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# High performance photocatalytic and thermoelectric two-dimensional asymmetrically ordered Janus-like MXene alloys†

Zicong Marvin Wong, Dangi Deng, Wen Shi, Gang Wu, D Teck Leong Tan \*\* and Shuo-Wang Yang \*\*

MXenes are two-dimensional transition metal carbides, nitrides, and/or carbonitrides which display extensive versatility in their elemental composition for tuning and optimization of their functional properties. In this work, by using a multiscale approach of density functional theory calculations, cluster expansion method, and Monte Carlo simulations, we systematically investigate the  $Ti_{2(1-x)}Mo_{2x}CO_2$  alloy system of two-dimensional  $Ti_2CO_2$  and  $Mo_2CO_2$  MXenes with 0 < x < 1. From the alloy formation energies, we identify the stable ground-state and report the possible existence of an asymmetrically ordered Janus-like MXene alloy TiMoCO2. This alloy is theoretically validated to be stable thermodynamically, mechanically, and thermally, with a high order-disorder phase transition temperature of  $\sim 640$  K. Importantly, TiMoCO<sub>2</sub> is a semiconductor with a band gap of 0.60 eV and its valence band maximum and conduction band minimum are located on opposite sides of the MXene monolayer. As such, the asymmetric configuration generates an intrinsic dipole moment, which realigns the conduction and valence band edges, permitting photocatalytic redox water-splitting to favorably occur with a higher solar absorption efficiency in the visible-infrared region due to the smaller band gap. From the transport calculations, we propose the TiMoCO2 MXene alloy to exhibit high room temperature n- and p-type power factors of 49.8 μW cm<sup>-1</sup> K<sup>-2</sup> and 15 μW cm<sup>-1</sup> K<sup>-2</sup>, respectively, outperforming other related experimentally synthesized Ti-Mo based MXene alloys and even similar to that of p-doped SnSe which is one of the best performing thermoelectric materials. All these indicate the stable asymmetrically ordered Janus-like TiMoCO2 MXene alloy to be a promising high performance water-splitting photocatalyst and thermoelectric material, paving the way for more stable Janus-like MXenes with interesting and outstanding functional properties to be discovered.

# 1. Introduction

MXenes are a large family of two-dimensional (2D) layered transition metal carbides, nitrides, and/or carbonitrides discovered in

Institute of High Performance Computing, Agency for Science, Technology and Research, 1 Fusionopolis Way, #16-16 Connexis, 138632, Singapore. E-mail: yangsw@ihpc.a-star.edu.sg, tantl@ihpc.a-star.edu.sg

† Electronic supplementary information (ESI) available: Structures of the 3 different fully O-terminated MXenes; electrostatic potential profile for the Janus-like ground-state TiMoCO2 MXene alloy; projected electronic band structures of the Janus-like ground-state TiMoCO2, and its analogous conjectural ZrMoCO2, HfMoCO2, TiCrCO2, and TiWCO2 MXene alloys; table of structures of various TiMoCO2 MXene alloys with different O-sites and their constituent MXenes Ti<sub>2</sub>CO<sub>2</sub> and Mo<sub>2</sub>CO<sub>2</sub>, their relative energies with respect to the constituents obtained via density functional theory, and the Bader charges of the respective elements; table of calculated DFT and alloy formation energies of bare TiMoC MXene with respect to its constituent Ti2C and Mo2C MXenes; and table of band gap and the respective contributions and directional effective hole and electron masses of the valence band maximum and conduction band minimum for Janus-like ground-state TiMoCO2, and its analogous conjectural ZrMoCO2, HfMoCO2, TiCrCO2, and TiWCO2 MXene alloys. See DOI: 10.1039/d0ma00391c

2011, with the structural formula  $M_{n+1}X_n$  (where M is transition metal, and X is C and/or N, with n = 1, 2, 3, or 4). They are typically derived from their MAX phase precursors by the selective chemical etching of the A-group elements such as Al or Ge,<sup>4</sup> although other synthesis approaches are feasible as well.5,6 Although post-chemical etching results in MXenes to be mostly terminated with a mixture of functional groups, e.g., O, OH, and F, a recent study has shown that exclusively O-terminated MXenes, i.e.,  $M_{n+1}X_nO_2$ , can be attainable via post-treatment under vacuum and subsequent exposure to O<sub>2</sub> gas at slightly elevated temperatures.<sup>7</sup> Owing to the compositional tunability, elemental flexibility, 2D crystallinity, excellent electrical conductivity, and hydrophilicity, MXenes have been under the spotlight for a wide spectrum of research spanning from energy,3 environment,8 and even biomedical applications. This rich chemistry exhibits the versatility of MXenes in the design of materials.

As the world becomes increasingly conscious towards climate change and sustainability, alternative green energy sources such as hydrogen production from photocatalytic water-splitting have become heavily sought for as a clean fuel. As such, MXenes have

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been garnering intense research interest in photocatalytic watersplitting. 10-12 Typically, a practical water-splitting photocatalyst has to fulfill 3 main requirements: (1) band gap > 1.23 eV such that the band edges straddle water redox potentials; (2) high solar photon absorption; and (3) a low electron-hole recombination rate. 13,14 Moreover, 2D materials like MXenes display high specific reaction surface and the low dimensionality minimizes the distances that photogenerated electrons and holes have to migrate to the reaction interface, reducing the possibility of electron-hole recombination and enhancing the photocatalytic performance. Currently, only a handful of MXenes are theoretically identified to exhibit semiconducting behaviors, e.g., Ti<sub>2</sub>CO<sub>2</sub>,  $Zr_2CO_2$ ,  $Hf_2CO_2$ , and  $Sc_2CT_2$  (where T = O, F, OH). Unfortunately, these MXenes possess either insufficient band gaps, misaligned band edges, or weak solar absorption, which compromise their photocatalytic performance.<sup>16</sup> Our previous work has suggested stoichiometric engineering of the disordered alloy of Ti<sub>2</sub>CO<sub>2</sub> and  $Zr_2CO_2$ , i.e.,  $Ti_{2(1-x)}Zr_{2x}CO_2$  where x = 0.2778, to show enhanced photocatalytic capabilities.<sup>17</sup> However, it is possible that the random distribution of Ti and Zr in the M-sites could lead to disorder-based scatterings which may impede the mobility of the photogenerated electrons and holes. 18 Therefore, ordered semiconductor MXene alloys with reduced disorder-based scatterings would be desirable.

