Rational Design of Two-Dimensional Transition Metal Carbide/Nitride (MXene) Hybrids and Nanocomposites for Catalytic Energy Storage and Conversion

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This is the author’s manuscript of the article published in final edited form as:

ABSTRACT

Electro-, photo-, and photoelectrocatalysis play a critical role toward the realization of a sustainable energy economy. They facilitate numerous redox reactions in energy storage and conversion systems, enabling the production of chemical feedstock and clean fuels from abundant resources like water, carbon dioxide, and nitrogen. One major obstacle for their large-scale implementation is the scarcity of cost-effective, durable, and efficient catalysts. A family of two-dimensional transition metal carbides, nitrides, and carbonitrides (MXenes) has recently emerged as promising earth-abundant candidates for large-area catalytic energy storage and conversion due to their unique properties of hydrophilicity, high metallic conductivity, and ease of production by solution processing. To take full advantage of these desirable properties, MXenes have been combined with other materials to form MXene hybrids with significantly enhanced catalytic performances beyond the sum of their individual components. MXene hybridization tunes the electronic structure towards optimal binding of redox active species to improve intrinsic activity, while increasing the density and accessibility of active sites. This review outlines recent strategies in the design of MXene hybrids for industrially relevant electrocatalytic, photocatalytic, and photoelectrocatalytic applications such as water splitting, metal-air/sulfur batteries, carbon dioxide reduction, and nitrogen reduction. By clarifying the roles of individual material components in the MXene hybrids, we provide design strategies to synergistically couple MXenes with associated materials for highly efficient and durable catalytic applications. We conclude by highlighting key gaps in the current understanding of MXene hybrids to guide future MXene hybrid designs in catalytic energy storage and conversion applications.

KEYWORDS: MXenes, hybrid material, electrocatalysis, photocatalysis, water splitting, CO₂ reduction, N₂ reduction, energy conversion, batteries, energy storage
The advancement of our energy economy using cost-effective, accessible, and environmentally responsible means is critical to ensure a sustainable supply to meet the burgeoning global energy demand.\textsuperscript{1,2} In this regard, catalysts play a critical role in enabling efficient energy conversion of earth-abundant resources such as water, carbon dioxide, and nitrogen to clean fuels and chemical feedstock precursors such as hydrogen, hydrocarbons, and ammonia.\textsuperscript{3–5} Durable and active catalysts are also required to facilitate long-term energy storage in H\textsubscript{2} fuel cells and batteries of varying architectures.\textsuperscript{6–8} Presently, precious platinum group metals (PGMs) such as Pt, their alloys, and related structures (PtIr, IrO\textsubscript{2}, RuO\textsubscript{2}, etc.) are widely regarded as the most effective catalysts for the hydrogen evolution reaction (HER), oxygen reduction reaction (ORR), and oxygen evolution reaction (OER), respectively.\textsuperscript{9–11} The widespread commercialization and adoption into energy storage and conversion devices however, can only be realized with alternative earth-abundant and cost-effective catalysts with high catalytic activity and stability.\textsuperscript{3,4}

Since their discovery in 2011, more than 30 different compositions belonging to a family of two-dimensional (2D) transition metal carbides, nitrides, and carbonitrides (MXenes) have been successfully synthesized, with many other stable compositions predicted theoretically.\textsuperscript{12–14} MXenes have attracted significant attention as promising catalysts due to their unique combination of physical and chemical properties.\textsuperscript{13–15} They possess high metallic electrical conductivity with hydrophilicity similar to graphene oxide, and mechanical strength of transition metal (TM) carbides/nitrides. MXenes can be easily scaled, because of their solution processible top-down synthesis, into stable colloidal solutions or flexible films for energy applications (electrocatalysts, batteries, and supercapacitors) and beyond (flexible electronics, sensors, and electromagnetic shielding).\textsuperscript{13,14,16} In addition to their high metallic conductivity to facilitate efficient electronic charge transport at the electrode-electrolyte interface, the large-area 2D basal planes of MXenes
are HER-active\textsuperscript{17–19} unlike the widely studied semiconducting 2H-MoS\textsubscript{2} with HER activity restricted only to their edges.\textsuperscript{20} Separate theoretical calculations have also predicted a variety of MXenes for efficient photocatalytic (PC), electrocatalytic (EC), and photoelectrocatalytic (PEC) reactions such as the hydrogen evolution (HER), oxygen evolution (OER), oxygen reduction (ORR), carbon dioxide reduction (CO\textsubscript{2}RR), and nitrogen reduction (N\textsubscript{2}RR) reactions.\textsuperscript{21–25} MXenes have a general formula of M\textsubscript{n+1}X\textsubscript{n}T\textsubscript{x} where n = 1–4, M = early transition metal such as Ti, V and Mo, X = C and/or N and T\textsubscript{x} = surface termination groups such as –O, –OH, –F, and –Cl, which are formed on the outer M basal plane during the synthesis process.\textsuperscript{26–28} For consistency, a standardized notation of M\textsubscript{n+1}X\textsubscript{n}T\textsubscript{x} will be used to describe MXenes throughout this review.

MXenes are most commonly produced through the top-down separation of layered MXene sheets from their parent layered ternary TM carbide MAX phase materials ("A" = group 13 or 14 elements such as Al, Si, and Ga) by aqueous wet selective etching and removal of A-group atom layers.\textsuperscript{12,27} The A-group atom layers are preferentially etched away as the M-A bonds binding the MXene layers together in the MAX structure are chemically more active than the stronger M-X bonds within the MXene sheets.\textsuperscript{29–31} This scalable top-down etching process to produce MXenes is most commonly conducted under ambient conditions with gentle heating using hydrofluoric acid (HF) directly, or HF produced \textit{in-situ} from F-containing etchants.\textsuperscript{32–35} After etching away the A-group atom layers, the exposed outer M basal planes of the etched MXene sheets are functionalized with hydrophilic surface termination groups (T\textsubscript{x}) such as –O, –OH, –F and –Cl due to the aqueous F- or Cl-containing etchants.\textsuperscript{14,31,36} The surface chemistry of MXenes can be controlled to some extent by the selective wet etching conditions or through post-etch processing to optimize binding toward specific electroactive species for efficient catalysis.\textsuperscript{18,37} Subsequent delamination and exfoliation separates the weakly bound MXene sheets into single- or few-layered MXenes using
intercalants,\textsuperscript{38,39} exposing large MXene surface areas for catalysis and to serve as a conductive support for co-catalysts.

Most recently, organic and inorganic materials such as TM compounds (carbides, phosphides, chalcogenides, oxides, \textit{etc.}), layered double hydroxides (LDHs), metal-organic frameworks (MOFs), carbon-based (C-based) materials such as graphitic carbon nitride (g-C\textsubscript{3}N\textsubscript{4}) and carbon nanotubes (CNTs), quantum dots (QDs), metallic alloys, and polymers have been combined with MXenes to form MXene hybrids and composites.\textsuperscript{40–45} These MXene hybrids exhibit significantly enhanced catalytic activities and stability beyond the sum of their individual components, due to synergistic coupling between MXenes and the secondary materials. The strong and intimate coupling increases the density of catalytically active sites, and optimizes these active sites toward binding of electrochemically active species for efficient catalysis.\textsuperscript{40–45} The hybrid mix can perform a variety of functions such as a large-area, electronically conductive support, electronic structure modulator, and co-catalyst,\textsuperscript{40–45} as we discuss in this review. Indeed, MXene hybrids have garnered significant traction since 2014, with the number of papers published doubling year-on-year, and more than half (168) of the publications on MXene hybrids in 2019 (312) focused on catalysis and energy storage (\textbf{Figure 1a}). The breakdown of the many different secondary materials integrated with MXenes to form MXene hybrids also illustrates the versatility of MXene hybrids to meet the demands of a wide spectrum of applications (\textbf{Figure 1b}), especially in the many important catalytic reactions of interest (\textbf{Figure 1c}).

In this review, we summarize recent strategies in the design of MXene hybrids for industrially relevant EC, PC, and PEC applications in energy storage (metal-air/sulfur batteries catalyzed by OER, ORR, and sulfur redox reactions) and conversion (water splitting catalyzed by OER and HER). We clarify the roles of each material component in the MXene hybrids for their
intended applications to provide a set of rational design rules to synergistically couple MXenes with the relevant secondary materials for highly efficient and durable catalysis. The feasibility of MXene hybrids for important emerging catalytic applications such as CO_2RR and N_2RR is then discussed. The review concludes by highlighting key gaps in the current fundamental understanding of MXene hybrids and provides an outlook to inform forthcoming MXene hybrid designs in catalytic energy storage and conversion applications.

**Figure 1.** (a) Graphical representation of the total publications to date (*March 2020*) on MXene hybrids for various applications on Web of Science. Color filled radial bars represent different applications, while uncolored radial bars are the projected number of publications for the rest of 2020. (b) Spider plot survey of all MXene hybrid publications to date, illustrating the type of MXene hybrids most used for specific applications. For example, 31% of all published MXene hybrid papers for catalysis used MXene/TM oxides hybrids. (c) Spider plot survey of all MXene hybrid publications for catalysis to date, illustrating the type of MXene hybrids most used for various catalytic reactions of interest. For instance, MXene/TM compound hybrids account for 32% of all HER-related papers using MXene hybrids. Raw data used for each plot is provided in Tables S1a–S1c respectively.
DESIGN STRATEGIES FOR MXENE HYBRIDS AND COMPOSITES

The layered nature and variety in surface $T_x$ groups of MXenes\textsuperscript{30} afford a high degree of tuneability in their properties\textsuperscript{13–15} and hence, strong compatibility with a wide range of secondary materials to produce MXene hybrids with high catalytic performances. MXene hybrids with precise and well-formed interfaces have been reported to improve charge transfer,\textsuperscript{46–48} mitigate volumetric expansion in batteries\textsuperscript{49,50} and resist secondary material agglomeration through confinement effects.\textsuperscript{51–53} In this review, we define a MXene hybrid/composite as a material comprising MXenes and at least one other non-MXene material, either organic or inorganic. This is a more general term than MXene heterostructures, which are produced when a material is grown onto MXenes in a repetitive layered manner (see Vocabulary for full definition of MXene hybrid/composite and MXene heterostructures).

A growing number of MXenes have been integrated with non-MXene secondary materials to form MXene hybrids, realizing unique material properties and enhanced performance. These MXenes include, but are not limited to, Ti$_2$CT$_x$,\textsuperscript{54} Ti$_3$C$_2$T$_x$,\textsuperscript{55,56} Ti$_3$CNT$_x$,\textsuperscript{56} V$_2$CT$_x$,\textsuperscript{57,58} V$_4$C$_3$T$_x$,\textsuperscript{59,60} Nb$_2$CT$_x$,\textsuperscript{61,62} Nb$_4$CT$_x$,\textsuperscript{62} Mo$_{1.33}$CT$_x$,\textsuperscript{63} Mo$_2$CT$_x$,\textsuperscript{64–66} Mo$_2$TiC$_2$T$_x$,\textsuperscript{67} and Ta$_4$C$_3$T$_x$.\textsuperscript{68} Since rapid electronic charge transfer is crucial to efficient electrocatalysis, Ti$_3$C$_2$T$_x$ is most extensively incorporated into MXene hybrids for electrocatalytic applications, due to their high electrical conductivity (up to 15000 S cm$^{-1}$)\textsuperscript{69} as compared to other MXenes. The wide diversity in MXene-compatible secondary materials (TM compounds, supramolecular structures, C-based materials, and metals), coupled with the many different synthetic methods to form these hybrids (Figure 2), illustrates the boundless potential for MXene hybrids and composites to meet the various demands in energy-related applications and beyond.\textsuperscript{40–44}
**Figure 2.** Three different MXene types combined with secondary materials to form MXene hybrids. Cyan, dark grey and yellow spheres represent M, X, and Tₓ respectively. Common processing methods to fabricate the MXene hybrids and their catalytic applications are also listed.

The first step to designing efficient and durable MXene hybrids for energy storage and conversion is the deliberate choice of secondary material and the processing method. Highly interconnected charge transfer networks for electronic transport, a large-area/hierarchical/porous network for mass transfer of electroactive species, and an abundance of catalytically active sites, are all highly desirable properties in MXene hybrids for catalytic applications.⁴⁰-⁴⁴ Additionally, a
strongly and chemically coupled MXene hybrid synergistically increases the hybrid’s intrinsic activity, while secondary material dislodgement or degradation during electrocatalysis is prevented, especially at large applied potentials. MXene hybrids are synthesized through three main routes: (1) in-situ conversion of the MXenes’ surface structure through oxidation (e.g. $\text{Ti}_3\text{C}_2\text{T}_x$ to $\text{TiO}_2$) or sulfidation (e.g. $\text{Mo}_2\text{TiC}_2\text{T}_x$ to $\text{MoS}_2$), (2) reactive chemical formation of secondary materials on MXenes, using methods such as hydro/solvothermal synthesis, solution processing, or deposition, and (3) non-reactive assembly of secondary materials by drop-casting and adsorption processes. In this section, we summarize the common and unique features for each of these three synthetic routes for MXene hybrid formation, highlighting their suitability, advantages, and shortfalls (and how they have been overcome).

**In-situ conversion of MXene surfaces to form MXene hybrids.** The layered structure of MXenes leaves the TM atomic layers exposed on their outer basal planes (left schematics in Figure 2). These TM layers are highly oxophilic and hence prone to surface reactions such as oxidation (e.g. Ti to $\text{TiO}_2$ in $\text{Ti}_3\text{C}_2\text{T}_x$) or sulfidation (e.g. Mo to $\text{MoS}_2$ in $\text{Mo}_2\text{TiC}_2\text{T}_x$). In these processes, the MXenes’ surface is transformed, either partially or fully, to form a second material. We thus refer to this process as an in-situ conversion of the MXenes’ surface. The advantage of surface conversion is clear: the second material is strongly integrated with MXenes, forming a hybrid interface. Since the MXene surfaces are directly involved in the conversion, no additional TM source is required for the secondary material growth. Consequently, this process is self-limited by the MXenes’ surface exposure, minimizing the issue of excessive secondary material nucleation and growth that would otherwise restrict access of reactants to catalytically active sites.

