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## Structural engineering of MXenes for enhanced magnesium ion diffusion: a computational study<sup>†</sup>

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The unique layered structure and tunable surface terminations of MXenes play a critical role in  $Mg^{2+}$  storage and diffusion dynamics. This study systematically investigates the behavior of  $Mg^{2+}$  in  $Ti_3C_2O_2$  and its nitrogen-doped derivatives through theoretical calculations. In  $Ti_3C_2O_2$  monolayers,  $Mg^{2+}$  exhibits a high diffusion barrier of 0.81 eV due to strong electrostatic interactions. However, AA-stacking reduces this barrier to 0.32 eV by introducing staggered active sites. The instability caused by interlayer O–O repulsion is mitigated by modulating the N/O ratio ( $Ti_3C_2O_{1.78}N_{0.22}$ ), resulting in a diffusion barrier of 0.27 eV. Transition metal substitution further optimizes performance, as exemplified by  $Nb_3C_2N_2$ , which achieves an ultralow barrier of 0.23 eV through weakened N–N covalency and enhanced metal–N interactions. Voltage analysis reveals that  $Nb_3C_2N_2$  possesses dual functionality as both cathode (4.00 V) and anode (0.64 V), contrasting with the anode-specific behavior observed in Ti-based MXenes.

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### Introduction

With the rapid depletion of fossil fuels, sustainable energy storage solutions grow increasingly vital.<sup>1</sup> In the past few decades, rechargeable lithium-ion batteries (LIBs) have dominated portable electronics and energy storage systems through their high energy density and cycling stability.<sup>2,3</sup> However, natural scarcity and dendritic growth risks of LIBs<sup>4–6</sup> necessitate alternative solutions.<sup>7</sup> Rechargeable magnesium ion batteries (MIBs) emerge as a promising alternative to LIB, offering high energy density and superior electrochemical stability.<sup>8</sup> Mg is the fifth most abundant element in the Earth's crust,<sup>9</sup> enabling substantial cost savings compared to Li. The melting temperature of Mg is measured at 923 K, much higher than that of Li (453.7 K), making MIBs the safer option for working at high temperatures like in aviation. Divalent  $Mg^{2+}$  delivers doubled volumetric capacity (3833 vs. 2046 mA h cm<sup>-3</sup>) and dendrite-free operation with low reduction potential (−2.37 vs. SHE), and is expected to achieve reversible deposition in multiple electrolytes.<sup>8,10–12</sup>

However,  $Mg^{2+}$  (0.72 Å) has a similar ionic radius to Li<sup>+</sup> (0.76 Å).<sup>13</sup> The large charge/radius ratio of  $Mg^{2+}$  leads to strong electrostatic interaction with the electrode framework, resulting in

sluggish ion diffusion.<sup>14,15</sup> Previous research on MIB electrode materials mainly focused on the transition metal–oxygen/sulfide, polyanionic compounds, and other two-dimensional materials. In transition metal oxides/sulfides systems (e.g.,  $\alpha$ -V<sub>2</sub>O<sub>5</sub>,<sup>16</sup>  $\alpha$ -MoO<sub>3</sub>,<sup>17</sup> Ti<sub>2</sub>S<sub>3</sub>/TiS<sub>2</sub> (ref. 18)), strong  $Mg^{2+}$ -framework electrostatic interactions induce structural transformations that elevate  $Mg^{2+}$  diffusion barriers to 0.9–1.30 eV. By contrast, Chevrel phases and polyanionic compounds exhibit reversible  $Mg^{2+}$  (de)intercalation with mitigated electrostatic interactions.<sup>19</sup> But even for polyanionic materials with good ion mobility, their energy barrier in MIBs is usually higher than 0.6 eV, such as  $Mg_{0.25}FePO_4$  (1.03 eV) and  $NaV_2O_2(PO_4)_2F$  (0.78 eV).<sup>20</sup> Moreover, for the olivine-type  $Mg_xMnSiS_4$ , the strong Mn–S covalent interaction weakens the electrostatic interactions between  $Mg^{2+}$  and the host material ions, thereby enhancing  $Mg^{2+}$  diffusion compared to its oxide counterpart  $Mg_xMnSiO_4$ . However, the diffusion barrier for  $Mg^{2+}$  in  $Mg_xMnSiS_4$  remains as high as 0.76 eV.

Transition metals-based layered materials have stable structures and excellent electrical conductivity.<sup>21</sup> More importantly, the layered space provides a broad diffusion channel for  $Mg^{2+}$ . Thus, 2D materials possess a lower diffusion barrier than the above materials, such as Si<sub>2</sub>BN (0.08–0.35 eV), TiSe<sub>2</sub> (0.88 eV), VSe<sub>2</sub> (0.346 eV) and VS<sub>2</sub> (0.593 eV).<sup>22,23</sup> As a layered material, MXenes have garnered significant attention in electrochemical energy storage due to their electrical conductivity, large specific surface area, and stable layered architecture for the guest ions.<sup>16,24,25</sup> Lukatskaya *et al.* showed that  $Ti_3C_2T_x$  can provide ample accommodation to accommodate cations, such as Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, Mg<sup>2+</sup> and Al<sup>3+</sup>, which produce large volumetric capacitance.<sup>26</sup> Xie *et al.*<sup>15</sup> confirm that MXene monolayer cation

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storage occurs in a comparatively low voltage window, which could be used as anodes for rechargeable batteries. Currently, about 40 MXenes have been synthesized experimentally and hundreds of new MXenes are simulated theoretically.<sup>27–30</sup> It is crucial for MIB to investigate and enhance the diffusion performance of Mg ions within MXene materials.<sup>31,32</sup> Moreover, traditional research on MXene batteries typically focuses on ion storage mechanisms under the default stacking configuration. In fact, the existence of different stacking types is an important feature of layered materials. The most typical example is the layered oxide cathode of potassium-ion batteries. Their different stacking structures can form different potassium-ion storage environments, such as edge-sharing octahedral (O3) or face-sharing prismatic (P2). O3-type materials have high structural stability and can effectively inhibit the migration of transition metals into the K layer, thereby reducing structural degradation during the cycling process. P2-type materials exhibit excellent rate performance due to their unique prismatic coordination environment, which provides shorter K ion migration paths and lower energy barriers. Obviously, stacking type has a significant impact on the storage and migration of ions.<sup>33</sup> However, research on the impact of stacking configurations on the MXene electrochemical properties and the corresponding modulation mechanisms is still lacking.