Likewise, another abundant but inefficiently utilized energy source is waste thermal energy, for which, thermoelectric (TE) materials can be exploited for conversion into useful electrical energy. The efficiency of thermoelectric materials can be evaluated using a dimensionless figure-of-merit, defined as  $zT = \frac{S^2 \sigma}{\kappa} T$ , where S is the Seebeck coefficient,  $\sigma$  is the electrical conductivity,  $\kappa$  is the total thermal conductivity, and T is the temperature. As such, a good performing thermoelectric material should possess a high power factor  $S^2\sigma$  and low  $\kappa$ , although these two parameters are usually negatively correlated.<sup>19</sup> Recently, a few Mo-based MXene flexible thin films, e.g., Mo<sub>2</sub>C, Mo<sub>2</sub>TiC<sub>2</sub>, and Mo<sub>2</sub>Ti<sub>2</sub>C<sub>3</sub>, have been experimentally investigated for their thermoelectric performance.<sup>20</sup> The Mo<sub>2</sub>TiC<sub>2</sub> film displays the highest  $S^2\sigma$  of ~3.09  $\mu$ W cm<sup>-1</sup> K<sup>-2</sup> at 800 K (and  $\sim 0.5 \,\mu\text{W cm}^{-1}\,\text{K}^{-2}$  at 300 K) among these thin films. While these values are not exceptional, one can exploit the compositional versatility of MXenes to further optimize, tune, and create a better thermoelectric material.

As such, we wonder whether it is possible to surpass these limitations to discover a novel alloying approach to make MXene photocatalysis and thermoelectric properties better for sustainability applications. Herein, we investigate the substitutional alloy MXene Ti<sub>2(1-x)</sub>Mo<sub>2x</sub>CO<sub>2</sub> and attempt to predict its synthesizability, noting that its thicker counterparts, Mo<sub>2</sub>TiC<sub>2</sub> and Mo<sub>2</sub>Ti<sub>2</sub>C<sub>3</sub>, have been successfully synthesized and are well-studied.<sup>20-22</sup> Using a multiscale approach of density functional theory (DFT) calculations, cluster expansion (CE) method, and Monte Carlo (MC) simulations, <sup>23–27</sup> we screened through a total of more than 9 million different Ti<sub>2(1-x)</sub>Mo<sub>2x</sub>CO<sub>2</sub> MXene alloy configurations with the 3 different types of O-termination. We then systematically map out their structure-stability relationship by validating the thermodynamically stable alloy ground-state(s), followed by the assessment

of their suitability as water-splitting photocatalysts and thermoelectric materials, using their respective figure-of-merit derived from electronic, optical, and thermal calculations. Thus, for the first time, we report the possible existence of an asymmetrically ordered Janus-like (or 2-faced) MXene alloy TiMoCO2. This alloy is thermodynamically, mechanically, and thermally stable with a high orderdisorder phase transition temperature of  $\sim$ 640 K. In essence, the TiMoCO<sub>2</sub> MXene allov is a semiconductor with a band gap of 0.60 eV. The asymmetric structure gives rise to an intrinsic dipole moment leading to band edge realignment due to the valence band maximum (VBM) and conduction band minimum (CBM) situating on the opposite surfaces of the MXene monolayer, which in turn straddles water redox potentials. Photocatalytic redox water-splitting can therefore occur with a much higher solar absorption efficiency, reaching above  $10^4$ – $10^5$  cm<sup>-1</sup> in the visible-infrared region. Its room temperature intrinsic hole and electron mobilities are evaluated to be 133.61-142.39 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> and 19.65-29.99 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> respectively, surpassing some 2D monolayer transition metal dichalcogenides (TMDs). Moreover, we determine the TiMoCO2 MXene alloy to exhibit high room temperature n- and p-type power factors of 33.7–49.8  $\mu$ W cm<sup>-1</sup> K<sup>-2</sup> and ~15  $\mu$ W cm<sup>-1</sup> K<sup>-2</sup>, respectively, which outperform other related experimentally synthesized Ti-Mo based MXene alloys and are even comparable to that of p-doped SnSe, which is one of the best performing thermoelectric materials. All these indicate the stable asymmetrically ordered Janus-like TiMoCO<sub>2</sub> MXene alloy to be a promising high performance water-splitting photocatalyst and thermoelectric material. Our work paves the way for more Janus-like MXenes with interesting and outstanding properties to be discovered.

