub>2</sub> are consistent with those of **P1-CNPs**, and the high-resolution XPS spectra of Ti 2p showed four deconvoluted peaks (Fig. S10†) with binding energies of 464.8 eV and 459.0 eV corresponding to Ti–O 2p<sup>1/2</sup> and Ti–O 2p<sup>3/2</sup> and 461.2 eV and 455.3 eV corresponding to Ti–C 2p<sup>1/2</sup> and Ti–C 2p<sup>3/2</sup>, respectively. This result is in agreement with the HRTEM ones described above.

### Photocatalytic mechanism and the band structure

To further understand the enhanced photocatalytic performance of the **CM<sub>n</sub>** series composites compared to **CM0**, we conducted a series of characterization studies. The Mott–Schottky analysis was first performed to evaluate the flat-band potential

and Fermi levels of **CM0** and **CM1** prepared from ISC-25-4'-NH<sub>2</sub>.<sup>45–47</sup> As shown in Fig. 8a, the flat-band potentials of **CM0** and **CM1** were calculated to be  $-0.87$  V and  $-0.26$  V (*versus* NHE, pH = 7), which were directly treated as Fermi levels since the surface Fermi levels are very close to the flat-band potential.<sup>48,49</sup> To further determine the valence band (VB) and  $E_g$ <sup>50</sup> of **CM0** and **CM1**, XPS valence band spectroscopy (VB-XPS) and solid-state UV-vis diffuse reflectance spectroscopy were also carried out. From the VB-XPS results (Fig. 8c and d), the VB positions of **CM0** and **CM1** were determined to be 1.44 eV and 0.96 eV relative to the Fermi level,<sup>51</sup> which indicates that the VB positions of **CM0** and **CM1** are 0.57 V and 0.70 V (*versus* NHE, pH = 7), respectively. The band gaps of **CM0** and **CM1** were found to be 3.27 eV and 3.26 eV, respectively, according to Fig. 8b. The corresponding conduction bands (CB) edges of **CM0** and **CM1** were calculated to be  $-2.70$  V and  $-2.56$  V (*versus* NHE, pH = 7), respectively, by combining the  $CB = E_g - VB$  equation.<sup>52</sup>

Based on the above analysis, we proposed a photocatalytic mechanism for illustrating the higher PHE performance of the **CM<sub>n</sub>** series composites, as shown in Fig. 9. After **P1-CNPs** combined with the Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets to form a binary complex, the Fermi level of **CM1** shifted upwards relative to **CM0**. The Mott–Schottky curve in Fig. 8a shows that **P1-CNPs** are n-type semiconductors and their flat-band potential is much more negative than that of the Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets, so the close contact between the two components in **CM1** results in the transfer of electrons from the **P1-CNPs** to the Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> nanosheets. This is evidenced by the flat-band potential correction ( $-0.10$  V *vs.* NHE) in **CM1** compared to **CM0**, suggesting that the Fermi level of **P1-CNPs** in **CM1** decreases upon their binding to Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>. A similar situation was reported in the studies of Jakob<sup>53</sup> and Ran.<sup>46</sup> Furthermore, the immobilized positive charges in the **P1-CNPs** are near the **P1-CNPs**/



**Fig. 8** (a) Mott–Schottky plots of ISC-25-4'-NH<sub>2</sub> derived **P1-CNPs** (*i.e.* **CM0**) and **CM1** composites; (b) solid-state UV-vis diffuse reflectance spectra of **CM0** and **CM1**; (c and d) valence band XPS spectra of **CM0** (c) and **CM1** (d).

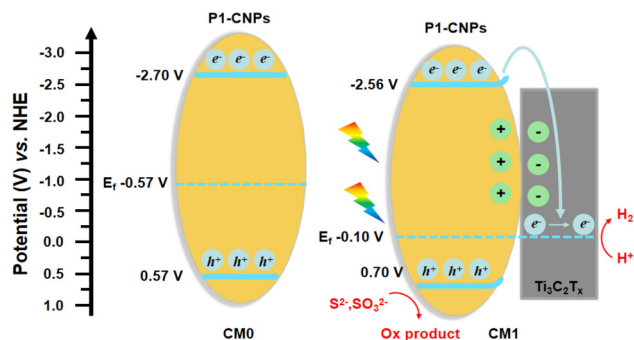


Fig. 9 Proposed schematic diagram of the photocatalytic mechanism for CM0 and CM1 under light irradiation.

$\text{Ti}_3\text{C}_2\text{T}_x$  interface, where a space charge layer (or internal electric field) is formed and the CB and VB in the **P1-CNPs** are bent “upwards”, thus forming a Schottky junction between them. Under full-spectrum solar irradiation, electrons are excited from the VB of the **P1-CNPs** to the CB, and due to the reduced thickness of the space charge layer in the **P1-CNPs** particles, photogenerated electrons in the CB can still migrate through the “upward” bent CB to the Fermi level of  $\text{Ti}_3\text{C}_2\text{T}_x$ , leaving photogenerated holes in the VB of **P1-CNPs**. The previously mentioned formation of Schottky junctions can act as electron traps for trapping photogenerated electrons without blocking the transfer of electrons from **P1-CNPs** to  $\text{Ti}_3\text{C}_2\text{T}_x$ .<sup>54–56</sup> After the transfer of electrons to the  $\text{Ti}_3\text{C}_2\text{T}_x$  nanosheets through the above pathway, the excellent electrical conductivity of  $\text{Ti}_3\text{C}_2\text{T}_x$  allows electrons to quickly shuttle to the surface. As a result, based on the excellent HER capabilities of  $\text{Ti}_3\text{C}_2\text{T}_x$ , the **CMn** series composites achieved more efficient PHE performance compared to pure **P1-CNPs**.

## Conclusions

In summary, we proposed a strategy for disulfide bond cleavage to prepare discrete MCSCs with multiple active/inactive modified functional groups. Furthermore, three models with different structures were tested for their dispersibility, binding strength with MXene nanosheets, and photocatalytic hydrogen generation performance. The results showed that the amino-modified clusters had better performance than ligand-free type ones. This work provides insights for the subsequent design and synthesis of other types of discrete MCSCs with active functional groups, and is expected to have long-term implications for the functional expansion of metal chalcogenide cluster-based nanomaterials.

## Author contributions

Jin Wu: Investigation, writing – original draft, and writing – review & editing. Qiang Fu and Zixin Wu: TEM image testing. Peipei Sun: data curation. Xing Zhu and Ying Wang: HRTEM

and SAED testing. Ning Chen: funding acquisition. Tao Wu: Writing – review & editing, supervision, project administration, and funding acquisition.

## Conflicts of interest

The authors declare no competing financial interests.

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