This work investigates the  $Mg^{2+}$  storage and diffusion mechanisms in MXene-based materials, emphasizing the interplay between structural configurations and electrochemical performance. Based on integrating stacking engineering, functional group modulation, and transition metal substitution strategies, we elucidate the fundamental principles governing  $Mg^{2+}$  behavior, such as mitigating electrostatic interactions, optimizing interlayer covalent bonding, and modulating electronic structures. These findings demonstrate that hierarchical structural design-stacking control, doping, and metal substitution-significantly enhances  $Mg^{2+}$  kinetics and stability, advancing high-performance MIB electrodes.

## Method

All first-principles density functional theory (DFT) calculations are performed using the Vienna *Ab initio* Simulation Package (VASP)<sup>34–37</sup> based on Perdew–Burke–Ernzerhof's (PBE) generalized gradient approximation (GGA-PBE).<sup>38,39</sup> The plane wave truncation energy is set to 550 eV. The valence electron selection for pseudopotential elements is as follows: Mg-3s<sup>2</sup>3p<sup>0</sup>, O-2s<sup>2</sup>2p<sup>4</sup>, C-2s<sup>2</sup>2p<sup>2</sup>, N-2s<sup>2</sup>2p<sup>3</sup>, Cl-3s<sup>2</sup>3p<sup>5</sup>, S-3s<sup>2</sup>3p<sup>4</sup>, F-2s<sup>2</sup>2p<sup>5</sup>, Ti-3d<sup>2</sup>4s<sup>2</sup>, V-3d<sup>3</sup>4s<sup>2</sup>, Zr-4d<sup>2</sup>5s<sup>2</sup>, Nb-4d<sup>4</sup>5s<sup>1</sup>. Considering the strong correlation system of transition metal D-orbital electrons, GGA + *U* method is adopted in the calculation, and the *U* values of Ti, Nb, V and Zr are set as 2.5 eV, 2.0 eV, 2.5 eV and 2.0 eV respectively, which are obtained from the reference of element *U* values given by Materials Studio software. Conjugate gradient method was used to optimize the lattice structure. The energy convergence standard was set to 10–6 eV per atom and the force convergence standard was set to 0.01 eV Å<sup>–1</sup>. In addition, this work adopts the CI-NEB method<sup>40–42</sup> to simulate the diffusion performance of  $Mg^{2+}$  in single/multilayer  $M_3C_2T_x$  ( $M = Ti, V, Zr, Nb$ ,  $T = O, N$ )

in a  $3 \times 3 \times 1$  supercell model. Including the beginning and end positions before and after diffusion, a total of 7 insertion points are set, and the convergence standard is that the force of each insertion point is less than 0.03 eV Å<sup>–1</sup>.

## Results and discussion

### Structure and properties of bulk phase $Ti_3C_2O_2$

The  $Ti_3C_2$  structure is stacked by the Ti–C–Ti–C–Ti layer. In each layer, the atoms are arranged in hexagons. During the preparation of MXene, the –OH in the etched aqueous solution will spontaneously adsorb on the surface, forming –O terminated groups and water. Based on the type of molten salt and subsequent treatment conditions, other terminated groups, such as –Cl, –I, –Br, –S, –Se and –N could also be adsorbed on the surface of MXene layers. Among these function groups, previous theoretical studies mainly considered –O adsorbed in MXene electrode materials. Because divalent  $O^{2-}$  with a larger chemical activity has more affinity for the cation exposed framework. The adsorbed atoms inherit the hexagonal arrangement of the Ti–C–Ti–C–Ti layer, which distributes above the middle Ti atoms (Fig. 1).

Generally, the synthesized MXene materials are bulk rather than monolayer. According to the relative position of adjacent layers, the MXene structure has three possible stacking types, including AA (Table S1†), ABC<sub>1</sub> (Table S2†) and ABC<sub>2</sub> (Table S3†) types (Fig. 2a). For ABC<sub>1</sub> and ABC<sub>2</sub> types, terminal O is directly opposite to the outer Ti and C in the adjacent  $Ti_3C_2O_2$  layer, respectively. The AA-stacking structure is characterized by the alignment of the terminal O groups of adjacent  $Ti_3C_2O_2$  layers. According to the calculated phonon spectra, the phonon band for AA-stacking appears at –0.40 THz, which indicates a slight kinetic instability (Fig. 2b). By contrast, the phonon band structures without imaginary frequencies suggest that ABC<sub>1</sub> and ABC<sub>2</sub>-stacking configurations are more likely to exist in nature. Moreover, our calculated relative energies show that the  $Ti_3C_2O_2$  bulk with ABC<sub>1</sub>-stacking is the most stable configuration (Fig. 2c). In comparison, the AA and ABC<sub>2</sub> types have relatively higher energy of 0.30 and 0.09 eV, respectively. Such relative stability of ABC<sub>1</sub>-stacking is also applies consistently to the case of –Cl and –F terminate.

### Adsorption of $Mg^{2+}$ in $Ti_3C_2O_2$

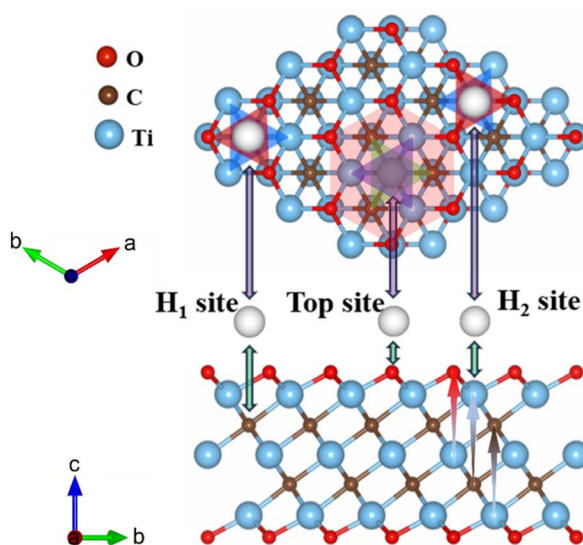
Each surface of the  $Ti_3C_2O_2$  layer has three possible  $Mg^{2+}$  active sites named H<sub>1</sub>, H<sub>2</sub> and T, which are above the C, Ti and O atoms, respectively (Fig. 1a). H<sub>1</sub> and H<sub>2</sub> sites in two triangular hollow regions formed by three O and three Ti atoms, respectively. While the T site has only one coordinated O atom. To determine the most favorable  $Mg^{2+}$  active site, the  $Mg^{2+}$  binding energy  $E_b$  on the  $2 \times 2 \times 1$   $Ti_3C_2O_2$  monolayer was calculated through the following formula:

