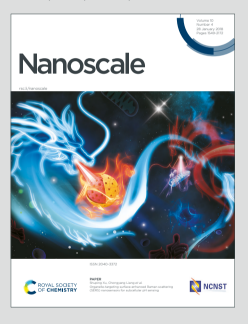




Nanoscale

Accepted Manuscript

This article can be cited before page numbers have been issued, to do this please use: S. Kohli, G. Rathee, I. Jha, L. Phor, H. sable and V. Chaudhary, *Nanoscale*, 2025, DOI: 10.1039/D5NR02336J.



This is an Accepted Manuscript, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about Accepted Manuscripts in the <u>Information for Authors</u>.

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal's standard <u>Terms & Conditions</u> and the <u>Ethical guidelines</u> still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this Accepted Manuscript or any consequences arising from the use of any information it contains.



Strategically Engineering 2D MXenes-based Advanced Adsorbents for Sustain MR023363
Wastewater Remediation of Dyes

Sahil Kohli¹**, Garima Rathee²*, Indrani Jha¹, Lakshita Phor³, Harsh Sable⁴, Vishal Chaudhary⁵*

¹Department of Chemistry, School of Engineering and Technology, Manav Rachna International Institute of Research and Studies, Faridabad, Haryana-121004, India.

²Department of Chemical Engineering, Universitat Politècnica de Catalunya, Barcelona, Spain.

³Department of Physics, University Centre for Research and Development, Chandigarh University, Gharuan, Mohali 140413.

⁴Sharda School of Allied Health Sciences, Sharda University, Greater Noida, U.P. (201306), India.

⁵Centre for Theoretical Physics and Natural Philosophy, Nakhonsawan Studiorum for Advanced Studies, Mahidol University, Nakhonsawan, 60130, Thailand.

*Correspondence:

- sahilkohli41@gmail.com (S.K.), sahil.set@mriu.edu.in
- Chaudhary00vishal@gmail.com (V.C.)

*Equal authorship: SK and GR

View Article Online

DOI: 10.1039/D5NR02336J

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Nanoscale Accepted Manuscript

The exponential growth of the global population in the digital era has accelerated urbanization and industrialization, leading to severe water pollution from the discharge of toxic dyes into aquatic ecosystems. Two dimensional (2D) MXene-based nano-adsorbents have recently emerged as promising candidates for developing sustainable wastewater remediation technologies due to their tunable physicochemical properties, including a high negative zeta potential, a large specific surface area, exceptional adsorption capacity, superior electrical and thermal conductivity, hydrophilicity, and a rich surface chemistry. Strategic optimization approaches for MXenes, encompassing interlayer spacing modification, surface engineering, stoichiometric tuning, morphological control, bandgap engineering, membrane fabrication, hybridization, and functionalization, have significantly enhanced their adsorptive performance and dye removal efficiency for practical wastewater treatment applications. This comprehensive review examines the latest advances in MXene-based nano-adsorbent engineering and their implementation as key components in wastewater treatment strategies for efficient dye removal from industrial effluents, providing fundamental insights into dye-MXene interactions to elucidate underlying removal mechanisms. The review highlights the sustainable characteristics of MXene-based nano-adsorbents, including their dye removal capacity, regeneration potential, recyclability, catalytic efficiency, and enhanced physicochemical properties, while addressing critical challenges, such as toxicity concerns, biocompatibility issues, and scalability limitations, that currently hinder their translation from the laboratory to the market. Innovative solutions are proposed through the integration of digital-age technologies, particularly artificial intelligence and machine learning approaches, with the implementation of these recommendations facilitating the establishment of MXenebased nano-adsorbents as sustainable alternatives to conventional commercial adsorbents. This

aligns with the UN Sustainable Development Goals and contributes to the principles of Mariota Continuous Health, promoting global welfare.

Keywords: Nanomaterials; Adsorbents; MXenes; Wastewater remediation; Dyes.

View Article Online 1. Introduction DOI: 10.1039/D5NR02336J

As industrialization progresses rapidly to cater to the needs of coming generations, there is improper discharge of toxic pollutants from industry settings, resulting in serious global One Health threats, including water pollution and damage to human health. In accordance with World Health Organization (WHO), environmental contamination impairs the human respiratory, neurological, and immune systems, making people more susceptible to serious diseases, including monkeypox, coronavirus disease (COVID-19), and cancer. As a result, legislators, environmentalists, researchers, and industrialists have expressed concern about the need for environmental rehabilitation. It can be accomplished by monitoring and managing various contaminants (air, water, radiation, and solid) using specific methodologies. 1-6 Chemical water contaminants are classified as organic and inorganic pollutants. The former comes primarily from human activity, such as sewage plants, agricultural runoff, and industrial effluents. Dye compounds are among the organic pollutants, while radionuclides and heavy metal ions are the most frequent inorganic pollutants found in water. These contaminants are mostly discharged by paper, paint, and textile factories. Most of these industries frequently discharge their effluent into water bodies, which can lead to several issues, including decreased solar penetration into the marine ecosystem, genotoxicity, carcinogenicity, and adverse effects on human wellness, the habitat, and marine life.⁷⁻⁹ Harmful molecules of dye in water leads to mutagenicity, and disruption of main human tissues such as liver, kidney, reproductive system, and central nervous system. 10 As a result of industrial and human activity, dyes are regarded as dangerous pollutants since they can be found in surface water. 11 To protect the environment, it is crucial to eliminate these harmful toxins from the aquatic environment.¹²

Various technologies have been established in the past few years to effectively remove dye pollutants from the water, including advanced oxidation, ^{13–15} catalytic ozonation, ¹⁶ flocculation and coagulation, ¹⁷ Fenton oxidation and electrocoagulation, ¹⁸ ion exchange, ¹⁹ membrane

separation,²⁰ and photo-catalysis.^{21,22} These approaches have certain drawbacks Figury Article Online including the production of contaminated sludge, difficulties with disposal and handling, scaling up sophistication problems, high energy consumption, increased colouration of effluents, financial constraints, intricate mechanisms, clogged membranes, restricted flow rates, and production of byproducts.²³

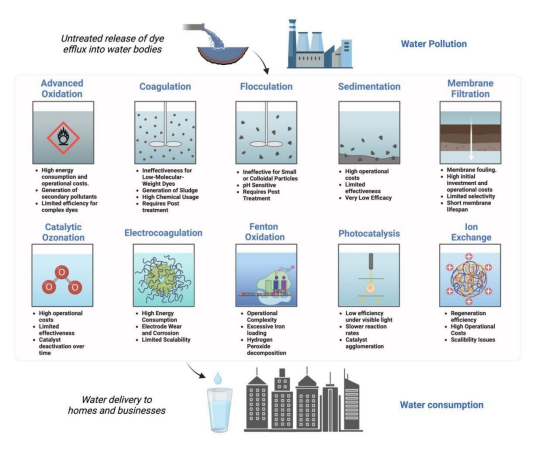


Figure 1. Drawbacks of traditional methods employed for dye removal

Among these methods, adsorption is considered as one of the effective substitute techniques for de-colorizing wastewater owing to its easy handling, affordability, sustainability, and quick water treatment.^{24–31} Furthermore, during the regeneration process, adsorption stops the creation of hazardous chemicals, which results in secondary contamination. For efficient adsorption, adsorbents with appropriate functionality and large surface area are usually utilized for the adsorption process. Few porous materials MOF, activated carbon, zeolites, chitosan,

and kaolinite) were built as adsorbents for environmental contaminants. 32,33 Hydrogel material subject on the subject of the contaminants and kaolinite on the subject of the subject of

are also used as a superior adsorbent for the elimination of water-based contaminants.^{34,35}

The research community has shown substantial interest in 2D materials owing to their distinct chemical and physical properties, which set them apart from their bulk counterparts. This difference became particularly evident after the groundbreaking discovery of graphene through mechanical exfoliation into single layers.³⁶ Following the graphene's discovery, various 2D materials, including metal oxides, metal hydroxides, hexagonal boron nitrides, and TMDs, have been identified and demonstrated their capability across various applications.^{37–41} These 2D materials are low-dimensional materials possessing enormous surface areas and have been recognized as the best adsorbents for a variety of contaminants. Examples of these materials include carbon-based nanomaterials such as carbon nanotubes (CNTs) and graphene, ordered mesoporous silica, and phosphorenes. 42-44

A number of nanomaterials have lately shown great potential in water purification due to their notable physicochemical properties. Among them, MXenes are an innovative family of cuttingedge nanomaterials with promising applications in water purification and material science. They are also referred to as 2-D transition metal carbides, carbonitrides or nitrides. MXene's unique design, fine structure, remarkable stability, notable oxidation and chemical resistance, high conductivity, and environmental friendliness have piqued the curiosity of researchers recently. 45 They are ideal candidates for use in wastewater treatment applications due to their enhanced surface area and availability of several functional moieties, such as -OH and -O, on the MXene surface. 46 Their varied chemical composition, photo-catalytic capabilities, 2D layered architecture, and active metallic hydroxide sites have made these materials stand out as excellent candidates for use in water purification systems.⁴⁷ The MXenes-based materials are also very useful in the adsorption of various organic dyes which are toxic to the

environment. Additionally, MXenes serve as raw materials for the creation of nanostructive colling MXene membranes, demonstrating beneficial usage in wastewater treatment methods. 48,49

This review article addresses the potential use of MXenes, 2D transition metal carbides and nitrides in improving dye adsorption for wastewater treatment. MXenes are characterized by their unique surface chemistry, increased surface area, and superior conductivity, which positions them a prime choice to eliminate hazardous dyes from industrial effluents. In this article, the advancements in MXene-based adsorbents, including the physicochemical features, adsorption mechanisms that improve the efficacy of MXenes in dye removal operations, and the design and techniques of synthesis for these adsorbents are discussed. Additionally, it highlights recent progress in integrating MXenes into composite materials and membrane systems to optimize performance. Furthermore, the regeneration, recyclability, challenges and recommendations are also addressed for a sustainable environment. This review explores the viability of MXenes as an eco-friendly and sustainable approach for water pollution, paving the way for future investigation and expansion in sustainable wastewater management technologies.

2. Engineering structural and physicochemical properties of MXenes and related nanosorbents employed for dye removal

2.1. Fundamental structure of MXenes and their nanoadsorbents

MXenes are derived from the MAX phases, which are ternary compounds of carbides or nitrides. They follow with the common formula $M_{n+1}AX_n$, where M signifies a transition metal (Cr, Ti), A corresponds to an element either from the group 13 or 14 of the periodic table (e.g., Si, Al), X indicates nitrogen or carbon and n is typically 1, 2, or 3. These phases are characterized by densely arranged M layers, with X atoms occupying the octahedral sites within the layered hexagonal MAX phases. However, strong acids can easily etch away the A-layer due to the greater strength of M-X bonds compared to M-A bonds. After etching, the

resulting MXene can be expressed as $M_{n+1}X_nT_x$, where T signifies surface terminal groups with color of R023363 as -OH, -O, and -F, and so on, and x is termination number. This 2-D material, which shares similarities with graphene, was given the name "MXene" because of its structure. The suffix "-ene" is taken from the word "graphene," while "MX" is derived from the material's structure. Additionally, there are a lot of edges and void spaces between the various layers, which help adsorbates to diffuse and allows integration with nanomaterials or polymers. Examples of widely used MXenes are Mo_2CT_x , T_2CT_x , $T_3C_2T_x$, and so on. 50

The fundamental MAX phase and MXenes have the same hexagonal crystal structure. Typically, MAX phases are expressed as $M_{n+1}AX_n$ (n=1-3), which yields M_2AX (211), M_3AX_2 (312), and M_4AX_3 (413) (**Figure 2a**). ^{51,52} The "X" atoms fill the octahedral spaces created by the arrangement of "M" atoms of transition metal organizing into octahedra in all MAX phases, resulting in M_6X interspersed with "A" element sheets and M_6X overlapping with "A" element sheets. ⁵³ The only thing that separates the (211), (312), and (413) phases are the numbers of the M atom layers between A atom layer. Conclusively, the MAX phase consists of alternative layers of MX and A. Ti_3AIC_2 has a unit cell that resembles a hexagon and a five-atomic sheet structure. Among the five atomic sheets, three Ti sublayers containing carbon atoms are located at the "octahedral interstitial sites," and a reactive Al layer links the two adjacent layers. The surface terminating groups zone, the intralayer skeleton, and the interlayer make up the structure of $Ti_3C_2T_x$. ⁵⁴

Figure 2. (a) The structural arrangement of MAX phases and their associated MXenes. Reproduced with permission from Ref.⁵² Copyright (2013) John Wiley & Sons. (b) Various examples of different MAX phases.

2.2. Physicochemical properties of MXenes and their nanosorbents

The prominent properties of MXene include its tunable high Young modulus, band gap, and thermal and electric conductivities. Notably, the hydrophilic surfaces of MXenes, characterized by strong metallic conductivities, differentiate themselves from most 2D materials, such as graphene. Finally, to optimize the properties and performance of their applications, various strategies can be employed, including (i) composition adjustments (such as formation of solid solution and variation of "M" and "X" elements), (ii) surface functionalization (achieved through thermal and chemical treatments), and (iii) structural and morphological modifications. The primary features of the MXenes family are outlined below.

2.2.1. Mechanical attributes of MXenes for effective adsorption

The mechanical properties of MXenes are crucial for their effectiveness as adsorbents for dye removal. Young's modulus, a critical indicator of mechanical strength, assesses a material's rigidity and its ability to resist elastic deformation. MXenes are widely recognized for their high Young's modulus values, generally spanning from tens to hundreds of gigapascals (GPa).

For example, the Ti₃C₂Tx MXene has Young's modulus of approximately 330 GPa_Dwhile of Ro₂₃₃₆J

MXenes, such as V_2C and Nb_2C , also exhibit similarly high values. ^{61}MX ene membranes with 7 nm tip radius, demonstrate an effective Young's modulus and a tensile strength of 333 ± 30 GPa and 17.3 ± 1.6 GPa respectively. Additionally, the Young's modulus of Ti_3C_2 is 502 GPa, as determined through molecular dynamics simulations. 62,63 The layer thickness significantly affects the tensile strength of MXene. Luo *et al.* demonstrated that with the increase in thickness of the MXene film from 2.30 to 17.0 μ m, the elastic modulus decreases from 17 GPa to 8 GPa, and the tensile strength reduces from 61 MPa to 36 MPa. 64 In addition to possessing a high Young's modulus, MXenes exhibit robust physical properties. MXenes exhibit tensile strengths typically ranging from several hundred to several thousand megapascals (MPa), indicating considerable robustness. For example, $Ti_3C_2T_x$ MXene is reported to possess a strength of approximately 400 MPa. 65 Liang *et al.* demonstrated that addition of 3 wt% MXene enhanced the fracture mechanical Strength of ceramics by 7.6 MPa·m¹/², marking a 36% increase compared to unmodified ceramics. 66

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

This mechanical robustness is coupled with excellent flexibility due to their layered structure, which allows MXenes to bend and deform without compromising their structural integrity. Such properties are advantageous in wastewater treatment, where maintaining functionality under different conditions is essential. Flexibility is another essential aspect of mechanical behavior. MXenes have a layered structure that imparts a degree of flexibility, allowing them to bend and distort. The specific mechanical properties of MXenes can vary based on factors such as composition, doping, production methods, and the existence of defects or functional groups. Owing to their versatility, MXenes are appealing prospects for application in the adsorption of dyes.⁶⁷

2.2.2. Electronic and electrical properties aiding to effective dye removal

MXenes have distinct electronic and electrical properties that greatly improve their efficacy discrete or the discrete distinct electronic and electrical properties that greatly improve their efficacy discrete or their electric and electronic properties. MXenes can modify two of their primary attributes of interest: their electric and electronic properties. MXenes metallic or semi-metallic conductivity allows for quick electron transfer processes, which can increase dye-adsorbent surface interactions. This conductivity facilitates charge transfer, promotes electrostatic interactions, and may improve the adsorption kinetics of anionic or cationic dyes. Experimentally, the pressed MXene discs exhibited electrical conductivities comparable to those of multi-layered graphene (with resistance varying from 22 to 339 Ω , based on the chemical formula and 'n' index) and surpassed those of reduced graphene oxide (rGO) and CNTs. 52,68 Furthermore, the values of resistivity were found to increase with the functional groups and number of layers. 55,69,70 The calculated conductivities typically exhibit higher values than those measured experimentally. 71

The electrical conductivity of Ti₃C₂Tx ranged from 850 to 9880 S/cm, which can be attributed to variations in (i) defect concentration, (ii) surface functional groups, (iii) delamination yield, (iv) d-spacing between MXene flakes, and (v) lateral diameters influenced by the etching technique used. 72,73 Generally, reduced HF content and etching durations produce MXenes with less flaws and higher lateral dimensions, resulting in enhanced electronic conductivity (for instance, bigger flake sizes yielded conductivities five times greater than those of smaller MXenes). 74 Furthermore, environmental humidity may affect their conductivities, 75 indicating potential uses for relative humidity sensing materials. 76,77 An effective method to improve electrical characteristics is surface modification using alkaline and thermal treatments. They demonstrate conductivities that rise by up to two orders of magnitude. This augmentation results from the modification and/or elimination of functional moieties and embedded molecules. 78–80 Also, the functional groups present on the surface might result in localized electronic states, which boost adsorption capacity *via* enhanced chemical interactions. The

2.2.3. Optical attributes aiding to effective dye removal efficacies

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

MXenes have unique optical characteristics that make them ideal for dye adsorption applications. Their metallic conductivity and configurable bandgap enable efficient interaction with light, making them ideal for photocatalytic reactions. The functional moieties present on the MXene's surface can also affect light absorption, allowing for selective interactions with certain dyes based on their absorption spectra. Furthermore, the layered structure of MXenes can provide plasmonic effects, which increase the local electromagnetic fields around the material, enabling dye adsorption. This combination of optical features not only improves dye removal effectiveness but also provides opportunities for the progress of new materials for environmental remediation and sensing applications. 52,81

2.2.4. pH-Responsive surface chemistry of MXenes and its role in selective dve removal

MXene surface charge behavior and zeta potential variations across different pH conditions represent a critical yet underexplored aspect that significantly influences dye selectivity in adsorption applications. The surface terminations of MXenes (primarily -OH, -O, and -F groups) exhibit pH-dependent protonation and deprotonation behaviors, leading to dynamic changes in surface charge density and zeta potential values. Under acidic conditions, MXenes typically exhibit positive zeta potentials due to the protonation of surface hydroxyl groups, whereas alkaline environments promote deprotonation, resulting in increasingly negative surface charges. This pH-responsive behavior directly governs the electrostatic interactions between MXene surfaces and ionic dyes, where cationic dyes show enhanced adsorption at higher pH values when MXene surfaces are negatively charged, and anionic dyes preferentially

adsorb under acidic conditions when surfaces carry positive charges. Despite its fundamental icle online importance for rational design of selective dye removal systems, systematic investigations correlating MXene surface charge characteristics with dye selectivity mechanisms across comprehensive pH ranges remain limited, representing a significant knowledge gap that hinders optimization of MXene-based water treatment technologies. 60,82,83

2.2.5. Adsorptive attributes of MXenes and nanosorbents make them ideal adsorbents

The adsorption properties influence the performance of MXene-based adsorbent in removing colors from mixtures. Factors such as hydrophilicity, specific surface area (SSA), toxicity, porosity, uniformity, and various functional groups on MXene adsorbent surface are key determinant in adsorption of dye.

2.2.5.1. Enlarged specific surface area for high adsorption

Adsorption is often a surface process; thus, as an adsorbent's SSA increases, the number of pollutants absorbed also increases. By increasing surface area, several more active sites might be accessible for the likely adsorption of dyes. Usually, a large SSA provides additional active sites dedicated to the adsorption of pollutants, leading to a higher adsorption capacity. These nanomaterials' large SSA and negative surfaces of MXenes have made it possible for them to demonstrate possible adsorption capabilities.⁸⁴

2.2.5.2. Abundant surface functional groups for better adsorption

The functional groups on MXene-derived adsorbents, which are readily available, can significantly contribute to the removal of dyes from aquatic environments. This is due to the robust binding interactions between the molecules of the cationic dye and the negatively charged functional moieties on the MXene adsorbents. These interactions frequently lead to an increase in the amount of dye that is taken up by the adsorbents. Similarly, the existence of surface-bound positive functional groups on MXene-derived adsorbents promotes the adsorption of anionic dyes through electrostatic interaction (attractive) between the oppositely

charged species. Since different surface functional moieties gives different chemical via dicte online physical features of MXenes, their availability plays significant part in the utilization of MXene-based adsorbents. Additionally, the molecular intercalations or cations of tiny organic interlayer spacing also enhance the material's capacity for adsorption for MXene adsorbents. The increased interlayer space can explain this; consequently, increased analyte exposure occurs with interlayer gaps and functional divisions. 85

2.2.5.3. Significant wettability aiding efficient adsorption

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Another aspect that substantially affects an adsorbent surface's power for adsorption is its wettability. The wettability of MXene-based adsorbents can significantly influence selective dye uptake. Wet etching technique utilizing (in situ) hydrogen fluoride is the most widely used synthetic method for producing MXenes. This procedure gives the surface of the 2D sheets a spectrum of O and OH containing moieties, making them hydrophilic. Ref. As a result, MXenes show remarkable alignment with liquid procedure in an aqueous media, underscoring their increased capability for forming material hybrids and managing wettability in systems. Essentially, a liquid droplet's ability to wet on the MXene surface depends on the interaction at the solid-liquid phases. Some of the additional factors that affect this interaction are electrostatic interactions, H-bonding, and van der Waals interactions.