Most surface conversions involve the treatment of MXenes with gases such as $\text{O}_2$ or $\text{CO}_2$ for oxidation, or $\text{H}_2\text{S}$ for sulfidation. These processes are typically conducted at elevated
temperatures to form highly crystalline secondary materials on MXene surfaces.\textsuperscript{67,83} Hence, there exists a delicate balance between the extent of MXene conversion, and preserving the MXenes’ structural integrity that can be compromised in oxidizing environments at high temperatures.\textsuperscript{83} The range of secondary materials that can be grown through MXene surface conversion is also limited by the chemical identity and reactivity of the MXenes’ TM surface exposed. For instance, the Ti surface of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} implies that only Ti-based secondary materials can be grown through this conversion route. In this regard, we anticipate that disordered solid solution MXenes such as (Ti,V)\textsubscript{2}CT\textsubscript{x} and (Mo,V)\textsubscript{4}C\textsubscript{3}T\textsubscript{x}, denoted by the chemical formula (M',M'')\textsubscript{n+1}C\textsubscript{n}T\textsubscript{x}, will be interesting for growth of secondary materials through this surface conversion route, since they possess two different TMs at the surface in a random distribution.\textsuperscript{84}

Thermally-assisted oxidation is an efficient one-step method to produce MXene/TM oxide hybrids.\textsuperscript{41,43} MXene surface oxidation is most commonly employed in Ti-based MXenes to form TiO\textsubscript{2} photocatalysts,\textsuperscript{55} although oxidation can be also applied to V, Nb, and Mo-based MXenes to form their respective oxides.\textsuperscript{58,62,82} \textit{In-situ} oxidation is achieved through flash oxidation (1150 °C, 30 s),\textsuperscript{55} prolonged oxidation from calcination (350–650 °C),\textsuperscript{85–87} hydro/solvothermal processes (120–220 °C),\textsuperscript{47,78,88–91} or CO\textsubscript{2} oxidation (500–900 °C).\textsuperscript{62,83} Control over the MXenes’ surface T\textsubscript{x} groups prior to oxidation is crucial, since different surface terminations have resulted in morphological differences in the oxides formed. Li \textit{et al.} reported that calcining untreated Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} with –O, –F, and –OH terminations resulted in larger anatase TiO\textsubscript{2} nanoparticles (NPs) with exposed (001) planes, compared to the smaller and more spherical anatase TiO\textsubscript{2} NPs formed by calcining Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} with primarily –OH terminations.\textsuperscript{87} With precise control and optimization over the heat treatment conditions, a variety of functional TiO\textsubscript{2}/MXene hybrids can be formed with high PC activity toward water splitting, CO\textsubscript{2}RR, and N\textsubscript{2}RR.\textsuperscript{87,89}
For example, Ti$_3$C$_2$Tx has been oxidized to varying degrees to produce hybrids ranging from Ti$_3$C$_2$Tx/TiO$_2$ (partial MXene surface oxidation) to C/TiO$_2$ (complete MXene oxidation). While controlled partial oxidation retained some of the Ti-C layered structure from the MXene precursor, complete oxidation degraded the entire MXene structure. The latter resulted in a Ti-O structure supported by C-C bonded layers, in the form of TiO$_2$ on 2D carbon sheets (C/TiO$_2$). The morphology and phase of the resulting TiO$_2$ NPs was also highly dependent on the treatment conditions. This is important as the identity and quantity of exposed facets, together with the distribution and phase of TiO$_2$ formed, directly influences PC activity. Ghassemi et al. utilized in-situ environmental transmission electron microscopy (TEM) and Raman spectroscopy to elucidate the oxidation mechanism of Ti$_3$C$_2$Tx under various conditions (Figure 3a). In flash oxidation, the exposed Ti basal planes serve as nucleating sites to form a thin 2D anatase TiO$_2$ film. Subsequently, the 2D TiO$_2$ nuclei grew vertically to form 3D TiO$_2$ anatase NPs from the outward migration of inner Ti layer atoms. Conversely, a slower heating process yielded rutile TiO$_2$ NPs without the formation of the intermediate anatase phase. Experimentally, anatase TiO$_2$ was formed from partial Ti$_3$C$_2$Tx oxidation at lower temperatures, while rutile TiO$_2$ was mostly formed through near-complete oxidation of Ti$_3$C$_2$Tx, often resulting in degradation of the Ti-C MXene interior.

MXene/TM chalcogenides hybrids have also been formed through in-situ sulfidation of MXene surfaces. Although TM chalcogenides are HER-active, they are less electrically conductive than MXenes. Thus, MXene/TM chalcogenides hybrids benefit from (1) significantly improved charge transfer kinetics due to the metallic MXenes, and (2) an increased number of active sites with TM chalcogenides acting as co-catalysts in the hybrid structure. In ordered double TM (Mo, Ti) MXenes such as Mo$_2$TiC$_2$Tx and Mo$_2$Ti$_2$C$_3$Tx, Mo atoms preferentially...
occupy the surface (outer) basal planes. This led to a follow-up study on the in-situ sulfidation of the Mo-rich surface of \( \text{Mo}_2\text{TiC}_2\text{T}_x \) to form the \( \text{MoS}_2@\text{Mo}_2\text{TiC}_2\text{T}_x \) hybrid. After an initial liquid mixing process to incorporate sulfur on and within the \( \text{Mo}_2\text{TiC}_2\text{T}_x \) sheets, heat treatment in an inert Ar atmosphere with additional elemental sulfur converted the surface Mo-O motifs to 2H-MoS\(_2\) to form the final \( \text{MoS}_2@\text{Mo}_2\text{TiC}_2\text{T}_x \) hybrid (Figure 3b). The \( \text{Mo}_2\text{TiC}_2\text{T}_x : \text{MoS}_2 \) ratio was also tuned from 1 : 0.62 and 1 : 2.19 by adjusting the treatment temperature from 500 to 700 °C. Furthermore, the intimate interfacial relationship was evident in the stacking of thin 2H-MoS\(_2\) on the MXenes (Figures 3c and 3d). This 2H-MoS\(_2\) on MXene hybrid exhibited an increased specific capacity when used as a lithium ion anode, which was attributed to the incorporation of MoS\(_2\). We speculate that ordered double TM MXenes with a Mo-rich surface, such as \( \text{Mo}_2\text{TiC}_2\text{T}_x \) and \( \text{Mo}_2\text{TiC}_3\text{T}_x \), might be better suited for sulfidation compared to single TM MXenes (\( \text{Mo}_2\text{CT}_x \)), due to the presence of a conductive inner Ti layer after surface Mo sulfidation to MoS\(_2\).

**Figure 3.** In-situ conversion of MXene surfaces to form MXene hybrids. (a) Oxidation mechanism of \( \text{Ti}_3\text{C}_2\text{T}_x \) to form \( \text{TiO}_2 \) under different oxidation conditions. Adapted with permission from ref 96. Copyright 2014 Royal Society of Chemistry. (b) In-situ sulfidation of \( \text{Mo}_2\text{TiC}_2\text{T}_x \) and cross-sectional TEM image of the resulting \( \text{MoS}_2@\text{Mo}_2\text{TiC}_2\text{T}_x \) hybrid heat treated at (c) 500 °C and (d) 700 °C. Adapted with permission from ref 67. Copyright 2018 Wiley-VCH.
In-situ reactive formation of secondary materials onto MXenes to form MXene hybrids. Compared to in-situ conversion of MXene surfaces, this approach allows for the incorporation of wider material compositions, independent of the MXenes’ TM surface identity. Here, MXenes can be viewed as substrates, not reactants, for secondary material growth. The main advantage of in-situ reactive formation is the preservation of the MXenes’ integrity in the synthesis process, as virtually any precursors (cationic, anionic, or metallic) can be utilized for the nucleation and growth of the secondary material. With this process, it is possible to form hybrids with high secondary material to MXene ratios as reported for Bi$_2$WO$_6$/Ti$_3$C$_2$Tx (Bi$_2$WO$_6$ : MXene layer height ≈3:1)$^{101}$ or MoSe$_2$/Ti$_3$C$_2$Tx.$^{102}$ Thus, this strategy is most useful if the secondary material is catalytically active, since large quantities can be grown on the large surface area of MXenes.

Typical synthesis routes available for this strategy include hydro/solvothermal synthesis,$^{47,72,103–105}$ solution processing,$^{73}$ and deposition (physical or electro/photo-induced).$^{74–79}$ Most of these synthetic routes, except physical deposition, rely on the van der Waals$^{53}$ or ionic/electrostatic$^{106}$ interactions between the negatively charged MXene surfaces (from the T$_x$ groups) and positively charged secondary material precursors.$^{41}$ After initial adsorption, the secondary materials are formed in-situ onto the MXenes’ surface through chemical processing with additional reactants and/or heat treatment.$^{101,107}$

Hydro/solvothermal synthesis exploits the interactions between the liquid solvent, mineralizer (e.g. base), and precursor moieties at elevated temperatures and pressures, mimicking the processes occurring at the Earth’s crust.$^{108}$ This synthesis route is a simple and scalable way to synthesize many different types of secondary materials (Figure 2) in various morphologies,$^{109–111}$ and is versatile enough to yield solid solution mixtures$^{112}$ or different material allotropes$^{113}$ by carefully tuning the processing conditions. Although hydrothermal synthesis is typically directed
at bulk material synthesis, heterogeneous nucleation and epitaxial films can be achieved by adding suitable substrates.\textsuperscript{114,115} A major challenge in this route is aggressive corrosion by water molecules or –OH moieties at elevated temperatures and pressures, which may induce undesired damage to the MXenes.\textsuperscript{70} This can be partially alleviated by replacing water with non-aqueous solvents, thus the term solvothermal.\textsuperscript{116} Wu et al. proposed the addition of glucose to the solvent to protect MXenes from oxidative damage.\textsuperscript{70} During the reaction, glucose molecules were adhered on MXene surfaces \textit{via} hydrogen bonding and converted to a simple hydrocarbon coating (Figure 4a), which was subsequently converted to carbon with thermal treatment. The carbon coating was shown to improve charge transport characteristics and prevent secondary material aggregation.\textsuperscript{117}

Solution processing at ambient pressure is an alternative to hydro/solvothermal treatment, since the gentler reaction conditions can reduce unwanted MXene damage.\textsuperscript{73} To achieve controlled growth, metallic precursors are first adsorbed onto MXenes \textit{via} a mixing process. Thereafter, anionic precursors are introduced to chemically react and form the desired secondary materials on the MXene surface. An additional annealing step is usually required to complete the reaction and enhance the integration of the secondary material on MXenes.\textsuperscript{118–120} A variety of inorganic materials such as TM oxides,\textsuperscript{121} phosphides,\textsuperscript{122} chalcogenides,\textsuperscript{118} nitrides,\textsuperscript{123} MOFs,\textsuperscript{124} perovskites,\textsuperscript{125} and LDHs\textsuperscript{109,126} have been hybridized with MXenes this way (Figure 2). Yu et al. prepared FeNi-LDH@Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} hybrids by the co-precipitation of Fe\textsuperscript{3+} and Ni\textsuperscript{2+} cations on negatively charged Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} sheets for EC OER (Figure 4b).\textsuperscript{126} The abundance of –OH and –F surface groups facilitated the initial electrostatic nucleation and subsequent growth of LDH sheets on MXenes with minimal oxidation. Additionally, secondary materials that are initially electrostatically adsorbed and formed on MXene surfaces, can be chemically converted to other more catalytically active materials.\textsuperscript{119,120} For instance, zeolitic imidazolate framework-67 (ZIF-67)
self-assembled on Ti$_3$C$_2$Tx was etched to NiCo-LDH by ultrasonication after the addition of Ni$^{2+}$ precursors. Then, heat treatment and sulfidation of the mildly OER-active NiCo-LDH/Ti$_3$C$_2$Tx yielded NiCoS/Ti$_3$C$_2$Tx MXene hybrids with significantly enhanced OER activity (Figure 4c).\textsuperscript{120}

**Figure 4.** *In-situ* formation of secondary materials on MXenes to form MXene hybrids. (a) Carbon nanoplating to resist MXene oxidation during hydrothermal treatment. Adapted with permission from ref \textsuperscript{70}. Copyright 2017 Wiley-VCH. (b) *In-situ* electrostatic self-assembly of FeNi-LDH/Ti$_3$C$_2$Tx hybrid. Adapted with permission from ref \textsuperscript{126}. Copyright 2017 Elsevier Ltd. (c) X-ray diffraction (XRD) spectra showing the conversion of ZIF-67/Ti$_3$C$_2$Tx to NiCoS/Ti$_3$C$_2$Tx hybrid. Adapted with permission from ref \textsuperscript{120}. Copyright 2018 American Chemical Society.

Finally, MXene hybrids can be formed via various deposition methods. Deposition methods are generally non-invasive, controlled, and a more conformal way of fabricating MXene hybrids and heterostructures as compared to the random assembly methods of *in-situ* conversion and self-assembly. Secondary materials have been deposited on MXene surfaces through atomic layer deposition (ALD),\textsuperscript{79,127–130} electro-,\textsuperscript{76,77,131,132} or photodeposition.\textsuperscript{78,133} MXenes have been used as substrates for ALD and chemical vapor deposition (CVD) of secondary material precursors to grow TM oxides,\textsuperscript{79} metal NPs,\textsuperscript{127} CNTs,\textsuperscript{128} and inorganic NPs.\textsuperscript{129} These techniques promote strong interfacial bonding due to *in-situ* hybrid formation at elevated temperatures,\textsuperscript{128,129} which
promotes charge transfer for electrocatalysis.\textsuperscript{130} While ALD and CVD are useful for atomic-scale studies at the MXene hybrid interface,\textsuperscript{65} these methods may not be viable to produce hybrids for electrocatalysis at larger scales due to the cost and difficulty in scaling up.

MXene hybrids comprising TM phosphides,\textsuperscript{74} oxides,\textsuperscript{131} metals,\textsuperscript{76} and C-based materials\textsuperscript{77,132} have also been formed through electrodeposition. Electrodeposition can be used to produce MXene hybrid coatings or electrodes without the use of binders or additives, which can potentially hinder charge and mass transport during electrocatalysis.\textsuperscript{132} In one study, the conductive Ti$_3$C$_2$Tx film was directly used as a cathode for the deposition of secondary material precursors.\textsuperscript{76} Alternatively, other electrodeposition studies disperse both the MXenes and secondary material precursors in the electrolyte, and deposit the hybrid onto a conductive electrode such as nickel foam,\textsuperscript{131} graphite paper\textsuperscript{132} or nickel plate.\textsuperscript{74} Specifically, –OH surface groups on Ti$_3$C$_2$Tx have been demonstrated to improve the adsorption of Sn$^{4+}$ and Pb$^{2+}$,\textsuperscript{134,135} which presents the possibility of tuning the T$_x$ composition to selectively favor the electrodeposition of specific metallic precursors. Complementary to electrodeposition, photodeposition was used to reduce metallic ions to deposit metallic NPs such as Cu\textsuperscript{78} and Pt\textsuperscript{133} \textit{in-situ} onto MXene surfaces. Photodeposition possesses an advantage over other deposition methods in that it can be used to target specific areas or facets where photo-generated electrons accumulate.\textsuperscript{136} However, the light intensity must be controlled carefully, as very intense light irradiation has been shown to oxidize and degrade MXenes.\textsuperscript{137,138}

Non-reactive assembly of secondary materials onto MXenes to form MXene hybrids.

Secondary materials with a positive surface charge can also be easily combined with MXenes to form MXene hybrids through physical processes such as drop-casting\textsuperscript{80} and adsorption.\textsuperscript{56} This method also relies on the same electrostatic/van der Waals self-assembly principle as the previous
method (reactive chemical formation of secondary materials on MXenes). However, in this non-reactive method, the secondary materials are in the final forms prior to integration with MXenes. Hence, the MXenes are typically not subjected to high temperature treatment steps such as hydro/solvothermal treatment for secondary material integration. This method is most suitable for ambient-stable secondary materials that can be easily solution-processed.

Adsorption of C-based materials such as g-C$_3$N$_4$, CNTs, graphene oxide (GO), and reduced graphene oxide (rGO) is commonly used for the assembly of MXene/C-based material hybrids. CNTs are often mixed with MXenes by sonication, while g-C$_3$N$_4$, GO, and rGO are electrostatically adsorbed. MXene hybrids with CNTs, g-C$_3$N$_4$, and GO/rGO are then traditionally filtered to yield free standing films for electrode applications and capacitors. After the adsorption process, MXene hybrids with CNTs and GO/rGO can be further freeze-dried to form conductive aerogel networks for capacitors. Generally, MXene/g-C$_3$N$_4$ hybrids promote electron transfer at the interface to augment the catalytic activity of g-C$_3$N$_4$, while CNTs, GO, and rGO on MXenes serve as additional conductive and/or structural supports. Other secondary materials such as TM oxides, phosphates, chalcogenides, nitrides, MOFs, and LDHs can also be combined with MXenes this way.