# 2. Methodology

## Cluster expansion (CE) method and Monte Carlo (MC) simulations

According to the CE formalism, 28-30 a particular alloy configuration, o, on a fixed lattice, can be described using a set of occupational variables. For this work, a pseudo-binary  $Ti_{2(1-x)}$ Mo<sub>2x</sub>CO<sub>2</sub> substitutional MXene alloy is generated on the metallic sublattice with Ti and Mo, while ensuring the composition of C and O on the anion sublattice to remain unvaried. As such, for a given  $\varrho$ , its respective formation energy  $E_f$  corresponds its stability at 0 K to its most stable O-terminated MXene constituents, fcc-Ti<sub>2</sub>CO<sub>2</sub> and hcp-Mo<sub>2</sub>CO<sub>2</sub>:

$$E_{f}(Ti_{2(1-x)}Mo_{2x}CO_{2},Q) = E(Ti_{2(1-x)}Mo_{2x}CO_{2},Q)$$
$$- (1 - x)E(Ti_{2}CO_{2}) - xE(Mo_{2}CO_{2})$$
(1)

where  $E(Ti_2CO_2)$ ,  $E(Mo_2CO_2)$ , and  $E(Ti_{2(1-x)}Mo_{2x}CO_2,\varrho)$  are the total energies per atom of Ti<sub>2</sub>CO<sub>2</sub>, Mo<sub>2</sub>CO<sub>2</sub>, and a particular g of the Ti<sub>2(1-x)</sub>Mo<sub>2x</sub>CO<sub>2</sub> alloy, respectively. This configurationdependent  $E_f$  is described via the CE formalism<sup>31–33</sup> where we can obtain the effective cluster interactions (ECIs) between the alloyants, Ti and Mo. To perform CE, we utilize the Thermodynamic Tool-Kit (TTK) code, 17,22,23,30,32-41 the fundamental principles of which have been methodically elaborated in our previous works for other MXene alloys. 17,40,41 In essence, for

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each alloy configuration (of each fcc, hcp, and fcc + hcp O-terminated MXene, *i.e.*, 3 "different" systems), we construct an initial CE Hamiltonian by fitting the ECIs from a learning set of DFT-calculated  $E_f$  of  $\sim 100$  configurations comprising up to 20-atom  $\mathrm{Ti}_{2(1-x)}\mathrm{Mo}_{2x}\mathrm{CO}_2$  MXene supercells. This is followed by screening of over 3 million alloy structures with unique configurations (up to 70-atom supercells) to obtain their corresponding CE energies. Alloy configurations with  $E_f$  close to or at the ground-state hull would be verified via DFT and added to the learning set to generate an updated CE Hamiltonian for the ensuing iteration(s). The iterative process is ended and the CE is considered to be properly truncated when either no new ground-states are predicted and/or the CE  $E_f$  values are reproduced precisely within < 0.003 eV atom $^{-1}$ .

Utilizing the properly truncated CE, we perform MC simulations via the TTK code to survey the equilibrium Ti–Mo distribution in the alloys at the ground-state hull composition across different temperatures. A simulation cell of  $24 \times 24 \times 1$  times the size of the primitive unit cell of  $Ti_2CO_2$  and  $Mo_2CO_2$  is used. As such, a series of canonical (fixed chemical potential and temperature) MC runs with  $30\,000$  sampling steps per run is conducted to assess the Ti–Mo distribution, and to predict the order–disorder phase transition temperature of the alloy at ground-state hull composition.

#### 2.2 First-principles calculations

Density functional theory (DFT) calculations<sup>42</sup> are performed using the Perdew-Burke-Ernzerhof (PBE) exchange correlation based on the generalized gradient approximation (GGA)<sup>43,44</sup> as implemented in the Vienna ab initio Simulation Package (VASP). 45,46 Electron-ion interactions are described via the projector augmented-wave (PAW) method.<sup>47</sup> The plane-wave cut-off energy is set to 500 eV. During structural relaxation, the volume of the unit cell is fixed while the cell shape is allowed to change. A vacuum spacing of ~15 Å is ensured in the out-of-plane direction to minimize spurious interactions between periodic images. All atomic coordinates are therefore fully relaxed until the Hellmann-Feynman forces on each atom are < 0.001 eV  $\mathring{A}^{-1}$ . A dipole correction in the direction normal to the MXene plane is included for all the calculations. 48 Monkhorst-Pack k-point meshes<sup>49</sup> with a density of  $\sim 40$  k-points per Å<sup>-1</sup> inclusive of the  $\Gamma$ -point are used during structural optimizations, whereas finer meshes of  $\sim 160$  k-points per  $\text{Å}^{-1}$  are utilized for detailed electronic density-of-states (DOS) and electronic band structure calculations. To achieve an accurate description of the band gaps, the band-edge positions, and the dielectric functions of the ground-state alloy, the Heyd-Scuseria-Ernzerhof (HSE06)50,51 hybrid functional is utilized. The effective electron and hole masses are determined by the parabolic fitting of the conduction band minimum (CBM) and valence band maximum (VBM), respectively, along selected k-point paths in the in-plane directions.<sup>52</sup>

To investigate the elastic properties, the elastic tensor is determined using the stress-strain relationship via six finite distortions of the lattice to derive the elastic constants.<sup>53</sup> The ionic contributions are determined from the inversion of the

ionic Hessian matrix which is evaluated from finite differences and multiplied with the internal strain tensor.  $^{54}$ 

*Ab initio* molecular dynamics (AIMD) simulations are employed to evaluate the stability of the ground-state MXene alloy at finite temperatures; the simulations are carried out with a 5  $\times$  5  $\times$  1 supercell at 300 K and 600 K with the Nosé–Hoover thermostat<sup>55,56</sup> for 20 ps with a time step of 1 fs.