2.2.5.4. Improving stability for practical prospects of nanosorbents

One of the main problems researchers face with MXenes is its stability. Density functional tight-binding calculations, supported by formation and cohesive energy studies as well as ab initio simulations, have demonstrated that MXenes are stable. Specifically, hydroxylated MXene derivatives with varying atomic distributions of nitrogen (N) and carbon (C) represent stable phases with favorable thermodynamic properties. However, under real reaction conditions, the stability of MXenes and their surface topologies remains unknown. Transition metal-containing layered MXenes have a remarkable combination of excellent metallic

conductivity and rich, adaptable surface chemistry. MXenes become thermodynamically includes the conductivity and rich, adaptable surface chemistry. metastable as a result of this. Additionally, it is anticipated that MXenes' larger surface energy and low resistance to oxidation in oxygenated aqueous environments, even at room temperature, will lessen the electrical conductivity and passivate the reactive-interface.⁸⁸ Furthermore, after storing delaminated Ti₃C₂T_x MXene for two weeks, MXenes usually undergo oxidation, such as MXene titania layer oxidation to anatase TiO2, which results in a color shift to off-white from greenish-black.⁸⁹ Ti₃C₂T_x, on the other hand, was shown to be stable in an argon or hydrogen environment up to 1473 K, demonstrating that it is able to maintain its stability even at higher temperatures.⁶⁹ Thermal stability of MXenes (Ti₃C₂T_x) powders is demonstrated to be as low as 200 °C in oxygen, and as high as 800 °C in argon atmosphere. At 200 °C in oxygen, MXenes surface oxidizes to produce amorphous carbon and anatase TiO₂.^{69,90} Initially, the surfaces of MXene oxidize to form anatase crystals, preventing further oxidation. Then, as a result of the anatase's enhanced O₂ diffusion, the leftover MXene was gradually oxidized at higher temperatures until the anatase converted all of the MXene to CO₂, TiO₂, and rutile.⁶⁹

2.2.5.5. Toxicity related to MXene nanosystems

The toxicity profile of titanium-based MXenes presents a complex landscape requiring comprehensive evaluation across multiple biological systems and temporal scales. While nontoxic degradation products including C, N, and Ti breakdown into CO₂, N₂, and Ti compounds that are safe in trace amounts,⁹¹ the true group toxicity of MXenes remains challenging to ascertain due to their varied compound classes and significant heterogeneity in X or M elements and surface chemistry (Tx).92 Ti₃C₂Tx demonstrates concentration-dependent cytotoxicity with differential effects on normal versus cancerous cell lines. Studies reveal higher toxicity against cancerous cells (A549 and A375) compared to normal cell lines (MRC-5 and HaCaT). 93,94 Neural stem cell studies indicate no observable adverse effects at 12.5

μg/mL but significant cytotoxicity at 25 μg/mL with compromised cell membrane integrity. ^{95cle Online}
The primary toxicity mechanisms involve oxidative stress pathways generating reactive oxygen species (ROS), leading to cellular membrane damage and mitochondrial dysfunction. ⁹⁶
Surface termination chemistry significantly influences MXene biocompatibility profiles.
Fluorinated Ti₃C₂Tx synthesized via conventional HF etching exhibits enhanced cytotoxicity due to fluoride ion release, which promotes ROS generation. ⁹⁷ Halogen-free Ti₃C₂Tx produced through NaOH-based etching demonstrates superior biocompatibility with no noticeable cytotoxicity even at high concentrations. PEGylation alters protein corona formation and cellular uptake mechanisms, with PEGylated MXenes showing modified biocompatibility profiles. ⁹⁸

Environmental persistence studies reveal Ti₃C₂Tx stability in aquatic media for at least 4 days at concentrations of 5-20 mg/L, with no dissolution detected over 42 days. 99 Aquatic organism studies using zebrafish embryo models indicate LC₅₀ values of 257.46 µg/mL, classifying Ti₃C₂Tx as "practically non-toxic" under acute exposure conditions. 100 However, chronic exposure effects including developmental abnormalities and behavioral changes remain poorly characterized. Metabolomic analysis in Daphnia magna reveals metabolic disruption at potential ecosystem-level impacts. 101 sublethal concentrations. indicating environmental behavior involves complex transformation pathways influenced by water chemistry and redox conditions. Aggregation processes in natural waters alter bioavailability, while photodegradation generates titanium dioxide nanoparticles with unknown ecological implications.⁹⁹ Oxidative degradation over weeks to months may produce metabolites with different toxicity profiles compared to parent materials.

Comprehensive safety assessment using Safe and Sustainable by Design (SSbD) framework suggests Ti₃C₂Tx can be safe and sustainable when properly designed and applied.⁹⁹ Life cycle assessment (LCA) studies identify synthesis impacts as major environmental concerns,

particularly titanium precursor production and hazardous chemical usage. 102 Green, synthers a sticle Online

alternatives using electrochemical etching reduce environmental impact but require systematic toxicity evaluation. Current regulatory frameworks lack MXene-specific guidelines, creating knowledge gaps for commercial applications. Standardized testing protocols addressing unique physicochemical properties are needed for comprehensive risk assessment. Risk mitigation strategies include surface engineering for reduced toxicity, biocompatible coating systems, and real-time monitoring for environmental release detection. Critical research needs include longterm exposure studies, mechanistic toxicology investigations, and ecosystem-level impact assessments. Predictive toxicology models incorporating machine learning approaches can accelerate safety evaluation while reducing animal testing requirements. Integration of advanced characterization techniques will provide mechanistic insights essential for safe-bydesign approaches.

3. Methods for the synthesis of MXenes and their nanosorbents for dye removal

Most MXenes originate from MAX phase, in which A atoms are embedded in MX layers while M and X atoms positioned at hexagonal crystal's centre and apex, respectively. After eliminating A atoms from MAX phases, MX layers could be maintained by utilizing this "laminar" structure. However, unlike graphene, which rely on weak van der Waals forces to preserve the structure, the strong interlayer bonding in MXenes prevents mechanical exfoliation, making it impossible to produce 2D MXenes through this method. 103 Numerous methods have been introduced so far to prepare MXenes, which have numerous advantages and disadvantages (Figure 3).

View Article Online DOI: 10.1039/D5NR02336J

Methods for preparing 2D Mxenes

Advantages and Disadvantages

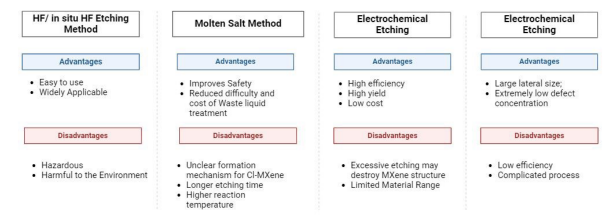


Figure 3. Advantages and Disadvantages of various methods of 2D Mxenes.

3.1. Top-down approaches

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Utilizing precursors is a key part of the top-down technique for the synthesis of MXenes. Although ScAl₃C₃, Hf₃(Al, Si)₄C₆, and Zr₃Al₃C₅, are examples of non-MAX structures, Ti₃AlC₂ and Ti₂AlC are examples of structures that are frequently found in MAX. At the moment, MAX is the precursor that is used the most frequently in the production of MXene. In MAX phase, strength of M-X bonds is greater than that of M-A bonds. Multi-layered MXenes are produced as a result of the breaking of the M-A connection in MAX during the synthesis process. This also results in element A being etched away from the parent structure. These multilayered MXenes are subsequently delaminated to yield single-layered MXenes. MXene's surface produces terminal groups like -OH, -F, and -O, to stabilize its structure. Mechanical stripping is commonly used in top-down methods of synthesis for 2D materials. Mechanical stripping alone is ineffective for breaking the precursor material's M-A bonds. Consequently, chemical stripping is the method of choice for the production of MXenes, whereas mechanical stripping is solely utilized for the production of single-layered MXenes by

the manufacturer. Various etching methods can disrupt M-A bonds: electrochemical attallice online base, fluoride, and lewis acid molten salt.

3.1.1. HF/in-situ HF-etching method

The two primary methods for preparing MXene are HF and in situ HF-etching (Figure 4a). 104 The loosely layered accordion-like MXene structure was initially described by Gogotsi and colleagues, who selectively etched Al from bulk Ti₃AlC₂ in HF solution (50 wt%). ¹⁰⁵ HF broke the Ti-Al bonds in the Al layers, dissolving them and releasing a high volume of H₂. As a result, the reaction started off with a powerful bubbling phenomenon. Additionally, a large number of functional moieties, like -F, -OH, and -O moieties, were joined to the surface of Ti atoms, giving rise to high hydrophilicity and distinctive electrochemical properties. Following this, HF etching was used to successfully exfoliate Ti₂CT_x, V₂CT_x, and Mo₂CT_x. ^{106–108} It should be mentioned that the MAX phase's structural characteristics determine the etching conditions. Ti₂CT_x only needed to be treated for 10 hours in 40-wt% HF, but Nb₂AlC needed 90 hours of etching time in 50-wt% HF. The outcomes were verified using theoretical approaches. The calculation findings confirmed that, in order to etch Al from Nb₂AlC rather than Ti₂AlC, a longer period and greater concentrations of HF solution are required. This is due to the fact that the bond energy of Nb-Al, which is 1.21 eV, marginally greater than Ti-Al, which is 0.98 eV respectively. 109 It is important to note, though, that an overly aggressive etching process will produce more surface flaws, which could adversely affect the characteristics of MXene sheets. 110 Successful Al layer delamination significantly reduces the interlayer force, facilitating the easy separation of MXene layers from their neighbouring ones.

Many attempts have been made to create safer and milder production methods for HF due to its significant toxicity and high danger. Using a combination of HCl and LiF rather than HF solution allowed for production of few-layer MXene with fewer surface flaws and greater interlayer spacing in 2014.¹¹¹ H⁺ and F⁻ were liberated during the etching process to create HF

3.1.2. Molten salt etching method

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Both in situ HF and HF etching are effective methods for generating carbides MXene, although they perform poorly when etching nitrides MXene. Two concepts were put out to illustrate the challenges involved in manufacturing nitrides MXene. Initially, the computational results indicated that $Ti_{n+1}N_n$ had a lower cohesive energy than $Ti_{n+1}C_n$, indicating lower stability. The energy required to form $Ti_{n+1}N_n$ is likewise greater than $Ti_{n+1}C_n$, indicating that the Al atoms were firmly bound in Ti_{n+1}AlN_n structure. 115 Gogotsi et al. used a combination of molten salt as etchant to produce nitrides MXene to get around these issues. 116 In specific, Ti₄AlN₃ was initially mixed with molten salt (29 wt% LiF, 59 wt% KF, 12 wt% NaF) in mass ratio of 1:1, and heated at 550 °C for 30 minutes in Ar atmosphere. After that, more washing with H₂SO₄ and deionised water was needed to eliminate the fluorides that contained aluminum. The prominent broad peak (002) of Ti₄N₃T_x shifted from 7.6° to 6.3°, as corroborated by XRD data, showing an extended interlayer distance. The EDX results also showed that no Al atom was seen, indicating that nitrides MXene was successfully etched using molten salt.

View Article Online

DOI: 10.1039/D5NR02336J

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Li and colleagues explained an alkali hydrothermal treatment process that yields high-purity multilayer MXene without the use of fluorine. 117 Al(oxide) hydroxides can dissolve completely at high temperatures and concentrations of NaOH. Additional delamination using DMSO or TMAOH together with ultrasound approach can aid in the production of a few-layer MXene with smaller diameters. 118 In contrast to fluorine-assisted approaches, there are more Ocontaining groups rather than -F groups dispersed on the surface, which could improve the electrochemical operation. Treatment with alkali prevents the production of HF. However, harsh environments might damage the internal structure and increase surface flaws. There is another method of creating MXene sheets besides conventional wet chemical etching, which is electrochemical etching. Yang and colleagues created an electrochemical etching technique using an organic system mixture of NH₄Cl and TMAOH. ¹¹⁹ The counter electrode and working electrode were made of two bulk pieces, and only working electrode was etched. Chloride ions functioned as the Ti-Al bond breaker at a constant applied potential of 5 V, and the NH₄OH that was produced assisted in extending the edge. Compared to the conventional HF-etched MXene, the exfoliated sheets had an average lateral size of nearly 2 μm and were primarily single or double layers. Furthermore, Ti₃C₂T_x sheets generated through electrochemical etching often have a stacked architecture that is comparable to that of bulk Ti₃AlC₂ without exhibiting any obvious expansion. This is due to the fact that the reaction process does not include the release of a significant amount of gas. It should be mentioned that the outcome of etched MXene is also greatly influenced by the choice of electrolytes. For instance, electrochemical treatment in solutions containing NaCl and HCl invariably produces amorphous carbon, which obstructs additional etching. 120 Sun and colleagues discovered that when MAX was treated with 2 M HCl aqueous electrolyte, a three-layer structural composite was produced. ¹²¹ The components of this hybrid were unetched MAX, carbon-derived carbines recorded RAZ (CDCs), and MXene, which required additional purification in order to get pure MXene sheets. Above all, using particular etching techniques such as molten salt etching, HF etching, and other fluorine-free approaches, MXenes could be extracted from bulk MAX phases by exploiting the difference in intensities between M–X and M–A bonds. It should be mentioned that the M–A bond energies had a significant impact on the etching conditions, with higher etching times and etchant concentrations needed for stronger M–A bond. Furthermore, the etching rate and reaction conditions are significantly influenced by the size of MAX particles. ¹⁰⁵ Attrition milling the MAX powder beforehand could significantly cut down on etching time while maintaining overall yield.

3.2. Bottom-up approaches

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Apart from the etching approach described above, various alternative bottom-up techniques have been developed to build ultrathin 2D highly crystalline MXene particularly for Mo₂C materials. These techniques include chemical vapor deposition (CVD) method, template method, PEPLD.^{122–126}

3.2.1. Chemical vapor deposition method

CVD was utilized by Ren *et al.* to successfully generate ultrathin two-dimensional α-Mo₂C crystals, with few nanometers thickness and lateral dimensions of up to 100 μm (Figure 4c). This procedure required the use of methane over bilayer substrate made of copper and molybdenum foils. Further, this CVD process has been expanded to produce ultrathin TaC and WC crystals from additional transition metals such as Ta and W. Notably, this approach has the benefit of producing MXenes with large lateral diameters and few flaws, which substantially simplifies the analysis of their intrinsic properties.¹²⁵ Despite the fact that the fabrication of MXene monolayers has not yet been achieved through this method, it presents the possibility of future research and advancement in bottom-up approach for synthesis of

MXene. MXene thickness and form can be controlled very precisely by the use of Clyphological continuous of the other hand, this procedure calls for highly specialized machinery and individuals who have received adequate training. 105

3.2.2. Plasma-enhanced pulsed laser deposition method

The PELPD system is a step up from traditional pulsed laser deposition systems. The very initial ultrathin films of Mo₂C synthesized using PELPD, made with methane plasma as the source of carbon, interacted with Mo vapor produced by pulsed laser. Utilizing sapphire substrate that was heated upto 700 °C, this reaction was carried out in order to generate films of high-quality with thickness that can be altered by modifying the pulse rate of laser.^{122,123}

3.2.3. Template method

To create 2D MoN, a salt-templated technique was employed. In this process, hexagonal 2D NaCl crystals coated with MoO₃ were first prepared, and then NH₃ was used to ammoniate them at 650 °C (**Figure 4d**). As a result, 2D MoN@NaCl powder, in which MoN species combined with the NaCl matrix, were successfully synthesized. 2-D nanosheets of MoN were produced by removing the NaCl using deionized water afterward. The high yield of the template approach is its main benefit over CVD. This method also provides the capacity to precisely regulate the surface functional moieties. It is crucial to recognize that energy input is necessary for the precursor to be converted into nitride or carbide, which adds to energy consumption. 124

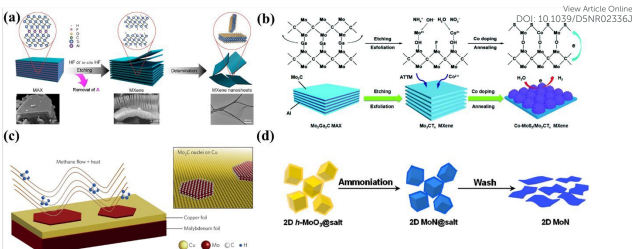


Figure 4. (a) Illustration of the process of etching and delamination of MXene. Reproduced with permission from Ref.¹⁰⁴ Copyright 2019, Elsevier. b) The schematic diagram for synthesizing the Co–MoS₂/Mo₂CT_x nanohybrids. Reproduced with permission from Ref.¹⁰⁶ (c) CVD growth of large-area α-Mo₂C thin layer on Cu substrate. Reproduced with permission from Ref.¹²⁶ Copyright 2015, Springer Nature. (d) Schematic of synthesis of 2D MoN. Reproduced with permission from Ref.¹²³ Copyright 2015, American Chemical Society.

4. Rational engineering of MXene-based materials for enhanced dye removal performance

The rational engineering of MXene-based materials for enhanced dye removal performance represents a systematic approach to designing and optimizing these advanced adsorbents through targeted modifications that address specific limitations and enhance desired properties. As the field of MXene-based water treatment technologies has evolved, researchers have developed sophisticated strategies that go beyond the use of pristine MXenes to create engineered materials with superior performance characteristics. These rational design approaches are guided by a fundamental understanding of structure-property relationships, adsorption mechanisms, and the specific requirements for effective dye removal from aqueous environments. The engineering strategies encompass four primary categories: surface

functionalization to introduce particular chemical interactions, interlayer engineering resolution to introduce particular chemical interactions. optimize structural accessibility, composite formation to achieve synergistic effects, and morphological control to enhance practical implementation. Each of these approaches addresses different aspects of the dye adsorption process, from molecular-level interactions to macroscopic material handling and system integration. A comparative analysis of published literature reveals several vital aspects when examining the efficiency of various MXene-based materials in dye adsorption (Table 1). The information includes adsorption capacity or removal efficiency, experimental conditions, and the type of adsorption isotherms and kinetics models applied for removing different dyes from the aqueous environment using MXene-based materials. This comprehensive evaluation demonstrates that rational engineering approaches have successfully achieved remarkable improvements in adsorption performance, with engineered MXene-based materials exhibiting significantly enhanced adsorption capacities representing orders of magnitude enhancement compared to pristine MXenes. The systematic design and optimization of these materials not only advance our fundamental understanding of MXene chemistry and physics but also provide practical pathways toward the development of commercially viable water treatment technologies capable of addressing the growing global

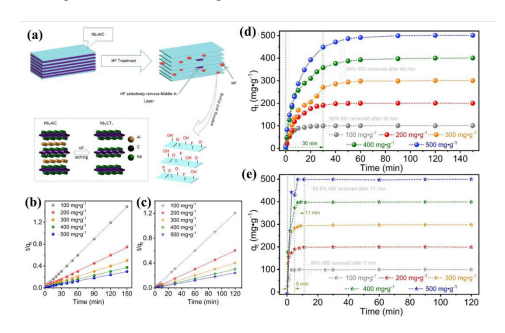
4.1. Surface Functionalization Strategies

challenge of dye-contaminated wastewater.

Surface functionalization is a key method for improving the adsorption capabilities of MXenes by changing their surface chemistry and charge properties. The natural surface terminations of MXenes (-F, -OH, =O) create reactive sites for various chemical modifications, allowing the addition of specific functional groups that can greatly enhance their selectivity, capacity, and stability for dye removal. These methods aim to modify the surface features of MXenes without significantly altering their layered structure, thus maintaining their unique 2D properties while adding new functionalities. Surface functionalization techniques include chemical

modifications through covalent bonding of organic groups, alkali treatments to change suffice online terminations, organic coupling reactions to attach complex molecular structures, and surfactant modifications to adjust surface charge and hydrophilicity. Each approach provides distinct benefits, such as boosting electrostatic interactions, hydrogen bonding, π - π interactions, and other molecular interactions that affect dye adsorption. Successful surface functionalization not only improves adsorption performance but also enables the creation of pH-responsive materials, selective adsorbents for specific dyes, and systems with better regeneration ability, making them highly useful for practical water treatment.