$$E_b = E_{Ti_3C_2N_2Mg_x} - E_{Ti_3C_2N_2} - xE_{Mg} \quad (1)$$

where  $E_{Mg_xTi_3C_2O_2}$ ,  $E_{Ti_3C_2O_2}$  and  $E_{Mg}$  are the total energy of  $Ti_3C_2O_2Mg_x$ ,  $Ti_3C_2O_2$  and one atom of metal Mg. The calculation results show that the H<sub>1</sub> site exhibited a more negative  $E_b$  (–2.16 eV) than H<sub>2</sub> (–1.36 eV) site. While the  $Mg^{2+}$  on T site migrates to



(a)



(b)

Binding energy: -0.68 eV -1.36 eV -2.16 eV

The Valence: +0.84 +1.41 +1.55

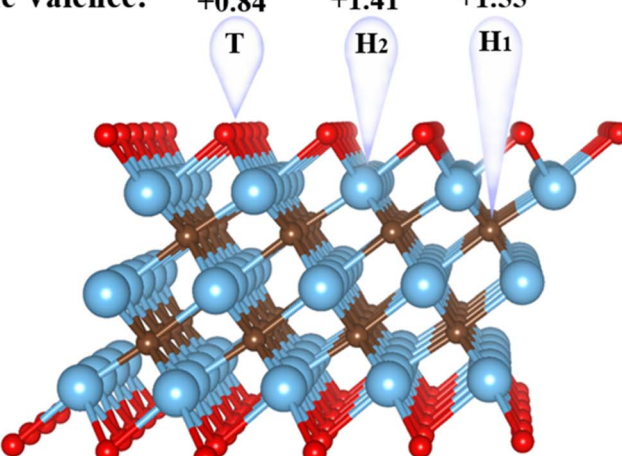


Fig. 1 (a) The top and side views of the three  $Mg^{2+}$  adsorption sites, namely,  $H_1$ ,  $H_2$  and  $T$  site on monolayer  $Ti_3C_2O_2$ . (b) Binding energy and valence states of  $Mg^{2+}$  at  $H_1$ ,  $H_2$  and  $T$  sites.

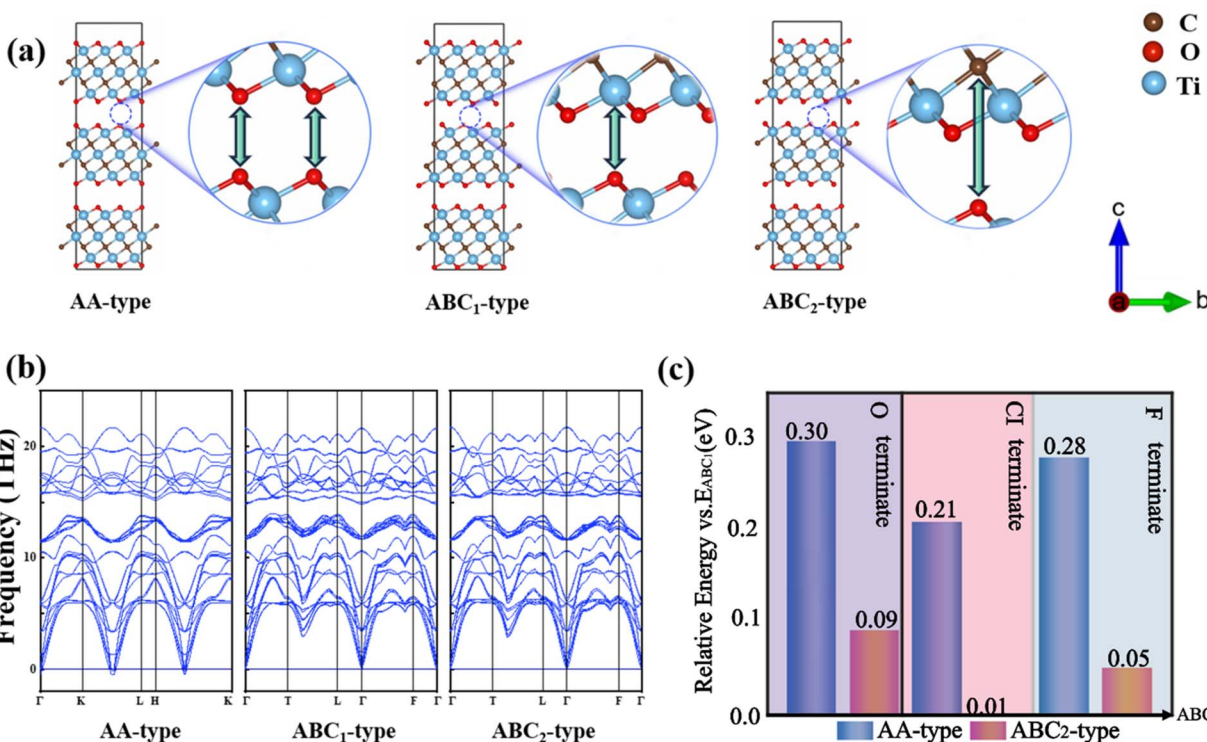


Fig. 2 (a) The side-views of the three stacked  $Ti_3C_2O_2$ , namely  $ABC_1$  type,  $AA$  type and  $ABC_2$  type. (b) The phonon band structures of  $AA$ -,  $ABC_1$ - and  $ABC_2$ -stacked configuration. (c) The relative energy of  $ABC_1$ - and  $ABC_2$ -type vs. the energy of  $AA$ -type for  $Ti_3C_2O_2$ ,  $Ti_3C_2Cl_2$  and  $Ti_3C_2F_2$ , respectively, in left to right order.