MXene hybrids have also been formed by drop-casting MXenes onto the secondary material (or vice versa). For example, TiO$_2$ nanorods were grown hydrothermally on a fluorine-doped tin oxide (FTO) substrate, and then coated with Ti$_3$C$_2$T$_x$ through solution drop casting to produce the Ti$_3$C$_2$T$_x$/TiO$_2$/FTO photoanode for PEC water splitting. While drop-casting is a less-intensive process, post-processing steps such as annealing may be required to optimize MXene-secondary material interfacial chemical bonding and charge transfer.
Summary, specific challenges, and outlook for processing of MXene hybrids. We have described how MXene hybrids can be formed through (1) in-situ conversion by using MXene surfaces as reactants, (2) direct chemical growth of secondary materials on MXenes, and (3) deposition of secondary materials on MXenes. The processing method employed for MXene hybrids formation is highly dependent on their intended catalytic application. For instance, PC applications require semiconducting photo-absorbers, which can be formed via in-situ oxidation or hydro/solvothermal treatment of Ti-based MXenes. In contrast, oxide formation on MXenes is detrimental to electrocatalysis, as oxides are less conductive and limit charge transport properties, so a self-assembly process may be more appropriate. MXene hybrids should also be carefully designed to optimize chemical coupling at the MXene-secondary material interfaces toward catalytic activity enhancement. For example, strong chemical coupling between Ti₃C₂Tx with g-C₃N₄ afforded a highly OER-active hybrid. A strongly coupled MXene hybrid also reduces the probability of hybrid disintegration during electrocatalysis. Finally, the surface chemistry of MXene Tₓ groups play an important role in determining the morphology, phase, and growth characteristics of secondary materials in the various synthesis methods. Since the presence of termination groups on MXene surfaces is dependent on the synthesis processes, it is thus crucial to carefully select the route of MXene synthesis, as well as the subsequent treatment steps, to develop MXene hybrids with high catalytic activity and structural stability.
MXENE HYBRIDS FOR WATER SPLITTING REACTIONS

MXenes’ unique set of physical and electronic properties make them excellent candidates for catalytic water splitting. High hydrophilicity and solution processability allow MXenes to be robust and durable in aqueous electrolytes such as acids, bases, and seawater. Metallic electronic conductivity enables MXenes to facilitate efficient electronic charge transport at the electrode-electrolyte interface as a large-area conductive 2D substrate when coupled to other highly active but less conductive co-catalysts. This combination of properties circumvent the limitations of conventional C-based supports such as graphene whose hydrophobicity restricts their use in aqueous electrocatalysis, and GO whose conductivity is reduced because of hydroxyl functionalization. Hybridization also creates additional catalytically active motifs at the interface and modulates the electronic structure to augment intrinsic catalytic activity. With precise control over the MXene hybrid’s interfaces and growth of secondary materials, bifunctional non-PGM MXene hybrid couples have outperformed full PGM couples (Pt/C as HER cathode, RuO₂ or IrO₂ as OER anode) in both activity and stability. Here, we highlight highly active and non-PGM HER, OER, and bifunctional water splitting MXene hybrid catalysts, clarifying the roles played by each material component in the overall catalyst design. For the reader’s benefit, a summary table of performance metrics for various MXene hybrids in EC HER/OER and PC/PEC HER is provided in Tables S2 and S3 respectively.

MXene hybrids and composites for electrocatalytic (EC) HER. MXenes have garnered significant attention after they were theoretically predicted and then demonstrated to be HER-active. Unlike other 2D materials like 2H-MoS₂ that exhibit higher catalytic activity on the edges, the 2D basal planes of MXenes are more HER active, allowing for more efficient catalyst utilization. HER activity in bare MXenes was reported experimentally using Mo₂CTₓ and
Ti$_2$CT$_x$ by Seh et al., with the –O T$_x$ surface groups determined as active sites on the MXenes’ basal planes.\textsuperscript{17} Due to the modest EC HER activity of bare MXenes, initial efforts were directed at improving their HER activity through physically increasing the density of electrochemically active surface area (ECSA) by nanostructuring,\textsuperscript{157,158} or by modulating the MXene electronic structure to achieve a more thermoneutral $^*\text{H}$ adsorption free energy ($\Delta G_{\text{H}} \to 0$), a common predictor for HER activity.\textsuperscript{154–156} Effective binding energy tuning may be achieved through T$_x$ control,\textsuperscript{18,154} and doping with other TMs\textsuperscript{127,159–162} and/or non-metals (such as N, S, and P).\textsuperscript{163–165}

The effect of TM doping in changing $^*\text{H}$ binding energy was recently illustrated by Djire et al. using scanning electrochemical microscopy (SECM) on Ti$_4$N$_3$T$_x$ MXenes.\textsuperscript{19} SECM is a unique tool capable of generating high-resolution local conductivity and EC activity maps of the catalyst surface. In the feedback mode, ferrocene (Fc), a redox mediator, is included to probe the surface conductivity. A higher tip current ($i_T$) will be observed on a more conductive surface due to a faster rate of Fe$^+$ reduction. As the MXene flakes are placed on top of a conductive substrate like indium tin oxide (ITO), $i_T$ contrast also demarcates the boundaries of the MXene flake edges. The HER activity can be mapped by applying a negative bias to the substrate, referred to as the substrate generation/tip collection (SG/TC) mode.

Alloying Ti$_4$N$_3$T$_x$ with V, which produced V-alloyed Ti$_4$N$_3$T$_x$ (denoted as y M V-Ti$_4$N$_3$T$_x$, where y = molar concentration of aqueous VOSO$_4$ dopant used for alloying), did not improve the surface conductivity, as shown by the negligible $i_T$ in 0.3 M V-Ti$_4$N$_3$T$_x$, similar to bare Ti$_4$N$_3$T$_x$ (Figures 5a and 5c). However, the HER activity in V-alloyed Ti$_4$N$_3$T$_x$ increased very significantly compared to the bare MXenes (Figures 5b and 5d). Improved conductivity in V-Ti$_4$N$_3$T$_x$ was also be achieved by increasing the concentration of VOSO$_4$ dopant used (Figure 5e) and was accompanied by a slight improvement in HER activity (Figure 5f). This study by Djire et al. is
experimental proof of basal plane HER activity in MXenes, as inferred from the lower \( i_T \) around the flake edges in the SG/TC mode. Furthermore, it demonstrates that the conductivity and HER activity in MXenes may be decoupled, possibly due to the self-gating phenomenon, reported previously on the basal planes of a variety of semiconducting 2D materials, including 2H-MoS\(_2\).\(^{166}\)

**Figure 5.** SECM feedback mode and SG/TC mapping of (a, b) bare Ti\(_4\)N\(_3\)T\(_x\) and V-alloyed Ti\(_4\)N\(_3\)T\(_x\), using (c, d) 0.3 M or (e, f) 1.0 M aqueous VOSO\(_4\) dopant for alloying. Adapted with permission from ref \(^{19}\). Copyright 2020 WILEY-VCH.

Coupling MXenes with other HER-active co-catalysts such as TM chalcogenides,\(^{102,167}\) phosphides,\(^{168–170}\) carbides,\(^{151,171}\) and QDs (metallic alloys and black phosphorus, BP) are common ways to enhance HER activity.\(^{52,53}\) Here, MXenes serve as a large-area conductive and mechanical support to facilitate charge transfer, act as a HER co-catalyst, and modulate the hybrid’s electronic structure to optimize for H\(^+\) adsorption and HER activity. With significantly higher HER activities than the sum of their individual material components, intimately coupled MXene hybrids benefit from synergistic HER enhancement to achieve PGM-like activities.
Coupling MXenes with TM chalcogenides for EC HER was reported by Wu et al. in a hierarchical MoS$_2$/Ti$_3$C$_2$Tx@C architecture (Figure 4a). The MXene support constrained 2H-MoS$_2$ to grow vertically, maximizing ECSA and mass transport of gaseous products (Figure 6a). The strong interfacial and chemical relationship between Ti$_3$C$_2$Tx and 2H-MoS$_2$ was verified by the Mo (MoS$_2$)-C (MXene) X-ray photoelectron spectroscopy (XPS) peak, which would be absent in a physical mixture of Ti$_3$C$_2$Tx and 2H-MoS$_2$. With a small charge transfer resistance of 1.28 $\Omega$, MoS$_2$/Ti$_3$C$_2$Tx@C required a small overpotential ($\eta_{f=10}$) of 135 mV at a current density of -10 mA cm$^{-2}$ in 0.5 M H$_2$SO$_4$, 217 mV smaller than the MoS$_2$/rGO@C hybrid when rGO is used as the substrate (curves 1 and 2, Figure 6b, see Vocabulary for a definition of overpotential and Tafel slope). The nanohybrids were also durable in acid for 20 h of operation at -130 mV (vs. reversible hydrogen electrode, RHE), due to the protective carbon nanoplating surrounding the hybrid. Additionally, coupling other TM chalcogenides such as MoS$_2$, MoSe$_2$, NiS$_2$, NiSe$_2$, and VS$_2$ to various MXenes have also achieved synergistically enhanced HER activity.

TM carbides and phosphides have also been integrated into MXene hybrids for EC HER. TM phosphides will be discussed in the next sub-section due to their HER and OER bifunctionality. A collaborative catalytic interface was formed between the cobalt-doped $\beta$-Mo$_2$C co-catalyst, Ti$_3$C$_2$Tx, and N-doped carbon (NC) to enhance electrical conductivity and expose ECSA, while improving H$^+$ and water absorption kinetics for efficient HER in a wide pH range, including in seawater. Theoretical calculations revealed that the Co$_x$Mo$_{2-x}$C/NC interface reduced the H$^+$ absorption barrier to 0.05 eV, and the water dissociation barrier to 0.78 eV. Notably, the water dissociation barrier was lower than Pt (111) at 0.89 eV. With accelerated water dissociation and HER kinetics, the optimized Co$_{0.31}$Mo$_{1.69}$/MXene/NC hybrid exhibited Pt-like HER activity with an $\eta_{f=10}$ and Tafel slope of 75 mV and 32 mV dec$^{-1}$, respectively, in 1 M KOH.
(curve 2, Figure 6c), outperforming the individual β-Mo₂C and MXene components (curves 4 and 6, Figure 6c). This corroborated with a separate report on Mo₂C/Ti₃C₂Tₓ@NC, whose exchange current density ($j_0$) in acid was four-fold higher than Mo₂C@NC without the MXene support.¹⁷¹ These works demonstrate the synergistic co-operation between material components, with β-Mo₂C and NC providing favorable H⁺ adsorption sites,¹⁵¹ while MXenes facilitated charge transport.¹⁷¹

QDs have also been anchored on MXenes to modulate the electronic structure or alter the adsorption kinetics of intermediates. Du et al. modified the Ti₃C₂Tₓ surface with Ni/Co alloyed QDs to reduce the overly strong Ti-H interaction.¹⁷⁷ Nb was also doped into Ti₃C₂Tₓ to raise the Fermi level to the conduction band, improving the electronic conductivity of the Ti₃C₂Tₓ support. The optimized Ni₀.₉Co₀.₁@Ti₂.₅Nb₀.₅C₂Tₓ hybrid showed $\eta_{f=10}$ of 43 mV for HER, a significant improvement compared to the 516 mV required for Ti₂.₅Nb₀.₅C₂Tₓ without the alloyed QDs. Critically, the HER enhancement was intrinsic, given that the HER improvement trends were still valid after normalization for ECSA. Separately, BP QDs were decorated on Ti₃C₂Tₓ to similarly reduce Ti₃C₂Tₓ surface binding to H⁺ from -1.01 eV to as low as -0.17 eV (Figure 6d).⁵³ Beneficially, the MXene support physically confined the growth of BP to discrete 0D QDs, which prevented agglomeration, and maximized ECSA and mass transport for electrocatalysis.⁵²⁵³
Figure 6. MXene hybrids for EC water splitting. (a) TEM image and (b) polarization curves of MoS2/Ti3C2Tx-MXene@C, showing vertical growth of MoS2 on Ti3C2Tx. Adapted with permission from ref 70. Copyright 2017 WILEY-VCH. (c) Polarization curves of Co9Mo2-xC/MXene/NC. Adapted with permission from ref 151. Copyright 2019 WILEY-VCH. (d) Free energy diagram of BP QDs/Ti3C2Tx. Adapted with permission from ref 53. Copyright 2018 Royal Society of Chemistry. (e) Polarization curves and (f) Constant current (-10 mA cm⁻²) stability test of FeNi-LDH/Ti3C2Tx-MXene. Adapted with permission from ref 126. Copyright 2017 Elsevier Ltd. (g) Assembly and (h) Scanning electron microscope (SEM) image of 3D CoP-MXene. Adapted with permission from ref 169. Copyright 2018 American Chemical Society.

**MXene hybrids and composites for electrocatalytic (EC) OER and water splitting.**

Compared to HER, OER is a more complex reaction as it involves a four-electron transfer process and multiple reaction intermediates. Achieving efficient and durable OER often require electrocatalysts with intricate heterostructuring and design.178 Although bare MXenes are not efficient OER catalysts,45 they can serve as electrically conductive supports, modulate the
electronic density of their coupled OER co-catalyst, and even create additional OER-active motifs to greatly enhance OER activity.\textsuperscript{44,139} For instance, coupling Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} with g-C\textsubscript{3}N\textsubscript{4} created a highly OER-active Ti (MXene)-N\textsubscript{x} (g-C\textsubscript{3}N\textsubscript{4}) motif at the interface for PGM-like OER activity.\textsuperscript{139} Many OER-active materials such as TM chalcogenides,\textsuperscript{102,120} oxides,\textsuperscript{131,179} nitrides,\textsuperscript{180} phosphides,\textsuperscript{168–170} MOFs,\textsuperscript{107} LDHs,\textsuperscript{126,153,181} C-based materials,\textsuperscript{139,182} and QDs\textsuperscript{53,183} have been coupled with MXenes to augment OER activity. More significantly, these strategies have yielded non-PGM MXene hybrids that have outperformed PGM-based electrocatalysts for EC OER and water splitting.

