Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: *J. Mater. Chem. A*, 2025, **13**, 24038

Computational discovery of a novel double transition metal nitride MXene and its applications as an anchoring and catalytic material in Li-Se batteries†

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Double transition metal (DTM) nitride MXenes offer enhanced electrical conductivity and tunable catalytic properties compared to conventional carbide-based MXenes. In this study, we employed first-principles density functional theory (DFT) calculations to discover and computationally validate a novel DTM nitride MXene, Nb₂TiN₂, derived from its MAX phase precursor and investigated its potential as an anchoring material (AM) for Li-Se batteries. This newly proposed MXene expands the compositional landscape of DTM nitrides and opens new avenues for functional material design. We performed a comprehensive analysis of the thermodynamic and electronic properties of Nb2TiAlN2, and the MAX phase precursor to Nb₂TiN₂ to assess its structural stability and exfoliation potential. Exfoliation energy calculations confirmed the feasibility of synthesizing Nb2TiN2 from Nb2TiAlN2. We then explored the functionalized form, Nb₂TiN₂S₂, evaluating its capability to serve as an effective anchoring material (AM) in Li-Se batteries by analyzing the reaction mechanisms and kinetics of the selenium reduction reaction (SeRR). Our results indicate that Nb₂TiN₂S₂ exhibits a strong binding affinity for lithium polyselenides (Li₂Se_n), effectively suppressing the shuttle effect. Gibbs free energy calculations for the rate-limiting step of the SeRR reveal favorable kinetics and reduced reaction barriers. Overall, this study provides a detailed evaluation of the structural and electronic properties of a newly proposed DTM nitride MXene and its Sfunctionalized derivative and the catalyzing effect of Nb2TiN2S2 in accelerating the reaction kinetics in Li-Se batteries. These findings underscore the potential importance of the further exploration of MXenes to address current challenges in high-performance Li-Se batteries.

Received 20th April 2025 Accepted 22nd June 2025

DOI: 10.1039/d5ta03099d

rsc.li/materials-a

1. Introduction

Over the past decade, graphene has grabbed global attention as a groundbreaking two-dimensional (2D) material, demonstrating remarkable potential across diverse fields such as energy storage, electronics, and catalysis.¹⁻⁴ This significant achievement has fueled a widespread interest in the search for alternative 2D materials that could further revolutionize these domains. Among these emerging materials, MXenes—a family of transition metal carbides, nitrides, and carbonitrides—have gained substantial recognition for their exceptional electrical

conductivity, hydrophilic nature, mechanical flexibility, and

MXenes originate from a broader class of layered ternary carbides, nitrides, and carbonitrides known as MAX phases.⁶ These materials follow the general formula $M_{n+1}AX_n$, where M represents a transition metal, A is a group 13 or 14 element, and X denotes carbon (C) or nitrogen (N). The unique structure of MAX phases allows them to be selectively etched to produce MXenes, which follow the formula $M_{n+1}X_n$. A key distinguishing feature of MXenes lies in their synthesis process, which involves

dispersion ability.^{5,6} Their high specific surface area and the rich presence of surface functional groups make them highly adaptable for a broad range of applications, including supercapacitors, catalysis, sensors, energy storage systems, and optoelectronics.⁷⁻¹¹ Today, MXenes stand as one of the most expansive and rapidly growing families of 2D materials, offering unmatched versatility and tunability for next-generation technological advancements. The unique combination of metallic conductivity, chemical stability, and functional surface chemistry has positioned MXenes as a promising class of functional materials.

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d5ta03099d

the selective removal of the A-group element, such as aluminum, from MAX phases. 12,13 This process is typically achieved using aggressive etching agents like hydrofluoric acid (HF) or ammonium bifluoride (NH4HF2), which effectively dissolve the A-layer while preserving the underlying $M_{n+1}X_n$ layers.¹⁴ As a result, the final structure of MXenes can be represented by the formula $M_{n+1}X_nT_x$, where T_x denotes surface terminations (such as -OH, -F, or -O groups) that form during the etching process. The composition and surface chemistry of MXenes play a crucial role in determining their physical, chemical, and electronic properties. 5,15 The ability to tune surface terminations and modify the structural composition opens up new opportunities for MXenes, particularly in electrochemical energy storage, catalysis, sensing, and electronic devices.

MXenes can be broadly categorized into two primary types based on their composition: carbide MXenes (c-MXenes) and nitride MXenes (n-MXenes).16 Among the broader family of MXenes, n-MXenes have attracted significant attention due to their superior electrical conductivity, enhanced mechanical strength, and improved catalytic performance compared to conventional c-MXenes.17-19 Previous studies have demonstrated that when comparing c-MXenes and n-MXenes with the same metal composition, n-MXenes exhibit superior electrical conductivity and enhanced capacitive properties. For example, Zhang et al.17 reported that n-MXenes exhibit higher in-plane Young's moduli and electrical conductivity compared to their c-MXene counterparts. This enhancement was attributed to the smaller lattice constants and reduced monolayer thickness of n-MXenes, which contribute to their superior mechanical and electronic properties. Another study indicated that substituting carbon with nitrogen in the same MXene composition results in a lower point of zero charge, which enhances the material's metallic characteristics and increases its capacitance. 19 Molecular dynamics (MD) simulations further reinforced these findings, revealing that the Young's modulus of n-MXenes is significantly higher than that of c-MXenes.18 This enhanced mechanical strength, combined with their exceptional electrical conductivity and distinctive surface chemistry, enables n-MXenes to be utilized in a broad range of applications.20 Beyond mechanical and electrical performance, n-MXenes have also been explored for their magnetic properties. Studies on Mn₂NT_x suggest that 2D n-MXenes hold great promise for spintronic applications, making them potential candidates for next-generation magnetic devices.21 Additionally, Ti₄N₃T_x has been identified as a highly efficient electrocatalyst for the hydrogen evolution reaction (HER) as it exhibits a unique combination of metallic and semiconducting properties that result in enhanced catalytic activity.22 By employing density functional theory (DFT) calculations, Fan et al.23 demonstrated 2D V₂N/V₂NT₂ monolayers functionalized with S as a potential S cathode host for Li-S batteries.