Li and co-authors effectively synthesized Nb₂CT_x MXene with a particular surface area by etching Nb₂AlC with hydrofluoric acid (**Figure 5a**). The produced Nb₂CT_x MXene is highly effective at adsorbing methylene Blue (MB) and methyl orange (MO) dyes, achieving maximum capacity exceeding 500 mg/g. For starting concentrations of 100 and 200 mg/g, ~99% MO dye may be removed in ~30 min (**Figure 5 d and e**). However, MB dye removal requires just 5 min of contact time. Furthermore, the adsorption process of Nb₂CT_x MXene is governed by a pseudo-second-order kinetics model (**Figure 5 b and c**), with chemisorption at the surface absorption active sites serving as the rate limit.¹²⁷



The elimination of MB and congo red (CR) dyes from wastewater was accomplished by Li *et al.* by the fabrication of AA-alk-MXene-based adsorbent. This was accomplished by attaching acrylic acid (AA) to alkalized single or several layered MXene nanosheets. The influence of pH, temperature, concentration of dye, contact time and AA dosage on adsorption was examined. The process of adsorption used the pseudo-second-order kinetic and langmuir isotherm adsorption model. The highest adsorption capacity of a AA-modified sample (2 ml) for MB and CR was found to be 193.92 mg/g and 264.46 mg/g, respectively. Furthermore, hydrogen bonding, electrostatic attraction, along with interlayer force, are likely the primary driving forces behind the adsorption mechanism of AA-alk-MXene (Figure 6 a and b). 128

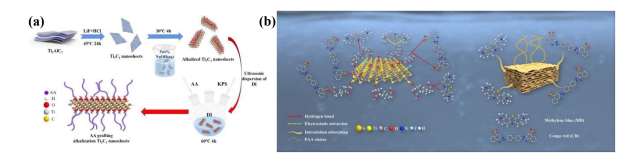


Figure 6: (a) Schematic Diagram for preparing AA-alk-MXene; (b) The adsorption mechanism diagram of AA-alk-MXene for MB and CR. Reproduced with permission from Ref. 128 Copyright 2022, Elsevier.

By modifying Ti_3C_2 to incorporate sulfonic groups through aromatic coupling-diazotization and incorporate sulfonic groups through aromatic coupling-diazotization and the sulfonic groups through a sulfonic groups through the sulfonic group groups through the sulfonic group group groups through the sulfonic group Wei et al. achieved the fabrication of 2D material MXenes. This was accomplished by removing the Al layer from Ti₃AlC₂ through a simple process. A further investigation into the adsorption behavior of functionalized Ti₃C₂ against MB was conducted under various experimental conditions, including pH, solution temperature, initial MB concentration, and contact time. The results demonstrated that Ti₃C₂-SO₃H achieved an adsorption performance of 111.11 mg/g. Kinetic and isotherm analyses discovered that the pseudo-first-order and the Langmuir isotherm adsorption models were appropriate for explaining the experimental data. The deposition of MB onto adsorbent surfaces was endothermic, while dye adsorption is most effective when the aqueous solution is alkaline (Figure 7). 129

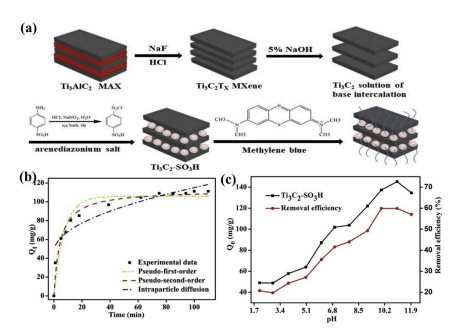


Figure 7: (a) Schematic diagram for the preparation of Ti₃C₂–SO₃H for MB removal. (b) The kinetic curve of Ti₃C₂-SO₃H adsorption MB. (c) Effect of pH; on the adsorption of MB onto Ti₃C₂–SO₃H. Reproduced with permission from Ref. ¹²⁹ Copyright 2019, Elsevier.

Kelishami et al. designed a cetyltrimethylammonium bromide (CTAB)-modified multi-layered Ti₃C₂Tx MXene (CMM) from Ti₃AlC₂ precursor and was employed to remove MO (Figure 8

a and b). This material adsorbent was resulted from an electrostatic combination between the resultance cationic surfactant solution (CTAB) and negatively charged $Ti_3C_2T_x$ nanosheets. The accessible active sites were exposed as a result of this, which resulted in an increase in the spacing between the nanosheets and an increase in adsorption efficiency. The adsorbents were analyzed by using a few analytical techniques, and the impact of various parameters (contact time, pH, loading adsorbent, and initial concentration of dye) was investigated. The results of this study demonstrated that MO adsorbs CMM to its maximum capacity at an adsorbent dosage of 0.83 g/L, a contact time of 90 minutes, and a solution pH of 3. The results of the adsorption experiment were most accurately represented by the pseudo-second-order kinetic model ($R^2 = 0.9924$) and the Langmuir isotherm ($R^2 = 0.9990$). Approximately 213.00 mg/g was the maximum amount of adsorption capacity that MO possessed. There is a possibility that the MO adsorption mechanism on CMM involves many interactions, including electrostatic adsorption, π -cation interactions, and hydrogen bonding. 130

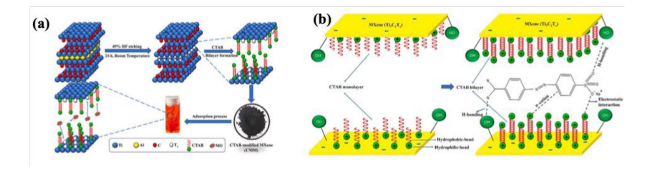


Figure 8. (a) Illustration diagram for the intercalation of CTAB in MXene (CMM) for the adsorption of MO; (b) CTAB monolayer and bilayer formation on MXene (Ti₃C₂T_x) sheets and adsorption mechanisms of MO by MXene sheets. Reproduced with permission from Ref.¹³⁰ Copyright 2024, Elsevier.

4.2. Interlayer Engineering and Spacing Modification

Interlayer engineering and spacing modification represent critical strategies for optimizing Marchanical Str structural architecture of MXenes to enhance their dye adsorption capabilities. The layered nature of MXenes, with interlayer spaces typically ranging from 0.98 to 1.5 nm in their pristing form, provides unique opportunities for structural manipulation to accommodate different dye molecules and improve accessibility to adsorption sites. These approaches focus on controlling the interlayer distance through various techniques that either expand, maintain, or systematically modify the spacing between MXene layers. Interlayer expansion techniques involve the insertion of molecules or ions that physically separate the layers, creating larger void spaces for enhanced accommodation of dye molecules. Ion intercalation methods utilize cations or anions to penetrate between layers, causing structural swelling and increased surface accessibility. Delamination strategies aim to entirely or partially separate multi-layered MXene structures into individual or few-layer nanosheets, maximizing the available surface area. Pillaring approaches involve the insertion of rigid molecular or ionic species that act as structural supports, maintaining controlled interlayer distances while preventing restacking. These interlayer engineering strategies are particularly effective because they directly address the accessibility limitations inherent in layered materials, enabling larger dye molecules to penetrate the interlayer galleries and access previously unavailable adsorption sites. The controlled modification of interlayer spacing also facilitates size-selective adsorption, improves mass transfer kinetics, and enhances the overall adsorption capacity by creating a three-dimensional network of accessible adsorption sites throughout the material structure. Gogotsi et al. tested Ti₃C₂T_x stacked sheets bound by hydrogen bonds and/or van der Waals interactions for the adsorption of MB dye. Ti₃C₂T_x showed adsorption capacity of 39 mg/g for MB. The adsorption of MB on Ti₃C₂T_x was most closely with Freundlich isotherm model. ¹³¹ Through the use of alkali solution, ZhengMing et al. described a straightforward method for increasing the interlayer gap of Ti₃C₂T_x while modulating the surface functional groups of the

material (Figure 9a). 132 Using LiOH, the approach is able to enhance the spacing between Ro23361

layers by 29% in Ti₃C₂T_x MXene. Additionally, the modification of functional groups involves the transformation of -F into -OH. NaOH-Ti₃C₂T_x and LiOH-Ti₃C₂T_x MXenes are able to adsorb MB more quickly compared to other types of MXene adsorbents. Notably, NaOH-Ti₃C₂T_x has the maximum ability to adsorb MB, capacity of 189 mg/g, attributed to the combination of intercalation adsorption and surface adsorption of the MXene (Figure 9b). The High-resolution transmission electron microscopy (HRTEM) images (Figure 9c, d, and e) reveal significant structural modifications following alkaline metal ion intercalation in LiOH-Ti₃C₂T_x, NaOH-Ti₃C₂T_x, and KOH-Ti₃C₂T_x systems. The intercalation process involves alkaline metal ions (Li⁺, Na⁺, K⁺) inserting between MXene layers, causing interlayer spacing expansion through electrostatic interactions and hydration shell formation. Structural expansion mechanisms include: direct ion insertion where alkaline cations occupy interlayer galleries, creating physical separation between Ti₃C₂T_x sheets; hydration-induced swelling as intercalated ions attract water molecules, forming hydration shells that further increase interlayer distance; and surface functionalization changes where alkaline treatment converts surface -F terminations to -OH groups, enhancing hydrophilicity and water uptake. The measured interlayer spacings show progressive expansion: LiOH-Ti₃C₂T_x (~1.22-1.65 nm), NaOH-Ti₃C₂T_x (~1.16-2.0 nm), and KOH-Ti₃C₂T_x (~1.40-1.73 nm), correlating with ionic radius differences (Li⁺ < Na⁺ < K⁺) and hydration characteristics. This structural expansion directly contributes to enhanced adsorption capacity by increasing accessible surface area and creating larger pore channels for dye molecule accommodation, explaining the superior MB

adsorption performance observed in alkaline-modified MXene systems.

Nanoscale Accepted Manuscript

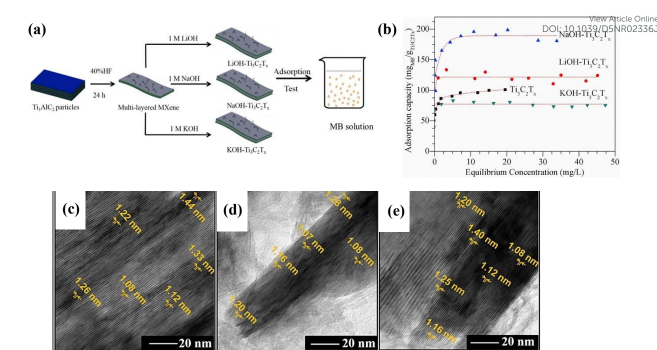


Figure 9: (a) Illustration diagram for the fabrication of different Alkali- $Ti_3C_2T_x$ towards adsorption of MB dye. (b) Adsorption isotherms of MB on $Ti_3C_2T_x$, LiOH- $Ti_3C_2T_x$, NaOH- $Ti_3C_2T_x$ and KOH- $Ti_3C_2T_x$. HRTEM images of (c) LiOH- $Ti_3C_2T_x$, (d): NaOH- $Ti_3C_2T_x$ and (e): KOH- $Ti_3C_2T_x$. Reproduced with permission from Ref. Copyright 2017, Elsevier.

Yu and co-workers developed a novel hydrothermal method utilizing less toxic etching agents such as NaBF₄, HCl for the synthesis of Ti₃C₂ MXene (*h*-Ti₃C₂). Because of slow-release mechanism during hydrothermal, h-Ti₃C₂ has a higher lattice parameter *c*, and a longer interlayer distance, as well as a better SSA than t-Ti₃C₂. The hydrothermal etching method avoids high-concentration HF and is more effective in synthesizing Ti₃C₂ flakes. Furthermore, the etching method can be used for other MXene compositions, such as Nb₂C and *h*-MXenes shows superior adsorption performance for MB, with a capacity of 24 mg/g.¹³³ Cagnetta and colleagues synthesized Ti₃C₂T_x MXene layers pillared with terephthalate using an innovative MC etching approach that involved exfoliating a titanium MAX phase with a small volume of concentrated hydrofluoric acid using a high-energy ball mill. The obtained material has a

greater SSA and strong adsorption capability of MB of 209 mg/g because of wider interlar advantage of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong adsorption capability of MB of 209 mg/g because of wider interlar and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and strong additional capability of MB of 209 mg/g because of wider and wider space between free carboxylate groups of terephthalate and MXene sheets. Also, dye adsorption with the material is best explained by pseudo-second-order model. 134

Yang et al. produced a suspension of Ti₃C₂T_x MXene nanosheets using the intercalation and delamination of multi-layered Ti₃C₂T_x utilizing TMAOH. The prepared material showed excellent adsorption of cationic MB dye. The highest adsorption capacity, noted at 318 K, reached 1026 mg/g. This performance best fits with pseudo-second-order kinetic and Freundlich isotherm models. The mechanism of adsorption is mainly attributed to the ion exchange and electrostatic attraction for MB removal. 135 By intercalating GO into an Alk-MXene layer, Li and the authors were able to create a novel AMXGO absorbent. This absorbent demonstrated a high level of efficiency in the elimination of MG and anionic CR (Figure 10). The results of the FTIR, XRD, and SEM investigations showed that the AMXGO absorbent possesses a characteristic 3D layer-by-layer structure with plentiful oxygen-bearing groups, and that its heat stability has greatly improved. Based on the results of the BET analysis, it was determined that the AMXGO1 adsorbent possesses a greater SSA (16.686 m²/g) and pore volume (0.04733 cm³/g). It was discovered that the performance of adsorption was dependent on the mass ratio of Alk-MXene to GO, the starting dye concentration, pH, contact time, and temperature. The AMXGO1 absorbent, which had a mass ratio of 3:1, had the maximum capacity to adsorb 1111.6 mg/g of MG and 1133.7 mg/g of CR, and it had dye removal rates that were greater than 92%. Both pseudo-second-order kinetic and Freundlich isotherm models are utilized to describe the adsorption behaviour of AMXGO1 for both CR and MG. 136

Figure 10: Diagram for the preparation of AMXGO. Reproduced with permission from Ref. ¹³⁶ Copyright 2024, Elsevier.

4.3. Composite Formation Strategies

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Composite formation strategies represent a versatile and powerful approach to developing advanced MXene-based adsorbents by synergistically combining the unique properties of MXenes with complementary materials to achieve enhanced dye removal performance. These strategies leverage the principle that the integration of different materials can create synergistic effects that surpass the individual performance of constituent components, resulting in composites with superior adsorption capacities, improved selectivity, enhanced stability, and additional functionalities such as magnetic separability or pH responsiveness. The composite formation approaches can be systematically categorized based on the nature of the secondary materials integrated with MXenes. MXene-organic composites involve the combination of MXenes with organic polymers, biomaterials, or carbon-based materials, creating hybrid systems that benefit from the conductivity and surface chemistry of MXenes while gaining enhanced mechanical properties, biocompatibility, or specific functional groups from the

organic components. MXene-inorganic composites integrate MXenes with metal oxide State online

hydroxides, or other inorganic materials, often resulting in enhanced adsorption capacity through increased surface area, additional adsorption sites, or new interaction mechanisms such as magnetic separation capabilities. Multi-hybrid systems represent the most sophisticated approach, combining MXenes with multiple different material types to create complex architectures that integrate the advantages of various components, such as combining organic polymers for flexibility, inorganic nanoparticles for functionality, and MXenes for conductivity and surface area. These composite formation strategies not only enhance the fundamental adsorption properties but also address practical considerations such as material recovery, regeneration efficiency, mechanical stability, and cost-effectiveness, making them particularly attractive for real-world water treatment applications where multiple performance criteria must be simultaneously optimized.

Peng *et al.* effectively synthesized MXene-based core-shell composites, MXene-COOH@(PEI/PAA)_n, using layer-by-layer technique. Furthermore, these nanocomposites were studied using spectral and morphological techniques, revealing that they contain more reaction sites and mesoporous structures. The produced composites were able to efficiently adsorb MB dye after approximately 200 minutes, showing that the generated composites might be used as a highly efficient adsorbent. The observed data are well matched with pseudo-second-order model and show a significant correlation coefficient of R²> 0.99, indicating that the core-shell composites have outstanding adsorption capabilities.¹³⁷

The PHGC/MXene that was created by Zhang *et al.* (**Figure 11 a and b**) exhibited remarkable a strong selectivity and cycle stability for ionic dyes across a range of pH levels. PHGC/MXene shown exceptional adsorption selectivity and great pH-responsiveness in the mixed dyes system. It was able to adsorb MB at a pH of 11.0, and methyl blue (AB93) at pH 2.0, achieving maximum adsorption capacities of 555.56 mg/g and 207.47 mg/g, respectively. The adsorption

kinetics for these dyes were found to be in agreement with the two-level kinetic model that was dicte online proposed, while Langmuir model demonstrated a good fit for adsorption isotherms being studied. It was determined that electrostatic adsorption and hydrogen bonding were the primary processes responsible for the adsorption process, with van der Waals forces also making a contribution. Despite undergoing 12 rounds of desorption and regeneration, the clearance rates for MB and AB93 remained more than 90%. 138

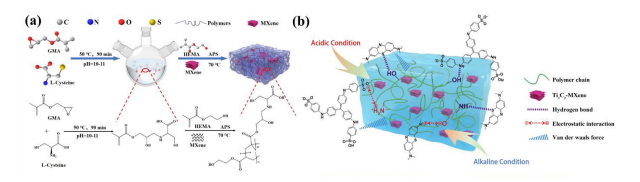


Figure 11. (a) Schematic preparation of PHGC/MXene hydrogel; (b) Adsorption mechanism diagram of PHGC/MXene for AB93 or MB. Reproduced with permission from Ref.¹³⁸ Copyright 2019, Elsevier.

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

The amino-functionalized bacterial cellulose/Ti₃C₂T_x MXene (ABC/MX) composites were developed by Wen and his colleagues by the utilization of an electrostatic self-assembly technique that comprised modifications of polydopamine (PDA) and polyethyleneimine (PEI) (Figure 12a). Based on the findings of this research, the effective insertion of amino groups strengthens the interfacial contacts between nanofibers of BC and nanosheets of Ti₃C₂T_x, it also enhances the number of active sites for adsorption. With a maximal ability to adsorb 1103.7 mg/g of CR, the results demonstrate that the composite possesses a remarkable removal efficiency.¹³⁹ Using a chemical etching approach, Wang *et al.* created a few-layer MXene. Subsequently, PEI was grafted onto the surface of the MXene with the assistance of glutaraldehyde-assisted crosslinking (Figure 12b). The adsorption properties of water-soluble

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

dyes were extensively studied. The results indicated that grafting PEI may improve Microscopic explains a successful as adsorption capacities. The modified MXene showed a remarkable ability to adsorb 909.1 mg/g of MO. **Figure 12c** reveals the binding mechanism between the components of the MPEI composites. The pseudo-second-order kinetic and Langmuir isothermal adsorption model are the best appropriate to the adsorption processes of MO, which are spontaneously endothermic and chemisorption. 140

Wang and co-authors designed a novel $CoFe_2O_4/CS$ composite-supported onto alk-MXene magnetic adsorbent *via* hydrothermal and self-assembly approach (**Figure 12d**). Adding $CoFe_2O_4/CS$ to alk-MXene can significantly increase the capacity to adsorb anionic and cationic dyes. The prepared material showed high adsorption capacity upto 1333.9, 537.6, and 2095.9 mg/g for rhodamine B (RhB), malachite green (MG), and CR dyes, respectively. The higher adsorption performance of the composites is attributed to synergistic effect of hydrogen bonding, π - π interaction, and electrostatic interaction.¹⁴¹

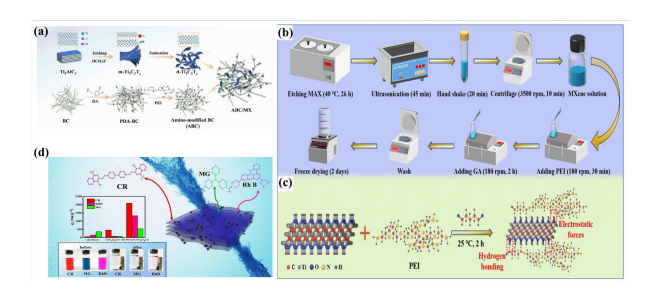
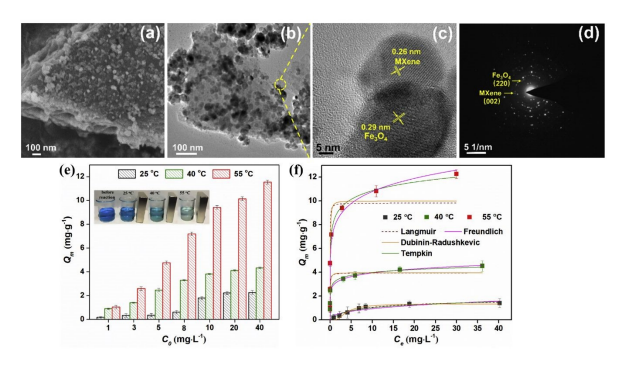


Figure 12: (a) Schematic diagram of the preparation of ABC/MX composite. Reproduced with permission from Ref.¹³⁹ Copyright 2024, Elsevier. (b) Synthesis procedures of the MPEI composites and (c) Diagram of the inter-component microscopic combination mechanism in MPEI composites. Reproduced with permission from Ref.¹⁴⁰ Copyright 2023, Royal Society of

Zhang *et al.* used an in-situ growing technique to create a new 2D MXene coated with Fe₃O₄. The prepared material was analyzed for MB dye adsorption at different temperatures. This material exhibits superparamagnetic characteristics and a typical 2D lamellar structure. The elimination of MB occurred as endothermic process, as evidenced by maximum elimination capacity of 11.68 mg/g and 91.93% decolorization efficiency at 55 °C, considerably surpassing the performance at lower temperatures. Additionally, at high temperatures (40 and 55 °C), the adsorption isotherm demonstrated that the model which best fits for removal method of MB is Freundlich isotherm, but at low temperatures (25 °C), the Langmuir isotherm fit best. Through electrostatic attraction and hydrogen bonding at high temperatures, Ti-OH groups on the material's surface enhance MB decolorization. At 25 °C, surface adsorption by electrostatic interaction aids in the elimination of MB (Figure 13).¹⁴²

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM



Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Figure 13: (a) SEM, (b) TEM, (c) HRTEM, (d) SAED pattern of 2D-MX@F@Qfricle Online nanocomposites. (e) The removal capacity of MB over 2D-MX@Fe₃O₄ at 25, 40 and 55 °C. (f) Adsorption isotherm of 2D-MX@Fe₃O₄ for MB removal. Reproduced with permission from Ref. 142 Copyright 2019, Elsevier.