The elucidation of the phonon dispersions and the electronphonon couplings is performed using density functional perturbation theory (DFPT)<sup>57,58</sup> under PBE-GGA as implemented in the PHONON package of QUANTUM ESPRESSO. 59,60 Here, a set of optimized norm-conserving Vanderbilt (ONCV) pseudopotentials from PseudoDojo<sup>61</sup> are used to replace core electrons, along with a plane-wave cut-off energy of 100 Ry and a well-converged and uniform  $12 \times 12 \times 1$  k-point mesh for the Brillouin zone sampling, followed by the interpolation of the phonon dispersion and electron-phonon matrix elements into well-converged ultrafine  $240 \times 240 \times 1$  k-point and  $400 \times 400 \times 1$  g-point meshes with maximally localized Wannier functions<sup>62,63</sup> as implemented in Electron-Phonon coupling using the Wannier function (EPW)<sup>64</sup> code for the evaluation of the transport properties for an accurate description of the carrier mobilities and thermoelectric performance via the self-energy relaxation time approximation (SERTA)<sup>65</sup> within the framework of the Boltzmann transport equation (BTE). On the basis of the BTE, we evaluate the electron velocity matrix using first-order k-derivatives of the Hamiltonian via EPW, followed by the subsequent determination of the electrical conductivity  $\sigma$  and Seebeck coefficient S as elaborated according to published works.66,67

## Results and discussion

#### 3.1 Alloy screening and formation stability

A fully O-terminated MXene M<sub>2</sub>CO<sub>2</sub> can adopt 1 of the 3 possible geometries of O atoms, i.e., fcc, hcp, or fcc + hcp (fcc on one surface, and hcp on the other) (Fig. S1, ESI†). While our previous and other published works have shown Ti2CO2 and Mo2CO2 to preferentially adopt the fcc and hcp geometry, respectively, 15,17,41 we have considered all 3 geometries in our screening of alloys lest there are thermodynamically MXene alloys with any of these 3 possible geometries. As such, Fig. 1 illustrates the formation energies  $E_f$  of more than 9 million different alloy configurations derived from eqn (1) for the fcc, hcp, and fcc + hcp O-termination geometries, using the constructed ECIs by the CE method. The most stable configurations (at 0 K) are known as ground-states, which form a ground-state hull (dashed-dotted line) across compositions, implying the theoretical lower bound of  $E_f$  for the  $Ti_{2(1-x)}Mo_{2x}CO_2$  alloy. On the other hand, states or configurations above the ground-state hull are accessible only at finite temperatures. Accordingly, from Fig. 1, only 2 stable alloy ground-states at x = 0.25 (Ti<sub>1.5</sub>Mo<sub>0.5</sub>CO<sub>2</sub>) and x = 0.5 (TiMoCO<sub>2</sub>) can be observed, of which Ti<sub>1.5</sub>Mo<sub>0.5</sub>CO<sub>2</sub> and TiMoCO<sub>2</sub> occur as fcc and fcc + hcp O-terminated MXenes, respectively. On the grounds that only TiMoCO2 displays semiconducting character which is crucial for photocatalytic water-splitting and will be Paper Materials Advances

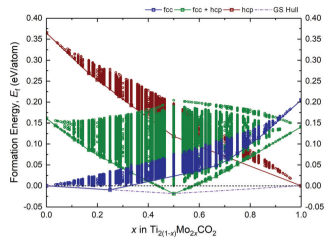


Fig. 1 Plot of formation energy  $E_f$  versus composition for  ${\rm Ti}_{2(1-x)}{\rm Mo}_{2x}{\rm CO}_2$  MXene. Each point represents a particular alloy configuration whose relative stability is indicated by its  $E_f$  with respect to its most energetically stable fcc O-terminated  ${\rm Ti}_2{\rm CO}_2$  and hcp O-terminated  ${\rm Mo}_2{\rm CO}_2$  constituent MXenes. Alloys with fcc, hcp, and fcc + hcp O-termination geometry are indicated in blue, red, and green, respectively. The solid line refers to the local 'ground-state' hull which joins the lowest energy structures across compositions for each corresponding O-termination geometry; whereas the dashed–dotted line is the overall ground–state (GS) hull which joins the lowest energy structures across compositions regardless of the geometry of O-termination.

revealed later, whereas  $Ti_{1.5}Mo_{0.5}CO_2$  is metallic, we focus on the in-depth investigations of the  $TiMoCO_2$  MXene alloy in the following sections.

#### 3.2 Asymmetrically ordered Janus-like TiMoCO<sub>2</sub> MXene alloy

3.2.1 Structural properties and stability. TiMoCO<sub>2</sub> possesses an asymmetrically ordered Janus-like structure in which the atomic layers are stacked in the sequence O(fcc)-Ti-C-Mo-O(hcp) as illustrated in Fig. 2a. Due to this asymmetric structure, the TiMoCO<sub>2</sub> MXene alloy has the space group P3m1 (no. 156) (instead of  $P\bar{3}m1$  (no. 164) like most other  $M_2CO_2$  MXenes) and possesses no inversion symmetry. The configuration of the M-sites (M-C-M') in this MXene alloy is rather intriguing because from our previous works on other Ti-based MXene alloy systems  $(Ti_{2(1-x)}M_{2x}CO_2$ , where M = Zr, Hf, V, Nb, and Ta), <sup>17,41</sup> alloys with such a Janus-like configuration typically possess the most positive  $E_f$  and if ground-states with x = 0.5 exist, it would likely occur with in-plane ordering. Analogously, even experimentally synthesized 2D TMD MoSSe alloys with a Janus-like configuration (S-Mo-Se)<sup>68,69</sup> have been theoretically evaluated to be energetically less favorable than the disordered counterpart. 70-72 The origin of this stability in TiMoCO<sub>2</sub> can be attributed to the appropriate O-termination geometries for the respective Ti (fcc) and Mo (hcp) sides resulting in Bader charges to be similar to those of O, Ti, and Mo in their thermodynamically stable constituent MXenes Ti<sub>2</sub>CO<sub>2</sub> (fcc) and Mo<sub>2</sub>CO<sub>2</sub> (hcp) (Table S1, ESI†). Moreover, we have further verified that even the bare TiMoC MXene alloy without O-terminations is stable by 81.8 meV per atom with respect to the constituent Ti<sub>2</sub>C and Mo<sub>2</sub>C MXenes (Table S2, ESI†). Accordingly, our TiMoCO<sub>2</sub> MXene alloy with its

favorable  $E_{
m f}$  and thermodynamic stability would have enhanced synthesizability.