the adjacent  $H_1$  site after structural optimization. In the case of fixed  $Mg^{2+}$  at a specific  $xy$  coordinate, the binding energy at the  $T$  site is only  $-0.68$  eV. This indicates that a low coordination degree is not favorable for  $Mg^{2+}$  storage. According to the Bader charge analysis, the valence states of  $Mg^{2+}$  on  $H_1$ ,  $H_2$  and  $T$  site

was  $+1.55$ ,  $+1.41$  and  $+0.84$ , respectively (Fig. 1b). Obviously,  $Mg^{2+}$  at the  $H_1$  site is closer to its theoretical standard valence state, which is more conducive to its binding. When the  $Ti_3C_2O_2$  layers are stacked to form the bulk, the  $Mg^{2+}$  active site is provided by both upper and lower  $Ti_3C_2O_2$  layers. All possible

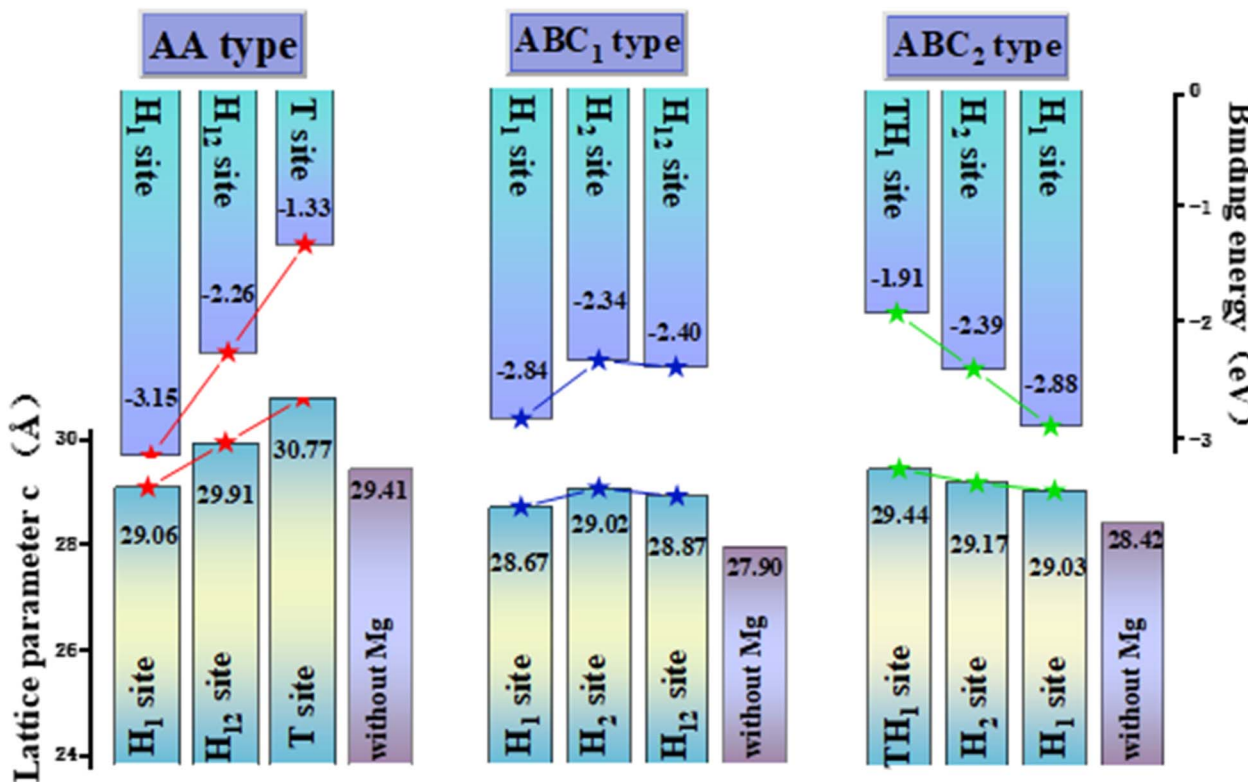


Fig. 3 The binding energy of  $Mg^{2+}$  at each adsorption site of  $Ti_3C_2O_2$  and the change trend of lattice parameter  $c$ .

adsorption sites of  $Mg^{2+}$  in the three 42 configurations are taken into account. For the AA-stacking type, the two  $Ti_3C_2O_2$  layers form  $H_1$ ,  $H_{12}$  and T active sites for  $Mg^{2+}$ , where the “ $H_{12}$ ” site means the combination of the  $H_1$  and  $H_2$  sites from two adjacent  $Ti_3C_2O_2$  layers, respectively.  $ABC_1$ - $Ti_3C_2O_2$  forms  $H_1$ ,  $H_2$  and  $H_{12}$  sites.  $ABC_2$ - $Ti_3C_2O_2$  forms  $TH_1$ ,  $H_1$  and  $H_2$  sites. For all these three stacking configurations, the  $H_1$  site exhibited stronger binding energy than the other sites (Fig. 3), which is consistent with the  $Mg^{2+}$  adsorption behavior for  $Ti_3C_2O_2$  monolayer. Among the three configurations, the  $H_1$  site in AA-stacked structure exhibited the most negative adsorption energy, which indicated that the AA-stacking is more favorable adsorptive configuration for  $Mg^{2+}$ . According to the optimized crystal, the lower the binding energy, the smaller the lattice parameter  $c$ . This suggests that the strong coulomb attraction between the O atoms and the adsorbed  $Mg^{2+}$  can mitigate volume expansion during ion storage. Moreover, it is evident that the structural energy difference between the  $ABC_1$ - and  $ABC_2$ - $Ti_3C_2O_2$  is very small (Fig. 2c and 4). Furthermore, during structural optimized,  $Mg^{2+}$  adsorbed  $ABC_2$ - $Ti_3C_2O_2$  will relax into  $ABC_1$  type through the sliding of adjacent  $Ti_3C_2O_2$  layers. Therefore, the  $ABC_2$  configuration is improbable to form under practical conditions and will hence not be considered in subsequent studies.