Eltaweil et al. fabricated Ti₃C₂T_X/NiFeMn-LDH@Gel composites in order to remove organic CR dye synthesized via cross-linking reaction by incorporating Ti₃C₂T_X MXene and NiFeMn-LDH into gelatin (Figure 14a). The results shown that the adsorption ability of Ti₃C₂T_X/NiFeMn-LDH@Gel towards CR was significantly enhanced by raising the Ti₃C₂T_X content in the matrix to 10%. With a q_{max} value of 588.24 mg/g, the Freundlich model offered the best explanation for adsorption of CR. Moreover, pseudo-second-order model was determined to be best for studying the adsorption kinetics of CR accompanied. The remarkable recyclability over multiple cycles confirms the composite's sustainability. 143 Using a macroporous polymeric support, Yang et al. were able to effectively generate a flexible, rigid, and porous MXene/COF hybrid flake. This flake is characterized by the presence of three 2D COF flakes that are bridged covalently on the surface of the MXene by the process of in-situ growth. COFs (TpTAPB, TpBD, TpPa) were synthesized via Schiff-base reactions using 1,3,5-triformylphloroglucinol with 1,3,5- tris(4-aminophenyl) benzene, benzidine and p-phenylenediamine monomers (Figure 14b). Further, interlayer of MXene (Ti₃C₂T_x) nanosheets expands adjacent interlayer spacing and introduces numerous sieving pores when COFs are intercalated into it. The optimized MXene/TpTAPB hybrid membrane when utilized for eliminating organic dyes, it exhibits a removal efficiency of 96.4% for AB93, 98.2% for CR, 97.2% for MO and 98.7% for chrome black T.144

Wang *et al.* developed alk-MXene/ZIF composite, with small ZIF particles in situ synthesized on interlayer and on the alk-MXene surface (**Figure 14c**). In the meantime, stable intercalation

framework is produced, allowing dye molecules to transfer mass more quickly. Adding profice online particles increases the adsorption functional groups and SSA in alk-MXene/ZIF composite.

The composites achieved excellent adsorption values, with maximum capacity of 7111.3 and 539.7 mg/g for MG and CR, respectively. The impact of coexisting ions, temperature, concentration, contact time, and pH on the adsorption performance were studied. Adsorption kinetics studies demonstrated that dye adsorption onto the composite surface is consistent with Elovich and pseudo-second-order kinetic models. Additionally, the adsorption mechanism of adsorbent is mainly due to chemisorption. 145

By integrating few-layer MXene and hydrophilic biomaterial DASNP, followed by immobilization of DASNP onto few-layer MXene through dialdehyde-based cross-linking technique, Li et al. were able to generate a novel environmentally friendly composite material known as DSP-M (**Figure 14d**). The findings indicate that DSP-M had outstanding adsorption efficiency for both adsorbates. Further, Langmuir maximum adsorption capacity for monomer adsorption for RhB was measured to be 678.19 mg/L, while the capacity for CR was measured to be 754.41 mg/L. These results were in agreement with the pseudo-second-order, intraparticle diffusion, and Langmuir models. For the purpose of conducting additional research on the adsorptive mechanism, the statistical model of physics was utilized. The DSP-M characteristics post-adsorption and findings of the simulation suggest that the RhB and CR removal by DSP-M occurred predominantly through physical adsorption mechanism involving van der Waals, electrostatic force, and hydrogen bonding. 146

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Wang and co-authors designed a ZnS/CuFe₂O₄/MXene (ZSCFOM) composites featuring ternary heterostructures *via* solvothermal approach to efficiently adsorb azo dyes. The composite primarily achieved azo dyes adsorption via electrostatic interactions and hydrogen bonding, with maximum capacity to adsorb 377 mg/g for DBM and 390 mg/g for DBRN. The

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

pseudo-second-order and Langmuir model were used to characterize the adsorption property sticle Online of ZSCFOM, suggesting that adsorbate forms a monolayer on the ZSCFOM surface. 147

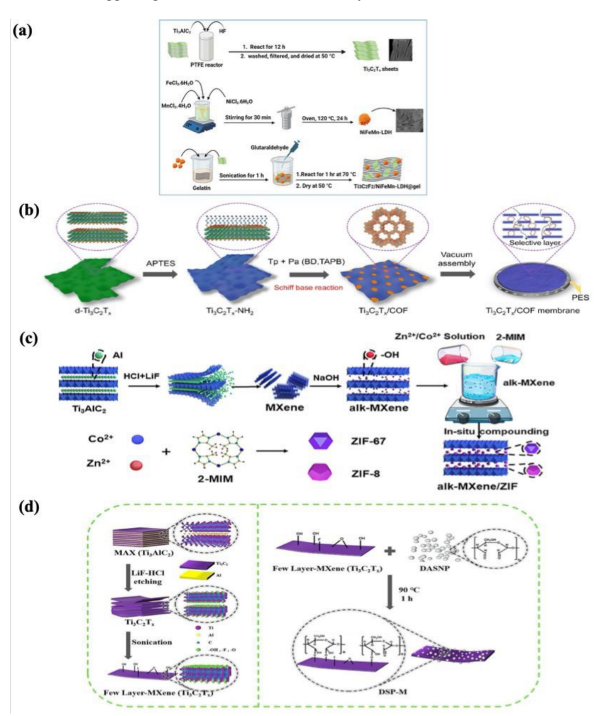


Figure 14: (a) A schematic presentation for the synthesis of Ti₃C₂T_X/NiFeMn-LDH@Gel composite. Reproduced with permission from Ref.¹⁴³ Copyright 2024, Elsevier. (b) Schematic illustration of the separation mechanism of MXene/COF in the dye wastewater purification process. Reproduced with permission from Ref.¹⁴⁴ Copyright 2024, Elsevier. (c) Schematic

preparation process of alk-MXene/ZIF composite. Reproduced with permission from Ref. 45 cle Online Copyright 2024, Elsevier. (d) Schematic illustration for the step-wise synthesis of DSP-M composite. Reproduced with permission from Ref. 146 Copyright 2023, Elsevier.

Moreover, green methods have also been adopted to design MXene-based nano adsorbents. Ijaz *et al.* developed an efficient and rapid Fe₂O₃/BC/MXene composite by functionalizing *Shorea faguetiana* biochar with Fe₂O₃ and MXene for selective adsorption. XPS and FTIR revealed that the Fe₂O₃/BC/MXene composites had more surface functional groups (F-, C=O, CN, NH, and OH-) than the original biochar. The characterization revealed that the intended modified BC had a high MB adsorption capacity in wastewater. At 293 K, MB showed a q_m value of 899.03 mg/g. The adsorption followed a pseudo-second-order model (R² = 1) and the Langmuir isotherm. Surface electrostatic forces and hydrogen bonding helped to remove MB more efficiently. This demonstrates monolayer adsorption on the Fe₂O₃/BC/MXene composite, which is regulated by chemical adsorption. ¹⁴⁸

An effective, fast, and selective adsorption of MB was achieved by Imtiaz *et al.* through the

An effective, fast, and selective adsorption of MB was achieved by Imtiaz *et al.* through the development of Ni₃(HITP)₂/MXene/CS. This was accomplished by functionalizing chitosan biopolymer with MXene and 2D MOFs (Ni₃(HITP)₂). The TEM image clearly shows irregularly-shaped MOFs and spherical CS particles adhered to the MXene surface. At 298 K, MB had a maximum adsorption capability of 424.99 mg/g, 400.59 mg/g at 308 K, and 305.80 mg/g at 318 K. The adsorption of MB on Ni₃(HITP)₂/MXene/CS composite followed the pseudo-second-order kinetic and Langmuir models. The composite has -OH, -F, -O, and N groups, which allow for effective MB absorption *via* hydrogen bonding, electrostatic attraction, and complexation interaction.¹⁴⁹

4.4. Morphological and Structural Control

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Morphological and structural control strategies focus on engineering the macroscopic discolaring discolaring the macroscopic discolaring the macroscopic discolaring discolaring the macroscopic discolaring discolaring the macro architecture and three-dimensional organization of MXene-based materials to optimize their performance in dye adsorption applications. Unlike surface functionalization or interlayer modifications that primarily alter chemical properties or nanoscale structure, these approaches target the overall physical form, porosity, and architectural arrangement of MXene materials to enhance mass transfer, improve accessibility, and facilitate practical implementation in water treatment systems. The controlled manipulation of morphology enables the creation of materials with tailored pore structures, enhanced mechanical properties, and optimized flow characteristics that are essential for real-world applications. Membrane fabrication represents a critical approach for creating selective barriers with controlled permeability and high surface area-to-volume ratios, enabling efficient dye removal while maintaining structural integrity under operational conditions. Aerogel formation involves the development of ultralightweight, highly porous three-dimensional networks that maximize surface accessibility while providing excellent adsorption kinetics through interconnected pore channels. Nanofiber integration focuses on incorporating MXenes into fibrous architectures that combine high surface area with excellent mechanical flexibility and easy handling characteristics. Threedimensional architecture design encompasses the creation of complex hierarchical structures that integrate multiple length scales, from nanoscale MXene sheets to macroscopic frameworks, enabling optimized mass transfer pathways and enhanced adsorption site utilization. These morphological control strategies are particularly valuable because they address critical practical considerations such as pressure drop in flow-through systems, ease of material recovery and regeneration, mechanical durability under operational stresses, and scalability for industrial applications. By controlling the overall architecture and form factor of MXene-based adsorbents, these approaches bridge the gap between fundamental material

properties and practical implementation requirements, enabling the development of efficiency included in the development of efficiency in the efficiency in the development of efficiency in the development of efficiency in the development of efficiency in the efficiency in the development of efficiency in the ef

Purkayastha and co-author used vacuum filtering to produce 2D Ti₃C₂T_X MXene on porous PAN membrane, demonstrating its effective adsorption of MB. MX-PAN had an adsorption effectiveness of 85%, compared to 12% achieved by PAN membrane. Adsorption efficiency increased dramatically after manufacturing PAN membrane along with Ti₃C₂T_X MXene nanoflakes. Shang *et al.* combined SA with d-Ti₃C₂T_X nanosheets (MXene), then employed electrospinning and subsequent Ca²⁺-mediated crosslinking for production of various SA/MXene NMs. The impact of MXene in NMs on MB adsorption ability were examined. SA/MXene NMs exhibited the ability to adsorb 440 mg/g of MB at the optimal MXene concentration comprising 0.74 wt.%, surpassing electrospun SA NMs, pristine MXene, or SA/MXene composite beads with equivalent MXene content. Moreover, the optimal SA/MXene NMs demonstrated high reusability. The Langmuir and pseudo-second-order model were in good agreement with the results, indicating monolayer adsorption aligned with the mechanism of chemical adsorption. 151

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

By using cross-linking approaches to incorporate PEI and amino-functionalized Ti₃C₂T_x into SA aerogel matrix, Wang *et al.* created the MXene/PEI-modified SA aerogel. The plentiful active PEI groups, together with the reduction capabilities of MXene, considerably enhance the adsorption capacity of 3568 mg/g for CR, attributed to the robust electrostatic attraction along with the synergistic effects of intercalation and surface adsorption. The results showed that adsorption process of CR aligned well with pseudo-second-order kinetic and Langmuir isotherm model. The mechanical strength of the aerogel was greatly enhanced by the double-network structure composed of polymeric PEI and SA, allowing for easy recycling without secondary contamination, with only a slight reduction in capacity following five cycles.¹⁵²

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM.

The innovative nanocomposite hydrogel of ZIF-8@IL-MXene/Poly(N-isopropylacry/lany) and Contine Online (NIPAM), which was fabricated by Xiong and his co-authors, is capable of successfully adsorbing crystal violet (CV) from wastewater (Figure 15a). The formation of composite hydrogel IL-MXene/PNIPAM was accomplished through in-situ polymerization by grafting IL onto MXene surface and subsequently introduced into solutions of NIPAM monomer. This process was completed in a single step. Utilizing in-situ formation of ZIF-8 on pore walls of composite hydrogels, ZIF-8@IL-MXene/PNIPAM were produced. The nanocomposite hydrogel displayed maximum adsorption capacities for CV of 325.03 mg/g under ambient temperature, with the reducing to 91% following five cycles of adsorptiondesorption. The results are in accordance with Freundlich and pseudo-second-order kinetics models, based on several interactions between molecules of adsorbent and hydrogel. 153 By self-assembling CNTs, CTAB and MXene nanosheets, Ding et al. were able to synthesize a unique CMC three-dimensional composite. In order to determine whether or not the produced material could adsorb AO7, CR, and MO the material was tested. In comparison to CNTs and MXene, the CMC composite had a greater capacity for adsorption of AO7, CR, and MO with maximum capacity to adsorb 367.9, 628.5, and 294.2 mg/g, respectively. As the temperature increased, the adsorption capabilities for AO7 and MO by CMC composite reduced, whereas the adsorption capabilities of CR increased. When it comes to the process of adsorption, the hydrogen bonding, π - π electron-donor-acceptor, and electrostatic interactions were all crucial contributors. The energy distribution of sites analysis revealed that the CMC composite possessed more adsorption active sites than MXene and CNTs. As a result, CMC composites possessed better adsorption capabilities for the anionic dyes that were being studied. It is possible that the changes in adsorption patterns for the anionic dyes can be attributed to the availability of the adsorption sites as well as the changes in the heterogeneity of the site on the CMC surface. Based on the findings of the approximate site energy distribution analysis, it was

discovered that the CMC composite has a greater number of adsorption active sites composite of R023363 to MXene and CNTs. This indicates that the CMC composite possesses a superior adsorption capability for specific anionic dyes. The amount of accessible adsorption sites and the change in site heterogeneity on the CMC surface are two factors that can be attributed to the differences in adsorption behaviors that occur between three anionic dyes from different compounds (Figure 15b-f).¹⁵⁴

Wang and co-authors developed an innovative approach to modify the electrospun polylactic acid fibrous membrane, improving its adaptability for treatment of wastewater under challenging conditions. PLA fibre membranes were initially coated with PDA, followed by deposition of MXene facilitated by PDA (**Figure 15g**). The as-prepared SC-PLA/PDA/MXene membranes proved effective for MB adsorption, with maximum capacity to adsorb 434.8 mg/g. The MB adsorption by the membrane aligns closely with the pseudo-second-order kinetic and Langmuir models. The above circumstance also indicates that chemisorption predominates in the MB elimination process. In essence, the membrane and MB molecules exhibit π - π interaction and electrostatic attraction. ¹⁵⁵

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Xing *et al.* fabricated TiVCT_X MXene/graphene nanosheet-based aerogels, named TiVCT_X/GAs, *via a* simple self-assembly hydrothermal process, followed by freeze-drying **(Figure 15h)**. Combining bimetallic MXene and graphene aerogel in TiVCT_X/GAs provides a remarkable broad-spectrum dye removal capability from wastewater. TiVCT_X/GAs effectively adsorb different dyes, with capacities of 319.67, 229.97, 303.45, and 217.87 mg/g for MB, CR, RB, and MO, respectively. Kinetics and thermodynamics studies confirm that the process of adsorption follows pseudo-second-order kinetic and Langmuir adsorption models.¹⁵⁶

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

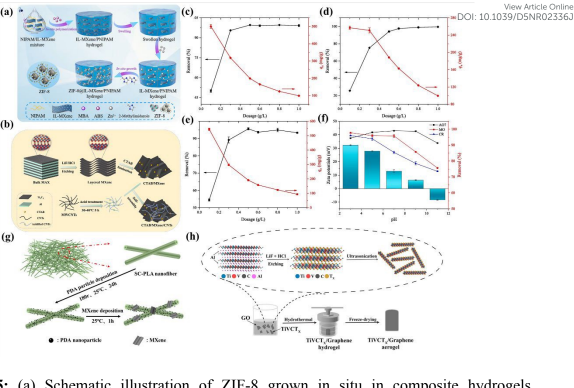


Figure 15: (a) Schematic illustration of ZIF-8 grown in situ in composite hydrogels. Reproduced with permission from Ref.¹⁵³ Copyright 2024, Elsevier. (b) Scheme of the preparation process of the CMC composite. (c), (d) & (e) Effect of adsorbent dosage on the removal efficiency and adsorption capacity (q_e) of AO7, MO, and CR. (f) effect of pH on the adsorption and zeta potentials of the CMC composite at different pH values. Reproduced with permission from Ref.¹⁵⁴ Copyright 2024, Elsevier. (g) Fabrication procedures of composite fibers. Reproduced with permission from Ref.¹⁵⁵ Copyright 2024, Elsevier. (h) Preparation Process of TiVCT_X/Gas. Reproduced with permission from Ref.¹⁵⁶ Copyright 2024, Elsevier.

MXene/carbon foam (MCF) hybrid aerogel was synthesized by Li and co-authors using Ti₃C₂T_X-MXene and MF. Both cationic and anionic dyes were used to investigate the removal properties of the MCF. Several different methods of characterization were utilized in order to examine the physicochemical properties of MCF. Additionally, statistical physics models were utilized in order to gain a deeper comprehension of dye adsorption mechanism that it possesses. MCF was shown to have outstanding adsorptive efficiency against the harmful dyes CR and

MB, with adsorption capacities of 647.75 mg/g and 356.97 mg/g respectively. According MBCO23361 the findings, these capacities were better fitted for Langmuir and pseudo-second-order kinetic models. In addition, the saturated multilayer model was utilized in order to explore the molecular process that underlies the adsorption systems of MCF-MB and MCF-CR components. Through a process that involved multi-anchoring (303 and 318 K), single connection (333 K), and mixed adsorption orientation, the findings of the numerical simulation demonstrate that MCF adsorbed the MB molecules in the form of dimers. In contrast, CR molecules were adsorbed as monomers through a process that included mixed adsorption orientation (333 K), multi-anchored and pure non-parallel adsorption directions (303 K and 318 K), and adsorption directions that were not parallel to one another. 157

Electrostatic self-assembly of 2D MXene nanosheets and biomass-activated carbon was used by Li *et al.* to create a heterostructure adsorbent that they referred to as CMAC composite. This was accomplished in the presence of a CTAB. This method prevented the re-stacking of MXene nanosheets, which resulted in a reduction in the multilayer plate structure of MXene and an increase in the layer spacing. This resulted in an increase in exposure of accessible active sites, which made adsorption performance even more significantly improved. As a consequence of the tests, it was shown that CMAC possessed an exceptional adsorption efficiency for CR, with adsorption capacities of 1264.032 mg/g. The Langmuir, intraparticle diffusion and pseudo-second-order kinetic models suit the dye adsorption data well. The adsorption mechanism is attributed to the combination of hydrogen bonding, physical adsorption, and electrostatic interactions.¹⁵⁸

Li *et al.* developed a straightforward and eco-friendly foaming technique in order to produce a macroporous cellulose nanocrystals (CNC)/MXene/polyvinyl alcohol (PVA) (C–CMP) foam that possessed exceptional adsorption capabilities. Glutaraldehyde was utilized as a crosslinker to enhance the adsorption effectiveness of MB dye. The adsorption experiment revealed that

MB has a maximum adsorption capacity of 239.92 mg/g. The adsorption performance control of the c

Page 50 of 119

Table 1: Comparison of different MXene-based materials for adsorption of various dyes.