Recently, double transition metal MXenes (DTM-MXenes) have surfaced as another special class of 2D materials in the realms of advanced composites, energy storage, and catalysis, complementing the mono-metallic MXene counterparts.²⁴⁻²⁶ This focus can be attributed to their unique structure, which integrates two distinct transition metals instead of one. DTM-

MXene has a formula of $M'_{n+1}M''_{n+1}X_n$, where M' and M" are two different transition metals. A prior study has revealed that compared to single metal MXene Ti₄C₃, DTM-MXene Ti₂Ta₂C₃ exhibits higher open circuit voltage and capacity to store lithium(Li) than Ti₄C₃.²⁷ The mechanical properties of DTM-MXenes have also been investigated, with Mo₂TiC₂ showing an exceptionally high Young's modulus of 361 GPa, significantly surpassing that of Ti₃C₂ MXene.²⁸ This enhanced mechanical strength reinforces their potential use in structural applications where durability and flexibility are crucial. Liu et al. studied the electrochemical characteristics of Ti₂NbC₂T_r as an anode material in Li-ion batteries.29 They reported a superior specific capacity and long cycling stability with Ti₂NbC₂T_x, which outperformed its single metal counterparts, Ti₃C₂T_x. Similar behavior was also noted for the Ti_rTa_{4-r}C₃ MXene. ²⁶ All of these studies underscore the potential of DTM-MXene in various applications. Despite these advancements, most research on DTM-MXenes has primarily focused on c-MXene systems, with little to no studies investigating DTM n-MXenes. It can be hypothesized that DTM-n-MXene may exhibit superior performance by utilizing the characteristics of both n-MXene and DTM-MXene.

The lithium selenium (Li-Se) batteries have recently emerged as an alternative to Li-S batteries.30,31 Se has a high theoretical gravimetric capacity (678 mA h g^{-1}) and volumetric capacity (3268 mA h cm⁻³). 32-35 Most significantly, Se has higher electrical conductivity $1 \times 10^{-3} \text{ S m}^{-1}$ (about 20 orders of magnitude than S), which can result in increased capacity utilization and rate capability.36 Unfortunately, the dissolution of soluble intermediate lithium polyselenides (Li₂Se_n) into the ether-based electrolyte during the electrochemical cycling results in a low battery capacity and inhibits the cycle performance of Li-Se batteries.37 Additional constraints like sluggish reaction kinetics and irreversibility also hinder their practical realization. To overcome these issues, for Li-Se systems, several strategies have been reported, such as confinement of Se in porous carbon, adsorption of polyselenides by metal oxide, insertion of carbon interlayer to adsorb soluble polyselenides, and design of nanoporous or nanofibrous Se structure. 38-40 To achieve the desired performance in Li-Se batteries, the use of CoSe₂-porous carbon composites, 41 graphene-selenium hybrid microballs,42 and various metal organic frameworks (MOF) are found to mitigate, but not completely eliminate the dissolution of polyselenides. It has been observed that nitrogen-doped carbon structures, due to their porous character, facilitate an ion transport channel and better restrict Se inside the host. 41,43 Han et al. demonstrated that the electrical and ionic conductivities can be enhanced, and the dissolution of polyselenides can be suppressed by probing the hierarchical structure created by the fusion of graphene and 3D porous carbon nanoparticles.44 The synthesized Fe-CNT/TiO2 catalyst through hightemperature sintering of a carbon nanotube-decorated MOF and MXene nanoarchitecture enhances Se hosting and demonstrates exceptional battery performance with high capacity retention and efficiency across various cycling rates. 45 Se incorporation into the microporous carbon nanofibers via vacuum and heat treatment enhances the anchoring behavior

and provides uniform distribution of the selenides on the carbon matrix.⁴⁰ Despite extensive research on Li–Se batteries, challenges such as sluggish electrode kinetics and polyselenide shuttle effects remain unresolved. This highlights the need for further research into alternative anchoring materials (AMs) for Li–Se batteries to improve the polyselenides adsorption capacity and enhance the reaction kinetics.

This study focuses on the development of a novel DTM-n-MXene, Nb₂TiN₂, from its precursor MAX phase. To achieve this, we conducted a comprehensive computational study using first-principles DFT simulations to analyze the stability and electronic characteristics of MAX phases and the formation of Nb₂TiN₂ MXene. Our study began with an in-depth investigation of the stability of the Nb2TiAlN2 MAX phases, which was further extended to the evaluation of the exfoliation process by calculating the exfoliation energy. To assess the practical energy storage potential of the newly developed Nb₂TiN₂ MXene, we investigated its adsorption behavior and reaction kinetics in Li-Se batteries. Specifically, we studied sulfur-functionalized Nb₂-TiN₂S₂, evaluating its ability to stabilize Li₂Se_n and facilitate Se reduction reactions (SeRR). The adsorption mechanisms of Li₂Se_n on Nb₂TiN₂S₂ and the kinetics of SeRR are all investigated using DFT simulations.

Calculation methodology

Spin polarized plane wave DFT calculations were carried out using the Vienna ab initio Simulation Package (VASP).46 The exchange-correlation energy was described using the Perdew-Burke-Ernzerhof (PBE) functional within the framework of the generalized gradient approximation (GGA).47 The PBE functional was selected for this study because of its demonstrated ability to accurately capture the electronic structure of MXenes and their interactions with lithium polychalcogenides in Lichalcogen batteries. 48-52 In addition, PBE offers a practical advantage with its significantly lower computational cost compared to hybrid functionals such as HSE06, which makes it well-suited for exploring larger system sizes and multiple adsorption configurations. The Projector Augmented Wave (PAW) method was employed to explicitly treat the interaction between valence and core electrons, enabling an accurate representation of the electronic structures by including the effect of core electrons on valence electron density. The planewave energy cutoff of 520 eV is used. The electronic structure minimization was performed using the blocked Davidson iteration scheme,46 and van der Waals (vdW) interactions were accounted for using the empirical DFT-D3 dispersion correction.53 We used the conjugate gradient method to optimize the geometry while letting the atoms relax until the residual force was less than 0.05 eV \mathring{A}^{-1} and the energy convergence reached 1 \times 10⁻⁴ eV. In order to sample the Brillouin zone integration for the electronic structure computations and atomic relaxations, the Monkhorst-Pack grid scheme with $8 \times 8 \times 1$ k-point was used for the calculations related to MAX phase and MXene unit cell and $5 \times 5 \times 1$ k-point was used for the Li₂Se_n adsorption calculations on the MXene substrate. To obtain the stable configuration of the MAX phase, the structure was fully relaxed,

allowing for any adjustments in both shape and volume until the forces on atoms and the stress tensor components were minimized. The integration of the Brillouin zone was carried out using the tetrahedron method with Bloch corrections. Bader charge analysis was employed to quantify the charge population on the polyselenides and MXenes. We calculated the charge density difference based on the expression $\rho_{\rm b} = \rho_{\rm absorbed\ state} - (\rho_{\rm absorbate} + \rho_{\rm AM})$, where $\rho_{\rm absorbed\ state}$, $\rho_{\rm absorbate}$, and $\rho_{\rm AM}$ refer to the charge transfer of Li₂Se_n adsorbed AM, the isolated polyselenides, and the AM, respectively.