To further validate the stability of the ground-state configuration of TiMoCO2, a series of canonical MC runs, AIMD simulations, and phonon calculations are performed. Fig. 2b and c show, respectively, a plot of average energy  $U_{\text{avg}}$  (of the MC simulation cell) and another plot of heat capacity  $C_v$ , both versus temperature T. The ordered configuration at the end of the MC runs corresponds exactly to the configuration of the DFT- and CE-determined ground-state TiMoCO2 MXene alloy (Fig. 2a). Moreover, the order-disorder phase transition temperature, which is the peak in the plot of  $C_v$  versus T, is determined to be  $\sim$ 640 K, implying that the ordered configuration is thermodynamically stable under room conditions ( $\sim$  300 K). Besides, AIMD simulations in Fig. 2d carried out at 300 K and 600 K (below its order-disorder phase transition temperature) show the energies to fluctuate about constant values, with negligible structural deviations and no structural reconstructions after annealing for 20 ps. In addition, the phonon dispersion in Fig. 2e along the high-symmetry path of the Brillouin zone exhibits no negative (imaginary) frequencies, suggesting the dynamic stability of the TiMoCO2 ground-state configuration.

Likewise, the mechanical stability of the TiMoCO<sub>2</sub> MXene alloy is also examined. A (hexagonal) 2D MXene possesses 3 independent non-zero elastic constants which are calculated to be:  $C_{11} = 322.22$  N m<sup>-1</sup>,  $C_{12} = 103.06$  N m<sup>-1</sup>, and  $C_{66} = 109.58$  N m<sup>-1</sup>. These elastic constants fulfill the Born stability criteria according to Born–Huang's lattice dynamical theory for hexagonal 2D structures,  $^{73,74}$  *i.e.*,  $C_{11} > 0$ ,  $C_{11} - C_{12} > 0$ , and

 $C_{66} = \frac{C_{11} - C_{12}}{2}$ , indicating that the TiMoCO<sub>2</sub> MXene alloy is mechanically stable. Using these elastic constants, we evaluate the in-plane Young's modulus  $Y_{\rm s}$  and the Poisson's ratio  $\nu$  according to the following expressions:  $^{75,76}$   $Y_{\rm s} = \frac{C_{11}^2 - C_{12}^2}{C_{11}}$ 

and  $\nu = \frac{C_{12}}{C_{11}}$ , which give 289.26 N m<sup>-1</sup> and 0.32, respectively. We note that using the same approach, we obtain  $Y_{\rm s}$  values of 243.02 N m<sup>-1</sup> and 241.04 N m<sup>-1</sup> for the constituent MXenes Ti<sub>2</sub>CO<sub>2</sub> and Mo<sub>2</sub>CO<sub>2</sub>, indicating that the alloying would produce a mechanically stronger material. Moreover, this value of  $Y_{\rm s}$  is much higher than those of 2D TMDs such as monolayer MoS<sub>2</sub> (171 ± 11 N m<sup>-1</sup>)<sup>77</sup> and WS<sub>2</sub> (177 ± 12 N m<sup>-1</sup>),<sup>77</sup> and slightly lower than that of graphene (324 ± 13 N m<sup>-1</sup>),<sup>78</sup> which is the strongest 2D material. This implies the TiMoCO<sub>2</sub> MXene alloy to overall possess high mechanical strength as well. With the thermodynamically favorable  $E_{\rm f}$ , dynamically stable phonon dispersion, thermally stable MC and AIMD simulations, and favorable elastic constants, we believe that the experimental realization of this material is highly feasible.

3.2.2 Photocatalytic properties. Fig. 3a illustrates the electronic band structure of the TiMoCO<sub>2</sub> ground-state configuration with a band gap of 0.60 eV. The band gap is indirect with the valence band maximum (VBM) located on the  $\Gamma$  point, whereas the conduction band minimum (CBM) resides on the

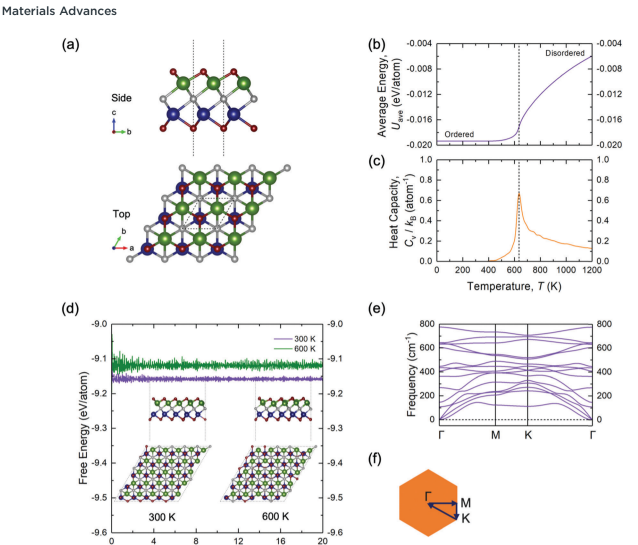


Fig. 2 (a) Side and top views of the Janus-like ground-state  $TiMoCO_2$  MXene alloy. The red, green, blue, and grey spheres correspond to O, Ti, Mo, and C, respectively. Plots of the (b) average energy  $U_{avg}$  and (c) heat capacity  $C_v$ , both versus the temperature T for the  $TiMoCO_2$  MXene alloy. (d) Plot showing the variations in the free energy of the ordered Janus-like ground-state  $TiMoCO_2$  MXene alloy at 300 K and 600 K with respect to the AIMD simulation time. The insets show the side and top views of the final crystal structure of the  $TiMoCO_2$  MXene alloy at 300 K and 600 K after 20 ps. (e) Phonon dispersion along the high-symmetry path of (f) the hexagonal Brillouin zone for the Janus-like ground-state  $TiMoCO_2$  MXene alloy.