Heterostructured and hybrid materials are often subject to lattice mismatch and strain at the interface, which has been exploited in phase engineering to alter electronic states near the interface and stabilize metastable material phases.\textsuperscript{184,185} Li \textit{et al.} stabilized a mixture of metastable 1T (23%) and stable 2H (77%) MoSe\textsubscript{2} on Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} as a bifunctional HER and OER electrocatalyst.\textsuperscript{102} The 1T/2H MoSe\textsubscript{2}/MXene hybrid (HER $\eta_{j=10} = 95$ mV) outperformed their individual components and other phases (2H MoSe\textsubscript{2}/MXene HER $\eta_{j=10} = 123$ mV) for both HER and OER. The non-PGM 1T/2H MoSe\textsubscript{2}/MXene couple also required a modest 1.64 V (vs. RHE) to maintain a current density of -10 mA cm\textsuperscript{-2} for 50 h while retaining a high 1T phase content. The activity improvement was attributed to the use of conductive MXenes to improve charge transfer and confine the 1T/2H MoSe\textsubscript{2} nanostructures to prevent aggregation. The intrinsic activity of MoSe\textsubscript{2} was also boosted by the abundance of MoSe\textsubscript{2} active sites, particularly from the more catalytically active and now stabilized 1T phase. This work illustrates the potential of using MXenes to stabilize metastable phases that are highly electrocatalytically active, as co-catalysts to achieve higher activities.\textsuperscript{102}

Although LDHs are excellent candidates for alkaline OER and water splitting, they are poor electrical conductors and aggregate easily.\textsuperscript{126,153,181} Previously, LDHs were used with C-based supports such as graphene to improve charge transport. These C-based supports, however, had to
be first functionalized to rGO/GO before they could be integrated with LDHs, compromising the support’s structural integrity and conductivity. Therefore, Yu et al. exploited the negatively charged surface $T_x$ groups of MXenes to electrostatically self-assemble FeNi-LDH nanostructures vertically on Ti$_3$C$_2$T$_x$ in a 3D array (Figure 4b). The vertical growth of LDH atop MXenes reduced repulsion and aggregation between positively charged LDH sheets while maximizing ECSA for catalysis. The benefits of strong chemical coupling within the hybrid was evident in their OER performance (Figure 6e) and stability (Figure 6f), which are far superior to the physical mixture (FeNi-LDH + Ti$_3$C$_2$T$_x$) or when the FeNi-LDH OER catalyst was hybridized with a less conductive substrate (FeNi-LDH/rGO) instead of MXenes. The reasoning for the enhanced OER is attributed to the increased oxidation state of Ni and Fe centers after hybridization, as shown in the +0.5 eV shift of the Ni and Fe 2p XPS peaks. A follow-up work integrated the FeNi-LDH/Ti$_3$C$_2$T$_x$ structure with a 3D nickel foam network to further maximize ECSA and mass transport, achieving a high current density of 500 mA cm$^{-2}$ at very low overpotentials for alkaline OER (300 mV), HER (205 mV), and bifunctional water splitting (1.75 V vs. RHE). Separately, extensive growth of Co-LDH on Ti$_3$C$_2$T$_x$ was successful in arresting Ti$_3$C$_2$T$_x$ surface oxidation during OER, which would have impeded both charge transfer at the MXene-LDH interface, and overall electronic conductivity in the hybrid.

The LDH-MXene hybrids can also be processed further to form bifunctional catalysts like phospho-sulfides. For example, FeNi-LDH/MXenes can be converted to 2D bimetallic Ni$_x$Fe$_{1-x}$PS$_3$ (MPS$_3$)/MXenes hybrid via thermal annealing with red phosphorus in a vacuum sealed ampule. The resulting MPS$_3$/MXenes hybrid possess a broken-up nanomosaic morphology with high ECSA, possibly due to the thermal stress. More importantly, the MPS$_3$/MXenes composition can be easily tailored by tuning the starting Fe : Ni ratio in the FeNi-LDH/MXenes precursor.
Interestingly, the catalytic reaction preference of MPS₃ hybrids depends on its Fe : Ni ratio – Ni₀.₉Fe₀.₁PS₃/Ti₃C₂Tₓ was found to be most proficient in catalyzing HER (ηᵢ=1₀ = 196 mV) in 1 M KOH, while Ni₀.₇Fe₀.₃PS₃/Ti₃C₂Tₓ exhibited excellent OER activity (ηᵢ=1₀ = 282 mV). The Ni₀.₇Fe₀.₃PS₃/Ti₃C₂Tₓ (OER) || Ni₀.₉Fe₀.₁PS₃/Ti₃C₂Tₓ (HER) couple can deliver -10 mA cm⁻² of complete water splitting at a lower potential (1.65 V vs. RHE) than the full-PGM IrO₂ || Pt/C reference standard (1.71 V vs. RHE). The MPS₃/MXenes couple was also stable for over 50 h at the said potential. This work demonstrates strain engineering for ECSA maximization, and the added flexibility of tuning multi-metal compounds to optimize toward different electrochemical reactions of interest as multi-functional electrocatalysts.

TM phosphides have also been integrated with MXenes as bifunctional HER and OER electrocatalysts.¹⁶⁸⁻¹⁷⁰ Xiu et al. reported a hierarchical CoP@3D Ti₃C₂Tₓ-MXene architecture to significantly enhance EC activity for water splitting.¹⁶⁹ After nebulizing MXene colloids containing cobalt salts into tiny droplets, the droplets were dried rapidly at 600 °C. Solvent evaporation induced an inward capillary force on the droplets to cause isotropic compression of MXene sheets into aggregation-resistant Co₃O₄@3D Ti₃C₂Tₓ-MXene hybrids (Figure 6g). Further phosphorization converted Co₃O₄ in the hybrids into OER-active CoP. The highly porous and interconnected 3D MXene architecture (Figure 6h) provided a large surface area for catalysis and facilitated long-range electrical charge transport. An efficient bifunctional HER and OER electrocatalyst, CoP@3D Ti₃C₂Tₓ-MXene required only 1.565 V (vs. RHE) to supply -10 mA cm⁻² of current density and remained stable (< 25 mV degradation) for 150 h at a high current density of -100 mA cm⁻². Combining TM phosphides with MXenes is also reported to shift the TM d-band center to higher energies, resulting in reduced occupancy of antibonding states from the TM and adsorbed O intermediate, for enhanced OH⁻ adsorption and OER.¹⁶⁸ Additionally, 1D CoP
nanorods grown on top of Ti$_3$C$_2$T$_x$ sheets was also reported to induce tensile stress and bending of Ti$_3$C$_2$T$_x$ to expose higher ECSA for improved OER activity.$^{170}$

**MXene hybrids and composites for photocatalytic (PC) and photoelectrocatalytic (PEC) water splitting.** Given the close alignment of the MXenes’ Fermi energy ($E_F$) position to the H$^+$/H$_2$ reduction potential,$^{187}$ efficient H$^+$ adsorption, and respectable EC water splitting performances as described earlier, MXene hybrids are a logical choice for PC/PEC water splitting to harvest sunlight as a renewable energy driving source. Unlike EC, PC reactions require semiconductor photo-absorbers, where electron-hole pairs are generated. Since most MXenes are metallic and not semiconducting,$^{152}$ other materials are necessarily drafted to harvest light and promote the separation of photo-generated carriers.$^{188}$ A few works feature MXene hybrids as multi-functional PEC HER, OER, and water splitting catalysts. Lin et al. reported a Ti$_3$C$_2$T$_x$/InGaN photo-anode with a 10-fold improvement in HER photocurrent density after MXene hybridization, verifying the high HER activity on the MXene surface.$^{189}$ Ti$_3$C$_2$T$_x$/TiO$_2$/NiFeCo-LDH was also tested for PEC OER, with the MXene and NiFeCo-LDH acting as the conductive support and OER catalyst, respectively.$^{181}$ Yu et al. reported Ti$_3$C$_2$T$_x$/TiO$_2$ as an OER photo-anode with a six-fold increment in applied bias photon-to-current efficiency compared to a bare TiO$_2$ photoanode.$^{80}$ Here, PEC OER occurred on the MXene surface, which served as a hole scavenger and OER active site.

As discussed previously, MXenes generally exhibit only modest OER activity and hence rely on hybridization with OER-active catalysts (such as TM chalcogenides$^{102}$ and LDHs$^{126,149}$), with the MXenes serving primarily as conductive supports and electronic modulators.$^{44,139}$ Thus, most PC/PEC applications of MXenes and their hybrids are directed at HER instead of OER to take advantage of MXenes’ HER activity. In this endeavor, MXenes, especially Ti-based MXenes, have
been employed as precursors to TM oxides with rich vacancies, which are widely considered as stable, efficient, and robust HER photocatalysts.\textsuperscript{83,97,100}

Yuan \textit{et al.} oxidized Ti$_3$C$_2$Tx in CO$_2$ at high temperatures to produce a 2D MXene-derived C/TiO$_2$ hybrid.\textsuperscript{83} Since the Ti and C atoms were already alternately arranged in the MXene structure (Ti$_3$C$_2$Tx), calcination is likely to result in alternating TiO$_2$ and 2D carbon sheets, while retaining the large-area 2D-layered structure of MXenes, which is suitable for PC HER (Figure 7a). After optimizing for calcination temperature and CO$_2$ flow rates, C/TiO$_2$-700-150 hybrid exhibited the highest PC H$_2$ generation rate of 24 \textmu mol g$^{-1}$ h$^{-1}$, representing an 89-fold enhancement from TiO$_2$. This was credited to the thin carbon layers facilitating charge transport, whose high electrical conductivity improved photo-carrier separation efficiency and reduced their recombination rate, in agreement with other works.\textsuperscript{100,190,191} The thin carbon layers also provided a high degree of transparency for visible light penetration to the TiO$_2$ photo-absorber. This method was also employed to synthesize S-doped TiO$_2$ NPs/C (Figure 7b) from S-doped Ti$_3$C$_2$Tx, whereby S doping further optimized H$^+$ binding energy for efficient HER.\textsuperscript{97} The strategy of using MXenes as precursors to synthesize MXene/TM oxide hybrids has also been applied to other MXenes, given the relative ease in oxidizing MXenes,\textsuperscript{192,193} such as in the oxidation of Nb$_2$CT$_x$ to Nb$_2$CT$_x$/Nb$_2$O$_5$,\textsuperscript{194} Mo$_2$CT$_x$ to Mo$_2$C/MoO$_x$,\textsuperscript{148} and V$_2$CT$_x$ to V$_2$CT$_x$/V$_2$O$_x$.\textsuperscript{57,195}

Apart from acting as TM oxide precursors, MXenes can also serve as efficient PC/PEC HER co-catalysts themselves to promote photo-generated charge carrier separation when coupled to semiconductors.\textsuperscript{133,196–199} The most widely studied MXene, Ti$_3$C$_2$T$_x$, has been applied as a PC HER co-catalyst and successfully coupled to a variety of semiconducting photocatalysts such as TM oxides,\textsuperscript{191,196} sulfides,\textsuperscript{124,187} perovskites,\textsuperscript{197,198} and g-$C_3$N$_4$.\textsuperscript{133,199}
For instance, Ran et al. investigated the use of Ti$_3$C$_2$Tx as a HER co-catalyst, with CdS acting as a photo-absorber for PC HER.\textsuperscript{187} The electrostatic self-assembly of CdS NPs on Ti$_3$C$_2$Tx NPs produced cauliflower-structured sub-microspheres (referred to as CTX, where X refers to the mass ratio of MXenes : CdS) with distinct and well-formed Ti$_3$C$_2$Tx/CdS interfaces (Figure 7c). An optimal 2.5 wt\% MXene incorporation (CT2.5) resulted in a substantial 136-fold improvement in H$_2$ production rate compared to bare CdS (CT0, Figure 7d). In fact, the strongly coupled Ti$_3$C$_2$Tx/CdS (CT2.5) hybrid outperformed other hybrids such as 2.5 wt\% Pt-CdS, NiS-CdS, Ni-CdS and MoS$_2$-CdS (Figure 7d). The outstanding PC HER activity of CT2.5 was attributed to a reduction in interfacial charge transfer resistance arising from intimate chemical coupling at the Ti$_3$C$_2$Tx/CdS interface, and high metallic conductivity of Ti$_3$C$_2$Tx. Furthermore, the photo-carrier separation efficiency was enhanced by deliberately matching the energy bands of Ti$_3$C$_2$Tx and CdS in a Schottky junction to trap photo-generated electrons and limit carrier recombination. This was observed in the suppression of both the radiative recombination photoluminescence (PL, $\lambda \approx$560 nm, Figure 7e), and defect level recombination (red-shifted tail from 650–700 nm in Figure 7e) peaks, resulting in a longer overall photo-carrier lifetime (Figure 7f). Finally, theoretical calculations revealed that –O terminations in Ti$_3$C$_2$Tx served as active HER sites due to their favorable $E_F$ position and efficient H$^+$ adsorption. Ti$_3$C$_2$Tx was also further hybridized with other semiconductor photocatalysts such as ZnS and Zn$_{0.8}$Cd$_{0.2}$S and similar trends in improved PC HER activity was observed. This work thus demonstrates the use of O-terminated Ti$_3$C$_2$Tx as efficient and durable PC HER co-catalysts for synergistic coupling with a variety of earth-abundant semiconductor photocatalysts (e.g. CdS, ZnS, Zn$_{1-x}$Cd$_x$S) to achieve significantly improved PC HER performances for PC water splitting.
Figure 7. MXene hybrids for PC HER. (a) SEM of Ti$_3$C$_2$Tx-derived 2D-layered C/TiO$_2$. Adapted with permission from ref $^{83}$. Copyright 2017 WILEY-VCH. (b) High resolution TEM (HRTEM) of Ti$_3$C$_2$Tx-derived S-TiO$_2$/C. Adapted with permission under a Creative Commons CC-BY license from ref $^{97}$. Copyright 2018 The Authors. (c) HRTEM of 2.5%-Ti$_3$C$_2$Tx-Cds hybrid (CT2.5). (d) PC HER activity of various co-catalysts hybridized with Cds. (e) PL and (f) time-resolved PL spectrum of bare Cds (CT0), and CT2.5 MXene hybrids. Adapted with permission under a Creative Commons CC-BY license from ref $^{187}$. Copyright 2017 The Author(s).

Summary, specific challenges, and outlook for EC, PC, and PEC water splitting. In this section, we reviewed the various materials hybridized with MXenes for EC/PC/PEC HER, OER, and water splitting. A summary table of performance metrics ($\eta$, Tafel slope, and turnover rate) for various MXene hybrids in EC HER/OER and PC/PEC HER is provided in Tables S2 and S3 respectively. In summary, MXenes play various critical roles: (1) providing a large-area conductive and structural support, (2) acting as a HER co-catalyst, (3) confining, stabilizing, and thus preventing the hybridized material from aggregating on the MXene support, and (4) inducing charge transfer to modulate the electronic band structure of the hybrid material to optimize $\text{H}^+$, $\text{OH}^-$, and water adsorption kinetics when strongly and chemically coupled to other materials.
Lattice strain induced at the MXene-secondary material interfaces can also alter electronic density and stabilize metastable co-catalyst material phases through strain and phase engineering. Apart from activity enhancement, MXene hybrids also benefit from improved stability in acidic and alkaline electrolytes (Figure 5f). A strong and chemically coupled MXene hybrid, especially at the hybrid interface, is important for prolonged cycling and operational stability, particularly at higher current densities. The NiFe-LDH/Ti₃C₂Tₓ/3D nickel foam hybrid was stable for 200 h of continuous water splitting in 1 M KOH at a high current density of 100 mA cm⁻² due to the strong chemical integration between the LDH, MXenes, and 3D nickel foam network. Conversely, Mo₂CTₓ/2H-MoS₂ hybrid suffered a significant ≈50% loss in operating current density after a short 8 h of HER. This was due to crack formation within the hybrid, and MoS₂ dislodgement during HER, possibly due to poor chemical integration between Mo₂CTₓ and 2H-MoS₂.