3. Results and discussion

3.1 Stability analysis of Nb₂TiAlN₂ MAX phase

DFT simulations were employed to investigate the structural and thermodynamic stability of the Nb₂TiAlN₂ MAX phase. The study began with the structural optimization of the MAX phase, starting from its initial configuration obtained from the Materials Project database.54 The optimized structure, depicted in Fig. 1a, represents the energetically stable configuration of the Nb₂TiAlN₂ MAX phase. This converged structure served as the basis for subsequent analyses, including assessments of thermodynamic stability, electronic structure, and mechanical properties. We used the Open Quantum Materials Database (OQMD)55 to identify the competing phases into which Nb2-TiAlN₂ can decompose. OQMD contains DFT-calculated thermodynamic and structural properties for over 1317811 structures. The Nb₂TiAlN₂ falls within the Nb-Ti-Al-N region of phase space. Our analysis revealed that Nb2TiAlN2 can decompose into AlN, TiN, Nb₂N, and Nb₃Al₂N, as shown in Fig. 1b. Although these phases do not represent the complete set of competing phases in the Nb-Ti-Al-N quaternary system, they collectively represent the lowest-energy ground state configuration that accurately captures the stoichiometric composition of Nb₂TiAlN₂. The isometric view of the Gibbs tetrahedron of the 4-D (Nb-Ti-Al-N) phase diagram generated via the OQMD database is shown in Fig. S1.† The phase diagram shows the composition of stable and unstable phases in the phase space. To evaluate the thermodynamic stability, the formation energy of the Nb₂TiAlN₂ MAX phase was calculated and compared with those of competing phases in the Nb-Ti-Al-N system. We found that MAX phase possesses low formation energy (-114.90 eV)compared to the total formation energy of its competing phases (-109.90 eV), which suggests that the MAX phase is energetically more favorable and thus likely to be the stable phase under standard conditions. Additionally, phonon dispersion calculations were performed to assess the dynamic stability of the MAX phases. The calculation was carried out using Density Functional Perturbation Theory (DFPT) and analyzed with Phonopy.56 The absence of imaginary frequencies, illustrated in Fig. 1c, confirms the structural stability of the phase. To further investigate the material's stability, ab initio molecular dynamics (AIMD) simulations were conducted at elevated temperatures.

The AIMD simulations were conducted using the canonical ensemble (NVT), where the system was maintained at a constant volume and temperature. The Nose–Hoover thermostat was applied to regulate the temperature at 1000 K. The AIMD

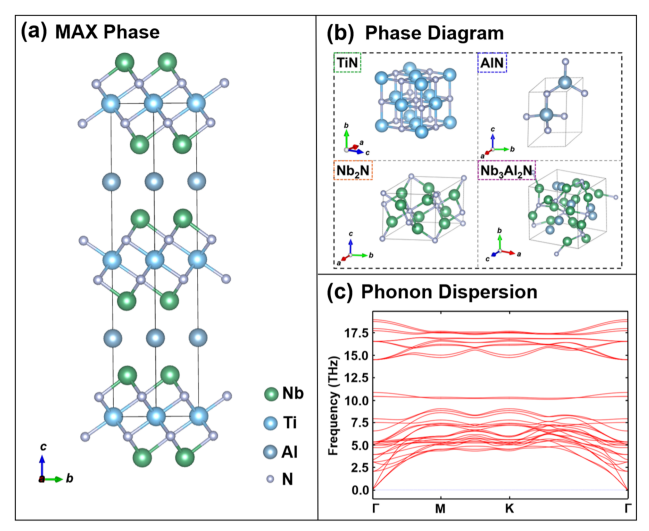


Fig. 1 The stable structures of (a) the Nb₂TiAlN₂ MAX phase; and (b) competing phases. (c) The phonon dispersion curve of the Nb₂TiAlN₂ MAX phase.

simulation was run for 10 picoseconds (ps) with a time step of 1 femtosecond (fs). Fig. S2† presents the distribution of total energy and temperature over the entire simulation duration. The results indicate that both energy and temperature exhibit minimal fluctuations, suggesting that the atomic structure remains stable under high-temperature conditions. Additionally, the initial and final structures of Nb2TiAlN2 in AIMD simulation, shown in Fig. S3,† clearly depicts that the structure remains stable at high temperature even after 10 ps. The absence of significant energy deviations or structural distortions reinforces the high thermodynamic stability of Nb₂TiAlN₂ at elevated temperatures. All of these findings indicate that Nb₂TiAlN₂ MAX phase demonstrates favorable thermodynamic and dynamic stabilities.

3.2 Electronic conductivity of Nb₂TiAlN₂ MAX phase

MXene's metallic nature essentially originates from the dlocalized electrons of the transition metal atoms. Previous studies showed that the metallic conductivity of MXenes and their functionalized forms allow electron diffusion between the

MXene surface and the lithium polysulfides during the redox reactions to achieve high capacity in the Li-S batteries. 50,57 This charge transfer also facilitates the chemical adsorption and catalytic conversion of higher-order polysulfides into lowerorder polysulfides on the MXene surface. As a result, the metallic nature not only promotes efficient redox kinetics but also improves the polysulfides immobilization within the cathode scaffolds to mitigate the shuttle effect through electronmediated interactions. 58 Li-Se batteries with Nb2TiN2S2 are also expected to exhibit similar trends. To gain a deeper understanding of the electronic characteristics of the Nb2TiAlN2 MAX phase, we performed density of states (DOS) analysis. Fig. 2 presents the calculated total DOS for the Nb₂TiAlN₂ MAX phase. Our results reveal a significant density of states at the Fermi energy level ($E_f = 0$ eV), which is a clear indicator of the metallic nature and high electrical conductivity of Nb₂TiAlN₂. To further analyze the electronic contributions from different atoms and orbitals of Nb₂TiAlN₂, we performed projected density of states (PDOS) analysis which offers a breakdown of the contribution of individual atomic orbitals to the overall electronic states of the