#### $Mg^{2+}$ diffuses in AA, $ABC_1$ and monolayer $Ti_3C_2O_2$

The rate performance of anode is determined by the kinetics of electronic and ionic diffusion. To further study the  $Mg^{2+}$

diffusion kinetic properties for the AA- and  $ABC_1$ -stacking structures, the  $Mg^{2+}$  diffusion barrier is evaluated in the  $3 \times 3 \times 1$   $Ti_3C_2O_2$  bulk/monolayer using the CI-NEB method. In each configuration,  $Mg^{2+}$  is simulated to migrate between two energetically favorable active sites (Fig. 4a). On  $Ti_3C_2O_2$  monolayer, the diffusion of  $Mg^{2+}$  between two adjacent  $H_1$  sites across  $H_2$  site (described as  $H_1 \rightarrow H_2 \rightarrow H_1$ ) has a relatively large barrier of 0.81 eV (Fig. 4d). Such a value is close to the  $Mg^{2+}$  barrier (0.72 eV) on the  $Ti_2NO_2$  monolayer reported by Wang *et al.*<sup>28</sup> According to that work, the univalent  $Li^+$ ,  $Na^+$ ,  $K^+$  diffusion barriers are lower than 0.3 eV, which suggests that the pathway of  $Ti_2NO_2$  monolayer is insufficient to provide an adequate coordination environment for the bivalent  $Mg^{2+}$ .

In the AA-stacking  $Ti_3C_2O_2$ , the  $Mg^{2+}$  barrier is significantly improved to 0.32 eV. The optimized AA-stacking structure shows that the adjacent  $H_{12}$  active sites provided by the upper and lower  $Ti_3C_2O_2$  surfaces are opposite, which can be described as “ $H_2 \rightarrow H_1 \rightarrow H_2$ ” (Fig. 4b). Notably, compared with the  $Ti_3C_2O_2$  monolayer,  $Mg^{2+}$  can only move from  $H_2 \rightarrow H_1$  through a single step, and can thereby transfer from one  $H_1$  active site to another. It can be seen that the periodic migration path of  $Mg^{2+}$  has been shortened by almost half. Besides, according to the CI-NEB method, the ionic migration barrier is obtained by subtracting the energy of the stable state (SS) from that of the transition state (TS), *i.e.*,  $E_{\text{barrier}} = E_{\text{TS}} - E_{\text{SS}}$ .<sup>43</sup> The contribution of metastable  $H_2$  site in AA- $Ti_3C_2O_2$  increases the  $E_{\text{SS}}$ , alleviating energy changes during  $Mg^{2+}$



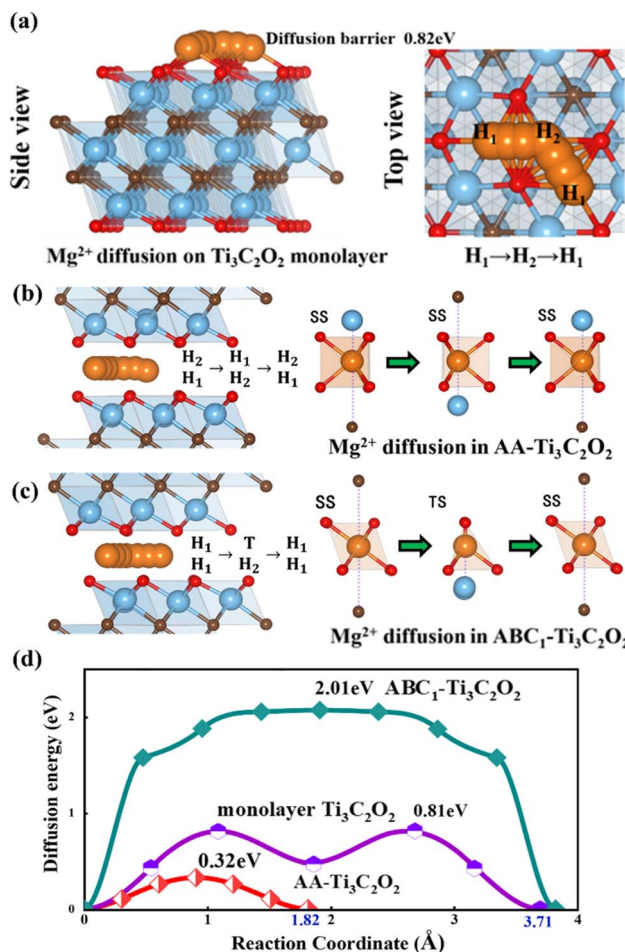


Fig. 4 (a) Diffusion path of magnesium on single layer  $\text{Ti}_3\text{C}_2\text{O}_2$  surface. (b) Diffusion paths of  $\text{Mg}^{2+}$  in AA type and (c)  $\text{ABC}_1$  type  $\text{Ti}_3\text{C}_2\text{O}_2$  structure, respectively. (d) Diffusion energy barrier of  $\text{Mg}^{2+}$  in AA type,  $\text{ABC}_1$  type and single layer  $\text{Ti}_3\text{C}_2\text{O}_2$ .

migration. In comparison, the  $\text{Mg}^{2+}$  diffusion channel in the  $\text{ABC}_1\text{-Ti}_3\text{C}_2\text{O}_2$  can be described as “ $\text{H}_1 \xrightarrow{\text{T}} \text{H}_2 \xrightarrow{\text{H}_1} \text{H}_1$ ” (Fig. 4c). The stable active site composed of two stable  $\text{H}_1$  sites lead to a low  $E_{\text{SS}}$  value. While the metastable  $\text{H}_2$  and exceptionally unstable  $\text{T}$  site significantly increases the  $E_{\text{TS}}$ . Such substantial energy disparity between  $E_{\text{TS}}$  and  $E_{\text{SS}}$  lead to a large diffusion barrier of 2.01 eV for  $\text{ABC}_1\text{-Ti}_3\text{C}_2\text{O}_2$ . To further investigate the influencing factors of  $\text{Mg}^{2+}$  diffusion barrier, the Bader charge of  $\text{Mg}^{2+}$  in stable states ( $e_s$ ) and transition states ( $e_t$ ) as well as the charge difference ( $\Delta e = |e_s - e_t|$ ) between them were calculated based on the CI-NEB image. The result showed that the  $\Delta e$  for AA and  $\text{ABC}_1$  types are 0.005 and 0.206e, which are consistent with their barrier order of 0.32, and 2.01 eV, respectively. The  $\text{Mg}^{2+}$  channel in  $\text{ABC}_1\text{-Ti}_3\text{C}_2\text{O}_2$  has large fluctuation on electron capture, which is also an important factors that cause the largest diffusion barrier.