Presently, non-PGM MXene hybrid couples have outperformed full-PGM couples (IrO₂ and RuO₂ for OER, Pt/C for HER) for EC water splitting due to the exceptional OER activity from the non-MXene OER-active components (TM phosphides, LDHs). From an activity standpoint, more work should thus be done on improving non-PGM MXene hybrids for HER, and this begins by working with Mo-based MXenes, which possess a higher intrinsic HER activity compared to Ti-based MXenes. Concurrently, further improvement in the OER competitiveness of MXene hybrids can be realized through the theory-guided development of MXenes with higher intrinsic OER activities such as nitride MXenes. Lastly, the use of operando and localized electrochemical/physical characterization techniques such as scanning electrochemical cell microscopy (SECCM) can elucidate more information on catalytic activity at different areas of the MXene hybrid, of which regions near and at the hybrid interface is of particular interest for further
These investigations are of particular importance to MXene hybrids for OER, since MXenes can be readily oxidized to TM oxides under the applied oxidizing potentials.
MXENE HYBRIDS AS ELECTROCATALYSTS FOR METAL-AIR AND METAL-SULFUR BATTERIES

Aside from their high HER and OER catalytic activity for water splitting, MXene hybrids are also similarly efficient in catalyzing the OER/ORR redox couple\textsuperscript{203,204} to serve as cathodes for metal-air batteries.\textsuperscript{180,205–207} With high metallic conductivity, MXene hybrids are well-positioned to facilitate rapid electronic transfer for sulfur redox reactions at the sulfur cathode in lithium-sulfur (Li-S) batteries as well.\textsuperscript{56,208} MXene hybrids have recently been demonstrated to possess a high binding affinity toward deleterious lithium polysulfide (LiPS) side products when used as a separator, extending the cycle life and reducing the capacity fade of Li-S batteries.\textsuperscript{209,210} In this section, we review the catalytic role of MXene hybrids towards realizing the industrial potential of metal-air and metal-sulfur batteries (see \textbf{Vocabulary} for full definition) as next-generation battery architectures to complement existing lithium ion batteries (LIBs).

\textbf{MXene hybrids and composites for electrocatalytic (EC) OER and ORR in metal-air batteries.} A post-LIB technological prospect, metal-air batteries comprise of design elements from both traditional batteries (using a metal anode) and fuel cells (using a cathode exposed to air).\textsuperscript{8} Metal-air batteries have attracted widespread interest due to their compact size, lightweight composition,\textsuperscript{211} and high theoretical energy density.\textsuperscript{212} For instance, Zn-air batteries have a theoretical energy density (1350 Wh kg$^{-1}$, excluding oxygen) about five times that of LIBs,\textsuperscript{212} and can be produced at significantly lower costs (<$10$ kW$^{-1}$ h$^{-1}$) than LIBs ($\approx$400–800 kW$^{-1}$ h$^{-1}$).\textsuperscript{213}

Unfortunately, the theoretical promise of metal-air batteries has yet to be practically realized due to their poor air cathode performance. The complex, multi-step, four-electron OER/ORR redox couple is severely hampered by sluggish kinetics as the rate-determining step.\textsuperscript{211}
Even though MXenes are not OER or ORR-active as previously discussed, MXenes have been utilized as conductive supports for other ORR-active electrocatalysts.\textsuperscript{182,206} Hybridizing MXenes with other materials also resulted in electronic structure modulation, and the creation of catalytic motifs to exhibit enhanced OER, ORR, and bifunctional OER/ORR activity comparable to PGMs such as Pt/C (for ORR) and IrO\textsubscript{2}/RuO\textsubscript{2} (for OER).\textsuperscript{123,139,180} Since MXene hybrids for EC OER have been described previously for water splitting, we focus our discussions here on MXene hybrids for EC ORR, bifunctional EC OER/ORR, and their use in Zn-air batteries.\textsuperscript{180,205–207} A summary of EC ORR performances for various MXene hybrids is provided in Table S2c.

Iron-nitrogen-carbon (Fe-N-C) materials are one of the most promising class of PGM-free ORR catalysts due to their highly active FeN\textsubscript{4} motifs.\textsuperscript{123,203,214} To further augment the intrinsic ORR activity of iron phthalocyanine (FePc), Li et al. hybridized FePc on Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} support and obtained a significant two-fold activity enhancement in FePc, which was also five times higher than Pt/C.\textsuperscript{123} Two-electron ORR was also virtually suppressed with a calculated electron number of \(\approx 4\) and an ultralow H\textsubscript{2}O\textsubscript{2} yield (<1\%). The authors posit that the negatively charged –OH/F T\textsubscript{x} groups of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} interacted strongly with the Fe\textsuperscript{2+} center through intermolecular van der Waals forces or hydrogen bonding to reduce the local Fe\textsuperscript{2+} electron density, which induced a transition to a higher spin state to optimize O\textsubscript{2} binding for ORR. A follow-up work synthesized Fe(phen)\textsubscript{2}/Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} hybrids through pyrolysis, using the MXene as a large-area 2D conductive support to prevent aggregation of the rugged Fe(phen)\textsubscript{2} coating.\textsuperscript{214} Separately, positively charged 2D Fe-N-C sheets were produced by decorating g-C\textsubscript{3}N\textsubscript{4} sheets with Fe NPs.\textsuperscript{203} By manipulating the Fe/g-C\textsubscript{3}N\textsubscript{4} precursor ratio, the positive zeta potential (from Fe\textsuperscript{2+}) per unit area of Fe-N-C sheets was adjusted to +30.4 mV to match the negative zeta potential (from MXenes’ T\textsubscript{x} groups) per unit area of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} sheets (-39.7 mV). This allowed both sheets to be electrostatically self-assembled...
in a 2D/2D layered superlattice-like heterostructure in aqueous solutions with enhanced ORR activity. This work can potentially be further extended to manipulate the positive sheet charges of other 2D materials using TM NP clusters to achieve the wet chemical, electrostatic self-assembly of 2D material/MXene layered heterostructures, circumventing the use of small-scale and energy intensive physical deposition methods such as CVD and ALD.

TM oxide\textsuperscript{207} and chalcogenide\textsuperscript{206} NPs have also been incorporated onto MXene scaffolds to synergistically enhance ORR activity for Zn-air batteries (Figure 8a). For example, 3D N-CoSe\textsubscript{2}/MXene architecture can be obtained with the aid of a poly(vinylalcohol) (PVA) cross-linker, followed by thermal annealing in NH\textsubscript{3}.\textsuperscript{206} The 3D architecture separates the MXene layers and prevents them from aggregation, thus maximizing ECSA and mass transport properties. Theoretical calculations revealed that N-CoSe\textsubscript{2}/MXene hybridization increases the activity at Co sites, by virtue of significant electron transfer from Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} to CoSe\textsubscript{2}, while N-doping reduces the thermodynamic barrier to the potential-limiting steps for both OER and ORR. This resulted in a highly efficient bifunctional OER/ORR electrocatalyst, with a higher specific capacity (751 mAh g\textsubscript{Zn}\textsuperscript{-1}) than the RuO\textsubscript{2}+Pt/C OER/ORR benchmark (690 mAh g\textsubscript{Zn}\textsuperscript{-1}) when discharged at 5 mA cm\textsuperscript{-2}. The discharge characteristics and maximum power density of N-CoSe\textsubscript{2}/3D Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} (142 mW cm\textsuperscript{-2} at 232 mA cm\textsuperscript{-2}) also outperforms the full PGM standard reference of RuO\textsubscript{2} + Pt/C (119 mW cm\textsuperscript{-2} at 168 mA cm\textsuperscript{-2}), as seen in Figure 8b. The high cycling stability (500 cycles, 166 h at 10 mA cm\textsuperscript{-2}) was attributed to the preservation of the 3D MXene architecture and strong integration of CoSe\textsubscript{2} NPs into the MXene network, which prevented CoSe\textsubscript{2} NPs from dislodging during cycling.
Figure 8. MXene hybrids as air cathodes for (a) Zn-air batteries. (b) Discharge and power density curves of N-CoSe$_2$/3D Ti$_3$C$_2$Tx cathode, compared against Pt/C + RuO$_2$. Adapted with permission from ref 206. Copyright 2019 American Chemical Society. (c) Charge-discharge polarization curves, (d) specific capacities (calculated with Zn mass consumed) and (e) galvanostatic charge-discharge cycling of NiFeMn nitride/N-Ti$_3$C$_2$Tx cathode, compared against Pt/C + RuO$_2$. Adapted with permission from ref 180. Copyright 2020 American Chemical Society.

Recently, fiber-shaped flexible Zn-air batteries were developed with molecularly thin trimetallic NiFeMn nitride sheets stabilized by Ti$_3$C$_2$Tx, exhibiting bifunctionality toward both OER and ORR. The stabilization effect was supported by theoretical calculations, while strong interfacial coupling between the nitride and Ti$_3$C$_2$Tx sheets was marked by the formation of ORR-active motifs (Ni/Fe/Mn-N-Ti) from the Ti (MXene)-N (trimetallic nitride) peak in the Ti 2p XPS spectrum. The ECSA for ORR/OER was maximized by stabilizing the 2D nitride sheets atop the MXene sheets, which facilitated efficient charge transport for electrocatalysis. This resulted in synergistic OER and ORR activity enhancement of the nitride/N-Ti$_3$C$_2$Tx hybrid after Ti$_3$C$_2$Tx incorporation, despite standalone Ti$_3$C$_2$Tx being practically OER and ORR-inactive. The potential difference between the OER $\eta_{10}$ and ORR half-wave potential ($E_{1/2}$) was smaller for the nitride/N-
Ti₃C₂Tex hybrid (0.68 V) than RuO₂ + Pt/C (0.80 V), illustrating the hybrid’s superior OER/ORR bifunctionality without the use of PGMs. When used in Zn-air battery applications, the bifunctional nitride/N-Ti₃C₂Tex air cathode operated with a smaller charge-discharge voltage gap (hence better rechargeability, Figure 8c), and higher specific capacity (Figure 8d, 630 mAh g⁻¹ Zn⁻¹) than RuO₂ + Pt/C. The hybrid also demonstrated long-term cycling stability for 120 cycles (120 h at 20 mA cm⁻²) with no appreciable voltage fade, in comparison to the large voltage increase and hence instability of the RuO₂ + Pt/C air cathode during the same cycling test (Figure 8e). After being incorporated into a unique fibrous design, the flexible Zn-air battery operated durably for more than 3 h under various mechanical deformation states for wearable technology applications.

Summary, specific challenges, and outlook for metal-air batteries. MXene hybrid metal-air batteries are still in their early stages of development, with predominantly Ti₃C₂Tex-based MXene hybrids being explored for Zn-air batteries. Theoretically, a high efficiency for MXene-derived catalyst supports has been predicted for Li-air batteries due to the MXenes’ ability to adsorb Li species to facilitate interfacial charge transfer and redox reactions.²¹⁵ We thus expect MXene hybrids to be explored for other metal-air chemistries such as Li, Na, Mg and Al, using other highly conductive MXenes apart from Ti₃C₂Tex.

Beyond metal-air batteries, the high ORR activity of MXene hybrids can also be translated to effective PC pollutant degradation and environmental remediation, an essential application that can be readily deployed in remote and less developed areas. For example, Ti₃C₂Tex/TiO₂,⁴⁷ Ti₃C₂Tex/Ag₃PO₄,¹²² and In₂S₃/TiO₂@Ti₃C₂Tex⁴⁸ hybrids have demonstrated very high activity and durability for PC removal of organic dyes and inorganic pollutants.⁴²,⁴⁴ By combining strategies for designing efficient water splitting photocatalysts in the previous section (maximizing photo-carrier separation and minimizing carrier recombination) with the strategies discussed here in
designing efficient ORR electrocatalysts, we anticipate further strides to be made in MXene hybrid photocatalysts for broad-based pollutant remediation applications too.

**MXene hybrids and composites for metal-sulfur batteries.** Li-S batteries have been touted as the next-generation energy storage device for decades due to their unique conversion mechanism and high theoretical energy density (2500 Wh kg\(^{-1}\)).\(^{216}\) During discharge, the Li anode is oxidized to Li\(^+\) ions which travel to the S cathode, reacts with it, and forms Li\(_2\)S through a series of lithiation steps and intermediate LiPS species \((n \text{ Li}^+ + S_8 + n \text{ e}^- \rightarrow \text{soluble Li}_2S_8 \rightarrow \text{Li}_2S_6 \rightarrow \text{Li}_2S_4 \rightarrow \ldots \rightarrow \text{solid Li}_2S)\). In reality, the majority of the intermediate LiPS species (Li\(_2\)S\(_4\) to Li\(_2\)S\(_8\)) dissolves readily into the electrolyte before reaching the insoluble Li\(_2\)S phase, resulting in a lower discharge capacity and rapid capacity decay.\(^{216}\) The soluble LiPS species can further diffuse to the Li anode to be chemically reduced to shorter-chain LiPS before diffusing back to the cathode to be re-oxidized in an internal short circuit/self-discharge (shuttling effect).\(^{216}\) Furthermore, the poor ionic and electrical conductivity of the S cathode, coupled with a large volumetric expansion in the cathode upon lithiation (\(~80\%)\), imposes structural stress and inhibits electrochemical performance at the S cathode.\(^{216}\) To address these issues, S cathodes have been hybridized with C-based materials to improve conductivity, while polymeric binders, TM oxides and chalcogenides have been incorporated to improve LiPS adsorption.\(^{216}\) Separator membranes, commonly coated with conductive polymers, TM oxides, or C-based materials, have also been placed in the electrolyte between the electrodes to capture the shuttling LiPS.\(^{216}\)

With a unique combination of metallic conductivity and polar T\(_x\) groups, MXenes have been employed as S-host materials for both the S cathode\(^{56,208}\) and separator,\(^{209,210}\) as they can chemically interact with LiPS and facilitate efficient electronic transport at the cathode.\(^{14}\) Experimentally, highly conductive MXenes (Ti\(_2\)CT\(_x\),\(^{56,217}\) Ti\(_3\)C\(_2\)T\(_x\),\(^{56,218}\) Ti\(_3\)CNT\(_x\),\(^{56}\) and V\(_2\)CT\(_x\))
have been investigated as cathode materials. Theoretical studies have also revealed various MXene structures as good binders for LiPS chemisorption: M$_2$X MXenes (Ti$_2$CT$_x$, Ti$_2$NT$_x$, V$_2$CT$_x$), M$_3$X$_2$ MXenes (Ti$_3$C$_2$T$_x$, V$_3$C$_2$T$_x$, Cr$_3$C$_2$T$_x$, Zr$_3$C$_2$T$_x$, Nb$_3$C$_2$T$_x$, Hf$_3$C$_2$T$_x$), and ordered double TM MXenes (Mo$_2$TiC$_2$T$_x$). Nazar’s group determined that the LiPS species are strongly bonded to the Ti atoms in Ti$_2$CT$_x$ through Lewis acid-base Ti-S bonding at the expense of Ti-OH (T$_x$) interactions. The effect of MXenes’ surface chemistry on LiPS affinity was elucidated by density functional theory (DFT) calculations which concluded that –O and –F T$_x$ groups act as stronger immobilizers than –F in restricting LiPS diffusion in conventional electrolytes. Additionally, N-doping of Ti$_3$C$_2$T$_x$ further improved charge transport capacity and the Lewis acid-base binding strength to LiPS species. Since the MXene surface chemistry and doping character can be directly tuned by the synthetic method, we expect future work to utilize theory-guided calculations to screen dopants and T$_x$ groups, to complement current synthetic methods and deliberately modulate the LiPS immobilization strength for a variety of MXenes to act as efficient S-host cathodes and separators for Li-S batteries.