K– $\Gamma$  valley. At first glance, this value of the band gap appears to fall short of the '1.23 eV band gap' prerequisite for typical redox water-splitting photocatalysis to be thermodynamically feasible. However, for TiMoCO $_2$  MXene alloys, this requirement can be overcome for reasons explained in the following paragraph. While a thicker absorption layer is typically required for an indirect band gap material, this indirect gap impedes the rate of electron–hole recombination, thereby increasing the probability for the holes and electrons to partake in the water-splitting redox reactions. <sup>79</sup>

Time (ps)

From the atomic projections of the electronic band structure and the charge density isosurface distributions (Fig. 3a–c), it can be observed that the VBM state is mainly composed of Mo-d<sub>z²</sub> and O-p<sub>z</sub> (at Mo side) orbitals, whereas the CBM state mainly originated from Ti-d<sub>xz</sub> orbitals. This distinction in the orbital contributions for both sides of TiMoCO<sub>2</sub> is the key factor

for allowing photocatalytic redox water-splitting to be thermodynamically feasible even with a small band gap. Due to the difference in the electronegativity between Ti (1.54) and Mo (2.16),80 and consequent Bader charge of Ti (+1.92e), Mo (+1.60e), and the fcc-terminated and hcp-terminated O (-1.08e and -0.93e, respectively), an intrinsic dipole moment of 0.232 D is generated with the electric field pointing from the Ti to Mo side. This dipole moment-induced electric field promotes photogenerated electron-hole separation and localizes the electrons and holes on Ti and Mo sides, respectively. As such, there exists an electrostatic potential difference of 1.53 eV as shown in Fig. S2 (ESI†) and Fig. 3d. Moreover, by aligning the band edge positions relative to the respective water redox potentials, i.e., the VBM to water oxidation potential (5.67 eV) w.r.t. the Mo side's vacuum level and the CBM to water reduction potential (4.44 eV) w.r.t. the Ti side's vacuum

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(d) (a) e\ E<sub>Vacuum-Ti</sub> (b) **CBM** O2/H2O Energy (eV) **VBM** 0 (e) 10<sup>6</sup> Absorption Soefficient,  $\alpha(\omega) (cm^{-1})$ 10<sup>5</sup> 10<sup>4</sup>

Fig. 3 (a) Projected electronic band structure of the Janus-like ground-state TiMoCO<sub>2</sub> MXene alloy via the HSE06 functional plotted along the highsymmetry path of the hexagonal Brillouin zone (see Fig. 2f). The size of each point corresponds to the relative magnitude of the respective elemental contribution to the band. The Fermi level is set to 0 eV at the VBM. The side view of the partial charge density of the (b) CBM (in green) and (c) VBM (in blue) of the TiMoCO<sub>2</sub> MXene alloy. The value of the isosurfaces is  $0.02 \text{ e Å}^{-3}$ . The red, green, blue, and grey spheres correspond to O, Ti, Mo, and C, respectively. (d) Band edge alignments of the VBM and CBM in blue and green solid lines, respectively, with regard to the water redox potentials in dashed lines, and their corresponding vacuum levels. The potential difference between the two sides of the TiMoCO<sub>2</sub> MXene alloy induced by the intrinsic dipole moment is denoted in red. For the actual electrostatic potential profile, please refer to Fig. S2 (ESI†). (e) Plot of the in-plane optical absorption coefficient  $\alpha(\omega)$  versus photon energy for the TiMoCO<sub>2</sub> MXene alloy. The visible wavelength region (1.65 to 3.10 eV) is shown as an iridescent rectangle for guide.

level, we observe in Fig. 3d that both the VBM and CBM straddle the water redox potentials, indicating that redox water-splitting photocatalysis is thermodynamically feasible albeit there is a band gap of 0.60 eV. The bending of the vacuum level along the direction of the internal electric field permits the reduction of the band gap requirement, i.e., difference between oxidation and reduction potentials of water, from 1.23 eV to 1.23 eV minus the value of the electrostatic potential difference between the two surfaces.81 Intriguingly, for our TiMoCO<sub>2</sub> MXene alloy with an electrostatic potential difference of 1.53 eV, the band gap requirement would therefore be 0 eV, indicating that so long as the VBM and CBM are located on the opposite sides of the MXene, the size of its band gap would not affect the thermodynamic feasibility of photocatalytic redox water-splitting.

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With the omission of the band gap prerequisite, the absorption properties of the TiMoCO<sub>2</sub> MXene alloy is investigated to elucidate its efficiency in solar conversion. Fig. 3e shows the absorption coefficient  $\alpha(\omega)$  derived from the calculated frequency-dependent complex dielectric function  $\varepsilon(\omega) = \varepsilon_1(\omega) +$ 

 $i\varepsilon_2(\omega)$  according to the relation:<sup>82</sup>  $\alpha(\omega) = \frac{\sqrt{2}\omega}{c}\sqrt{|\varepsilon(\omega)| - \varepsilon_1(\omega)}$ , with *c* being the speed of light in vacuum. It can be observed that the TiMoCO<sub>2</sub> MXene alloy exhibits strong absorption of a wide wavelength range from ultraviolet to visible to infrared with  $\alpha(\omega)$ reaching above  $10^4$ – $10^5$  cm<sup>-1</sup> in the visible-infrared region. Since this region accounts for majority of the incident solar energy, by implementation of the metric which integrates the product of the incident solar photon flux according to the AM1.5G solar

spectrum with the absorbance of the material up to its band gap's equivalent wavelength, as elaborated comprehensively in our previous work as well as other literature, 17,36,83 we obtain its maximum short-circuit current of 4.2 mA cm<sup>-2</sup>. This value is much larger than those of conventional photovoltaic materials such as Si, and also those of 2D materials like MoS<sub>2</sub> and WS<sub>2</sub>, with maximum short-circuit currents ranging from 0.1 to 3.6 mA cm<sup>-2</sup> (Table 1), which suggests TiMoCO<sub>2</sub> to be a potential photovoltaic material, in addition to its desirable visible-infrared absorbance for efficient photocatalysis.