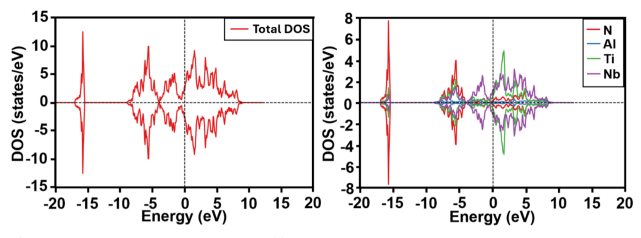


Fig. 2 Calculated band structure of the MAX phase. Total DOS is shown in the right plot whereas the contribution of individual orbital is presented in the left plot.

system. Our findings indicate that the d-orbitals of Nb and Ti play a dominant role in determining the electronic properties of Nb₂TiAlN₂. Both Nb-4d and Ti-3d states contribute significantly around the Fermi level. Additionally, the absence of an electronic band gap further confirms that Nb₂TiAlN₂ behaves as a metallic material rather than a semiconductor. The Al and N states contribute primarily to lower energy regions which indicates their minor role in direct conduction mechanisms but their potential influence on structural stability and bonding characteristics. The hybridization between the metal d-states and nonmetal p-states also suggests a complex bonding environment that supports both metallic and covalent interactions.

3.3 Exfoliation of Nb₂TiAlN₂ MAX phase

The exfoliation of MAX phases into their corresponding 2D MXenes involves the selective removal of Al layers through chemical etching, typically using hydrofluoric acid (HF) or other etching agents. The elimination of these Al layers disrupts the strong bonding network within the MAX phase and leads to the formation of weak van der Waals interactions between the remaining MXene layers. These weak interlayer interactions allow for the subsequent intercalation process, in which specific molecules or ions are introduced between the layers. The intercalants expand the interlayer spacing and facilitate the mechanical or chemical separation of individual MXene flakes and ultimately yield a 2D structure with enhanced flexibility and high surface area.⁵⁹

To investigate the feasibility of exfoliating the Nb_2TiAlN_2 MAX phase into its $2D\ Nb_2TiN_2$ MXene, we calculated the bond stiffness (k), which represents the resistance of a bond to deformation, and it is a crucial parameter for predicting the exfoliation behavior. It provides insight into which atomic bonds within the MAX phase are more susceptible to breaking during the exfoliation process. The calculated bond stiffness values for various atomic bonds in Nb_2TiAlN_2 are presented in Table 1. Our results suggest that the bond stiffness values for Nb–Al and Al–N bonds are significantly lower than those of Ti–N, N–N, and Nb–N bonds. This indicates that Al atoms are

weakly bonded to their neighboring atoms. The relatively strong bonding of Ti and Nb with nitrogen further supports the structural integrity of the remaining MXene layers after Al removal, allowing the formation of a stable 2D Nb₂TiAlN₂ MXene. To quantitatively assess the thermodynamic feasibility of exfoliation, we further calculated the exfoliation energy (E_{exf}) , which provides a measure of the energy required to separate the 2D MXene from the bulk MAX phase. The exfoliation energy was calculated using the equation $E_{\text{exf}} = (E_{\text{bulk MAX phase}} - E_{\text{2D-MXene}})$ $-E_{Al}/A$, where $E_{bulk\ MAX\ phase}$, $E_{2D\text{-}MXene}$, E_{Al} , and A denote the energy of bulk Nb₂TiAlN₂ MAX phase, the energy of 2D Nb₂TiN₂ MXene, the energy of Al atom in its bulk fcc phase and the surface area of Nb2TiN2 MXene flake, respectively. Our calculations yielded an E_{exf} value of approximately 0.20 eV Å⁻². Previous studies have reported that the critical exfoliation energy threshold is ~ 0.25 eV \mathring{A}^{-2} , with values below this threshold indicating a high probability of successful exfoliation. 60 Since our calculated exfoliation energy is well below this critical limit, this strongly suggests that the exfoliation of Nb₂-TiAlN₂ MAX phase into 2D Nb₂TiN₂ MXene is thermodynamically favorable.

3.4 Nb₂TiN₂ MXene and sulfur functionalization

The optimized geometry of the Nb₂TiN₂ MXene after the exfoliation from its MAX phase is illustrated in Fig. 3a and c. The Nb₂TiN₂ monolayer adopts a quint-layer configuration, where a central Ti atomic layer is symmetrically enclosed between two N layers, with Nb layers exposed on both the top and bottom surfaces. This structural arrangement follows the sequence of Nb-N-Ti-N-Nb. After the relaxation of the Nb₂TiN₂ monolayer, the optimized lattice parameters are found as a = b = 3.01 Å

 $\begin{tabular}{ll} \textbf{Table 1} & \textbf{Bond stiffness of different bonds present in Nb}_2 \textbf{TiAlN}_2 \ \textbf{MAX} \\ \textbf{phase} & \end{tabular}$

	Nb-Al	Ti-N	N-N	Nb-N	Al-N
Bond stiffness, k (N m ⁻¹)	609	1103	1111	1401	77

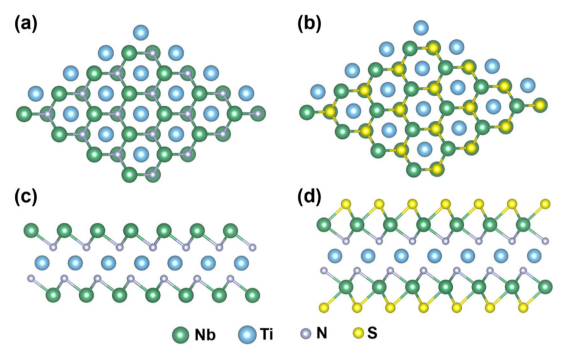


Fig. 3 Structure of optimized geometries of Nb₂TiN₂ MXene and its S-functionalized counterparts. (a–d) Represent the top view and the side view, respectively.

with a layer thickness of 5.08 Å. The bond lengths of Ti-N and Nb-N are observed as approximately 2.11 Å and 2.16 Å, respectively. The incorporation of functional groups provides additional knobs to tune the structural and electronic properties of Nb₂TiN₂ MXene. The introduction of functional groups such as -S, and -O can improve its hydrophilicity, conductivity, and interaction with other materials. 61,62 Additionally, controlled surface functionalization allows for tailoring the work function and charge transfer properties. Here, we incorporated S functionalization into the studied MXene and systematically evaluated its stability in comparison to the pristine phase. However, for functionalization, there are 3 possible adsorption sites (i) above the N atom (ii) above the Ti atom, and (iii) above the Nb atom. In our previous study, conducted for Mo₂TiC₂-MXene, we found that the C top site is energetically favorable and dynamically stable compared to the other two sites. 63 Following that, we applied the S-functionalization at all three sites, shown in Fig. S4,† and calculated their formation energies to determine the most favorable site for the functionalization. Interestingly, our result indicates that the N-top site of Nb₂TiN₂ MXene possesses relatively more negative formation energy (\sim -6.0 eV) compared to the Ti-top site(\sim -5.4 eV) and is the most stable configuration. However, it was found that the Nb-top site was structurally unstable for S-functionalization. These findings are in good agreement with our previous study.63 Fig. 3b and d depict the optimized configurations of the S-terminated Nb₂TiN₂ MXene. Notably, the bond length of Nb-S is measured at approximately 2.45 Å.