Beyond the high electrical conductivity and chemical affinity to LiPS, MXenes are also mechanically robust to accommodate the large volumetric changes in the S cathode during charge/discharge cycles. However, MXene nanosheets have a high tendency to self-restack, limiting the S/LiPS loading and restricting Li$^+$/electrolyte access at the S cathode. Thus, secondary materials have been integrated with MXenes as spacers to reduce MXene inter-layer re-stacking, while retaining the high electrical conductivity and LiPS binding affinity of MXenes. In this section, we highlight the use of MXene hybrids as S-host cathode and separator materials, illustrating how hybridization can augment the LiPS binding affinity to improve discharge capacity and impede capacity decay beyond that demonstrated by bare MXenes.
MXenes have been hybridized with TM chalcogenides\textsuperscript{224} and oxides\textsuperscript{105,208,225} to chemically interact with LiPS and expedite their catalytic conversion to Li$_2$S (Figure 9a). Zhang \textit{et al.} electrostatically self-assembled a 3D S/MnO$_2$@Ti$_3$C$_2$T$_x$ hybrid, taking advantage of the high surface area and conductivity of delaminated Ti$_3$C$_2$T$_x$ sheets, and the high chemical affinity of MnO$_2$ to LiPS.\textsuperscript{208} The 3D mesoporous conductive network enabled a high S loading of \textasciitilde70\% and could readily accommodate volumetric changes during charge/discharge cycles. Moreover, the intense O-Ti-S and O-S peaks in the O 1s XPS spectrum validated the strong MnO$_2$-MXene integration and LiPS retention ability of MnO$_2$. The S/MnO$_2$@Ti$_3$C$_2$T$_x$ cathode had a higher initial discharge capacity (1140 mAh g$_S^{-1}$, 0.05C) than the unhybridized components, which implies higher S utilization. The high cathode stability was demonstrated in the hybrid’s respectable specific capacity of 501 mAh g$_S^{-1}$ after 500 cycles at 1C, with a minor 0.06\% decay/cycle. Compared to the individual components (S/MnO$_2$ and S/Ti$_3$C$_2$T$_x$), the S/MnO$_2$@Ti$_3$C$_2$T$_x$ hybrid’s superior LiPS absorption was visually confirmed by the faintest coloration in the electrolyte after 4 h of cycling, indicating minimal LiPS dissolution into the electrolyte (Figure 9b).

Separately, the high LiPS binding affinity to other TM compounds such as TiO$_2$\textsuperscript{105,225} and MoS$_2$\textsuperscript{224} was validated experimentally. When used as symmetric cell electrodes in an electrolyte with Li$_2$S$_x$ species, Ti$_3$C$_2$T$_x$/1T-2H MoS$_2$-C hybrid afforded the highest redox current density, and hence the fastest rate of catalytic LiPS conversion, as compared to their separate components (Ti$_3$C$_2$T$_x$ and 2H-MoS$_2$) and the Ti$_3$C$_2$T$_x$/2H MoS$_2$-C hybrid (Figure 9c).\textsuperscript{224} This highlighted the importance of higher electrical conductivity (in 1T/2H-MoS$_2$) to facilitate electron transfer for LiPS redox conversion, as compared to using semiconducting 2H-MoS$_2$ for hybridization with MXenes. The 1T/2H-MoS$_2$ also contained a high concentration of edge defects and positively
charged S vacancies to bind strongly to negatively charged LiPS and facilitate electron transfer for rapid LiPS redox conversion.

Beyond TM compounds, C-based materials\textsuperscript{56,64,218} have also been hybridized with MXenes to further improve the cathode’s electronic conductivity\textsuperscript{56} for more efficient S utilization, and provide large surface areas to physically confine LiPS species.\textsuperscript{218} Interweaving Ti-based MXene nanosheets (Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}, Ti\textsubscript{2}CT\textsubscript{x}, and Ti\textsubscript{3}(CN)T\textsubscript{x}) with CNTs inhibited MXene inter-layer restacking to maximize Ti-S interactions for LiPS absorption, which led to stable cycling performances with high S loading of up to 5.5 mg cm\textsuperscript{-2} for high energy density Li-S batteries.\textsuperscript{56} Separately, Zhou \textit{et al.} treated Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} sheets with molten KOH to produce Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}/C hybrids with an expanded inter-layer spacing and significantly increased surface area (from 3.5 to 150 m\textsuperscript{2} g\textsuperscript{-1}).\textsuperscript{218} Hybridization with disordered C also reduced the charge transfer resistance by five-folds to accelerate LiPS conversion, increasing the calculated Li\textsubscript{2}S precipitation capacity from 95 to 226 mAh g\textsuperscript{-1}.\textsuperscript{218} Since TiO\textsubscript{2} was previously reported to improve LiPS binding efficiency,\textsuperscript{105,225} we predict that the partial oxidation of Ti-based MXenes to MXene/TiO\textsubscript{2}/C hybrids\textsuperscript{92} (similar to those for PC HER applications) may significantly enhance LiPS absorption and redox conversion kinetics, while providing a conductive, large-area support for LiPS deposition.

MXenes and their hybrids have also been incorporated in Li-S battery separators. As some MXenes are known to have high LiPS binding affinity, MXene-containing separators can immobilize parasitic LiPS shuttling between electrodes, thus limiting the internal short circuit effect and prolonging cycling stability.\textsuperscript{209,210,226} Similar to the cathode designs, MXenes have been hybridized with TM oxides (with high LiPS adsorption affinity)\textsuperscript{227} and C-based materials\textsuperscript{210,226} to act as efficient physical barriers at the separator facing the cathode. For example, Jiao \textit{et al.} produced TiO\textsubscript{2}@Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} hybrids from controlled partial oxidation of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}, to be mixed with
graphene and coated on a commercial Celgard polymer separator (Figure 9a).\textsuperscript{227} From the LiPS permeation test (Figure 9d), no significant darkening in the electrolyte was observed when the optimized (4 h oxidation) graphene-TiO\textsubscript{2}@Ti\textsubscript{3}C\textsubscript{2}Tx-polymer separator was used, in contrast to the untreated and graphene-coated separators. This validated the use of TiO\textsubscript{2} as LiPS adsorption centers, while MXenes provided a conductive large-area support to facilitate rapid LiPS conversion. When the separator was incorporated into a Li-S cell, only a minor 0.028\% capacity fade/cycle was observed over 1000 cycles at 2C (red curve, Figure 9e), with stable cycling also observed for over 200 cycles at a higher 7.3 mg cm\textsuperscript{-2} S loading.

**Summary, specific challenges, and outlook of metal-sulfur batteries.** The discovery of a strong Ti (MXene)-S (LiPS) bond,\textsuperscript{56} coupled with high metallic conductivity and mechanical flexibility, have enabled Ti-based MXenes to serve as large-area S-host materials in Li-S cathodes and separators. By hybridizing MXenes with other secondary materials with high affinity toward LiPS absorption, MXene hybrids have successfully enhanced LiPS capture and redox conversion kinetics to improve the discharge capacity and cycling stability. While not discussed in this section, MXenes have also been utilized as S-host materials for Na-S batteries,\textsuperscript{228} and have shown promise as alkali metal hosts to impede dendrite formation at the anode.\textsuperscript{229–232} We thus expect efforts to be made in the theoretical screening of potential MXene hybrid combinations, especially with conductive non-Ti-based MXenes, for LiPS binding affinity and catalytic conversion to Li\textsubscript{2}S. Combined with advanced synthetic designs in phase, defect, doping, and structural engineering of secondary materials on MXenes, future MXene hybrids could be applied in the cathode, anode, and separator to realize industrially relevant Li-S batteries with high discharge capacities, cycling stability, and S loading.
Figure 9. MXene hybrids as S cathodes and separators for (a) Li-S batteries. (b) Photographs of S/Mn$_2$O$_4$@Ti$_3$C$_2$T$_x$ S cathode during 4 h of charge-discharge (yellow colouration indicates LiPS leeching into the electrolyte). Adapted from ref 208. Copyright 2018 American Chemical Society. (c) Symmetric cell cyclic voltammetry (CV) cycling using the MXene hybrids for both electrodes. Adapted from ref 224. Copyright 2018 Wiley-VCH. (d) Visual LiPS permeation test with graphene-TiO$_2$@Ti$_3$C$_2$T$_x$-polymer separator (yellow/brown colouration on the right side of the cell indicates LiPS permeating through the separator) and (e) long-term cycling of graphene-TiO$_2$@Ti$_3$C$_2$T$_x$ separators (with different oxidation durations) at 2C. Adapted from ref 227. Copyright 2019 Wiley-VCH.
MXENE HYBRIDS FOR EMERGING EC, PC, AND PEC REACTIONS OF INTEREST

TM carbides and nitrides have a long history in catalyzing hydrogenation (or dehydrogenation) reactions due to their suitable surface interactions with the relevant reaction intermediates.\textsuperscript{233,234} With growing research interest in sustainable catalytic reactions, it was discovered that TM carbides and nitrides are also active in the catalytic conversion of abundant and stable moieties like CO\textsubscript{2} or N\textsubscript{2} to higher value-added chemicals.\textsuperscript{235} Sharing similarities to the bulk 3D crystalline TM carbide and nitride forms, MXenes are also anticipated to be catalytically active for CO\textsubscript{2} and N\textsubscript{2} activation due to their suitable adsorption sites\textsuperscript{236,237} and favorable adsorbate interactions.\textsuperscript{238} Having recently demonstrated promising nitrogen and carbon catalysis on MXenes,\textsuperscript{25,239} MXene hybrids have been predicted to expand the platform even further to enhance catalytic activity and catalyze other important reactions in the future. Here, we analyze the current strategies and pathways in which MXenes hybrids have been used to enhance important PC, EC, and PEC reactions of CO\textsubscript{2}, N\textsubscript{2}, and their related moieties.

**MXene hybrids and composites for electrocatalytic/photocatalytic (EC/PC) carbon dioxide reduction (CO\textsubscript{2}RR).** MXenes synthesized to date are similar to the bulk 3D crystalline non-MXene TM carbides and nitrides in terms of high electronic conductivity.\textsuperscript{152,240} Additionally, the MXenes’ basal planes are terminated with T\textsubscript{x} groups in ambient conditions or electrolytes.\textsuperscript{241} These T\textsubscript{x} groups usually comprise of electronegative elements (F, O, Cl) that can alter the MXenes’ effective work function\textsuperscript{242,243} and bring about unique MXene-intermediate molecule adsorbate interactions.\textsuperscript{23,244,245} In CO\textsubscript{2}RR, these unusual interactions render alternative thermodynamic pathways that were previously and commonly unfavorable on TM catalysts, feasible, providing a means to circumvent the scaled binding energy relations.\textsuperscript{246} Beyond providing active sites for CO\textsubscript{2}RR intermediates, Chen et al. described an interesting possibility of MXenes’ T\textsubscript{x} groups being
involved in the catalysis process. The DFT scheme in Figure 10a describes how H atoms in the T_x termination group can be “borrowed” by the intermediate during some of the proton-electron transfer steps, before being regenerated in the last three proton coupled electron transfer steps.

Despite encouraging theoretical simulations, EC CO_2RR catalysis by bare MXenes in aqueous electrolytes have yet to be reported, possibly due to the weaker CO_2RR intermediate-MXenes binding strength, and severe competition from HER as a side-reaction. Handoko et al. recently reported EC CO_2RR on Ti_2CT_x and Mo_2CT_x MXenes in a mixture of acetonitrile, water, and 1-Butyl-3-methylimidazolium tetrafluoroborate (BMIMBF_4). Formic acid (HCOOH) was the main product, with peak HCOOH Faradaic efficiency (FE_{HCOOH}, see Vocabulary for definition of FE) of 56% at -1.8 V (vs. standard hydrogen electrode, SHE) on Ti_2CT_x synthesized with KF-HCl (Figure 10b). Interestingly, higher activity and an earlier CO_2RR onset potential was observed for MXenes with lower –F T_x content. This observation is explained by DFT calculations, which revealed that the critical step involving the *COOH intermediate is more adversely affected by an increasing presence of –F T_x groups (Figure 10c).

As such, we believe that MXene hybrids can offer a viable value proposition. In addition to their excellent electronic conductivity, thin MXene layers would be a superior catalyst support choice compared to other C-based materials, due to stronger metal-support interactions. Furthermore, the possibility of intermetallic compound formation has been reported, revealing an active surface with a unique electronic structure that is more resistant to agglomeration. For example, Mo_2CO_2 MXene has been predicted as a good Pd support. Specifically, a more active Pd dimer species that allows for partial methane oxidation to methanol can be stabilized on Mo_2CO_2 MXenes.
In addition, MXenes have also been demonstrated to be suitable substrates for single atom catalysts (an emerging active frontier in heterogeneous catalysis\textsuperscript{251}) such as Pt, Ru and Rh, through a simple self-reduction reaction at room temperature.\textsuperscript{252} The single atom nature of PGMs substituting the Ti positions in Ti$_3$C$_2$T$_x$ MXene was demonstrated with extended X-ray absorption fine structure (EXAFS, \textbf{Figure 10d}) spectroscopy and high-angular annular dark-field (HAADF) scanning transmission electron microscopy (STEM) imaging (\textbf{Figure 10e}).\textsuperscript{252} In particular, Pt$_i$/Ti$_3$-C$_2$T$_x$ structures were reported to be active for CO$_2$ activation with amines and silane into formamide at 140 °C with near-complete conversion and selectivity.\textsuperscript{252}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure10.png}
\caption{MXenes for CO$_2$RR. (a) DFT scheme showing the possible involvement of T$_x$ atoms in CO$_2$RR to CH$_4$. Adapted with permission from ref.\textsuperscript{22} Copyright 2019 American Chemical Society. (b) Comparison of FE for CO$_2$RR to formic acid on Ti$_3$CT$_x$ and Mo$_2$CT$_x$. Error bars represent one standard deviation of three independent measurements. (c) Calculated free energy diagram at 0 V applied potential for CO$_2$RR to formic acid on Mo$_2$CT$_x$ MXenes with varying fractions of –F and –O surface T$_x$ groups. Purple, red, brown and white spheres represent Mo, O, C and H atoms respectively. Adapted with permission from ref.\textsuperscript{25} Copyright 2020 The Authors. (d) Two-dimensional wavelet transformation of Pt K-edge EXAFS (radial distance information in the y-axis and energy dependence k-space in the x-axis) and (e) HAADF STEM of Pt$_i$/Ti$_3$-C$_2$T$_x$, which clearly demonstrates the absence of Pt-Pt bond in the single atom Pt-on-Ti$_3$C$_2$T$_x$ MXene. Adapted with permission from ref.\textsuperscript{252} Copyright 2019 American Chemical Society.}
\end{figure}
We note that most experimental CO$_2$RR on MXenes or their hybrids were conducted using PC/PEC instead of EC. This is intriguing, as apart from the photophysical processes, PC should share many commonalities with EC. Discrepancies between the measured activities of the same TM catalyst (or co-catalyst) species in EC and PC approaches have been reported. These discrepancies can be partly attributed to the attenuated work function under steady state illumination and interfacial voltage loss, limited cathodic potential range (i.e. conduction band position), and temporal discontinuity between the photo-generated carrier lifetime (fs to ms) and reaction kinetics (ms to s). Besides impacting catalytic activity, we believe that these differences may enhance the selectivity in PC CO$_2$RR, as the diverse intermediate stability in CO$_2$RR and rate of competitive HER is sensitive to the applied potential.