MXene	Dye	Adsorption	Experimental	Isotherms/	Regeneration	Toxicity Evaluation
		capacity/	Conditions	kinetics		
		% Removal				
		efficiency				
$Ti_3C_2T_x^{131}$	MB	39	$C_0 (MB) = 0.05$	Freundlich	• Ti ₃ C ₂ T _x gradually oxidises to	No direct cytotoxicity or
			mg/ml, 25 °C, 20		TiO ₂ and Ti(OH) ₄ in water,	ecotoxicity tests were
			h		limiting reusability.	reported.
					• Structural changes (layer	• Possible formation of
					expansion and disorder) occur	TiO ₂ NPs and fluoride
					after 20 h in dye solution.	leaching may raise
					• Material is not stable for long-	concerns.
					term aqueous applications or	• Further studies needed
					storage.	to assess environmental
						and biological safety.
NaOH-Ti ₃ C ₂ T _x ¹³²	MB	189	25 °C, pH = 6–	Langmuir/PSO	No regeneration or reuse tests	• No cytotoxicity,
			6.5		were conducted.	ecotoxicity, or leaching
LiOH-Ti ₃ C ₂ T _x ¹³²	MB	121	25 °C, pH = 6–	Langmuir/PSO	Material performance	tests were included.
			6.5		stability over multiple cycles	Alkaline treatment may
KOH-Ti ₃ C ₂ T _x ¹³²	MB	77	25 °C, pH = 6–	Langmuir/PSO	not studied.	reduce fluoride (-F)
			6.5			groups, potentially

					Suggests need for future work	improving
					on recyclability.	biocompatibility.
						• Safety and
						environmental impact
						not evaluated.
<i>h</i> -Ti ₃ C ₂ ¹³³	MB	24	2 h	-	• No regeneration, desorption,	• No direct toxicity,
					or reuse cycles were tested or	cytocompatibility, or
					discussed.	environmental safety
					• Long-term stability of the	evaluations made.
					MXenes in dye removal	• Indirect benefit:
					applications not evaluated.	hydrothermal route
					• Future studies should assess	avoids toxic HF,
					recyclability to validate	reducing synthesis
					practical applicability.	hazards.
						• Leaching or stability of
						AlF ₃ by-products not
						assessed.
Nb ₂ CT _x ¹²⁷	MB	526.32	$C_0 = 100-500$	PSO	No regeneration/reusability	• No toxicity
			mg/L, 25 °C, pH		cycles tested.	(cytotoxicity or
			= 7, dose $= 1$		• Long-term performance	ecotoxicity) or leaching
			g/L, 700 rpm		stability and recyclability are	studies reported.
					not addressed.	• HF used for etching may
						pose safety concerns

Page 52 of 119

		1	T	T	ı	
					• Future studies should focus on	unless thoroughly
					desorption efficiency and	washed and neutralised.
					material durability.	• Environmental and
						biological safety remain
						unexplored.
PAA ₂ alk-MXene ¹²⁸	MB	193.92	$C_0 = 60-90$	Langmuir/ PSO	• The adsorbent AA2-alk-	No direct cytotoxicity or
			mg/L, 25–55 °C,		MXene maintained over 85%	ecotoxicity tests were
			pH = 2-10		dye removal efficiency after	reported in the study.
PAA ₂ alk-MXene ¹²⁸	CR	264.46	$C_0 = 60-90$	Langmuir/ PSO	five adsorption-desorption	• Use of mild reagents like
			mg/L, 25–55 °C,		cycles.	AA and NaOH suggests
			pH = 2-10		• Ethanol was used as the	potentially low toxicity,
					desorbing agent.	although this was not
					• Good reusability and	experimentally verified.
					structural stability.	Biocompatibility and
						environmental safety
						remain unexplored.
Ti ₃ C ₂ -SO ₃ H ¹²⁹	MB	111.11	T = 25 °C,	Langmuir/ PSO	• Regeneration or reuse	• Did not evaluate
			pH = 7,		experiments were not	toxicity, cytotoxicity, or
			dosage = 10 mg,		conducted.	leaching risks.
			$C_0 = 50 \text{ mg/L}$		• Long-term adsorbent stability	• Use of NaF and
					was not assessed.	arenediazonium salts
					• No adsorption—desorption	may pose safety
					cycles reported.	concerns.

						• Environmental impact
						and biocompatibility
						remain unexplored.
MXene-	MB	81.9672	$C_0 = 0-3 \text{ mg/L},$	Langmuir/ PSO	• The removal efficiency	• Environmental safety
COOH@(PEI/PAA) _n) ¹³⁷			25 °C, dose = 10		decreased from 85.6% to	and leaching of
			mg/50 mL		64.4% after 8 cycles.	components were not
					• The material showed good	assessed.
					stability and reusability.	• The synthesis avoids
						heavy metals, but
						toxicity remains
						unevaluated.
PHGC/MXene ¹³⁸	MB	555.56	$C_0 = 50-100$	Langmuir/ PSO	• PHGC/MXene hydrogel	• No in vivo or in vitro
			mg/L, 298-308		maintained over 90% removal	toxicity study was
			K, pH = 11		efficiency for AB93 and MB	reported.
					after 12 regeneration cycles.	• Environmental impact
					• NaHCO ₃ and HCl were	and cytocompatibility
					effective desorption agents for	were not evaluated.
					AB93 and MB, respectively.	• Safety validation is
					• The material showed excellent	lacking for biomedical
					reusability and economic	or large-scale
					regeneration.	applications.
cetyltrimethylammonium	МО	213	$C_0 = 10-500$	Langmuir/ PSO	• Adsorption capacity	• No direct toxicity
bromide-modified multi-			mg/L, 298 K,		decreased slightly from 25.52	assessment (e.g.,

Page 54 of 119

layered Ti ₃ C ₂ Tx MXene			pH= 3–12, dose		mg/g to 14.3 mg/g after 5	cytotoxicity,
(CMM) ¹³⁰			= 0-3.33 g/L		cycles.	ecotoxicity) was
					• Ethanol used for desorption	conducted in the study.
					demonstrated effective dye	• Implicit safety
					elution.	suggested via use of
					• CMM maintained	ethanol in desorption
					commendable reusability and	and absence of harmful
					structural integrity.	leachates.
ABC/MX composite ¹³⁹	CR	1103.7	$C_0 = 50-500$	Freundlich	• CR: High stability and	• No in vitro, in vivo, or
			mg/L, pH = 1-11,	model	reusability; >95% removal	ecotoxicity tests were
			300 K- 330 K,		retained after 5 cycles.	performed.
			dose = 0.4 g/L		• Cr(VI): Performance dropped	• However, authors
					significantly from 88.44% to	emphasised that:
					55.53% after 1 cycle.	Bacterial cellulose (BC)
						is a renewable,
						biodegradable, and non-
						toxic biopolymer.
						• Desorbed Cr(VI) is
						treated with Fe(II) to
						form precipitates,
						reducing secondary
						pollution risk.

						• Suggests low
						environmental burden
						under controlled usage
						but lacks direct toxicity
						proof
MXene-PEI ¹⁴⁰	MO	909.1	$C_0 = 120 - 300$	Langmuir/ PSO	• High stability and reusability	• No in vitro or in vivo
			mg/L, 328 K,		over 5 cycles:	toxicity tests.
			pH=3		MB: Removal decreased from	• Materials (MXene,
					$89.7\% \rightarrow 85.3\%$	chitosan, alginate) are
					Cr(VI): Removal decreased	described as
					from $93.4\% \rightarrow 87.2\%$	biocompatible and eco-
					• Structural Integrity:	friendly.
					Maintained porous	• No harmful by-products
					architecture and adsorption	or leaching reported.
					capacity.	• Indicates low
						environmental impact
						but lacks formal
						toxicological validation.
MX-PAN membrane ¹⁵⁰	MB	-	$C_0 = 10 \text{ mg/L}, 25$	-	• Reusability:	• No explicit cytotoxicity
			°C, pH= 7		After 5 adsorption–desorption	or environmental
					cycles:	toxicity tests conducted.
					Cr(VI): Removal decreased	• Components (MXene,
					from 91.7% to 86.3%	lignocellulose, CNF) are

Page 56 of 119

	1		1			
					MB: Removal declined from	natural or
					95.4% to 89.5%.	biocompatible.
					Retained adsorption capability	• Emphasis on green
					and structure, demonstrating	synthesis and
					excellent recyclability.	biodegradability.
						• Suggests low toxicity
						potential, but lacks
						empirical toxicology
						evidence.
SA/MXene nanofiber	MB	440	$C_0 = 50-220$	Langmuir/PSO	Performance Over Cycles:	No direct cytotoxicity or
membranes ¹⁵¹			mg/L, 98 K-323		• After 5 adsorption—desorption	environmental toxicity
			K, pH= 2-11,		cycles:	data.
			dose = 1-5 mg		Cr(VI): Decreased from 93.6%	• MXene and attapulgite
					→ 84.7%	considered relatively
					MB: Decreased from 91.2% →	biocompatible materials.
					85.2%	• Use of mild regenerants
					Observation: Slight decline	(ethanol, NaOH)
					but excellent reusability and	supports low
					structural robustness.	environmental burden.
						• Long-term stability of
						MXene against
						oxidation not discussed
						in toxicological terms.
	I	1	1	L	1	

Ti ₃ C ₂ T _X bound with	MB	209	pH=7, RT, 100	PSO	• Unique regeneration method:	• No cytotoxicity or
terephthalate (T-MX) ¹³⁴			mg/L		Spent T-MX (after MB	environmental safety
					adsorption) is reconverted into	data were presented.
					MAX phase by adding only Al	• Concerns noted:
					powder and applying	Potential contamination
					mechanochemical (MC) ball	from milling tools (Fe,
					milling.	alloying metals) under
					• The MAX phase is then re-	acidic HF etching.
					etched into MXene,	• These may bind to
					confirming the closed-loop	reactive Ti sites,
					recyclability of the material.	requiring further
					• Pollutants and terephthalate	toxicological
					pillars are carbonised and	investigation.
					reused as carbon source in the	• Materials used (MXene,
					MAX phase.	terephthalate) are
						otherwise not inherently
						toxic, and MC synthesis
						is solvent-free and
						greener.
TMAOH delaminated	MB	1026	$C_0 = 20-80$	Freundlich /	• Reusability was tested over 5	No direct cytotoxicity or
Ti3C2Tx MXene			mg/L, 318 K,	PSO	cycles using 1 M NaOH for	ecotoxicity tests were
nanosheets ¹³⁵			pH=6, dose =		desorption.	reported.
			0.01-0.06 g/L			

Page 58 of 119

alk-	MB	537.63	$C_0 = 17-50$	Langmuir/ PSO	The adsorbent demonstrated	• No toxicity assessment
MXene/CoFe ₂ O ₄ /CS ¹⁴¹			mg/L, 22–47 °C,		excellent recyclability with	or environmental
			pH = 2-12, dose		>80% removal efficiency	impact evaluation was
			= 10 mg/25 mL		maintained for RhB and MG	conducted in this study.
alk-	RhB	1333.86	$C_0 = 5-10 \text{ mg/L},$	Langmuir/ PSO	dyes after 5 cycles, while CR	• The paper focused
MXene/CoFe ₂ O ₄ /CS ¹⁴¹			22–47 °C, pH =		retained ~66% efficiency.	solely on adsorption
			2-12, dose = 10		Magnetic separation using	performance without
			mg/25 mL		NdFeB magnet enabled easy	investigating potential
alk-	CR	2095.9	$C_0 = 5-10 \text{ mg/L},$	Langmuir/ PSO	recovery from aqueous	environmental or health
MXene/CoFe ₂ O ₄ /CS ¹⁴¹			22–47 °C, pH =		solutions, with ethanol used	impacts of the
			2-12, dose = 10		as desorption agent for	synthesized materials.
			mg/25 mL		regeneration cycles.	
MXene (Ti ₃ C ₂)/Fe ₃ O ₄ ¹⁴²	MB	9.85	$C_0 = 1-40 \text{ mg/L},$	Freundlich	• No regeneration or	No toxicity assessment
			25–55 °C, dose =	isotherm	recyclability studies were	or environmental
			1 g/L		conducted in this research.	impact evaluation was
					• The study focused solely on	performed in this study.
					adsorption performance at	• The research
					different temperatures	concentrated on
					without investigating the	adsorption mechanisms
					reusability of the 2D-	and temperature effects
					MX@Fe ₃ O ₄ adsorbent.	without addressing
						potential health or
						environmental risks.

Ti ₃ C ₂ T _X /NiFeMn-	CR	588.24	$C_0 = 50-300$	Freundlich /	• The composite demonstrated	• A comprehensive
LDH@Gel ¹⁴³			mg/L, 298-328	PSO	excellent recyclability with	leaching test was
			K, pH = 3-9		74.94% and 71.72% removal	conducted using ICP-
					efficiency maintained for	MS technique to detect
					Cr(VI) and CR respectively	potential metal
					after 5 cycles.	leaching (Ni, Fe, Mn)
					• Regeneration was achieved	across pH range 3-9.
					using 1 M NaCl/methanol	Results showed almost
					solution with continuous	constant metal content
					stirring for 2 hours, followed	after adsorption
					by drying at 50 °C for 8 hours	processes, confirming
					before reuse.	the stability and safety
						of the composite for
						practical applications
						without significant
						metal leaching.
AMXGO ¹³⁶	MG	1111.6	$C_0 = 100-400$	Freundlich /	• The composite maintained	Zeta potential analysis
			mg/L, 298 K-328	PSO	>70% removal efficiency for	showed stable surface
			K, pH = 2-12		both MG and CR after 5	charge (-20.54 to -
					cycles.	52.00 mV, pH 2-12)
					• Regeneration used 5 mL	without harmful ion
					detergent + 100 mL DI water,	leaching.
	1	1	I	l	I	

					stirred for 3 hours, washed to	• XRD and FTIR
					neutral pH, then freeze-dried.	confirmed no structural
						degradation after
						multiple cycles,
						ensuring material
						safety.
MXene/COF ¹⁴⁴	MB	96.4%	effective		• The membrane maintained	• XRD analysis
			membrane area		>96% rejection and >243	confirmed no structural
			$= ~ 8.0 \text{ cm}^2$		L/m ² ·h·bar water permeance	degradation with d-
MXene/COF ¹⁴⁴	CR	98.2%	effective		after 45 hours of continuous	spacing remaining
			membrane area		operation.	stable at ~1.46 nm after
			$= \sim 8.0 \text{ cm}^2$		• No specific regeneration	extended operation.
MXene/COF ¹⁴⁴	MO	97.2%	effective		process described -	• Membranes showed
			membrane area		membrane showed excellent	excellent anti-swelling
			$= ~ 8.0 \text{ cm}^2$		stability without requiring	properties in pH 3-10
					regeneration protocols.	conditions, maintaining
						structural integrity
						without toxic leaching.
alk-MXene/ZIF	CR	539.7	298-318 K, pH=	Elovich/PSO	• The composite maintained	• XRD analysis
composites ¹⁴⁵			8		>85% removal efficiency for	confirmed stable
alk-MXene/ZIF	MG	7111.3	298-318 K, pH=	Elovich/PSO	all three pollutants (CR, TC,	crystalline structure
composites ¹⁴⁵			8		MG) after 5 cycles.	with no significant peak

Page 62 of 119

					• Regeneration achieved using anhydrous ethanol stirring for 2 hours, followed by deionized water washing and	shifts after air exposure for 45 days. • Membrane filtration tests showed >90%
					freeze-drying.	removal maintained after 5 cycles, demonstrating material safety without toxic leaching.
DSP-M ¹⁴⁶	RhB	678.19	$C_0 = 2-100$ mg/L, 30 °C-60 °C, pH= 7	Langmuir/ PSO	• The composite maintained stable performance with no specific regeneration protocol	• XPS analysis confirmed no harmful metal leaching or toxic
DSP-M ¹⁴⁶	CR	754.41	5–200 mg/L, 30 °C-60 °C, pH= 7	Langmuir/ PSO	described, showing structural integrity after multiple adsorption-desorption cycles. • Material stability confirmed through XRD and FTIR analysis showing no structural degradation after repeated use	byproduct formation during adsorption processes. • Statistical physics modeling confirmed physical adsorption processes (E < 8 kJ/mol) involving safe hydrogen bonding, electrostatic forces, and van der Waals interactions.

:475:(2):97:48:(3):97:48:(ı	Nanoscale		
Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 124/6:0 e of on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 124/6:0 e of on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 124/6:0 e of on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 124/6:0 e of on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 124/6:0 e of on 04 August 2025.							
ersity	ZnS/CuFe ₂ O ₄ /MXene	DBM	377	0.02–4 g/L, 12 h,	Langmuir/PSO	• Good reusability with no	• Successfully applied to
n Univ	(ZSCFOM) ¹⁴⁷			75 rpm, 20 °C		significant reduction in DBM	real environmental
/unna	ZnS/CuFe ₂ O ₄ /MXene	DBRN	390	0.02–4 g/L, 12 h,	Langmuir/PSO	removal rate after three	waters (tap, pond, river)
d by Y	(ZSCFOM) ¹⁴⁷			75 rpm, 20 °C		cycles.	with 100% removal of
ıloade						• Excellent thermal stability	0.05 g/L dyes.
Dowr						with minimal weight loss	Photocatalytic process
2025.						(0.23-0.38%) at 25-600 °C.	generates
ugust						• Magnetic separation	environmentally benign
04 A						capability for easy recovery	products through
hed or						using CuFe ₂ O ₄ component.	complete
Publis							mineralization.
							Magnetic properties
							enable easy separation
							and recovery,
							preventing secondary
							contamination.
	PHGC/MXene ¹³⁸	AB93	207.47	$C_0 = 50-100$	Langmuir/ PSO	• Excellent reusability with	• No comprehensive
				mg/L, 298-308		removal rates remaining	toxicity assessment of
				K, pH = 2		above 90% for AB93 and	the PHGC/MXene
						above 97% for MB after 12	composite material
						cycles.	itself was conducted.
						• Regeneration achieved using	• Effective removal of
						0.1 mol/L NaHCO ₃ solution	toxic anionic (AB93)

<u> </u>				I a 1704 101 17	
				for AB93 and 0.1 mol/L HCl	and cationic (MB) dyes
				solution for MB.	from contaminated
				• Minimal decrease in	water.
				performance (less than 3%	• Successfully tested
				compared to highest removal	across different pH
				rate) demonstrating excellent	ranges (2-12) without
				stability.	significant material
				Both acidic and alkaline	degradation.
				regeneration solutions proved	Non-toxic regeneration
				effective for respective dye	process using
				systems.	environmentally
					acceptable desorption
					agents (NaHCO ₃ and
					HCl).
					• No adverse effects on
					hydrogel structure
					during multiple
					regeneration cycles.
					Material demonstrated
					biocompatibility based
					on constituent
					components.
	<u> </u>	1	l	I	<u> </u>

ZIF-8@IL-	CV	325.03	$C_0 = 0.4-2 \text{ mg/L},$	Freundlich /	• Good reusability with	• Environmentally
MXene/PNIPAM ¹⁵³			25-45 °C, pH = 7	PSO	adsorption capacities	friendly synthesis and
					decreasing to 79%, 91%, and	desorption processes
					29% for 4-NP, CV, and Cu ²⁺	without organic
					respectively after five cycles.	solvents.
					• Desorption achieved via	• Successfully applied to
					volume phase transition in	multifunctional removal
					deionized water at 35-40 °C	of phenols, dyes, and
					(environmentally friendly	metal ions from
					process).	industrial wastewater.
					• Decline in Cu ²⁺ adsorption	• VPTT range 33-35 °C
					due to strong chelation bonds	enables safe
					difficult to break through	temperature-responsive
					phase transition.	desorption process.
						• No toxic byproducts
						reported during
						adsorption/desorption
						cycles.
AMXGO ¹⁵⁴	CR	1133.7	$C_0 = 100-400$	Freundlich/	• Good reusability with	• No significant effects
			mg/L, 298 K-328	PSO	removal efficiencies	from common cations
			K, pH = 2-12			

Nanoscale

Page 66 of 119

CMC composite ¹⁵⁴	AO7	367.9	$C_0 = 50-500$	Sips/PSO	gradually decreasing over five	(Na ⁺ , K ⁺ , Ca ²⁺ , Mg ²⁺)
			mg/L, 25-45 °C,		cycles.	and anions (Cl ⁻ , NO ₃ ⁻ ,
			pH = 3-11		• AO7 showed least decrease in	CO_3^{2-} , SO_4^{2-}) on dye
CMC composite ¹⁵⁴	МО	294.2	$C_0 = 50-500$	Sips/PSO	efficiency, followed by MO	removal.
			mg/L, 25-45 °C,		and CR after five cycles.	• Successfully tested in
			pH = 3-11		Gradual decrease attributed to	different water matrices
CMC composite ¹⁵⁴	CR	628.5	$C_0 = 50-500$	Sips/PSO	incomplete desorption and	(deionized, tap, and lake
			mg/L, 25-45 °C,		loss of surface functional	water).
			pH = 3-11		groups during regeneration	• Slightly reduced
					process.	performance in tap and
						lake water due to
						competing ions and
						organic matter.
						• Effective removal of
						carcinogenic,
						mutagenic, and toxic
						anionic azo dyes from
						contaminated water.
						• No specific toxicity
						assessment of the CMC
						composite material
						itself was conducted.