To gain deeper insights into the bonding characteristics and the effects of functionalization, we analyzed the electron localization function (ELF) for both the pristine and S-functionalized

Nb₂TiN₂ MXenes. ELF provides a quantitative measure of charge distribution within the material, ranging from 0 to 1, where ELF = 0 represents complete electron delocalization, characteristic of metallic or weakly bonded systems, while ELF = 1 corresponds to strong electron localization, typically observed in covalent or ionic bonding scenarios. Fig. 4a presents a comparative visualization of the charge density in Nb₂TiN₂ MXene before and after functionalization. In the pristine structure, a high electron density is observed around the N atoms, which highlights substantial charge transfer from Nb and Ti atoms to N. As a result, N plays a crucial role in governing the charge distribution, leading to strong Nb-N and Ti-N interactions. Upon S functionalization, notable changes in charge localization are observed. Due to increased charge transfer from the metal atoms, the S atoms exhibit an enriched electron cloud, and this charge transfer is stronger than the one observed on N in the pristine MXene. Due to its higher electronegativity and polarizability, S interacts more effectively with the underlying Nb and Ti atoms, forming bonds with a more pronounced covalent character. Additionally, the redistribution of charge density upon functionalization may influence the electronic properties, catalytic activity, and mechanical integrity, as electron-rich surface terminations can alter the material's work function and interaction with external species.

To assess the dynamic stability of pristine and Sfunctionalized Nb₂TiN₂ MXenes, we performed a phonon dispersion calculation, as presented in Fig. 4b. The phonon dispersion plot reveals that all vibrational modes exhibit positive frequencies across the Brillouin zone which confirms the dynamical stability of both the pristine and functionalized MXenes. Although a minor negative frequency is observed for

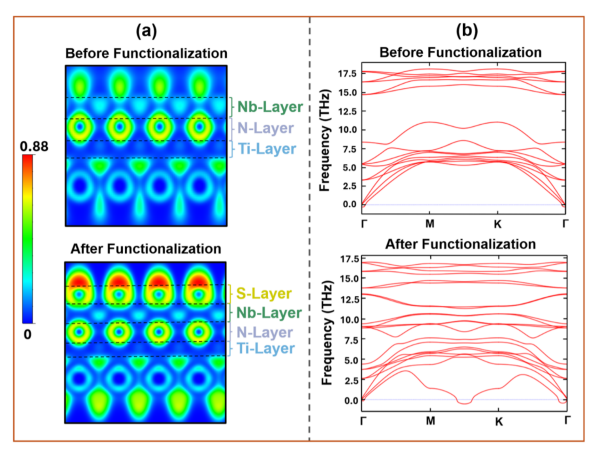


Fig. 4 (a) Electron localization function (ELF) and (b) phonon dispersion analysis of Nb₂TiN₂ MXene before functionalization and after functionalization.

functionalized case, it is insignificant and likely arises from numerical artifacts or computational approximations rather than indicating a true structural instability. We also performed AIMD calculations using an NVT ensemble with a Nose-Hoover thermostat at a temperature of 1000 K for 10 ps to evaluate the thermodynamic stability at high temperatures. The energy and temperature evolution with time shows insignificant fluctuations (Fig. S5 and S7†) and the identical initial and final structures were found to be highly stable for both pristine and Sfunctionalized MXene (Fig. S6 and S8†). The absence of substantial imaginary frequencies and rapid fluctuations of energy and temperature suggest that both materials maintain their structural integrity and thermodynamic stability in adverse conditions, making them promising candidates for further investigation in applications requiring stable MXene structures. As our simulations establish the stability of the Nb₂TiN₂S₂ MXene, we further explore its potential application in Li-Se batteries, specifically as an anchoring and catalyzing material to enhance battery performance.

3.5 Polysulfide adsorption on Nb₂TiN₂S₂

We first investigated the ability of the $Nb_2TiN_2S_2$ MXene to immobilize the otherwise soluble higher order polyselenides. Typically, the metal exposed surfaces of bare MXenes exhibit extremely high adsorption strength, which results in the

decomposition of polyselenides. To ensure reversible M–X battery operation, we incorporated surface modifications with S and studied their impact on the electrochemical performance as AMs.

The fundamental electrochemical processes in Li-Se and Li-S batteries share significant similarities.⁶⁴ During discharge, Li ions migrate from the anode and react with the Se cathode, forming intermediate Li₂Se_n. These intermediates undergo further reduction and ultimately yield the final discharge products, Li₂Se₂ and Li₂Se. To understand the anchoring behavior of Nb₂TiN₂S₂ for potential applications in metalchalcogenide batteries, we employed DFT simulations to study the adsorption of Se₈ and Li₂Se_n species. The interaction between the polyselenides and electrode surface plays a critical role in determining battery performance, particularly in mitigating the shuttling effect and improving the stability of Li-Se systems. In general, polychalcogenides interact with substrates through covalent bonding between metal sites and the surface S atoms of the MXene material. To identify the most energetically favorable adsorption configurations, we initially placed Li_2Se_n species in various positions on the Nb2TiN2S2 surface and performed structural optimizations. To quantify the anchoring strength of polyselenides on the pristine Nb₂TiN₂S₂ surface, we calculated the adsorption energies (E_{ads}) by employing the formula $E_{ads} = E_{Li_2Se_n} + E_{AM} - E_{Li_2Se_n+AM}$, where $E_{Li_2Se_n}$, E_{AM} , and