A commonly adopted hybrid combination for PC CO$_2$RR is MXenes coupled with a wider bandgap photo-absorber. MXenes made to date are metallic and some are predicted to have narrow bandgaps, thus hybridization with a wider bandgap photo-absorber is the only way to generate sufficient photo-carriers for catalysis. A notable example is the Ti$_3$C$_2$Tx/g-C$_3$N$_4$ hybrid which produced 2.24–5.19 and 0.04–0.05 µmol g$^{-1}$ h$^{-1}$ of CO and CH$_4$ respectively. Although g-C$_3$N$_4$ is a very good photo-absorber with high quantum yield even at visible wavelengths ($\lambda >$395 nm), it is limited by rapid charge recombination. The role of Ti$_3$C$_2$Tx in the hybrid is likely that of a charge separator, as their work function position is suitably positioned to accept electrons from the g-C$_3$N$_4$ conduction band, thus enhancing the photo-generated carrier lifetime (Figure 11a). The activity of Ti$_3$C$_2$Tx/g-C$_3$N$_4$ was further enhanced considerably by depositing Pd on the hybrid structure, which produced formate and methanol at a rate of 905 and 350 µmol g$^{-1}$ h$^{-1}$, respectively, in a PEC setting. Pd is a very interesting co-catalyst choice, because it is known to bind very strongly to adsorbed *CO in EC CO$_2$RR, which often leads to catalyst deactivation.
The authors attributed the high activity of the Ti$_3$C$_2$Tx/g-C$_3$N$_4$/Pd hybrid to electron trapping by protonic species on the Pd surface, but we speculate that the stronger binding may allow the key *CO intermediate to persist longer, allowing for photo-electron accumulation and transfer.

Depending on the relative band positions, hybridization of metals (i.e. MXenes) with semiconductors often results in a built-in Schottky junction. This increases the electronic transport resistance and it is thus generally undesired. However, in PC, the Schottky barrier junction may actually improve the overall carrier lifetime by minimizing the electron losses associated with back injections. One example of the adoption of this strategy is reported in Ti$_3$C$_2$Tx/CeO$_2$, which produced up to 40.2 µmol g$^{-1}$ h$^{-1}$ of CO.$^{265}$ The built-in junction may also exist in many other examples, especially in hybrids involving wide-bandgap n-type semiconductors such as TiO$_2$,$^{85}$ Bi$_2$WO$_6$,$^{101}$ or CsPbBr$_3$,$^{125}$ although this is hard to ascertain as the MXenes’ work function varies$^{243}$ depending on the actual T$_x$ composition in the reaction environment. Hybridization with these semiconductors generally yield modest CO$_2$RR activity (Table S4a), of up to 28 µmol g$^{-1}$ h$^{-1}$ products for Ti$_3$C$_2$Tx/P25/TiO$_2$. While a slightly higher activity was observed for Ti$_3$C$_2$Tx/CsPbBr$_3$ (32 µmol g$^{-1}$ h$^{-1}$ CO and 14 µmol g$^{-1}$ h$^{-1}$ CH$_4$), the perovskite’s (CsPbBr$_3$) instability in aqueous solutions necessitated the use of ethyl acetate as the electrolyte.$^{125}$
Figure 11. MXene hybrids for PC/PEC CO$_2$RR. (a) Transient PL spectroscopy showing a longer photo-generated carrier lifetime for Ti$_3$C$_2$T$_x$/g-C$_3$N$_4$ (10TC), compared to bare g-C$_3$N$_4$ (UCN). Adapted with permission from ref 259. Copyright 2020 Elsevier B.V. (b) Morphology of Ti$_3$C$_2$T$_x$/TiO$_2$ after controlled calcination of Ti$_3$C$_2$T$_x$ at 550°C. Adapted with permission from ref 85. Copyright 2018 Elsevier Inc. (c) TEM image of Ti$_3$C$_2$T$_x$ QD/Cu$_2$O and (d) HRTEM image of Ti$_3$C$_2$T$_x$ QDs, showing hexagonal units. (e) DFT calculated Fermi level and (f) band diagram of Ti$_3$C$_2$T$_x$ QD/Cu$_2$O. Adapted with permission from ref 121. Copyright 2018 WILEY-VCH.

TiO$_2$/Ti$_3$C$_2$T$_x$ is another popular MXene hybrid for PC CO$_2$RR, since TiO$_2$ can be naturally grown from in-situ conversion of Ti-based MXenes via partial oxidation. Under the right synthesis conditions, hierarchical hybrids were produced while retaining the conductive Ti$_3$C$_2$T$_x$ backbone (Figure 11b).$^{85}$ The in-situ grown TiO$_2$/Ti$_3$C$_2$T$_x$ hybrid, combined with Pd co-catalysts, is possibly the most active MXene hybrid for PEC CO$_2$RR, yielding a cocktail of products (formate, methanol, and ethanol) at a high total hydrocarbon yield of 1840 μmol g$^{-1}$ h$^{-1}$.$^{92}$ Separately, a different approach hybridized Ti$_3$C$_2$T$_x$ with Cu$_2$O (Figure 11c).$^{121}$ Instead of using large MXenes flakes directly, tiny Ti$_3$C$_2$T$_x$ QDs were prepared by rigorous sonication in inert argon. These MXene QDs were then treated with positively charged polyethyleneimine to aid their electrostatic self-assembly with negatively charged polystyrene sulfonate-coated Cu$_2$O nanorods. The theoretical Fermi level
of the hexagonal Ti$_3$C$_2$T$_x$ QDs was estimated to be significantly lower than the “bulk” Ti$_3$C$_2$T$_x$ sheets (Figures 11d and 11e), which resulted in more facile electron transfer from the Cu$_2$O photo-absorber, at a potential above the thermodynamic requirement for CO$_2$RR to methanol (-0.81 V vs. RHE, Figure 11f). The Ti$_3$C$_2$T$_x$ QD/Cu$_2$O hybrid supplied 78.5 µmol g$^{-1}$ h$^{-1}$ of methanol.

Beyond CO$_2$RR, CdS/Ti$_3$C$_2$T$_x$ was also shown to be active in the PC reduction of 4-nitroaniline to p-phenylenediamine (PPD), an important synthetic intermediate for dyes and aramid fibers. While the PC activity of CdS in reducing aromatic nitro-compounds is not surprising, hybridization with MXenes significantly enhanced their activity and photo-stability. It was proposed that hybridization with CdS formed a Janus-like structure, characterized by the strong metallic adsorption of Cd onto Ti$_3$C$_2$T$_x$, thus limiting the dissolution of CdS. Similar strategies may therefore be extended to other TM chalcogenides for enhancing CO$_2$RR activity.

**MXenes hybrids and composites for electrocatalytic/photocatalytic (EC/PC) nitrogen reduction (N$_2$RR).** The abundance of N$_2$ in the air has attracted many attempts to convert them to nitrate and ammonium compounds for agricultural applications. Combined with steam reforming of natural gas, the Haber-Bosch process is still widely used in modern ammonia production plants worldwide today. Although the Haber-Bosch process is an exothermic reaction overall, it still requires a massive energy input to drive the N≡N bond dissociation (941 kJ mol$^{-1}$) through the dissociative pathway over a suitable catalyst (Figure 12a).

EC/PC N$_2$RR offers an attractive alternative to the Haber-Bosch process. With the ability to be conducted under ambient conditions, N$_2$RR is potentially safer, cleaner, and a more sustainable way to produce NH$_3$ with a smaller plant footprint when coupled with sustainable energy inputs. Unlike the Haber-Bosch process with a kinetically (or thermodynamically,
depending on the substrate) limited N≡N dissociation step, PC/EC N$_2$RR preferentially undergoes the associative pathway, whereby the adsorbed N$_2$ molecule first undergoes stepwise protonation.$^{269}$ The associative N$_2$RR pathway diverges to three main pathways depending on the initial N$_2$ adsorption geometry (Figure 12b): alternating and distal protonation mechanisms operate when N$_2$ is adsorbed end-on, while the enzymatic mechanism operates after N$_2$ is adsorbed side-on.$^{272,273}$ There are also possibilities of pathway inter-mixing, depending on the thermodynamic and kinetic barriers.$^{274}$ Side-on N$_2$ adsorption is more favorable on the basal planes of bare M$_2$C MXenes,$^{24,275,276}$ M$_3$C$_2$ MXenes with group 14 “M” elements (Figure 12d),$^{277}$ and ordered double TM Mo$_2$TiC$_2$T$_x$ MXenes.$^{278}$ Conversely, distal adsorption is preferred on M$_3$C$_2$ MXenes with group 15 and 16 “M” elements.$^{277}$ However, it should be noted that bare MXenes are unlikely to be stable under N$_2$RR operating conditions,$^{24}$ and it is important to include the relevant T$_x$ species when performing theoretical calculations. The T$_x$ groups generally cause N$_2$ adsorption on MXenes to be more difficult,$^{24}$ although alternative adsorption sites can be provided by exposed “M” sites at the edges (Figure 12e),$^{279}$ or on single atom dopants.$^{274}$
Figure 12. N₂RR mechanistic study. (a) Dissociative N₂ reduction to NH₃. Adapted with permission from ref 269. Copyright 2016 Elsevier B.V. (b) Associative N₂ reduction to NH₃, with diverging pathways depending on the adsorption geometry of the N₂ molecule. Adapted with permission from ref 275. Copyright 2020 Royal Society of Chemistry. Schematics of side-on N₂ adsorption on (c) defective (101) TiO₂ surface with O-vacancy and on (d) Ti₃C₂Tₓ, MXene. Dark grey, red, light grey and blue balls represent C, O, Ti and N respectively. Adapted with permission from ref 280. Copyright 2020 Elsevier B.V. (e) Alternative end-on N₂ adsorption on the exposed Ti site on Ti₃C₂O₂ MXene edges. Gray, red, sky blue and blue balls represent C, O, Ti and N respectively. Adapted with permission from ref 279. Copyright 2018 Elsevier Inc.

Before discussing the catalytic N₂RR activity of MXenes and their hybrids, it is imperative that we discuss the best activity reporting practices. A recurring issue in most PC/EC N₂RR research is that the quantity of NH₃ product produced is very small, in the range of 10 to 1000 nmol,281 and thus very susceptible to contamination and false positives. Therefore, establishing a suitable NH₃ detection protocol with robust control measurements is vital. There are two commonly adopted methods to quantify the produced NH₃ in solution: (1) colorimetric assays, including Nessler’s reagent (Nessler) or Berthelot’s reagent (e.g. indophenol blue, IB), and (2) ion chromatography (IC). While both methods generally work well in detecting low concentrations of
NH₃ (<500 µg L⁻¹), any impurities that result in false strong absorption in the detection wavelength, such as methanol and carbon particles, must be avoided in colorimetric-based methods. We note that the IB-based colorimetric assays are unsuitable for use at >500 µg L⁻¹ NH₃, or in acidic media, due to higher deviations. Finally, rigorous experimentation, including statistical reproducibility and confirmation that the NH₃ originated from N₂RR, requires the use of isotopically labeled $^{15}$N₂. Detection of $^{15}$NH₃ can be achieved using mass spectrometry, infrared, or nuclear magnetic resonance (NMR) spectroscopy. For the reader’s benefit, we report the NH₃ detection method, and whether $^{15}$N isotope labeling control was performed in summary Table S4b.

Ti₃C₂Tₓ is the most commonly used MXene for PC/EC N₂RR due to their stability and close structural relationship with the popular TiO₂ photocatalyst. Ti₃C₂Tₓ has been reported to perform EC N₂RR with modest NH₃ yields (157–522 µmol g⁻¹ h⁻¹) and FE at relatively low cathodic potentials (-0.1 to -0.4 V vs. RHE), with electrochemical and structural stability. Investigations into MXene hybrids, especially through the use of single atom dopants, have predicted attractive N₂RR activity, as the single atom dopants provide more favorable N₂ adsorption sites and correspondingly lower activation barriers. A range of in-silico single atom modifications (boron and various TMs) have been studied, with Ru, Mo, Ti, and Fe commonly identified as promising dopants with the least negative N₂RR limiting potentials. Apart from calculating the lowest energy pathway from N₂ to NH₃, it is also critical to evaluate the formation energy and the stability of the hybrid through first principle calculations, or from more realistic kinetic stability studies.

Experimentally, Ti₃C₂Tₓ has been used to support metallic N₂RR electrocatalysts such as Ru (2321 µmol g⁻¹ h⁻¹ and 13% FE) and fine (<10 nm) Au NPs (1761 µmol g⁻¹ h⁻¹ and 18.3% FE), with up to 14 times higher N₂RR activity than bare Ti₃C₂Tₓ. Enhanced N₂ adsorption and
*N₂ coverage was a reason for the high catalytic activity on fine Au NPs/Ti₃C₂Tx, as shown in the desorption profile (Figure 13a). Despite numerous encouraging predictions of N₂RR on single-atom/MXene hybrids, only one example was reported experimentally, using Ru single atoms/Mo₂CTₓ. This hybrid electrocatalyst is highly N₂RR active, yielding 2382 μmol g⁻¹ h⁻¹ NH₃ at a significant 25.7% FE. The N₂RR active site was investigated using operando Ru K-edge X-ray absorption near edge structure (XANES). A slight increase in the Ru oxidation state to +3.56 was observed when N₂ was introduced without bias (at open circuit potential, OCP) compared to the baseline measurement in Ar (+3.27), attributed to the electron back-donation when the Ru-N₂ bond was formed. The oxidation state reduces back to +3.15 when a N₂RR-relevant potential (-0.3 V vs. RHE) was applied, indicating N₂RR. This phenomenon was also observed in the Fourier transformed-XANES (FT-XANES, Figure 13b). A discernible shift in the first neighbor peak to a shorter apparent distance of 1.51 Å was seen when N₂ was introduced, which reverted back to 1.56 Å during N₂RR, possibly due to Ru–N coordination during adsorption.

Beyond metallic catalysts, Ti₃C₂Tx was also combined with TM oxide catalysts such as MnₓOᵧ. Manganese oxides are N₂RR active, but exist in many structures with poor conductivity that limit their EC N₂RR performance. Hybridizing MnO₂ with Ti₃C₂Tx MXenes yielded up to 2003 μmol g⁻¹ h⁻¹ NH₃ (11.39% FE) and traces of N₂H₂ at -0.55 V vs. RHE. The superior N₂RR activity of Ti₃C₂Tx/MnO₂ was ascribed to multiple advantages provided by the Ti₃C₂Tx support: (1) prevention of MnO₂ aggregation, (2) facilitation of electron transfer to MnO₂ and avoidance of parasitic current loss, and (3) larger surface area with favorable adsorption and active sites.

As discussed earlier, Ti₃C₂Tx is a good precursor for in-situ conversion to Ti₃C₂Tx/TiO₂ hybrids. It is interesting that Ti₃C₂Tx/TiO₂ hybrids can display EC N₂RR activity with modest NH₃ yield (1545–1889 μmol g⁻¹ h⁻¹) and FE (8.42–16.07%) in aqueous solutions, despite potential
competition from HER and the poor conductivity of TiO$_2$. Expectedly, there are many more examples of Ti$_3$C$_2$T$_x$/TiO$_2$ hybrids reported for PC than EC N$_2$RR to utilize TiO$_2$ as a photocatalyst. The highest PC NH$_3$ yield using MXene hybrids so far is reported by Hou et al. (422 µmol g$^{-1}$ h$^{-1}$). This recent work demonstrated that the hybrid’s photo-response extends up to the near-infrared region. A significant yield (82 µmol g$^{-1}$ h$^{-1}$) was still detectable at 740 nm irradiation, corresponding to an apparent quantum efficiency (AQE) of 0.07% (Figure 13c). Liao et al. recently investigated the role of Ti$_3$C$_2$T$_x$ in enhancing TiO$_2$ N$_2$RR activity. In this work, the authors mixed P25 TiO$_2$ (80% anatase and 20% rutile phase) with Ti$_3$C$_2$T$_x$ and treated them in N$_2$ atmosphere. Interestingly, the hybridization process introduced significant O-vacancies in TiO$_2$, as indicated by the broadening of the Raman peak around 1500 cm$^{-1}$, and a strong low temperature electron spin resonance (ESR) signal (Figure 13d). The defective TiO$_2$ sites acted as an alternative adsorption site (Figure 12c) alongside Ti$_3$C$_2$ (Figure 12d), which enabled a higher coverage of adsorbed *N$_2$. At the optimum Ti$_3$C$_2$T$_x$ loading of 6%, the NH$_3$ yield of the hybrid was 10.74 µmol g$^{-1}$ h$^{-1}$ without any sacrificial reagents. The NH$_3$ yield was further enhanced to 43.44 µmol g$^{-1}$ h$^{-1}$ after 0.02 M CH$_3$OH was added as a sacrificial electron donor (Figure 13e), indicating that the anodic reaction (water oxidation) is still a significant bottleneck.