SC-PLA/PDA/MXene	MB	434.8	$C_0 = 0.5-2 \text{ g/L}$	Langmuir/ PSO	• Good cycling stability	• Biodegradable PLA-
membrane ¹⁵⁵					demonstrated over 10	based composite
					consecutive separation cycles	material providing eco-
					for oil/water separation	friendly alternative to
					• Water flux remained stable at	conventional polymer
					1429.0 L/(m ² ·h) even after 10	membranes
					cycles for E-N/W separation	• Eliminates potential
					• Oil flux maintained at	secondary pollution
					\sim 1862.0 L/(m ² ·h) for the first	from plastic debris
					4 cycles in E-W/N separation,	entering water systems
					then declined to 1563.3	• No comprehensive
					L/(m ² ·h) (84.6% retention)	toxicity assessment of
					after 10 cycles	the composite material
					• Alcohol regeneration	conducted
					treatment used between cycles	• Effective removal of
					with good recovery	toxic organic pollutants
					performance	(MB) from
					• Excellent reusability for	contaminated water
					simultaneous oil and MB	Biocompatible
					removal with water flux	polylactide substrate
					maintaining high levels after 6	reduces environmental
					reuse cycles	impact compared to

					Minor performance decline attributed to porosity blocking by residual oil during separation process	non-degradable polymer membranes • Polydopamine and MXene components generally considered biocompatible based on literature
Fe ₂ O ₃ /BC/MXene ¹⁴⁸	MB	899.03	C ₀ = 15-45 mg/L, 293-313 K, pH= 1-11	Langmuir/ PSO	 • Pb²+ sorption capacity: 99.91% → 90.99% after 5 cycles; MB: 98.63% → 89.47%. • Desorption efficiency: Pb²+: 96.8% → 78.32%; MB: 95.08% → 77.40% after 5 cycles. • Easy separation using external magnetic field. 	sustainable wastewater treatment. • Effectively removes
TiV _C T _X /GAs ¹⁵⁶	MB	319.67	298 - 338 K, pH= 1-13	Langmuir/ PSO		

TiV _C T _X /GAs ¹⁵⁶	RhB	303.45	298 K- 338 K,	Langmuir/ PSO	• Excellent cycling stability	• Leaching experiments
			pH= 1-13		demonstrated over 5	confirmed that
TiV _C T _X /GAs ¹⁵⁶	CR	229.97	298 K- 338 K,	Langmuir/ PSO	successive adsorption-	TiV _C T _x /GAs adsorbent
			pH= 1-13		desorption cycles.	does not produce
TiV _C T _X /GAs ¹⁵⁶	MO	217.87	298 K- 338 K,	Langmuir/ PSO	MB removal rate remained at	secondary pollution
			pH= 1-13		around 95.8% after five	during adsorption
					cycles.	process.
					• High absorption capacity	• Effectively removes
					maintained even after five	highly toxic organic
					cycles of absorption-	contaminants including
					squeezing for oil/solvent	dyes and antibiotic
					removal.	drugs from wastewater.
					• Easy regeneration through	• No comprehensive
					simple mechanical squeezing	toxicity assessment of
					due to excellent elasticity of	the composite material
					TiVCTx/GAs.	itself conducted.
						Biocompatible
						components generally
						considered safe for
						environmental
						applications.

Page 70 of 119 Nanoscale

MCF hybrid aerogel ¹⁵⁷	MB	356.97	$C_0 = 3-100$	Langmuir/ PSO	• Limited regeneration	• No comprehensive
			mg/L, 303 K-		assessment mentioned in the	toxicity assessment of
			333 K, pH = 11		study.	the MCF composite
MCF hybrid aerogel ¹⁵⁷	CR	647.75	$C_0 = 3-200$	Langmuir/ PSO	• No comprehensive cycling	material conducted.
			mg/L, 303- 333		stability evaluation provided.	• Focuses on removing
			K, pH = 6		• Brief mention that physical	highly toxic
					adsorption is beneficial for	contaminants: MB
					reversible processes and	causes DNA structure
					adsorbent regeneration after	damage, cancer risk,
					desorption.	heart rate acceleration,
					• Statistical physics modeling	shock, and tissue
					suggests physical interaction	necrosis.
					enables regeneration	• CR degradation
					potential.	produces benzidine, a
						recognized human
						carcinogen.
						• MCF designed as
						environmentally-
						friendly technology for
						hazardous dye
						elimination.
						• No secondary pollution
						production during

						adsorption process
						confirmed through
						leaching experiments.
Ni ₃ (HITP) ₂ /MXene/CS ¹⁴⁹	MB	424.99	10–400 mg/L,	Langmuir/PSO	• Five consecutive adsorption-	• No comprehensive
			298 K, pH= 2–9		desorption cycles conducted	toxicity assessment of
					for reusability evaluation.	the composite material
					• Composite maintained	conducted.
					excellent adsorption rate after	• Composite uses
					five cycles for both Pb(II) and	biocompatible chitosan
					MB.	biopolymer as base
					• Chemical adsorption nature	material.
					may pose challenges for	• No secondary pollution
					complete regeneration.	or leaching studies
					• Detailed regeneration	reported.
					protocols not clearly	
					presented in main text.	
MXene with biomass	CR	1264.032	$C_0 = 0-400$	Langmuir/PSO	Desorption and recyclability	• No comprehensive
activated carbon			mg/L, 25–75 °C,		studies conducted using	toxicity assessment of
(CMAC) ¹⁵⁸			pH = 6, dose =		ethanol as desorbing agent.	the CMAC composite
			10 mg/100 mL		• CMAC mixed with 100 mL	material conducted.
					ethanol and stirred at 25 °C	• Focus on removing
					for 24 hours.	highly toxic anionic azo
						dyes that are

Page 72 of 119

Nanoscale

					• Good adsorption effect	carcinogenic and
					maintained for all three dyes	mutagenic.
					after multiple desorption	• CMAC composite uses
					cycles.	biomass activated
					• No significant loss of	carbon as
					performance after several	environmentally
					cycles.	friendly base material.
					• Economic benefit and high	• No secondary pollution
					stability demonstrated for	or environmental impact
					wastewater treatment	studies reported.
					applications.	
macroporous cellulose	MB	239.92	pH=6, 338 K,	Langmuir/PSO	Adsorption-desorption	• No comprehensive
nanocrystals			250 mg/L		experiment conducted for	toxicity assessment of
(CNC)/MXene/PVA (C-					three cycles using 0.1 mol/L	the C-CMP composite
CMP) ¹⁵⁹					HCl for desorption.	material conducted.
					• MB removal efficiency	• Focus on removing
					maintained above 80% after	highly toxic MB which
					three cycles.	severely affects water
					• Good recyclability of C-CMP	transparency and
					composite foam confirmed.	dissolved oxygen.
					• Slightly decreased efficiency	• MB is difficult to
					attributed to strong interaction	degrade due to benzene
					between MB and C-CMP.	ring structure and

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 1246:0 e	Nanoscale		
3/2025			
on 8/23			
ersity	• Residual dye from previous	accumulates in	drainage
n Univ	cycles may affect subsequent	systems.	
r'unna	performance.	• Even 1	mg/L
d by y		concentration	severely
nloade		impacts water	quality.
Dowi		• Composite	uses
2025.		biocompatible	
vugust		materials:	cellulose
n 04 A		nanocrystals	and
o paq		polyvinyl alcol	nol.
Publis		• No envir	ronmental
		impact or s	secondary
		pollution	studies
		reported.	
	Co: concentration of dye		

5. Mechanism of adsorption fundamentals of MXenes-based nano-adsorbents for dyes

When it comes to water filtration, adsorption is among the most effective and economical technologies available. On the basis of variations in the adsorption forces, adsorption is typically divided into two categories: chemical adsorption and physical adsorption. Physical adsorption primarily occurs due to van der Waals forces, which are frequently observed in MXene due to material's large SSA. A number of different processes, including electrostatic interactions, coordination, ion exchange, and hydrogen bonding, are included in chemical adsorption. Due to the fact that Ti-C-O- or Ti-C-OH- groups on MXene have the ability to interact with a wide variety of cations, electrostatic contact is a frequent mechanism in the process of adsorption of MXene. Ion exchange or development of inner-sphere complexes is encouraged by the presence of a considerable number of functional groups and the stratified structure. There are a variety of factors that influence the adsorption mechanisms of MXenes, including the exact kind of MXenes, the properties of pollutants, and the natural environment of the aquatic environment. The adsorption mechanism for TiVCT_x/GAs, as proposed by Xing et al., is depicted in Figure 16a. Integrating H-bonding, π - π interactions, and electrostatic attraction significantly influences adsorption. The structural features of TiVCT_x/GAs, such as their porous structure with a large SSA and the abundance of adsorption active sites given by bimetallic layers of TiVCT_X and graphene sheets, also help to enhance the ability of TiVCT_X/GAs to adsorb MB dye. 156

Furthermore, Purkayastha and co-authors produced MX-PAN membranes that may bind MB molecules to both F and OH/=O terminating groups of MXene. The molecular structure of MB dye is shown in **Figure 16b**. The sulfur (S) or nitrogen (N) atoms in MB, carrying positive charge, classify it as cationic dye. After the aluminum atoms are removed *via* etching from Ti₃AlC₂ (parent powder), various functional groups (-F and -OH/=O) attach simultaneously on

surface of $Ti_3C_2T_X$, resulting in negative charge. These negative charges $[Ti-F]^{-1}$ and $[Ti^{V}_3C^{Article Online}_{DNR023363}]$ can be cationic species sensors. As shown in **Figure 16c**, MB molecules can interact with both F and OH/=O terminal groups. Therefore, a key mechanism for adsorption of MB dye is electrostatic interaction between positively charged MB molecules and negatively charged $Ti_3C_2T_X$ MXene. 150

Wang *et al.* explained that the adsorption of MG and CR on alk-MXene/ZIF composite could be ascribed to the hydrogen bonding, π - π , and electrostatic interaction. Additionally, the proposed adsorption mechanism is schematically illustrated in **Figure 16d.** ¹⁴⁵ To further understand the phenomenon of adsorption, Li *et al.* explained the adsorption process of AMXGO for CR and MG, as shown in **Figure 16e**. AMXGO dye adsorption involves various phenomena, including π - π conjugates, electrostatic force, intercalation adsorption, as well as pore adsorption. Among the most significant impacts is the electrostatic force, which is generated by the negative hydroxyl and carboxyl groups present on GO lamellae and Alk-MXene surfaces, which mix with cationic molecules. π - π conjugates have a considerable effect on CR adsorption because their structure contains six benzene rings, which is more than three in the structure of MG molecule. This also describes why AMXGO has superior adsorption efficacy toward anionic CR. Furthermore, the layer-by-layer structure of Alk-MXene and GO flakes in AMXGO favors intercalation adsorption of either CR or MG molecules. Furthermore, the AMXGO samples include numerous mesoporous pores, that ultimately promote pores adsorption of CR and MG. ¹³⁶

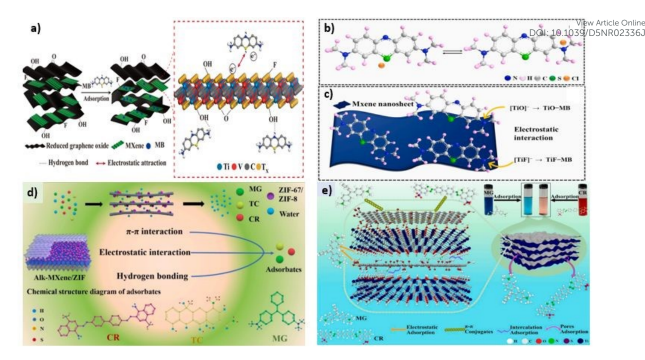


Figure 16: (a) Schematic Illustration of Adsorption Mechanism of TiVCT_X/Gas. Reproduced with permission from Ref. 156 Copyright 2024, American Chemical Society. (b) Molecular structure of MB. Reproduced with permission from Ref. 150 Copyright 2023, Elsevier. (c) Schematic illustration of electrostatic interaction between Ti₃C₂T_X MXene nanosheet and MB dye molecules. 150 Reproduced with permission from Ref. 150 Copyright 2023, Elsevier. (d) Schematic diagram on the adsorption mechanism of the alk-MXene/ZIF adsorbent. Reproduced with permission from Ref. 145 Copyright 2024, Elsevier. (e) Adsorption mechanism of AMXGO adsorbents for MG and CR. 136 Reproduced with permission from Ref. 136 Copyright 2024, Elsevier.

While the fundamental adsorption mechanisms have been identified, a comprehensive mechanistic framework reveals the relative contributions and synergistic interactions among different adsorption pathways in MXene-based systems.⁶⁰ Electrostatic interactions represent the primary driving force for cationic dye adsorption, with mechanism efficiency directly correlated to surface charge density and zeta potential values. Ti₃C₂T_x MXenes exhibit highly negative surface charges due to abundant -OH, -O, and -F terminations, creating strong

electrostatic fields capable of attracting cationic species over significant distance species over species ove Quantitative analysis reveals that electrostatic contribution accounts for the majority of total adsorption energy for MB and rhodamine B, while representing a smaller fraction for anionic dves like CR. pH-dependent studies demonstrate optimal electrostatic interactions occur at near-neutral pH, where MXene surface deprotonation maximizes negative charge density while maintaining structural stability. π - π stacking mechanisms show high selectivity toward aromatic dye molecules, with interaction strength proportional to aromatic ring number and electron density distribution. Computational studies using Density Functional Theory (DFT) calculations reveal significant binding energies for benzene ring interactions with Ti₃C₂T_x surfaces, with substantially higher energies for multi-ring systems like CR. Distance-dependent analysis shows optimal π - π interactions occur at specific separations, with parallel orientation providing maximum orbital overlap. MXene surface hybridization enhances π -electron delocalization, creating favorable interaction sites for aromatic dye molecules. 160 Hydrogen bonding exhibits remarkable complexity in MXene-dye systems, involving multiple donor-acceptor combinations and cooperative strengthening effects. Surface -OH and -O groups serve as hydrogen bond acceptors, while dye amino groups (-NH₂) and hydroxyl functionalities act as donors. 161 Spectroscopic analysis (FTIR and NMR) reveals moderate hydrogen bond strengths for individual interactions, with significantly enhanced cumulative effects for dyes with multiple bonding sites. Temperature-dependent studies show hydrogen bonding contributions decrease substantially at elevated temperatures, indicating entropydriven weakening of directional interactions. Ion exchange processes demonstrate high specificity toward charge-to-size ratios and hydration energies of exchangeable species. 162 Ti₃C₂T_x interlayer cations exhibit different exchange affinities for organic cations, with selectivity series: quaternary ammonium > primary ammonium > metal cations. Kinetic analysis reveals two-stage exchange processes: rapid surface exchange followed by slower

interlayer diffusion. Stoichiometric studies confirm equivalent exchange ratios for monovaled le confirme quivalent exchange ratios for monovaled le confirme quivalent exchange behaviors involving charge compensation mechanisms.

Physical adsorption through pore filling becomes dominant for microporous MXene derivatives and high molecular weight dyes. Pore size distribution analysis reveals optimal dye accommodation in mesopores, where molecular diffusion remains unrestricted while surface interactions are maximized. Molecular dynamics simulations demonstrate size-selective adsorption, with larger dye molecules showing reduced diffusion rates and surface-limited adsorption. Multiple mechanisms operate simultaneously in real adsorption systems, creating cooperative enhancement effects that exceed individual contributions. Electrostatic preconcentration brings dye molecules into proximity with MXene surfaces, facilitating secondary interactions including π - π stacking and hydrogen bonding. Computational analysis reveals cooperative binding energies significantly higher than additive individual contributions, indicating synergistic stabilization. Sequential mechanism analysis shows electrostatic interactions dominate initial adsorption, followed by π - π stacking at intermediate saturation and pore filling at high saturation. Solution pH dramatically influences mechanism prevalence and adsorption efficiency through surface chemistry modifications and dye speciation changes. 163

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Acidic conditions enhance protonation of MXene surface groups, reducing electrostatic attractions but increasing hydrogen bonding opportunities. Alkaline conditions maximize electrostatic interactions but may destabilize MXene structures through excessive deprotonation. Thermal analysis reveals differential temperature dependencies among adsorption mechanisms, with electrostatic interactions showing minimal temperature dependence, while hydrogen bonding decreases significantly at elevated temperatures. Multicomponent systems reveal mechanism-dependent selectivity patterns, with different dyes

competing for specific interaction sites, indicating moderate to intense competition depending rolling rolling moderate to intense competition depending rolling rol

6. Regeneration and recyclability of adsorbent

Adsorbents, such as MXene, used for dye adsorption should be recyclable and regenerative due to their practical and sustainable environmental applications. Several approaches have been established to recover the adsorption ability of dye-loaded MXene adsorbents for reuse. 134 Thermal regeneration comprises heating the adsorbents to encourage the desorption of dyes that have been absorbed. Chemical regeneration uses regenerants like alkalis, acids, organic solvents, or complexing agents to desorb the dyes from the adsorbent surface specifically. 164,165 Another approach is electrochemical regeneration, which uses an electrical potential to accelerate desorption with the advantage of selectivity and in situ regeneration. When the regenerant solution and dye-loaded adsorbents are mixed, and suitable desorption conditions are sustained, batch systems can perform these processes. The effectiveness of the regeneration is evaluated based on the amount of dye removed and the recovery of adsorption capacity. MXene adsorbents' long-term viability as cost-efficient and sustainable solutions for dye removal applications is partially due to their effective regeneration and recyclability. 83,166 For instance, the recyclability of Ti₃C₂T_X/NiFeMn-LDH@Gel was demonstrated after five times of recycling and it was observed the removal ratio of Ti₃C₂T_x/ NiFeMn-LDH@Gel for CR adsorption remained at 71.72%, displaying the effective regeneration property for Ti₃C₂T_X/NiFeMn-LDH@Gel as shown in Figure 17a. 143 Likewise, the cyclic stability of TiVCT_X/GAs, as illustrated in Figure 17b, highlights its remarkable performance in dye adsorption applications. Over five cycles of the absorption-squeezing process, the material

removing contaminants from wastewater. 156 The utilization of NaHCO3 solution as a desorption agent for PHGC/MXene/AB93 results in a negligible reduction in adsorption capacity of PHGC/MXene for AB93, even following 12 cycles of adsorption and desorption. The elimination rate is around 94% (Figure 17c), with variations not exceeding 1.5%. Upon utilizing hydrochloric acid solution as desorption agent for PHGC/MXene/MB, the PHGC/MXene's adsorption capacity for MB decreases by a minimal amount after 12 cycles of adsorption and desorption, while the efficiency of removal remains consistently higher than 97% (Figure 17d). The results suggest that the removal rate decreases by at least three percent throughout the course of the twelve cycles in comparison to its highest value. This suggests that PHGC/MXene hydrogel excels in both alkaline and acidic environments, demonstrating its versatility. In the case of PHGC/MXene/AB93 and PHGC/MXene/MB, respectively, solutions of sodium bicarbonate and hydrochloric acid demonstrate effective desorption characteristics. As a result, PHGC/MXene hydrogel is a stable, efficient, and renewable adsorbent that may be utilized for the treatment of wastewater that contains AB93 or MB. 138

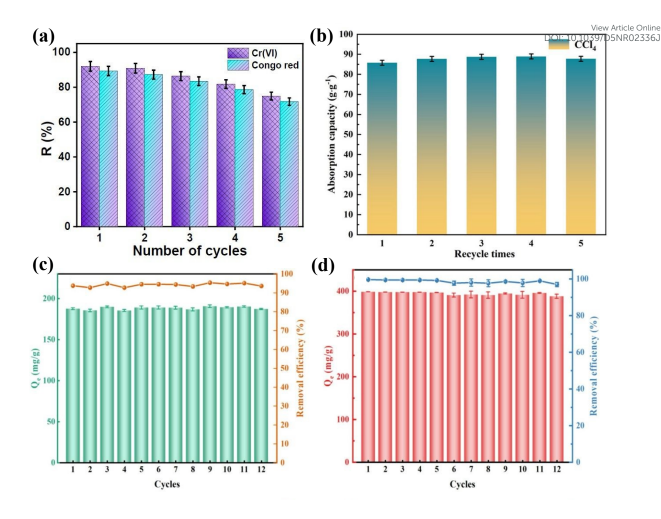


Figure 17: Reusability tests for (a) Ti₃C₂T_X/NiFeMn-LDH@Gel against Cr (VI) and CR. Reproduced with permission from Ref.¹⁴³ Copyright 2024, Elsevier. (b) TiVCT_X/Gas against CCl₄. Reproduced with permission from Ref.¹⁵⁶ Copyright 2024, American Chemical Society. Regeneration studies of (c) AB93 and (d) MB by PHGC/MXene. Reproduced with permission from Ref.¹³⁸ Copyright 2024, Elsevier.

Page 82 of 119

Table 2: Key strengths and limitations of Ti₃C₂T_x MXene for dye adsorption and photocatalysis

MXene	Positive Insights	Negative Insights
Ti ₃ C ₂ T _x ¹³¹	First report on MXene for dye removal & photocatalysis.	 Poor stability in water; oxidises to TiO₂.
	High MB adsorption via electrostatic interaction.	No toxicity or environmental safety data.
	Effective UV-driven dye degradation (MB & AB80).	 Moderate adsorption capacity (~39 mg/g).
	Strong binding (irreversible adsorption).	Complex mechanisms; needs further clarification.
	Good fit with Freundlich model (heterogeneous sites).	No real wastewater or reuse tests.
NaOH/LiOH/KOH-	Alkali treatment (NaOH, LiOH) increased interlayer spacing	No data on reusability or long-term performance.
$Ti_3C_2T_x^{132}$	(up to 29%).	Lack of toxicity and safety assessment.
	ullet Surface functionalisation (-F $ ightarrow$ -OH) improved adsorption	• Some variants (e.g. KOH–Ti ₃ C ₂ T _x) showed lower
	capacity.	capacity due to agglomeration.
	• NaOH-Ti ₃ C ₂ T _x showed highest MB adsorption capacity (189	Intercalation effects not uniform; morphology-
	mg/g).	dependent performance.
	Follows Langmuir model, indicating monolayer adsorption.	
	Faster dye removal rates compared to untreated MXene.	
h-Ti ₃ C ₂ ¹³³	Developed a fluoride-free, HF-free hydrothermal synthesis	No reusability or recyclability tests conducted.
	route using NaBF ₄ /HCl.	No quantification of maximum adsorption capacity
	• Resulting MXenes (h-Ti ₃ C ₂ , h-Nb ₂ C) had higher surface area	(only removal % shown).
	and better exfoliation.	Performance tested only for MB; limited scope (e.g.
	• Improved adsorption performance for MB over HF-etched	no heavy metals, real water).
	MXenes.	No mechanistic insight into selectivity or adsorption
		modelling (e.g. isotherms).