Photon Energy (eV)

Other than the appropriate band and promising absorption characteristics, the electron and hole mobilities of the TiMoCO<sub>2</sub> MXene alloy are also investigated to elucidate its efficacy in photocatalytic water-splitting. Herein, using the SERTA approach and accounting for electron-phonon scattering of all the phonon modes, <sup>65</sup> we obtain the room temperature intrinsic hole mobility  $\mu_p$  to be 142.39 and 133.61 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> along the x- and y-direction, respectively. These values show high isotropy in both x- and y-directions, which are consistent with the VBM being comprised of Mo-d<sub>z2</sub> and O-p<sub>z</sub> orbitals, and the consequent isotropic effective hole mass of  $0.23m_0$  along both x- and y-directions, indicating in-plane orientationindependent hole transport in either direction of TiMoCO<sub>2</sub>. In contrast, the room temperature intrinsic electron mobility  $\mu_n$ is anisotropic with 29.99 and 19.65 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> along the corresponding x- and y-directions, in accord with the contribution of the in-plane anisotropic Ti-d<sub>xz</sub> orbitals to the CBM with anisotropic effective electron masses of  $0.81m_0$  and  $1.31m_0$ along the respective x- and y-directions. It is evident that  $\mu_p$  is

Table 1 Figures-of-merit of the maximum short-circuit currents for photocatalytic water-splitting and power factors at 300 K for thermoelectric applications for TiMoCO<sub>2</sub> and other representative water-splitting photocatalytic and thermoelectric materials

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Material	Maximum short-circuit current <sup>a</sup> (mA cm <sup>-2</sup> )	Thermoelectric power factor (μW cm <sup>-1</sup> K <sup>-2</sup> )	
		n-doping	p-doping
TiMoCO <sub>2</sub>	4.2	33.7-49.8 <sup>a</sup>	16.0 <sup>a</sup>
$MoS_2$	3.6	$35.0^{88}$	_
$WS_2$	2.7	_	$2.8^{89}$
Si	0.1	_	_
$Mo_2TiC_2$	_	$0.5^{20}$	_
$Mo_2Ti_2C_3$	_	$0.1^{20}$	_
SnSe	_	_	$10.0 – 40.0^{90}$
<sup>a</sup> This work.			

larger than  $\mu_n$  as a result of lighter effective mass. Besides, the  $\sim 1$  order of magnitude difference between  $\mu_{\rm p}$  and  $\mu_{\rm n}$  promotes the effective spatial separation of the holes and electrons, which decreases the probability of photogenerated electronhole recombination.<sup>35,38</sup> To estimate the reliability of the calculated carrier mobilities with our approach,  $\mu_p$  and  $\mu_n$  values of widely investigated 2D MoS2 are analogously calculated to be 29.22 and 121.58  $\mathrm{cm}^2~\mathrm{V}^{-1}~\mathrm{s}^{-1}$ , respectively, which are close to their corresponding experimental measurements of  $\sim$  40–76 and  $\sim 60-150$  cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>.84-86 Evidently, TiMoCO<sub>2</sub> possesses a higher  $\mu_p$  than that of MoS<sub>2</sub>, suggesting its potential application as a hole transport material. Accordingly, the indirect band gap, dipole-induced electric field, appropriate band edge positions, high absorbance, and decent mobilities would favor photogeneration of the electrons and holes while bolstering effective spatial and charge separation of these photoelectrons and holes for effective water redox reactions, establishing the TiMoCO<sub>2</sub> MXene alloy to be an efficient water-splitting photocatalyst.

3.2.3 Thermoelectric properties. Meanwhile, from the calculated transport properties of the TiMoCO2 MXene alloy, we are able to further utilize these data to assess its potential thermoelectric performance. In the simulation of the doping effects, we achieve n- or p-type doping by assuming a shift of the position of the Fermi level into the conduction or valence bands, respectively, within the rigid band approximation. Fig. 4 shows the various thermoelectric quantities at room temperature, *i.e.*, Seebeck coefficient S, electrical conductivity  $\sigma$ , and power factor  $S^2\sigma$  with respect to carrier concentration N, which suggest n-doping of the TiMoCO<sub>2</sub> ground-state configuration to produce a better thermoelectric material with higher peak  $S^2\sigma$ than p-doping. These peak  $S^2\sigma$  values show varying degrees of isotropy corresponding to their band character, similar to their respective mobilities as described above. Moreover, compared with other representative thermoelectric materials as tabulated in Table 1, the peak n-doped  $S^2\sigma$  is  $\sim 2$  orders of magnitude larger than those of symmetrical Ti-Mo MXene alloys Mo<sub>2</sub>TiC<sub>2</sub> and  $Mo_2Ti_2C_3$  and  $\sim 1$  to 1.5 times larger than that of the  $MoS_2$ monolayer. However, the peak p-doped  $S^2\sigma$  is comparable to that of p-doped SnSe which is one of the best performing thermoelectric materials. It is important to also note that the