### Diffusion of $\text{Mg}^{2+}$ in AA- $\text{Ti}_3\text{C}_2\text{T}_x$ ( $\text{T} = \text{O}, \text{N}$ )

According to the above discussion, the adsorption and migration behavior of  $\text{Mg}^{2+}$  exhibits a pronounced preference for the

AA- $\text{Ti}_3\text{C}_2\text{O}_2$ . Regrettably, in layered stacking structures, the coulombic effect between layers significantly influence the stacking configuration. For the AA- $\text{Ti}_3\text{C}_2\text{O}_2$ , the O-O electrostatic repulsion from the adjacent layers leads to a preference for staggered arrangement in  $\text{Ti}_3\text{C}_2\text{O}_2$  bulk, thereby increasing the interlayer spacing and rendering the AA- $\text{Ti}_3\text{C}_2\text{O}_2$  unstable. One effective approach to mitigate electrostatic repulsion is to substitute bivalent O with monovalent halogens such as F or Cl. However, the introduction of F/Cl functional groups does reduce the energy difference between the AA- and  $\text{ABC}_1\text{-Ti}_3\text{C}_2\text{O}_2$ , yet the AA configuration remains relatively unstable (Fig. 2c). The remaining stacking control method is to utilize higher valence N as functional group, thereby converting electrostatic repulsion into covalent bonding through the addition of electrons.

Consequently, the O function group in AA,  $\text{ABC}_1$  and  $\text{ABC}_2$  configurations are replaced by N (Fig. 5a). The calculated relative energy revealed that the AA-type  $\text{Ti}_3\text{C}_2\text{N}_2$  become the most stable configuration. Taking AA- $\text{Ti}_3\text{C}_2\text{N}_2$  as the reference structure, the relative energies of  $\text{ABC}_1$  and  $\text{ABC}_2$  are 6.82 eV and 7.89 eV, respectively (Fig. 5b). Such large values indicate that the introduction of interlayer N-N covalent bonding significantly enhances the stability of the AA-stacking configuration. In the AA- $\text{Ti}_3\text{C}_2\text{N}_2$  system, the diffusion energy barrier of  $\text{Mg}^{2+}$  calculated using the CI-NEB method is 1.07 eV, which is 0.94 eV lower than that in the stable  $\text{ABC}_1$  configuration for  $\text{Ti}_3\text{C}_2\text{O}_2$ . Evidently, the electrostatic interaction between layers does not directly influence the migration of  $\text{Mg}^{2+}$ . Instead, it indirectly facilitates a favorable migration environment for  $\text{Mg}^{2+}$  by inducing structural changes. Despite this reduction, the barrier of 1.07 eV still represents a poor diffusion performance. This can be attributed to the strong bonding between nitrogen atoms in adjacent  $\text{Ti}_3\text{C}_2\text{N}_2$  layers, which results in reduced layer spacing and hinders the migration of  $\text{Mg}^{2+}$ . Consequently, it is crucial to determine an appropriate N doping concentration to maintain optimal layer spacing while controlling the MXene stacking type.

The calculation of the relative energies for AA and  $\text{ABC}_1$  configurations with varying N contents reveals that when the N

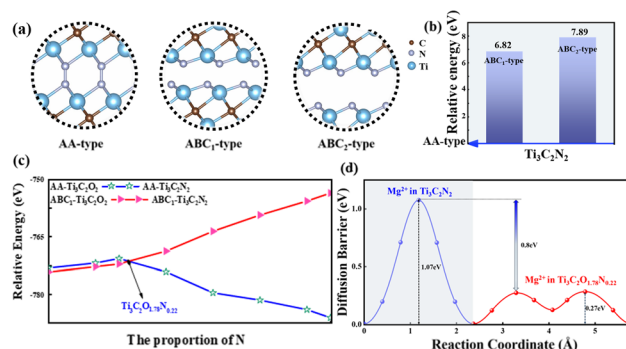


Fig. 5 (a) AA-,  $\text{ABC}_1$ - and  $\text{ABC}_2$ - $\text{Ti}_3\text{C}_2\text{N}_2$  interlayer structure. (b) The relative energy of  $\text{ABC}_1$ - and  $\text{ABC}_2$ - $\text{Ti}_3\text{C}_2\text{N}_2$  with respect to AA-type. (c) The relative energies of AA and  $\text{ABC}_1$  configurations with different N content. (d) Diffusion barrier comparison of  $\text{Mg}^{2+}$  in AA-stacking  $\text{Ti}_3\text{C}_2\text{N}_2$  and  $\text{Ti}_3\text{C}_2\text{N}_{1.78}\text{N}_{0.22}$  structures.

concentration reaches 11%, the stable phase MXene bulk transitions to the AA-stacking type, corresponding to the chemical formula  $Ti_3C_2O_{1.78}N_{0.22}$  (Fig. 5c). For this configuration, the  $Mg^{2+}$  diffusion barrier is decreased to 0.27 eV, which is significantly lower by 0.8 eV compared to  $Ti_3C_2N_2$  bulk. Notably,  $Mg^{2+}$  diffusion predominantly occurs near O functional groups rather than N functional groups (Fig. 5d). This can be attributed to the additional energy required for  $Mg^{2+}$  to overcome the interlayer N–N covalent bonds during diffusion, thereby increasing the migration energy barrier.