Ti ₃ C ₂ –SO ₃ H ¹²⁹	• Functionalisation with sulfonic groups increased adsorption capacity from 21.10 mg/g to 111.11 mg/g.	No regeneration or recyclability testing was conducted.
	 The material retained its adsorption efficiency after multiple cycles. Surface functionalisation introduced beneficial –COOH and – OH groups and increased interlayer spacing. 	generality of the findings.
PAA ₂ alk-MXene ¹²⁸	 Adsorbent showed high adsorption capacities of 264.46 mg/g for CR and 193.92 mg/g for MB. Adsorption followed the Langmuir isotherm and pseudosecond-order kinetics. 	 The study did not evaluate performance in real wastewater or multi-component dye systems. Toxicity or leaching of components was not assessed. Only two dyes (MB and CR) were tested, limiting the
Nb ₂ CT _x ¹²⁷	 Broader applicability: method extended to Nb₂C MXene synthesis. High specific surface area (~44.69 m²/g) and layered morphology support efficient adsorption. Very high adsorption capacities: MB = 526 mg/g, MO = 500 mg/g. Follows pseudo-second-order kinetics, indicating chemisorption at active sites. Faster adsorption for MB (99% removal in 5–11 min). Excellent efficiency for both cationic and anionic dyes. 	 No data on regeneration, recyclability, or real wastewater testing. Adsorption isotherms (Langmuir, Freundlich) did not fit well; mechanism modelling limited. Slight surface oxidation observed; long-term MXene stability not assessed.
	Larger interlayer spacing and higher BET surface area (44.6 vs. 8.9 m²/g). Broader and inhibitor model of a strended to NIL C. MYang.	• Adsorption only moderate (~24 mg/g for MB) compared to other MXenes.

Nanoscale

	 Adsorption followed Langmuir isotherm and pseudo-second-order kinetics. Adsorption capacity reached 723.35 mg/g with increased MB concentration. Surface chemistry and interlayer spacing were enhanced through diazonium modification. 	 The study only tested MB; no multicomponent or real wastewater systems. Safety and environmental aspects were not addressed. Adsorbent stability and scalability were not discussed.
MXene- COOH@(PEI/PAA) _n) ¹³⁷	 Adsorption followed pseudo-second-order kinetics and Langmuir isotherm (monolayer adsorption). Good selectivity and performance across three dyes (MB, ST, NR). Composites are easily separable and structurally stable. 	 Adsorption tested only in single dye systems, not in complex wastewater. Decline in adsorption capacity upon recycling may limit long-term use. Synthesis and LbL process may be time-consuming at industrial scale.
PHGC/MXene ¹³⁸	 PHGC/MXene hydrogel exhibited pH-responsive selectivity, with adsorption capacities of 555.56 mg/g for MB and 207.47 mg/g for AB93. High reusability and performance across acidic and alkaline conditions. The system demonstrated rapid adsorption and strong ionic dye interaction. 	 No real wastewater or multi-dye solution testing was conducted. Potential biodegradability or leaching behaviour of MXene was not assessed. Toxicity and environmental compatibility remain unexplored.
cetyltrimethylammonium bromide-modified multi- layered Ti ₃ C ₂ Tx MXene (CMM) ¹³⁰	 High adsorption capacity: up to 213.00 mg/g. Effective in a broad pH range, especially at pH 3. Adsorption follows Langmuir isotherm and pseudo-second-order kinetics. 	 Surface area decreased after CTAB modification (from 5.02 to 2.06 m²/g). No biological toxicity or environmental risk data provided.

Page 84 of 119

	 Mechanisms include electrostatic attraction, π–cation interaction, and hydrogen bonding. Maintains functionality in presence of common co-existing ions. 	Decreased adsorption efficiency after multiple reuse cycles.
ABC/MX composite ¹³⁹	 Electrostatic attraction, reduction, chelation, hydrogen bonding. Excellent selectivity and ion interference resistance. Thermally enhanced adsorption (endothermic; capacity ↑ with temperature). Stable for real-world application: Minimal impact from co-ions and long-term storage. 	 Cr(VI) regeneration is limited due to MXene oxidation. No cytotoxicity or leaching data; safety in biomedical or food applications not assured. Performance slightly affected by HPO₄²⁻ due to competitive adsorption. MXene's known susceptibility to oxidation may limit long-term use in oxidative environments. Complex DFT-supported adsorption models require experimental validation in multicomponent matrices.
MXene-PEI ¹⁴⁰	 Fast adsorption kinetics: Equilibrium within 20–60 min. Mechanisms: Electrostatic attraction, redox reaction (Cr(VI) → Cr(III)), π–π stacking, hydrogen bonding. Thermally and mechanically stable structure. Effective even in the presence of competing ions. Easy handling and reusability due to aerogel form. 	 No biological toxicity evaluation for MXene oxidation products or long-term exposure. Oxidation of MXene over cycles not deeply discussed. Desorption agents (ethanol, NaOH) could impact cost or scalability. No real wastewater or multi-contaminant system testing.

Page 86 of 119 Nanoscale

Nanoscale Accepted Manuscript

MX-PAN membrane ¹⁵⁰	Fast kinetics: Equilibrium achieved in 30–60 min.	No biological or environmental toxicity validation of
	• Mechanisms: Electrostatic interactions, hydrogen bonding, π – π	MXene degradation products.
	interactions, redox transformation.	Oxidation stability of MXene during long-term use
	Excellent mechanical strength and porosity from aerogel	not addressed.
	architecture.	Real wastewater validation not included.
	• Effective in presence of co-existing ions (Cl ⁻ , SO ₄ ²⁻ , CO ₃ ²⁻).	• Performance slightly decreased with coexisting PO ₄ ³⁻
	Scalable and sustainable fabrication method using wood waste	ions.
	and green chemistry.	
TMAOH delaminated	Exceptional adsorption capacity of 1026 mg/g at 318 K, among	Performance decreases in presence of multivalent
Ti3C2Tx MXene	the highest reported for MXenes.	cations (Ca ²⁺ , Al ³⁺) due to competition and
nanosheets ¹³⁵	Fast adsorption kinetics, reaching equilibrium in 10 minutes.	electrostatic shielding.
	Mechanism confirmed: electrostatic interaction and ion	Adsorption capacity decreases slightly after several
	exchange $(TMA^+ \leftrightarrow MB^+)$ dominate.	reuse cycles due to partial loss and delamination
	Highly dispersible suspension prevents restacking, ensuring	damage.
	active surface availability.	No real wastewater or biological testing provided,
	• High performance in MNM (membrane form): up to 96%	performance in complex matrices remains unproven.
	removal efficiency with good dye flux (52 L/m ² ·h at 0.6 mg/cm ²	Freeze-drying reduces performance, dry-state
	loading).	materials are less effective due to aggregation.
	• Good tolerance to pH range (2–10) and monovalent salts.	
alk-	Ultra-high adsorption capacities achieved: 2095.9 mg/g for CR,	Limited recyclability scope with significant capacity
MXene/CoFe ₂ O ₄ /CS ¹⁴¹	1333.9 mg/g for Rhodamine B, and 537.6 mg/g for Malachite	loss for anionic dyes (34% loss for CR vs <20% for
	Green, among the highest reported for similar adsorbents.	cationic dyes) indicating preferential performance
		toward cationic species.

	The system demonstrated versatility in removing both cationic	No environmental safety assessment conducted, and
		•
	and anionic dyes simultaneously with rapid kinetics	complex multi-step synthesis process involving
	(equilibrium within 60-120 minutes) and excellent magnetic	multiple components (alkalized MXene, CoFe ₂ O ₄ ,
	separability.	chitosan) may limit practical scalability and cost-
	Multiple synergistic mechanisms including electrostatic	effectiveness.
	interaction, π - π stacking, and hydrogen bonding provided	
	comprehensive dye removal capability with good structural	
	stability after cycling.	
MXene (Ti ₃ C ₂)/Fe ₃ O ₄ ¹⁴²	• Temperature-enhanced performance with 91.93% removal	Limited scope with testing only on single cationic dye
	efficiency at 55 °C compared to lower temperatures,	(MB) and no comprehensive evaluation across
	demonstrating excellent high-temperature adsorption capability	different dye types or real wastewater conditions.
	(11.68 mg/g maximum capacity).	No recyclability assessment conducted, leaving
	• The system exhibited superparamagnetic properties (20.3	questions about long-term performance, stability, and
	emu/g) enabling easy magnetic separation and recovery from	practical applicability, with potential concerns about
	aqueous solutions with good stability (zeta potential -48 mV).	magnetic material leaching and structural integrity
	Multiple adsorption mechanisms identified including	over multiple cycles.
	electrostatic attraction, hydrogen bonding (Ti-OH···N), and	
	surface interactions, with temperature-dependent mechanism	
	shifts from physisorption (25 °C) to chemisorption (55 °C).	
AMXGO ¹³⁶	 Ultra-high adsorption capacities: 1111.6 mg/g for MG and 	Limited real-world testing - only synthetic single-dye
ANVIAGO		
	1133.7 mg/g for CR, among highest for MXene-based	solutions, not actual industrial wastewater.
	adsorbents.	

Page 88 of 119 Nanoscale

Nanoscale Accepted Manuscript

	• Excellent versatility with >92% removal for both cationic and anionic dyes, plus selective separation capability.	Complex multi-step synthesis and long equilibrium times (24-48 hours) limit scalability and cost-
	 Multiple adsorption mechanisms with 3.5x surface area increase compared to pristine MXene. 	effectiveness.
ZIF-8@IL- MXene/PNIPAM ¹⁵³	 First hydrogel achieving simultaneous high-efficiency adsorption of phenols, dyes, and metal ions (198.40, 325.03, and 285.65 mg/g respectively). Excellent mechanical properties withstanding ten repeated compressions without damage. 100× increase in specific surface area through ZIF-8 incorporation. Novel ionic liquid grafting approach for MXene stabilization 	 Complex multi-step synthesis requiring precise control of ZIF-8 growth conditions and IL grafting. Significant decrease in Cu²⁺ regeneration efficiency (29% after 5 cycles) due to irreversible chelation. Limited scalability due to multiple synthesis stages and specialized materials (ionic liquids, ZIF-8 precursors). Reduced swelling ratios with ZIF-8 growth
SC-PLA/PDA/MXene membrane ¹⁵⁵	 Outstanding dual functionality combining excellent oil/water separation and organic dye adsorption capabilities. High flux performance with maximum water flux of 3009.2 L/(m²·h) and oil flux of 6397.9 L/(m²·h) for pure liquids. Exceptional MB adsorption capacity of 434.8 mg/g following Langmuir model. Excellent photothermal conversion ability enhancing separation efficiency under light irradiation (water flux: 9428.4 L/(m²·h), oil flux: 12498.1 L/(m²·h)). 	 Complex multi-step synthesis involving electrospinning, polydopamine self-polymerization, and MXene deposition requiring precise control. Limited long-term stability assessment under continuous industrial operating conditions. Gradual performance decline in oil flux during extended cycling due to irreversible pore blocking by residual oil.

	Superior amphiphilic properties enabling effective separation	Temperature and light dependency for optimal
	of different emulsion types (oil-in-water and water-in-oil).	performance may limit operational flexibility in
	Novel polydopamine-assisted MXene deposition strategy	varying environmental conditions.
	ensuring uniform distribution and stable attachment.	No comprehensive economic analysis or cost-
	Simultaneous removal capability for both oil and organic	effectiveness evaluation compared to conventional
	pollutants in single-step process.	separation methods.
	• Ultrahigh oil/water separation efficiency of 99.12% with	• Limited testing on real industrial wastewater with
	excellent selectivity.	complex compositions - most studies conducted using
	• Good mechanical properties with tensile strength above 4.5	synthetic solutions.
	MPa maintaining structural integrity.	Potential scalability challenges for large-scale
		industrial applications not thoroughly addressed.
		• Performance optimization requires specific
		conditions (pH, temperature, light irradiation) which
		may complicate real-world implementation.
		Incomplete removal during regeneration treatment
		leading to gradual accumulation of contaminants
		affecting long-term performance.
TiV _C T _X /GAs ¹⁵⁶	Outstanding broad-spectrum removal capabilities: 319.67,	Complex multi-step synthesis involving
	303.45, 229.97, 217.87, and 283.38 mg/g for MB, RhB, CR,	hydrothermal treatment, freeze-drying, and hazardous
	MO, and TCH respectively.	HF acid for MXene preparation.
	Superior performance compared to other adsorbents in	Limited long-term stability assessment with only 5-
	comparative analysis.	cycle regeneration study.
	I.	1

Nanoscale Page 90 of 119

- Excellent compatibility with harsh environments including wastewater containing multiple dyes/drugs and inorganic salts.
- Remarkable photothermal conversion ability reaching 95.0 °C within 1 min under sunlight irradiation.
- Large absorption capacity for oils and solvents ranging from 40-90 g/g.
- Spontaneous and endothermic adsorption process following Langmuir model and pseudo-second-order kinetics.

- Performance dependency on pH, temperature, and light conditions may limit operational flexibility.
- No comprehensive economic analysis or costeffectiveness evaluation provided.
- Mostly tested on synthetic wastewater rather than real industrial effluents.
- Endothermic nature requires higher temperatures for optimal performance which may increase operational costs.

Accepted Manuscript

Nanoscale

View Article Online DOI: 10.1039/D5NR02336J

7. Comparative insights into the advantages and limitations of $Ti_3C_2T_x$ MXene in dye adsorption and photocatalytic applications

A comprehensive analysis of Ti₃C₂T_x MXene variants and their performance characteristics is presented in **Table 2**, highlighting both positive insights and critical limitations across different modification strategies. Ti₃C₂T_x MXene represents a groundbreaking milestone as the first reported MXene system for dye removal and photocatalysis applications, establishing the foundation for an entire class of 2D materials in environmental remediation. The material exhibits exceptional electrostatic interaction capabilities with cationic dyes, particularly MB, by leveraging its inherent negative surface charge and rich surface chemistry to achieve a strong binding affinity. This electrostatic mechanism proves highly effective for irreversible adsorption, indicating robust dye-MXene interactions that prevent desorption under mild conditions.¹³¹

Photocatalytic versatility emerges as a distinctive advantage, with Ti₃C₂T_x exhibiting effective UV-driven dye degradation for both MB and AB80, demonstrating dual functionality beyond conventional adsorbents. The heterogeneous adsorption behavior, confirmed through excellent Freundlich model fitting, indicates the presence of multiple binding sites with varying energies, providing flexibility in accommodating diverse dye molecules. This multi-site adsorption mechanism contributes to the material's broad applicability across different dye classes and concentrations.¹³¹ Performance enhancement through strategic modifications demonstrates remarkable capabilities through surface functionalization strategies that dramatically amplify Ti₃C₂T_x performance. Alkali treatments using NaOH, LiOH, and KOH achieve remarkable improvements, with NaOH-treated Ti₃C₂T_x demonstrating an exceptional adsorption capacity of 189 mg/g for MB, representing a nearly five-fold increase over the pristine material. This enhancement stems from interlayer spacing expansion (up to 29% increase) and surface

chemistry modification through fluoride-to-hydroxyl group conversion (-F \rightarrow -OH), creating rolling color more favorable binding sites. ¹³² Fluoride-free synthesis approaches using NaBF₄/HCl represent a significant advancement, ¹³³ producing h-Ti₃C₂ with superior surface area (44.6 vs. 8.9 m²/g) and enhanced exfoliation characteristics. This environmentally safer synthesis route eliminates hazardous HF usage while achieving better performance for MB adsorption, addressing both environmental concerns and performance requirements. The Langmuir model compliance in modified systems indicates a transition to monolayer adsorption, suggesting a more uniform distribution of binding sites.

Chemical functionalization through sulfonic acid modification (Ti₃C₂-SO₃H) showcases dramatic capacity enhancement from 21.10 to 111.11 mg/g, with concentration-dependent performance reaching 723.35 mg/g under optimized conditions. This diazonium-mediated surface modification enhances both surface chemistry and interlayer spacing, creating highly active adsorption sites.¹²⁹ Surfactant modification using cetyltrimethylammonium bromide (CTAB) achieves 213.00 mg/g capacity while maintaining broad pH tolerance, particularly effective at pH 3.¹³⁰ Critical performance limitations and stability challenges reveal that oxidative instability represents the most significant limitation of Ti₃C₂T_x systems, with poor stability in aqueous environments leading to oxidation to TiO₂.¹³¹ This fundamental degradation pathway compromises long-term performance and material integrity, particularly problematic for continuous water treatment applications. The oxidation susceptibility extends across all Ti₃C₂T_x variants, indicating a materials-level challenge requiring protective strategies or alternative compositions.

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

The limited adsorption capacity in its pristine form presents significant competitive disadvantages, with the baseline $Ti_3C_2T_x$ achieving only ~39 mg/g, substantially lower than that of high-performing adsorbents.¹³¹ Even modified variants like h- Ti_3C_2 show moderate performance (~24 mg/g), indicating inherent capacity limitations that require extensive

modification for competitive performance.¹³³ Surface area reduction following certain colling modifications, such as CTAB treatment (5.02 to 2.06 m²/g), demonstrates trade-offs between functionalization and accessible surface area.¹³⁰ Mechanistic complexity across Ti₃C₂T_x systems requires further clarification,¹³¹ with multiple interaction pathways including electrostatic attraction, π-cation interactions, hydrogen bonding,¹³⁰ and ion exchange contributing to overall performance. This complexity makes performance prediction and optimization challenging, particularly in multi-component systems where competitive interactions may alter dominant mechanisms. Scalability and practical implementation barriers pose significant challenges, particularly due to the complexity of synthesis, which requires multi-step modifications. Alkali treatments,¹³² chemical functionalization,¹²⁹ and surfactant modifications¹³⁰ introduce additional processing steps, chemical consumption, and potential environmental concerns. The time-consuming nature of layer-by-layer assembly and precise control requirements for modification procedures limit industrial scalability and cost-effectiveness.¹³⁷

Regeneration limitations across Ti₃C₂T_x systems present economic and environmental concerns. Decreased adsorption efficiency after multiple reuse cycles, observed in CTAB-modified systems, indicates structural degradation or irreversible binding.¹³⁰ Desorption agent requirements (ethanol, NaOH) increase operational costs and waste generation, while incomplete regeneration leads to a gradual decline in performance. Real-world validation gaps represent critical knowledge deficits, with most studies conducted using synthetic single-dye solutions rather than complex industrial wastewaters. Performance degradation in the presence of multivalent cations (Ca²⁺, Al³⁺) and competitive anions highlights selectivity challenges in realistic water matrices. No comprehensive real wastewater testing limits practical applicability assessment.

Environmental and safety considerations highlight critical research gaps, with the assessing according to toxicity and biocompatibility representing urgent needs. No systematic evaluation of Ti₃C₂T_x's environmental impact or human health effects exists, while the toxicity of oxidation products, leaching behavior, and long-term environmental fate remain uncharacterized, raising concerns about its widespread deployment. Surface modification agents, such as CTAB and sulfonic acid groups, introduce additional chemical complexity, requiring safety evaluation. HF-based synthesis environmental concerns necessitate safer alternatives, with fluoride-free routes showing promise but requiring further optimization. ¹³³ Waste generation from synthesis processes and regeneration procedures requires a comprehensive life cycle assessment to evaluate their overall environmental impact. Challenges associated with disposing of spent MXene materials and their oxidation products necessitate a systematic investigation.

Future directions and innovation opportunities suggest that enhancing oxidative stability

represents the highest priority research direction, requiring protective coating strategies, alloy composition optimization, or control of operational conditions. Encapsulation approaches, such as those using protective polymer layers or composite formation with stable matrices, could address degradation issues while maintaining functional performance. Performance optimization strategies should focus on synergistic modification approaches combining interlayer engineering, surface functionalization, and morphological control. AI-guided design can accelerate composition-structure-property relationships discovery, enabling rational optimization of Ti₃C₂T_x variants for specific applications. Scale-up engineering focusing on continuous flow systems, automated synthesis procedures, and regeneration optimization will bridge the laboratory-to-industry gap. Economic feasibility studies incorporating material costs, processing requirements, and performance benefits are essential for commercial viability assessment. Comprehensive environmental impact evaluation through life cycle assessment

will guide sustainable development and regulatory compliance for $Ti_3C_2T_x$ -based with C_2T_x -based treatment technologies.