Alternatively, the oxidation of Ti$_3$C$_2$T$_x$ to TiO$_2$ can be fully completed, resulting in a unique MXene-derived TiO$_2$ structure with many Ti$^{3+}$ sites, which are inherently more active catalytically. These MXene-derived hybrids were used for EC and PC N$_2$RR. Using in-situ oxidation, a more complex MXene-derived hybrid of TiO$_2$@C@g-$C_3$N$_4$ was also produced from Ti$_3$C$_2$T$_x$ for PC N$_2$RR. This structure was demonstrated to be an efficient and stable N$_2$RR photocatalyst with an NH$_3$ production rate of 250.6 µmol g$^{-1}$ h$^{-1}$. The outstanding NH$_3$ production
performance was attributed to the abundance of surface defects, efficient electron-donation, suitable light harvesting, small transition impedance, and strong N₂ adsorption.

![Figure 13](image)

**Figure 13.** MXene hybrids for N₂RR. (a) N₂ temperature programmed desorption profile of Ti₃C₂Tₓ, <10 nm Au/Ti₃C₂Tₓ hybrid, and >200 nm Au/Ti₃C₂Tₓ hybrid. A broad peak around 130°C signifies N₂ physisorption, while the band closer to 280 °C represents N₂ chemisorption. Adapted with permission from ref 289. Copyright 2019 American Chemical Society. (b) FT-EXAFS spectra derived from the normalized *operando* Ru K-edge XANES spectra for single atom Ru-Mo₂CTₓ under various applied voltages vs. RHE in 0.5 M K₂SO₄. Adapted with permission from ref 290. Copyright 2020 WILEY-VCH. (c) AQEs for N₂ fixation on Ti₃C₂Tₓ/TiO₂ under monochromatic light irradiation. Adapted with permission from ref 86. Copyright 2020 Elsevier B.V. (d) ESR spectra of bare P25 and 6% Ti₃C₂Tₓ/P25 hybrid. (e) NH₃ yield after 1 h N₂RR on 0% (bare P25) and 6% Ti₃C₂Tₓ/P25 hybrid, in H₂O and with 0.2 M CH₃OH. Adapted with permission from ref 280. Copyright 2020 Elsevier B.V. (f) HRTEM of 15%-Ti₃C₂Tₓ/MIL-100(Fe) hybrid. Adapted with permission from ref 294. Copyright 2019 American Chemical Society.

Apart from N₂RR, NO oxidation is another pathway for nitrogen fixation and air pollution elimination, since NO accounts for 95% of all the NOₓ contaminations in air.²⁹⁴ Wang *et al.* reported the PC oxidation of NO by the Ti₃C₂Tₓ/MIL-100(Fe) MOF hybrid (Figure 13f). This work found that 58% NO was eliminated by the hybrid containing 15 wt% Ti₃C₂Tₓ, which was 2.8 and 4.1 times more efficient than the bare MIL-100(Fe) MOF and Ti₃C₂Tₓ MXene, respectively.
The enhanced NO oxidation was due to synergistic effects in the as-formed Schottky junction: Ti$_3$C$_2$Tx acted as an electron conductor and accumulated electrons on surface to promote O$_2$RR and form $\cdot$O$_2^-$. Both the generated $\cdot$O$_2^-$ and h$^+$ active species on the surface of MIL-100(Fe) then oxidized NO to NO$_3^-$ in the presence of water.

**Summary, specific challenges, and outlook for EC/PC CO$_2$RR and N$_2$RR.** Thus far, we have elaborated on how MXene hybrids have found their way into emerging catalytic reactions (of CO$_2$, N$_2$ and related compounds), functioning either as support or active materials. Although the use of MXenes in these emerging catalytic reactions is nascent and has yet to result in record performance metrics, it provides a variety of different reaction pathways and is a welcome addition as an alternative catalyst support choice. We predict that the role of MXenes in these emerging reactions will continue to grow as we discover more MXenes hybrids and learn more about their unique interactions with reaction intermediates and other co-catalysts. A summary table of performance metrics for various MXene hybrids in EC/PC CO$_2$RR and N$_2$RR is provided in Tables S4a and S4b respectively.

Specifically, we identify that the MXene/single atom hybrid is a worthwhile opportunity to pursue due to the many encouraging computational predictions but minimal experimental demonstrations. Single atom catalysis is a rapidly expanding field whereby methods of synthesizing and stabilizing these single atoms are quickly being developed.$^{251}$ Related to this point, mechanistic studies into MXene/single atom hybrids have not been established and is thus open for investigation, especially on the dynamic behavior of these single atom catalysts during catalytic reactions. Although the evidences so far suggest that single atom dopants are the active species, the catalytic action may also rely on the adsorbate interactions with the MXene catalyst support, for example through stepwise or cascading reactions. We foresee that *operando*
techniques capable of discerning transient catalyst state and surface adsorbates, such as resonant inelastic X-ray scattering and SECM, will play an important role here.

Another interesting research opportunity lies in bridging the gap in understanding between the PC and EC operating modes of CO$_2$RR and N$_2$RR on MXenes and their hybrids. We note that most direct experimental CO$_2$RR on MXene hybrids have been reported on PC compared to EC, possibly pointing to the direction of an indirect electron transfer driven reaction. In PC reactions, HER appears to be less severe a problem, even if these reactions (CO$_2$RR, N$_2$RR) are conducted in aqueous electrolytes. Although it is possible to run EC CO$_2$RR or N$_2$RR in non-aqueous electrolytes,$^{25}$ it is still important to improve our understanding on how to limit the HER side-reaction and run EC CO$_2$RR or N$_2$RR more economically in aqueous solutions. Some strategies for theoretical and experimental investigations are thus required (e.g. decoupling the lifetime of photo-generated carriers with the reaction dynamics). Additionally, studying how to tune the Fermi level of MXenes with different T$_x$ groups and the coupled co-catalyst is important to ensure the efficient photo-carrier movement towards the desired surface.

We also see a huge opportunity from MXene-derived materials in catalysis. Although this class of material does not fit the MXene hybrids category strictly, we recognize that the resulting MXene-derived materials behave differently and are much more active than those obtained using conventional methods. While many examples of TiO$_2$ hybrids have been derived from Ti-based MXenes, far fewer have reported using other MXenes. In particular, group 14, 15, and 16 TM MXene-derived materials could hold interesting properties for CO$_2$RR and N$_2$RR.
CONCLUDING REMARKS AND FUTURE OUTLOOK

In this review, we discussed the design rules behind MXene hybrids for catalytic energy storage and conversion applications. We summarized the different classes of secondary materials hybridized with MXenes and their synthesis methods, highlighting their differences. By carefully considering the MXenes’ properties (T\textsubscript{x} groups, basal plane defects, edge charges, and identity of the outer metal surface), a variety of secondary materials can be chemically coupled with MXenes to form MXene hybrids with varying architectures, from vertically-aligned nanomaterials on MXenes to lateral coverages of MXenes and even more complex 3D mesoporous structures.

In general, MXenes and their hybrids are susceptible to oxidative degradation in environments containing water and/or oxygen\textsuperscript{193} and under applied anodic potentials in aqueous media\textsuperscript{29}, both of which are common in catalysis. The MXenes’ instability should thus be thoughtfully considered when designing MXene hybrids for energy storage and conversion. This can be circumvented through recent advances in the synthesis (F\textsuperscript{-}-free\textsuperscript{295} and/or aqueous-free\textsuperscript{296,297}), storage,\textsuperscript{193,298} and stabilization (in thin films\textsuperscript{299} or colloids\textsuperscript{300}) of MXenes. These breakthroughs also offer greater flexibility and control of surface T\textsubscript{x} groups for targeted hybrid assembly. This can be translated to the preparation of stable MXenes and their hybrids for water-sensitive applications, such as non-polar organic composites, polymer inks for additive manufacturing, and organic electrolytes for supercapacitors and batteries\textsuperscript{301,302} and are also highly relevant for communities working on flexible electronics, biomedical, and electromagnetic applications.

Additionally, we expect directed efforts to be made beyond mono-metal MXenes to synthesize multi-metal MXenes such as ordered double MXenes (M’\textsubscript{2}MC\textsubscript{2}T\textsubscript{x} and M’\textsubscript{2}M\textsubscript{2}C\textsubscript{3}T\textsubscript{x}), which have already been theoretically predicted to possess high levels of HER activity.\textsuperscript{303} Adding
to the already-growing MXenes family, disordered double TM solid solution MXenes such as (Ti,V)$_2$CT$_x$ and (Mo,V)$_4$C$_3$T$_x$ are also attractive prospects for synthesis and characterization, given the ease of tuning their electrochemical and physical properties based on their compositions.$^{84,304}$

Another key challenge lies in the expansion of existing etching methods to ternary transition metal nitrides MAX materials to produce nitride MXenes, since only Ti$_2$NT$_x$ has been produced in this manner,$^{305}$ while Ti$_4$N$_3$T$_x$ has only been synthesized through harsh molten fluoride salt etching.$^{19,201}$ Other Mo-containing nitride MXenes are instead produced through ammoniating their corresponding carbide MXenes with NH$_3$ gas at a similarly high temperature of 600°C.$^{306}$ Additionally, closely related 2D MXene-like TM nitrides, such as MoN, have also been reported to be catalytically active for energy storage and conversion applications.$^{307–309}$ Investigating these layered 2D MXene-like TM nitrides can potentially reveal important insights into the predicted stability and chemical reactivity of MXene nitrides. In all, expanding the MXenes family can create more possibilities in advanced MXene hybrid design to realize theoretically predicted properties and competitive performances as multi-functional MXene hybrids and composites.

For each of the industrially relevant catalytic energy storage (metal-air/sulfur batteries) and conversion (water splitting, HER, OER, CO$_2$RR and N$_2$RR) applications, we clarified the roles of each material component in the MXene hybrids – a conductive support, co-catalyst, electronic structure modulator, photo-carrier generator and phase stabilizer, to name a few. More importantly, we described how strong chemical and electronic coupling within the hybrid can induce directed morphological growth, alter the electronic structure, and create more catalytically active motifs for synergistic enhancement in catalytic activity and stability. By combining T$_x$ control, doping and hybridization to modulate the electronic density and intrinsic activity of active site, coupled with nanostructuring to increase the density and accessibility of those active sites, many MXene hybrids
have outperformed their separate individual components in both catalytic activity and stability. We also provide application-specific challenges and our outlook at the end of each application section for readers working on those applications. A summary of key performance metrics for HER, OER, ORR, CO_2RR and N_2RR is provided in Tables S2–S4 for the reader’s convenience.

MXene hybrids and composites have achieved superior catalytic energy storage and conversion performances compared to traditional materials, and we anticipate further advances as the chemical and electronic nature of the MXene hybrid interfaces are elucidated using a powerful combination of *in-situ*/*operando* experiments and theoretical modelling.\textsuperscript{21,202,238} A better understanding of the dynamics at the MXenes’ hybrid surface and interface in the presence of reactive species, solvent, and applied potential is indispensable toward building a more realistic and multi-scale MXene hybrid model that incorporates the effects of electrochemical potential, solvation, and micro-kinetics into the current purely atomistic view of DFT, as recently demonstrated on metallic catalysts.\textsuperscript{310–312} So far, *operando* measurements demonstrated on bare MXenes\textsuperscript{19} have clarified our fundamental understanding of the localized electrochemical activity and electronic structure at the catalytic surface. For this purpose, it is not unreasonable to use MXene-like surrogate structures that can be synthesized using bottom-up approaches with atomic precision (CVD and ALD) for such fundamental studies.\textsuperscript{65,66,148}

Finally, efforts can be undertaken in the development of mild and scalable top-down MXene synthesis and hybridization techniques to (1) minimize MXene oxidation and mechanical damage during processing especially at high temperatures,\textsuperscript{192,193} (2) enable greater control over the chemical composition, morphology and distribution of the secondary material grown on MXenes, and (3) realize highly porous and 3D MXene hybrid networks for efficient mass and long-range charge transport to further enhance catalytic activity.\textsuperscript{169} Given the rapid rise in interest amongst
MXene hybrids and composites, we expect an increase in both experimental and theoretical efforts towards developing MXene hybrids for clean energy and related applications in the near future.
ASSOCIATED CONTENT

The authors declare no competing financial interests.

Supporting Information.

Detailed data entries (categorization of MXene hybrid publications, electrocatalytic HER/OER/ORR performances, photocatalytic HER performances, photo/electrocatalytic CO\textsubscript{2}RR/N\textsubscript{2}RR performances) is available free of charge in the Supporting Information online.

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

ACKNOWLEDGEMENT

This work was supported by the Singapore National Research Foundation (NRF-NRFF2017-04). H.-Y.J. acknowledges the National Natural Science Foundation of China (NSFC, No. 21703170) and Key Research and Development Program of Shaanxi (Program No. 2020GY-244). J.T. acknowledges the RS International Exchanges 2017 Cost Share Award (IEC\NSFC\170342) and the Leverhulme Trust (RPG-2017-122).
VOCABULARY

**MXene hybrid/composite**, a material fabricated by combining MXenes with another at least one other non-MXene material (metal, semiconductor or insulator) to form at least one interface. A hybrid/composite material is a more general classification than a heterostructure (see definition for heterostructure below).

**MXene heterostructure**, a material fabricated by growing at least one non-MXene material on MXenes in a repetitive layered manner, which forms more than one MXene-other material interface that repeats itself *i.e.* MXene-other material-MXene-other material-… and so on.

**(Photo)electrocatalyst**, a material that reduces the activation energy and overpotential (see definition for overpotential below) to drive an electrochemical reaction of interest, such as HER, OER, ORR, CO$_2$RR and N$_2$RR.

**Overpotential (at -10 mA cm$^{-2}$ current density) $\eta_{j=10}$**, the additional potential difference (voltage) beyond the thermodynamically determined redox potential to drive an electrochemical redox half-cell reaction at a specific current density.

**Tafel slope**, the amount of increase in overpotential required to increase the current density produced for an electrochemical redox half-cell reaction by a factor of ten.

**Metal-air/sulfur battery**, two different battery architectures for energy storage. Metal-air/sulfur batteries use a metal anode and an air/sulfur cathode. The redox half-cell reactions occurring at the cathode are OER/ORR for metal-air batteries and S-redox reactions for metal-sulfur batteries.
**Faradaic efficiency**, the ratio of the number of electrons used to produce the desired electrochemical product compared to the total number of electrons added into the system. For example, a 70% Faradaic efficiency in CO$_2$RR to CH$_4$ indicates that 70% of the electrons added are used to reduce CO$_2$ to CH$_4$ while the other 30% of electrons added are used to reduce CO$_2$ to other products.
TOC FIGURE (can be rescaled to 9 cm width × 4 cm height)
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