On the other hand, the strong interlayer N–N covalent bonding in the AA- $Ti_3C_2N_2$  structure is primarily responsible for the poor Mg diffusion kinetic property. Based on the such bonding situation, it can be proposed that enhancing the metal–N bond strength may weaken the interlayer N–N covalency, thereby improving ionic conductivity. In the periodic table, we have selected three transition metal elements, V, Zr and Nb, to substitute for Ti in AA- $Ti_3C_2N_2$ . V ( $3d^34s^2$ ), Nb ( $4d^45s^1$ ), and Zr ( $4d^25s^2$ ) each enhance N bonding through distinct mechanisms compared to Ti ( $3d^24s^2$ ). V and Nb provide additional valence electrons for covalent interactions, while Zr achieves stronger binding *via* reduced electronegativity-driven ionic attraction. After replacing Ti atoms in AA-stacking  $Ti_3C_2N_2$  with V, Zr and Nb, the dynamical stability of these four conformations are checked through examining the phonon band structures (Fig. 6a–d), which indicated that they are dynamically stable configurations with no states associated with imaginary frequencies.

#### Diffusion of $Mg^{2+}$ in AA- $M_3C_2N_2$ ( $M = Nb, V, Zr$ )

Subsequently, to analyze the diffusion behavior of  $Mg^{2+}$  in AA- $M_3C_2N_2$  ( $M = V, Zr$  and Nb), the  $Mg^{2+}$  migration pathways and corresponding energy barriers are investigated within  $3 \times 3 \times 1$  supercell. With a single  $Mg^{2+}$  intercalation into  $M_3C_2N_2$ , it was relaxed to the  $H_2$  site after structure optimization and the corresponding half of diffusion pathways were described as  $H_2 \rightarrow H_1$ , which showed that  $Mg^{2+}$  diffusion behavior in  $H_1 \rightarrow H_2$ , is consistent. The diffusion barriers of  $Mg^{2+}$  in  $V_3C_2N_2$ ,  $Zr_3C_2N_2$  and  $Nb_3C_2N_2$  had decreased compared with in  $Ti_3C_2N_2$  by 0.37 eV, 0.79 eV and 0.84 eV, respectively. Notably,  $Nb_3C_2N_2$  exhibits the smallest barrier of 0.23 eV. This indicates that N–N covalent binding is weakened more effectively by replacing Ti with Nb than with V and Zr (Fig. 6e). Overall, the  $Nb_3C_2N_2$  obtained by using the transition metal element Nb to replace Ti in AA- $Ti_3C_2N_2$  has the lowest energy barrier (Fig. S1†).

#### Theoretical voltage value of AA-stacking $Nb_3C_2N_2$

According to the above discussion, the favorite absorption site of  $Mg^{2+}$  is  $H_2$  site. Based on this, the formation energy of  $Nb_3C_2N_2Mg_x$  ( $0 \leq x \leq 1$ ) was calculated *via* the following equation to explore the thermodynamically stable Mg-containing phases,

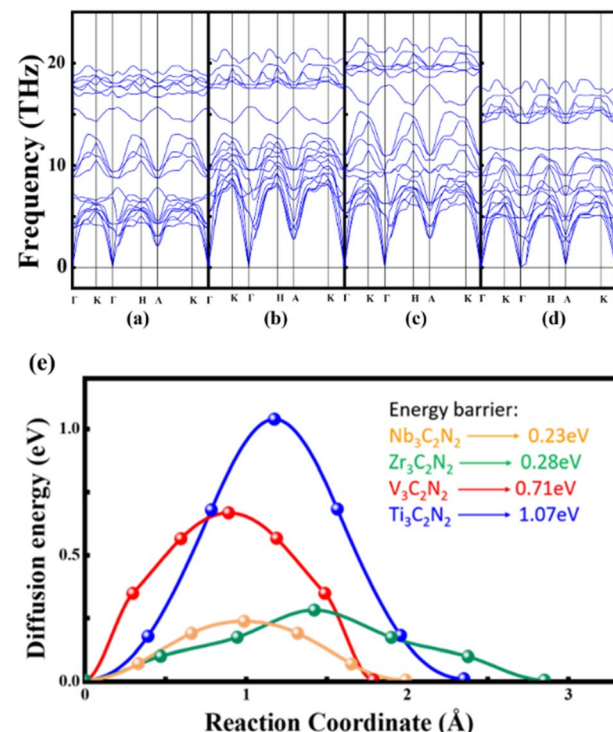


Fig. 6 Phonon spectra of (a)  $Nb_3C_2N_2$ , (b)  $Ti_3C_2N_2$ , (c)  $V_3C_2N_2$  and (d)  $Zr_3C_2N_2$ . (e) Diffusion energy barrier of  $Mg^{2+}$  in AA- $M_3C_2N_2$  ( $M = Ti, V, Zr, Nb$ ).

$$E_f = E_{Nb_3C_2N_2Mg_x} - \left(1 - \frac{x}{2}\right)E_{Nb_3C_2N_2} - \frac{x}{2}E_{Nb_3C_2N_2Mg_2} \quad (2)$$

where  $E_{Nb_3C_2N_2Mg_x}$ ,  $E_{Nb_3C_2N_2Mg_2}$  and  $E_{Nb_3C_2N_2}$  are total energies for  $Nb_3C_2N_2Mg_x$ ,  $Nb_3C_2N_2Mg_2$  and  $Nb_3C_2N_2$ , correspondingly,  $x$  is the  $Mg^{2+}$  adsorption degrees in  $Nb_3C_2N_2Mg_x$ , including 0, 0.25, 0.5, 0.75, 1, 1.25, 1.5, 1.75 and 2 (Fig. 7a). Our result suggests that the thermodynamically stable configurations are  $Nb_3C_2N_2Mg_{0.75}$ ,  $Nb_3C_2N_2Mg$  and  $Nb_3C_2N_2Mg_2$ , with their  $E_f$  points located at the bottom edge of the convex hull. Based on this, the open circuit voltage (OCV) of AA-stacking  $Nb_3C_2N_2$  with different  $Mg^{2+}$  adsorption degrees is calculated from the following formula,

$$V = \frac{1}{n} \left| \frac{2(E_{Nb_3C_2N_2Mg_{x_2}} - E_{Nb_3C_2N_2Mg_{x_1}})}{(x_2 - x_1)} - E_{Mg} \right| \quad (3)$$

where  $E_{Nb_3C_2N_2Mg_{x_2}}$  and  $E_{Nb_3C_2N_2Mg_{x_1}}$  are the total energies of  $Nb_3C_2N_2$  with different  $Mg^{2+}$  adsorption degrees  $x_1$  and  $x_2$ ,  $E_{Mg}$  is the energy of a Mg atom in metal Mg, charge value  $n = 2$  for  $Mg^{2+}$ . Interestingly, the calculated voltages contain two distinct plateaus at 4.00 V and 0.64 V, corresponding to the ranges of  $x = 0\text{--}0.75$  and  $x = 1\text{--}2$ , respectively (Fig. 7b). This results indicate that when a monolayer of  $Mg^{2+}$  is intercalated between the  $Nb_3C_2N_2$  layers, the binding affinity of  $Mg^{2+}$  to the  $Nb_3C_2N_2$  layers significantly decreases. Such a significant voltage difference indicates that the material can function as both the positive and negative electrode of the battery under varying ion concentrations. This behavior is analogous to that observed in