 $E_{\text{Li,Se,+AM}}$ denote the DFT energies of Li₂Se_n, AMs, and the polyselenide-adsorbed substrate. Additionally, we compared the adsorption characteristics of Nb2TiN2S2 MXene with graphene which was reported in a previous study on Li-Se battery.62 Fig. 5a presents the calculated adsorption energies of different Li₂Se_n species on Nb₂TiN₂S₂ and graphene. Our results indicate a progressive increase in the adsorption energy with increasing lithiation for Nb₂TiN₂S₂, demonstrating enhanced polyselenide anchoring as more Li atoms are incorporated. We compared the polyselenides absorption strengths on Nb2TiN2S2 with graphene as a representative carbonaceous material. The data for Li₂Se_n adsorption on graphene has been taken from our previous study.62 Fig. 5a exhibits a significant improvement in binding characteristics of Nb₂TiN₂S₂ over graphene substrate. Fig. 5b presents the relaxed atomic configurations of Li₂Se_n species adsorbed on the Nb2TiN2S2 surface. The optimized structures reveal that the Li₂Se_n species remain well-conserved during the adsorption process without undergoing significant structural or chemical degradation. The Se₈ molecules tend to adopt a nearly parallel orientation relative to the Nb₂TiN₂S₂ surface, with a minimum spacing of approximately 3.06 Å. For the other Li_2Se_n species the adsorption predominantly occurs at the bridging sites, where each Li atom establishes strong Li-S interactions with the nearby S atoms on the Nb₂TiN₂S₂ surface. This bridging configuration enhances the anchoring stability of the polyselenides, effectively preventing their dissolution and unwanted migration, which are critical challenges in Li-Se battery systems. The interaction with the Nb₂TiN₂S₂ substrate provides a strong anchoring performance while maintaining the structural integrity of the Li_2Se_n . To evaluate the possibility of polyselenide dissolution into the electrolytes, we compared our calculated adsorption energies with the adsorption energies

obtained for higher order polyselenides (Li₂Se₈, Li₂Se₆, and

Li₂Se₄) bound with the most common electrolytes solvents 1,3dioxalane (DOL) and 1,2-dimethoxyethane (DME) (shown in Fig. 5a), which were already reported in our previous work.62 Our findings suggest that Nb₂TiN₂S₂ substrate exhibits stronger binding characteristics for the polyselenides than DME and DOL, which indicates the capability of Nb₂TiN₂S₂ substrate to inhibit the dissolution of polyselenides into electrolytes, thereby limiting the shuttle effect in the Li-Se batteries.

To further understand the adsorption mechanism and establish a connection between the adsorption energies and underlying chemical interactions, we performed Bader charge analysis to quantify the charge transfer between Li₂Se_n species and the Nb₂TiN₂S₂ substrate. Our calculations reveal that Se₈ exhibited the lowest charge transfer, which aligns with its lower binding energy. The computed charge transfer values for different Li₂Se_n species are presented in Fig. 6a, where it has been observed that as the discharge process progresses, the quantity of electrons transferred from the polyselenides to the substrate increases. Positive charge transfer values indicate that charge is getting transferred from the polyselenides to the Nb₂TiN₂S₂ substrate. As a result, the lower-order polyselenides (Li₂Se₄, Li₂Se₂, and Li₂Se) are significantly prone to donate more charge to the AMs compared to higher-order species, which corroborates with their stronger interaction with Nb2TiN2S2. Higher-order polyselenides, which exhibit weaker charge transfer, also demonstrate lower binding energies, while lowerorder polyselenides show higher charge transfer and stronger adsorption.

To gain further insight into the nature of charge redistribution during adsorption, we also computed the average change in atomic charge of Li and Se atoms before and after adsorption. By comparing the atomic charge values in the gas phase (isolated Li₂Se_n molecules) and adsorbed phase (Li₂Se_n on

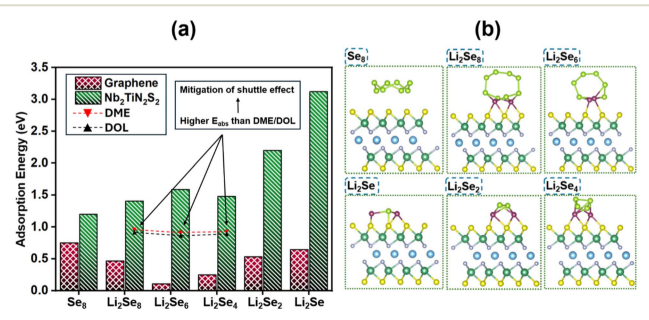


Fig. 5 (a) The calculated adsorption energies of Se_8 and Li_2Se_n adsorbed on graphene and $Nb_2TiN_2S_2$ (b) Optimized geometric configurations of Se₈ and Li_2Se_n adsorbed on $\text{Nb}_2\text{TiN}_2\text{S}_2$. Color code for atoms: Nb: forest green; Ti: sky blue; N: violate; S: yellow; Se: green; Na: navy blue; Li: purple. The data for graphene is taken from ref. 62.

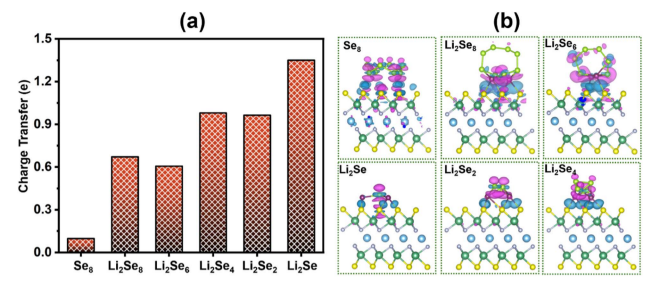


Fig. 6 (a) Computed charge transfer of Li_2Se_n adsorbed $Nb_2TiN_2S_2$. (b) Differential charge density (DCD) calculation of Se_8 and Li_2Se_n on Nb₂TiN₂S₂ substrate. The blue and pink colors define the charge accumulation and depletion, respectively.

Nb₂TiN₂S₂), we observed a notable trend: the charge transfer primarily originates from Se atoms rather than Li atoms. For example, the charge transfers of Li and Se were found -0.0168|e| and -1.317|e|, respectively, for Li₂Se adsorption. This finding is consistent with our previous research on Na-S battery systems, where we utilized Mo₂TiC₂S₂ MXene as an AM and observed a similar charge transfer mechanism.65 In both cases, the chalcogenide-substrate interaction is primarily facilitated by chalcogen atoms (S or Se), which play a crucial role in stabilizing the polychalcogenide species and preventing their dissolution.63 We also conducted the differential charge density (DCD) analysis to explain the charge transfer between the substrate and the adsorbates. The DCD analysis, shown in Fig. 6b, revealed a significant charge depletion around the Se atoms, confirming that Se acts as the primary electron donor in the adsorption process. Correspondingly, charge accumulation was observed on the Nb₂TiN₂S₂ substrate. This charge redistribution strengthens the interaction between the chalcogenide species and the substrate and develops the anchoring effect that prevents the dissolution of higher order polyselenides.