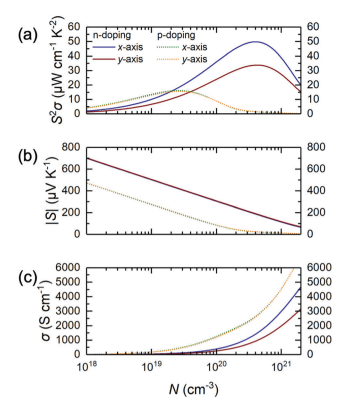


Fig. 4 Plots of thermoelectric quantities such as (a) power factor  $S^2\sigma$ , (b) Seebeck coefficient S, and (c) conductivity  $\sigma$  as a function of carrier density N, assuming an effective monolayer thickness of 7.2 Å, for the n- and p-doped Janus-like ground-state TiMoCO<sub>2</sub> MXene alloy at room temperature. Due to isotropy, the plots for p-doped thermoelectric properties for x- and y-directions are almost equal to and overlap with each other.

peak n- and p-doped  $S^2\sigma$  can be achieved at reasonably decent optimal doping concentrations of 4.0  $\times$  10<sup>20</sup> cm<sup>-3</sup> and 2.5  $\times$ 10<sup>19</sup> cm<sup>-3</sup>, respectively, assuming an effective monolayer thickness of 7.2 Å. On top of that, the softer ZA phonon mode (lowest phonon band in Fig. 2e) and the absence of inversion symmetry could potentially reduce thermal conductivity<sup>87</sup> which is crucial for maximizing thermoelectric figure-of-merit zT; and hence, entailing the TiMoCO2 MXene alloy to be a potential thermoelectric material upon optimal n- or p-type doping.

3.2.4 Further exploration in the design of functional materials. In addition, before concluding this work, it is imperative for us to propose possible approaches and alloying formulations to set a direction for further optimization and tuning of the functional properties of asymmetrically ordered Janus-like MXenes. As such, we consider permutations between elements in the same respective group as Ti or Mo in the periodic table, i.e., ZrMoCO2, HfMoCO2, TiCrCO2, and TiWCO2, since their valency is maintained albeit at different electron energy levels. Some preliminary results with regard to their electronic band structures are illustrated in Fig. S3 and their electronic properties are tabulated in Table S3 (ESI†). Upon Ti replacement with Zr or Hf, while the band gaps vary mildly and the effective hole masses increase slightly down the group, the main contribution to the CBM has transformed to Mo-d<sub>z2</sub> orbitals, indicating that since both the CBM and VBM are on the Mo's side, the

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photocatalytic water-splitting redox potential would not be realigned and the band gap would no longer permit redox water-splitting to occur simultaneously. In spite of this, the drastic reduction in the effective electron masses would suggest increased electron mobilities, potentially improving their n-type thermoelectric transport properties. On the other hand, substitution of Mo with Cr or W has resulted in significant decreases in the band gaps, for TiCrCO2 due to the CBM and VBM being on the Cr's side analogous to the situation described for ZrMoCO<sub>2</sub> and HfMoCO<sub>2</sub>, implying photocatalytic redox water-splitting to be thermodynamically unfavorable. However, the VBM and CBM of TiWCO2 still reside on the opposing sides of the MXene layer, which, coupled with the larger solar absorption from the reduced band gap and also a higher degree of photogenerated electron-hole separation due to larger effective mass variation between holes and electrons, could synergistically further enhance the photocatalytic performance. It is important to note that a caveat for these conjectural ZrMoCO<sub>2</sub>, HfMoCO2, TiCrCO2, and TiWCO2 MXene alloys is that their stability evaluation involving CE and DFT calculations is yet to be implemented and more meticulous investigations have to be made to validate their formation stabilities, synthesizability, and their photocatalytic water-splitting and thermoelectric performance. Regardless, we have showcased that such Janus-like asymmetrically ordered MXene alloys can give rise to intriguing and even contrasting properties, signifying that this would be a promising direction for tuning and optimization of MXenes for tailored functional properties.

## 4. Conclusions

In summary, through a systematic first-principles multiscale approach using DFT calculations, CE method, and MC simulations, we are able to identify a novel asymmetrical ordered Janus-like TiMoCO<sub>2</sub> MXene alloy ground-state by screening through the formation energies of more than 9 million different configurations of Ti<sub>2(1-x)</sub>Mo<sub>2x</sub>CO<sub>2</sub> substitutional alloy system with consideration of all 3 possible types of O-termination. This alloy is thermodynamically, mechanically, and thermally stable with a high order-disorder phase transition temperature of  $\sim 640$  K. Importantly, TiMoCO<sub>2</sub> is semiconducting with a band gap of 0.60 eV. It possesses an intrinsic dipole moment as a consequence of the asymmetric structure which realigns the conduction and valence band edges located on opposite sides of the MXene, permitting photocatalytic redox water-splitting to be thermodynamically favorable with enhanced solar absorption efficiency above 10<sup>4</sup>–10<sup>5</sup> cm<sup>-1</sup> in the visible-infrared region. As a thermoelectric material, it is found that TiMoCO<sub>2</sub> exhibits high room temperature n- and p-type power factors of 33.7-49.8  $\mu \text{W cm}^{-1} \text{ K}^{-2}$  and  $\sim 15 \ \mu \text{W cm}^{-1} \text{ K}^{-2}$ , respectively, surpassing other related experimentally synthesized Ti-Mo based MXene alloys and are even similar to that of p-doped SnSe which is one of the best performing thermoelectric materials. All these demonstrate the asymmetrically ordered Janus-like TiMoCO<sub>2</sub> MXene alloy to be a promising high performance water-splitting photocatalyst and thermoelectric material, paving

the way for more stable Janus-like MXenes with interesting and outstanding properties awaiting to be discovered.

# Conflicts of interest

The authors declare no competing financial interest.

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