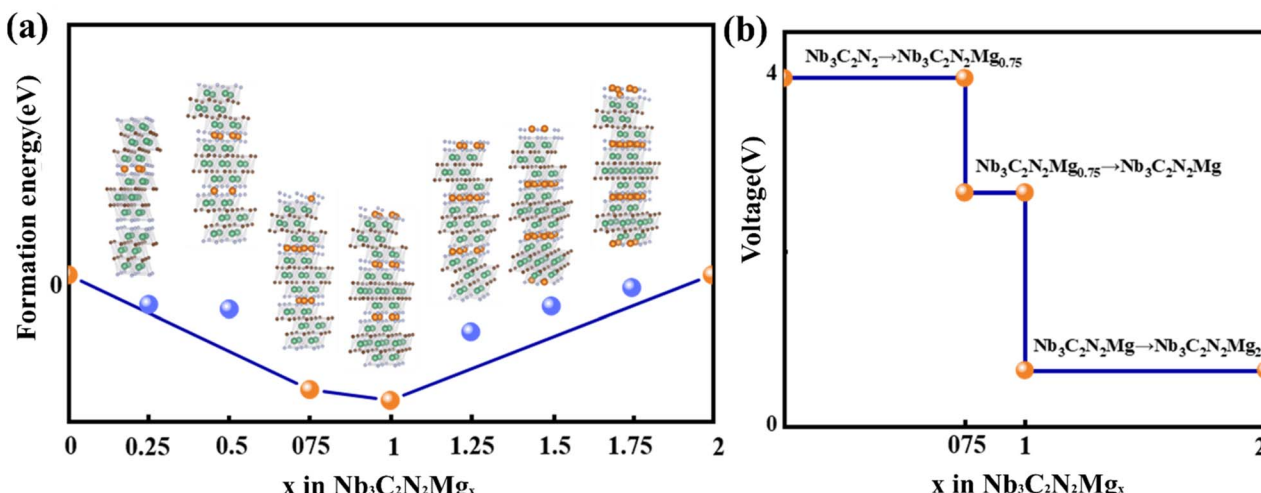


Fig. 7 (a) Formation energy and (b) theoretical voltage value of  $\text{Nb}_3\text{C}_2\text{N}_2\text{Mg}_x$ .

Prussian blue materials used in potassium-ion batteries.<sup>44</sup> By contrast, the voltage platforms presented by Ti-based MXene belong to the anode range (Fig. S2†). From a voltage stability perspective,  $\text{Ti}_3\text{C}_2\text{O}_2\text{Mg}_x$  and  $\text{Ti}_3\text{C}_2\text{O}_{1.78}\text{N}_{0.22}\text{Mg}_x$  exhibit three short platforms decreasing in the concentration of  $0 < x < 1$ . Although  $\text{Ti}_3\text{C}_2\text{N}_2$  possesses two stable platforms, the voltage of 0.02 V is too low for electrode, which is easy to transform the  $\text{Mg}^{2+}$  intercalation mechanism into electroplating.

## Conclusions

This study systematically investigates the  $\text{Mg}^{2+}$  storage and diffusion mechanisms in MXene materials, focusing on  $\text{Ti}_3\text{C}_2\text{O}_2$  and its nitrogen-doped derivatives. The strong electrostatic interaction between divalent  $\text{Mg}^{2+}$  and electrode frameworks leads to high diffusion barriers (0.81 eV in monolayers), which are mitigated by optimizing MXene stacking configurations and functional group engineering. AA-stacking  $\text{Ti}_3\text{C}_2\text{O}_2$  exhibits strong  $\text{Mg}^{2+}$  adsorption energy of  $-2.16$  eV and low diffusion barrier of 0.32 eV due to staggered active sites shortening migration paths. However, its instability from interlayer O–O repulsion necessitates structural modifications. Replacing O with N transforms stable  $\text{ABC}_1\text{-Ti}_3\text{C}_2\text{O}_2$  to AA- $\text{Ti}_3\text{C}_2\text{N}_2$  through enhanced interlayer N–N covalent bonding. However, excessive N doping increases the energy barrier (1.07 eV) due to decreased layer spacing. One effective approach is to control the N/O ratio, which balances interlayer spacing and covalent interactions, resulting in an energy barrier of 0.27 eV in the  $\text{Ti}_3\text{C}_2\text{O}_{1.78}\text{N}_{0.22}$  structure. Another strategy involves substituting transition metals into the MXene framework to modulate interlayer bonding. Introducing transition metals (V, Zr, Nb) into  $\text{Ti}_3\text{C}_2\text{N}_2$  weakens N–N covalency, with  $\text{Nb}_3\text{C}_2\text{N}_2$  achieving an ultralow barrier of 0.23 eV.  $\text{Nb}_3\text{C}_2\text{N}_2$  possesses dual voltage platforms (4.00 V and 0.64 V), enabling its dual role as cathode/anode, while Ti-based MXenes exhibit anode-specific behavior. These findings highlight that tailored MXene structures, through stacking control, nitrogen doping, and metal substitution, significantly enhance  $\text{Mg}^{2+}$  kinetics and stability,

positioning them as promising high-performance electrodes for next-generation Mg-ion batteries.

## Data availability

The datasets used and/or analyzed during the current study are available from the corresponding author upon reasonable request.

## Conflicts of interest

There are no conflicts to declare.

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