To understand the contribution between chemisorption and physisorption in the interactions between Li_2Se_n species and the substrates, we conducted further analysis to isolate the effect of vdW interactions. This analysis quantifies the contribution of vdW forces to adsorption by calculating the relative vdW ratio, which is determined by comparing the adsorption energy with and without vdW interactions. This was achieved by computing the adsorption energy for each polyselenide species using the formula,

$$E_{\mathrm{ads}}^{\mathrm{No\text{-}Vdw}} = E_{\mathrm{Li}_{2}\mathrm{Se}_{n}/\mathrm{Se}_{8}}^{\mathrm{No\text{-}VdW}} + E_{\mathrm{AM}}^{\mathrm{No\text{-}VdW}} - E_{\mathrm{AM}+\mathrm{Li}_{2}\mathrm{Se}_{n}/\mathrm{Se}_{8}}^{\mathrm{No\text{-}VdW}},$$

where $E_{\rm Li_2Se_n/Se_8}^{\rm No-VdW}$, $E_{\rm AM}^{\rm No-VdW}$ and $E_{\rm AM+Li_2Se_n/Se_8}^{\rm No-Vdw}$ represent the DFT energies of isolated Li₂Se_n/Se₈ molecules, the AM, and the polyselenide-adsorbed system without vdW interactions,

respectively. The relative contribution of vdW interactions was then calculated using the expression:

$$R = \frac{E_{\mathrm{ads}}^{\mathrm{vdW}} - E_{\mathrm{ads}}^{\mathrm{No\text{-}vdW}}}{E_{\mathrm{ads}}^{\mathrm{vdW}}}$$

The calculated VdW ratio values, as shown in Fig. 7, reveal a strong dependence of adsorption behavior on the degree of lithiation. Notably, Se₈ exhibits an exceptionally high vdW ratio, approaching nearly 100%, indicating that Se₈ adsorption is predominantly driven by vdW forces with minimal chemical interaction. This suggests that Se₈ remains weakly bonded to the substrate through non-covalent interactions, well corroborated with the observed weaker adsorption behavior on

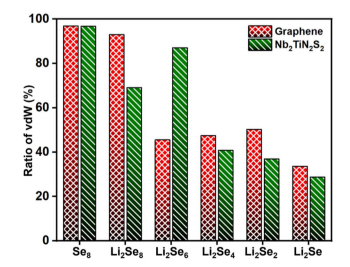


Fig. 7 The ratio of van-der Waals interactions of Se₈ and Li₂Se_n absorbed on graphene and Nb₂TiN₂S₂. The data for graphene is taken from ref. 62.

Nb₂TiN₂S₂. However, as lithiation progresses, the vdW ratio decreases significantly, indicating that chemical interactions become dominant with increasing Li content in the polyselenides. This trend is inversely related to the adsorption energy, further supporting the notion that as Li atoms interact with Se, they enhance the chemical binding by reducing the relative contribution of vdW interactions. Additionally, our analysis reveals that polyselenides exhibit stronger vdW interactions with graphene compared to Nb2TiN2S2. This finding indicates that while graphene exhibits weak physisorption behavior toward Li_2Se_n due to the non-polar nature of carbon, Nb₂TiN₂S₂ shows stronger chemical interactions with Lipolyselenides, making it a more effective material for anchoring Li_2Se_n species.

3.6 Electronic density of states of adsorbed Li₂Se_n

To ensure efficient redox reactions in Li-Se batteries, it is crucial that AMs maintain their intrinsic electronic conductivity following Li₂Se_n adsorption. A loss of conductivity upon adsorption could hinder electron transport, limiting the charge transfer kinetics and negatively impacting the overall electrochemical performance. To investigate this, we conducted DOS calculations for Nb₂TiN₂S₂ before and after Li₂Se_n adsorption to assess how adsorption affects the electronic characteristics of the substrate. The TDOS curve (Fig. S9†) confirms that pristine Nb₂TiN₂S₂ is metallic in nature, which aligns well with prior studies on functionalized MXenes. 62,63,66 The presence of a significant density of states at the Fermi level $(E_{\rm F})$ in the pristine system indicates high electronic conductivity which suggests Nb2TiN2S2 a potential host material for electrochemical applications. To further understand the contributions of different atomic orbitals, we analyzed the PDOS. As shown in Fig. S9,† the Ti-3d orbitals contribute the most to the electronic states near the Fermi level, while Nb-4d, S-3p, and N-2p states also make noticeable but relatively smaller contributions.

After the adsorption of Li_2Se_n , the PDOS plots (Fig. 8) reveal that a significant number of electronic states are still present at the Fermi level, confirming that Nb₂TiN₂S₂ retains its metallic conductivity even after polyselenides adsorption. This is particularly important because Se₈ is inherently insulating, and a conductive host material is essential to mitigate this limitation. In contrast to Se₈ adsorption, where the insulating characteristics are more pronounced, Li₂Se_n adsorption induces the emergence of adsorption peaks below the Fermi level, which correspond to charge transfer interactions between Li₂Se_n and the substrate. The PDOS further demonstrates that the contribution from Se-derived states is higher than that of S near the Fermi level, although this increase is negligible and does not significantly impact the overall conductivity of the system. This suggests that the electronic properties of Nb2TiN2S2 remain largely unaffected by Se-rich environments, further supporting its role as a stable, conductive AM.

Selenium reduction reaction (SeRR)

To further elucidate the SeRR kinetics occurring on the Nb2-TiN₂S₂ monolayer during the Li-Se battery discharge process, we investigated the overall reaction mechanisms based on the formation of Li₂Se from Se₈ and bulk Li. Understanding the thermodynamic feasibility of these transformations is crucial in assessing the electrochemical performance of Nb2TiN2S2 as an AM for Li₂Se_n. To quantify the driving forces of each step in the SeRR pathway, we computed the Gibbs free energy (ΔG) for each reaction step on the Nb2TiN2S2 substrate and compared the results with gas-phase data (Fig. 9). These free energy calculations allow us to determine the rate-determining step of the entire conversion process. Calculation details for this analysis were provided in our previous work.67

The reduction of Se₈ to Li₂Se₈ marks the first step in the Li-Se battery discharge process. Our calculations indicate that this initial reduction step is a spontaneous exothermic reaction for both the gas phase and the Nb₂TiN₂S₂-anchored systems. The exothermic nature of this step suggests that Li incorporation into Se occurs readily. The $Li_2Se_8 \rightarrow Li_2Se_6$ conversion is also found to be exothermic on both vacuum and AM. The third step, which involves the conversion of Li₂Se₆ to Li₂Se₄, is found to be endothermic, however the studied AM reduced the energy barrier for the formation of soluble Li2Se4 compared to gas phase conversion. Notably, in M-X (M = Li, Na, or K; X = S, Se) batteries, the M2X4 to M2X2 conversion is widely recognized as the rate-determining step in the overall discharge process. 65,68-70 Our results reveal that in gas phase, the Li₂Se₄ → Li₂Se₂ transformation exhibits a significant energy barrier of 1.25 eV, which could hinder the reaction kinetics and slow down the discharge process. However, when the reaction occurs on Nb₂-TiN₂S₂, the energy barrier is dramatically reduced to 0.201 eV, marking an 83.94% reduction in the reaction barrier compared to vacuum conditions. This substantial decrease in energy requirement evidence that the Nb2TiN2S2 substrate effectively accelerates the rate-determining step, enabling faster polyselenide conversion and enhancing the discharge reaction kinetics of Li-Se batteries. The improved Se reduction kinetics arises from the strong adsorption of polyselenides on Nb2-TiN₂S₂, which facilitates the reduction of higher order polyselenides to lower order polyselenides, owing to the electron transfer at the electrode-electrolyte interface. The sulfur functionalized Nb₂TiN₂ MXene provides abundant adsorption sites for polyselenides to suppress the shuttle effect by immobilizing the lithium polyselenides species (Fig. 5). At the same time, its high electrical conductivity enhances electron transfer during the adsorption process (Fig. 6a). The strong binding between polyselenides and the Nb2TiN2S2 surface, coupled with charge depletion from the polyselenide species toward Nb2TiN2S2 (Fig. 6b), accelerates the Se reduction reaction and lowers the energy barriers of intermediate steps. This effect is particularly pronounced in the $Li_2Se_4 \rightarrow Li_2Se_2$ conversion step compared to other stages (Fig. 9). Overall, these findings suggest that Nb₂TiN₂S₂ can effectively serve as both a catalytic and anchoring host to accelerate the conversion kinetics and improve the cycling performance of Li-Se batteries. A similar trend of acceleration in reduction kinetics was observed in our previous studies, where surface modification-based metalchalcogenides batteries exhibited significantly improved discharge reaction rates due to substrate-induced energy barrier

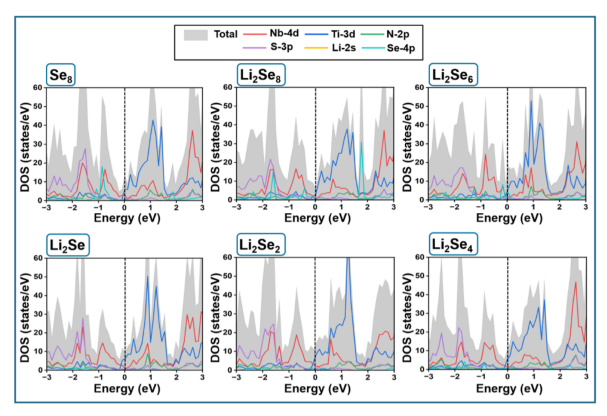


Fig. 8 PDOS of Se₈ and Li₂Se_n adsorbed on Nb₂TiN₂S₂. The Fermi level is denoted by vertical lines.

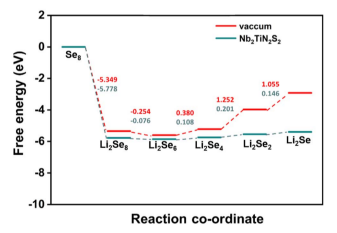


Fig. 9 Gibbs free energy diagram of selenium reduction reaction in vacuum and on $Nb_2TiN_2S_2$ for Li–Se systems.

reduction. 63,65,67,71 In conclusion, during discharge, the use $Nb_2TiN_2S_2$ as an AM proved to be effective for Li–Se batteries where Li–Se exhibited faster reduction reaction kinetics.

4. Conclusion

In this study, we discovered a new MXene, Nb₂TiN₂, and evaluated its structural and electronic properties, along with its potential as a cathode catalyst for Li–Se batteries. We started our investigation by analyzing the precursor Nb₂TiAlN₂ MAX

phase. With the relaxed MAX phase structure, we employed phonon dispersion and AIMD simulation to examine the dynamic stability and thermodynamic stability at high temperatures, respectively. Our finding suggests that the MAX phase exhibits excellent dynamic stability with zero negative frequency and thermodynamic stability even at 1000 K. The DOS analysis provides evidence of the presence of states at the Fermi level with the conducting property of the catalyst majorly arising from the Nb and Ti 'd' orbitals. The transformation of 2D MXene from the MAX phase was also examined with the calculation of exfoliation energy and bond stiffness of various bonds. Al-N and Nb-Al bonds are found to be significantly weaker than the other bonds, which is in favor of the exfoliation process. On the other hand, the exfoliation energy was also found within the critical value. We investigated the binding mechanism and reaction kinetics of Li2Se on the Nb2TiN2S2 substrate. Simulations revealed that Nb2TiN2S2 is effective in anchoring the Li_2Se_n to mitigate the shuttle effect. During the anchoring process, the charge transfer occurs from the Se atoms of the polyselenides to the AM material and the charge transfer increases as the order of polyselenides decreases. The DOS analysis of the polyselenides adsorption on AM indicates the presence of the metallic nature even after adsorption. According to the Gibbs free energy profile, Li₂Se₄ to Li₂Se₂ conversion is found to be the rate-determining step, and the inclusion of Nb2TiN2S2 lowers the energy barrier and is expected to accelerate the reaction kinetics significantly. In conclusion, the results demonstrate that Nb₂TiN₂S₂ MXene possesses

significant potential to advance the development of high-performance and sustainable Li-Se batteries, as well as broader catalytic systems. This investigation can be extended to other chalcogen-based systems such as Li-S, Na-S, and Na-Se batteries. Although the fundamental mechanisms share common features, there are also distinct differences depending on the chalcogen species and ion involved. On the other hand, the influence of different surface terminations such as -O, -F, and -OH on the catalytic performance of Nb₂TiN₂ MXene could offer another promising direction to explore. In our future work, we plan to utilize this MXene with various functional groups in these alternative battery chemistries to explore further insights into its catalytic functionality.

Data availability

The authors confirm that the data required to reproduce the findings of this study are available within the article and can be reproduced by density functional theory calculations.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work is partially supported by the National Science Foundation (Award No. CBET-2400109) and the Ralph E. Powe Junior Faculty Enhancement Award.

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