8. Benchmarking MXene-based adsorbents against established materials for sustainable dye remediation

To compare performance and demonstrate MXenes' potential in wastewater treatment, Table 3 provides a detailed quantitative comparison of 2D MXenes with well-known adsorbents, including MOFs, COFs, activated carbon, graphene-based materials, and CNTs for dye removal. The data, sourced from peer-reviewed studies between 2004 and 2024, highlights the outstanding adsorption abilities of MXene-based adsorbents under various conditions. TMAOH-Ti₃C₂Tx achieved the highest adsorption capacity of 1026 mg/g for MB, a 2.5 times increase over commercial activated carbon (400 mg/g) and a significant edge over GO (250-300 mg/g) and multi-walled CNTs (400 mg/g) in similar settings. 135,167 Likewise, MXene/biomass activated carbon composites showed excellent performance for CR, removing over 1400 mg/g, much higher than commercial activated carbon (500 mg/g) and functionalized CNTs (500 mg/g). 158,168 MXenes' top performance comes from their unique dual adsorption process that includes surface adsorption and intercalation, along with abundant surface functional groups (-OH, =O, -F), and adjustable surface chemistry that fosters strong electrostatic interactions with both cationic and anionic dyes. MXenes show quick kinetics, reaching equilibrium in 5-120 minutes versus 60-300 minutes for traditional adsorbents, and they can be reused with over 90% efficiency after multiple cycles. Although MOFs and COFs perform well in some cases, like TpStb-SO₃Na COF achieving high capacities for certain cationic dyes, MXenes offer greater versatility, faster kinetics, and wider applicability across diverse dye types and conditions, making them the most promising new adsorbents for sustainable wastewater treatment. 169

Table 3: Adsorption capacities of MXenes vs. conventional adsorbents for organic dyes

Nanoscale Accepted Manuscript

Adsorbent	Dye	Adsorption	Contact	pН	Temp (°C)	Initial Vi
Material		Capacity	Time			conc.
		(mg/g)				(mg/L)
	M	Kenes (2D Transit	ion Metal (_ Carbid	les)	
TMAOH-	MB	1026	Rapid	7	45	100
$Ti_3C_2Tx^{135}$						
Ti ₃ C ₂ Tx@Sodium	MB	969	60 min	10	25	300
Alginate ¹⁷⁰						
Ti ₃ C ₂ Tx/Carbon	MB	357	120 min	7	30	200
Foam ¹⁵⁷						
Ti ₃ C ₂ Tx/Carbon	CR	648	120 min	7	30	300
Foam ¹⁵⁷						
MXene/Biomass	CR	>1400	180 min	7	25	500
AC^{158}						
NaOH-Ti ₃ C ₂ Tx ¹³²	MB	189	120 min	7	25	100
Ti ₃ C ₂ Tx/Loofah	MB	175	60 min	7	25	200
Carbon ¹⁶⁸						
Ti ₃ C ₂ Tx/Loofah	CR	93	60 min	7	25	150
Carbon ¹⁶⁸						
	1	Activated	Carbon		1	1
Commercial PAC	MB	400	120 min	6-7	25	200
(Norit) ¹⁶⁷						
Commercial PAC	CR	500	120 min	6-7	25	250
(Norit) ¹⁶⁷						
Tea Seed Shell	MB	325	180 min	6-7	25	200
AC^{171}						
Water-activated	MB	149	5 min	6.5-	25	100
AC ¹⁷²				7		
Palm Kernel Shell	MB	15	60 min	7	25	100
AC^{173}						
Ashitaba Waste	CR	200-300	180 min	7	25	200
AC ¹⁷⁴						
Walnut Shell	MB	150-200	180 min	7	25	150
AC ¹⁷⁴						
		Graphene-bas	sed Materia	als		
Graphene Oxide ¹⁷⁵	MB	250-300	60 min	7	25	100

Graphene Oxide¹⁷⁴

Various COFs¹⁸¹

Rhodamine

60 min

7

25

100 View Article Online

200

200

250

150

200

100

200

100-300

Variable

200-400

200-250

9. Challenges, recommendations and future prospectives

MB

Although MXene-based materials are often employed to remove dyes in the environment, studies on these materials are still in their initial phases due to the numerous problems encountered by researchers. The theoretical concept of using MXene-based materials has expanded faster, but the experimental aspect remains slower. However, several challenging issues still need to be carefully considered. In order to realize their real-world relevance and commercial viability in large-scale water treatment applications, it will be necessary to take future steps toward scaling up MXenes and nanocomposites based on MXene.

120 min

7

25

Current MXene-based dye removal technologies operate at Technology Readiness Level (TRL) 3-4, characterized by laboratory-scale proof-of-concept and component validation in

controlled environments. 60 While fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adsorption mechanisms are well-junders to adjust on the fundamental adjust of the fund and performance optimization has been demonstrated in synthetic solutions, significant gaps remain in real-world validation, system integration, and scalable manufacturing processes. 182 Real wastewater performance presents the most critical challenge, as industrial effluents contain complex mixtures of organic contaminants, salts, suspended particles, heavy metals, and fluctuating pH that can drastically reduce MXene adsorption capacity and selectivity compared to single-component laboratory studies. Fouling mechanisms from biological wastes, organic matter, and inorganic scaling create surface deactivation and pore blockage, reducing recyclable capacity over operational cycles. 89 Cost and scalability barriers arise from expensive, hazardous synthesis methods using HF or LiF/HCl, with material costs of \$50-200/kg compared to \$1-10/kg for conventional adsorbents. 183 Environmental impact assessment through Life Cycle Analysis (LCA) reveals carbon footprints of 5-15 kg CO₂ equivalent per kg MXene, primarily from energy-intensive synthesis processes. 102 Advancement to TRL 6-7 requires pilot-scale demonstrations, comprehensive technoeconomic analysis, and development of green synthesis alternatives to establish commercial feasibility.

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Several vital areas require attention to fully harness the capabilities of MXene-based materials in dye removal and other applications:

1. Improvements in the etching procedure: Traditional etching methods for synthesizing MXenes often employ hazardous chemicals that can harm the environment. Therefore, exploring and creating more ecologically friendly techniques for the etching process is critical. This transition could make MXene production safer and more sustainable and increase its appeal for industrial applications in which environmental restrictions are growing more stringent.

- 2. **Research on toxicity**: Another critical area of exploration is the potential toxicity of the potential MXene-based materials. Understanding how these materials interact with the environment and human health is essential, especially for therapeutic applications. Comprehensive studies assessing the biocompatibility and environmental impact of MXenes will help establish safety protocols and regulatory frameworks, ensuring their responsible use in various applications, including medicine and environmental remediation.
- 3. **Surface characterization**: The surface characteristics of MXene-based materials play a significant role in their adsorption abilities. Gaining a deeper understanding of these properties, including surface area, functional groups, and morphology, can lead to modifications that enhance their performance in dye removal. Tailoring the surface chemistry of MXenes could optimize their interaction with various dye molecules, thereby improving their efficiency as adsorbents (**Figure 18**).
- 4. Lifespan of recycled materials and advanced integration strategies: Finally, further research is needed to evaluate the lifespan and reusability of recycled MXene-based materials in real applications. Understanding how these materials degrade over time and how they operate after several cycles of usage is critical to determining their feasibility in long-term applications. Investigating to create methods for the regeneration of MXenes could considerably improve their use in various applications, including wastewater treatment, energy storage, and sensing technologies. Integrating AI and ML can transform the development of MXene materials. These technologies may be utilized to forecast material degradation trends, enhance synthesis procedures, and formulate more effective recycling methods. ML models may evaluate massive datasets to discover trends and recommend ideal settings for material regeneration, therefore eliminating trial-and-error experimentation.

Nanoscale Accepted Manuscript

- 5. Performance in real wastewater: The majority of MXene research has been carried office online using artificial dye solutions in controlled lab settings. Real wastewater streams, on the other hand, are far more complicated and frequently include a combination of organic contaminants, salts, suspended particles, heavy metals, and fluctuating pH. These conflicting elements have the potential to drastically lower MXenes' adsorption capability and selectivity. Therefore, assessing MXene-based systems in realistic wastewater environments must be a top priority for future research in order to confirm their resilience, effectiveness, and longevity.
- 6. **Fouling and long-term stability:** MXenes are prone to fouling from biological wastes, organic matter, and other wastewater particles, which can result in surface deactivation, pore blockage, and decreased recyclable capacity. To ensure constant performance across several cycles, fouling must be addressed via surface functionalization, hybrid coating techniques, or integration with antifouling polymers. Additionally, MXene degradation under extended exposure to aquatic environments, particularly under variable redox conditions, remains a significant but little-studied problem.
- 7. Cost and scalability: The top-down etching techniques that use dangerous chemicals like HF or LiF/HCl are among the current MXene production methods that are expensive, dangerous, and challenging to scale. Additionally, the purification and post-synthesis delamination processes require a lot of energy and time. To move MXene manufacturing from laboratory to commercial scale, it will be essential to develop low-cost, scalable, and environmentally friendly synthesis methods, such as electrochemical etching, green exfoliation processes, and continuous flow systems.
- **8.** Environmental impact and sustainability: MXenes are intriguing prospects for green remediation solutions, but their total environmental impact must be carefully evaluated. To assess energy use, chemical use, emissions, and end-of-life disposal or recyclability, LCA

studies are crucial. To guarantee environmental and regulatory compliance, issues with rocassed leaching, nanoparticle release, and ecotoxicity should also be addressed using thorough risk assessment frameworks. Furthermore, performing a LCA is essential for identifying the environmental impacts of MXene-based materials throughout their life span, from synthesis to disposal. Such a technique guarantees that the materials are sustainable and effective, in agreement with the overall green chemistry initiatives. Evaluating LCA would make MXenes more attractive for firms embracing eco-friendly technology.

For biomedical and environmental applications, *in vivo* cytotoxicity analysis is significant in determining the toxicity of MXenes. Such tests would ensure the safer use of MXenes in real-world applications, such as sensing and wastewater treatment, and guarantee no risk to human health or ecosystems. Advanced characterization solutions including real-time monitoring capabilities, molecular docking studies, and computational simulation technologies, combined with emerging force measurement techniques and advanced X-ray methodologies, could significantly improve MXene system activity and reusability assessment. Force measurement approaches, particularly atomic force microscopy (AFM) and nanoindentation techniques, enable precise quantification of interfacial interactions between MXene surfaces and target molecules, providing critical insights into adsorption mechanisms at the nanoscale level. These force-based methodologies can map surface heterogeneity, measure adhesion forces, and characterize mechanical properties evolution during cycling processes, offering unprecedented understanding of structure-property relationships in MXene systems.¹⁸⁴

Although MXene-based materials hold great promise for environmental applications, addressing these challenges through focused research will be crucial for translating theoretical advancements into practical solutions. By improving synthesis methods, assessing toxicity, understanding surface properties, and evaluating lifespan, the full potential of MXenes can be realized, paving the way for innovative approaches to dye removal and other environmental

challenges. Despite years of investigation on MXene, these materials are still in their cartificiential phases of development. Standardization is still required to produce MXene, an attractive material with stable characteristics, including higher biocompatibility, uniform dispersion, prolonged strength, and higher efficiency compared to other 2D materials. 60 Industry-academia collaboration is essential for developing large-scale MXene-based materials, including encapsulated adsorbents, membranes, and electrodes, with increased hazardous removal efficiency. This will enable the production and deployment of low-cost MXene-based adsorbent solutions for wastewater treatment in residential and commercial areas.

Artificial Intelligence (AI) and Machine Learning (ML) technologies offer transformative potential for advancing MXene-based water treatment systems through multiple specific applications. 185 Machine learning algorithms can predict optimal MXene compositions and surface functionalization strategies by analyzing structure-property relationships from experimental datasets, with Density Functional Theory (DFT) calculations combined with ML models enabling screening of thousands of potential MXene variants to identify highperformance candidates before experimental synthesis. 186 Gaussian Process Regression and Random Forest algorithms have been successfully applied to predict adsorption capacities based on surface area, functional group density, and interlayer spacing parameters, ¹⁸⁷ while neural network models can optimize synthesis conditions, including etching time, temperature, and chemical concentrations to achieve target material properties. 188 Support Vector Machine (SVM) and Artificial Neural Network (ANN) models demonstrate capability in predicting dye removal efficiency based on water quality parameters including pH, ionic strength, competing ions, and dye concentrations, 189 with ensemble learning methods such as XGBoost and Random Forest showing superior accuracy in predicting breakthrough curves and saturation points for MXene-based filtration systems. 190

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

Real-time process optimization represents another critical application area, where adaptive a

lev most street on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM.

max max incompany with the street of the street

control algorithms integrated with IoT sensor networks enable dynamic optimization of treatment parameters. Parame

5G technology enables ultra-low latency communication between distributed sensor networks and central control systems, facilitating real-time optimization of multiple treatment units, while edge computing capabilities allow on-site ML inference for immediate decision-making without cloud dependency. Pot IoT sensors equipped with selective dye detection capabilities provide high-resolution monitoring of specific contaminants, enabling precision treatment protocols that automatically adjust based on real-time water quality parameters. Cloud-based platforms integrate multi-site data for comprehensive performance analysis and predictive modeling, with blockchain technology ensuring data integrity and traceability for regulatory compliance and performance verification. Specific implementation examples include predictive maintenance models that analyze pressure drop patterns, flow rate variations, and energy consumption trends to schedule optimal regeneration cycles, and multi-objective

optimization algorithms that balance treatment efficiency, energy consumption view fallow operational costs to minimize total system expenses. 197

Future prospects include quantum machine learning algorithms that may accelerate materials discovery by exploring vast chemical spaces more efficiently than classical computers, automated laboratory systems guided by AI algorithms conducting high-throughput synthesis and screening experiments with minimal human intervention, and integration of genomics data with ML models to predict ecological impacts and optimize MXene designs for environmental compatibility. Pederated learning frameworks enable collaborative model training across multiple treatment facilities while preserving data privacy and proprietary information, while augmented reality interfaces provide operators with real-time system status and maintenance guidance through smart glasses and mobile applications. Phis comprehensive AI/ML integration represents a paradigm shift toward intelligent water treatment systems that continuously learn, adapt, and optimize their performance, ultimately enabling more efficient, cost-effective, and sustainable MXene-based environmental remediation technologies capable of autonomous operation and predictive performance optimization across diverse water treatment scenarios.

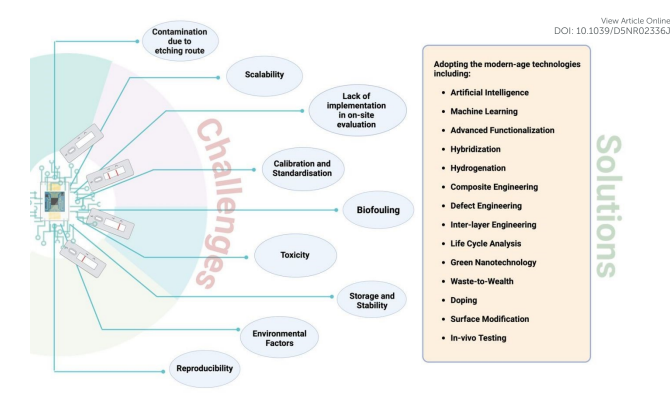


Figure 18. Challenges and Solutions for employing MXene-based materials in the adsorption of dyes

10. Conclusion

In conclusion, MXenes have demonstrated an emerging class of promising two-dimensional adsorbents for the adsorption of various dyes in an aqueous environment. There are various methods for synthesizing MXenes. Presently, the top-down method involving hydrofluoric acid etching is the most commonly used method. Although this approach is less expensive, there is more environmental harm risk. Their high surface areas, active surface terminal groups, hydrophilicity, and distinct combination of layered structures make their adsorption properties excellent. However, its oxidative and aggregative properties make effective adsorption and regeneration difficult. As a result, further modifications are needed. Surface modification, blending, and stripping are some modification methods for MXenes. Blending, in particular,

anoscale Accepted Manuscript

essential features of MXene materials.

This review has highlighted the substantial development in the design and fabrication of

MXene-based adsorbents, their adsorption capacity, their physicochemical properties, and

adsorption mechanisms of MXenes, including chemical, physical, and electrostatic interactions

for the removal of toxic dyes from the aqueous environment. Overall, MXenes are a cutting-

edge advance in water treatment, providing an alternative route to more sustainable and

effective solutions for dye-contaminated water. Furthermore, with the integration of modern

technologies such as IoT, AI, machine learning, and 5G communications, MXene-based

nanosorbents have the potential to be next-generation material-based intelligent environmental

technologies that form smart and connected societies and cities.

Abbreviation:

2D: Two dimensional

3D: Three dimensional

AA: Acrylic acid

AB93: Methyl blue

AI: Artificial intelligence

BC: Bacterial cellulose

BD: Benzidine

BET: Brunauer–Emmett–Teller

CDCs: carbon-derived carbides

CNT: Carbon nanotubes

COF: Covalent organic framework

CR: Congo Red

CTAB: Cationic surfactant solution

CVD: Chemical Vapor Deposition

DASNP: Dialdehyde starch nanoparticles

View Article Online

DOI: 10.1039/D5NR02336J

DBM: Direct brown M

DBRN: Direct black RN

DMSO: Dimethyl sulphoxide

EDX: Energy-dispersive X-ray

FTIR: Fourier-transform infrared spectroscopy

GPA: Gigapascals

GO: Graphene oxide

HRTEM: High-resolution transmission electron microscopy

IL: Ionic liquid

LCA: Life cycle assessment

LDH: Layered double hydroxides

MB: Methylene Blue

MG: Malachite Green

ML: Machine learning

MO: Methyl Orange

MOF: Metal-organic framework

MPa: Megapascals

NMs: Nanofiber membranes

PA: p-Phenylenediamine

PAN: Polyacrylonitrile

PDA: polydopamine

PEI: Polyethylenimine

PEPLD: Plasma-enhanced pulsed laser deposition

PLA: Polylactic acid

rGO: Reduced graphene oxide

RhB: Rhodamine B

ROS: Reactive oxygen species

SA: Sodium alginate

SAED: Selected area electron diffraction

SEM: Scanning electron microscopy

View Article Online

DOI: 10.1039/D5NR02336J

SSA: Specific surface area

SSbD: Safe and Sustainable by Design

TAPB: 1,3,5- tris(4-aminophenyl) benzene

TMAOH: Tetramethylammonium hydroxide

TP: 1,3,5-triformylphloroglucinol

UN: United Nation

XPS: X-ray photoelectron spectroscopy

XRD: X-ray diffraction

ZIF: Zeolitic imidazolate framework

Acknowledgements

Published on 04 August 2025. Downloaded by Yunnan University on 8/23/2025 12:46:06 PM

The authors would like to express their gratitude to the institutions and colleagues who provided valuable insights and feedback throughout the preparation of this review article.

Declaration of generative AI in scientific writing

While preparing this work the author(s) used Chatgpt and Claude to improve the readability and language of the manuscript. After using this tool/service, the author(s) reviewed and edited the content as needed and take(s) full responsibility for the content of the published article.

Conflict of Interests

The authors declare no conflicts of interest.

References

- A. Khosla, Sonu, H. T. A. Awan, K. Singh, Gaurav, R. Walvekar, Z. Zhao, A. Kaushik, M. Khalid and V. Chaudhary, 2022, preprint, DOI: 10.1002/advs.202203527.
- D. Pathania, S. Kumar, P. Thakur, V. Chaudhary, A. Kaushik, R. S. Varma, H. Furukawa, M. Sharma and A. Khosla, *Sci Rep*, DOI:10.1038/s41598-022-14984-3.
- 3 Sonu and V. Chaudhary, 2022, preprint, DOI: 10.1149/2754-2726/ac92ed.
- 4 S. Shekhar, A. K. Yadav, A. Khosla and P. R. Solanki, 2022, preprint, DOI: 10.1149/2754-2726/ac9227.
- 5 T. Suzuki, T. Hidaka, Y. Kumagai and M. Yamamoto, 2020, preprint, DOI: 10.1038/s41590-020-0802-6.

View Article Online

DOI: 10.1039/D5NR02336J

- 6 T. T. T. Toan and D. M. Nguyen, *ECS Sensors Plus*, 2022, **1**, 021604.
- 7 C. Valli Nachiyar, A. D. Rakshi, S. Sandhya, N. Britlin Deva Jebasta and J. Nellore, *Case Studies in Chemical and Environmental Engineering*, DOI:10.1016/j.cscee.2023.100339.
- J. Blair, G. Rathee, A. Puertas-Segura, L. M. Pérez and T. Tzanov, *Environ Res*, 2025, **279**, 121783.
- 9 G. Rathee, H. Chugh, S. Kohli, R. K. Gaur and R. Chandra, *Mater Adv*, DOI:10.1039/d2ma01081j.
- 10 Y. Zhou, J. Lu, Y. Zhou and Y. Liu, 2019, preprint, DOI: 10.1016/j.envpol.2019.05.072.
- E. F. D. Januário, T. B. Vidovix, N. de C. L. Beluci, R. M. Paixão, L. H. B. R. da Silva, N. C. Homem, R. Bergamasco and A. M. S. Vieira, 2021, preprint, DOI: 10.1016/j.scitotenv.2021.147957.
- Y. Pan, S. Sanati, R. Abazari, A. Jankowska, J. Goscianska, V. Srivastava, U. Lassi and J. Gao, *Coord Chem Rev*, 2025, **522**, 216231.
- S. X. Chin, K. S. Lau, S. Zakaria, C. H. Chia and C. Wongchoosuk, *Polymers (Basel)*, DOI:10.3390/polym14235165.
- P. V. Nidheesh, M. Zhou and M. A. Oturan, *Chemosphere*, DOI:10.1016/j.chemosphere.2017.12.195.
- 15 L. Bilińska and M. Gmurek, 2021, preprint, DOI: 10.1016/j.wri.2021.100160.
- 16 S. Khamparia and D. K. Jaspal, 2017, preprint, DOI: 10.1007/s11783-017-0899-5.
- 17 A. K. Verma, R. R. Dash and P. Bhunia, 2012, preprint, DOI: 10.1016/j.jenvman.2011.09.012.
- 18 S. Thakur and M. S. Chauhan, 2018.
- 19 M. M. Hassan and C. M. Carr, 2018, preprint, DOI: 10.1016/j.chemosphere.2018.06.043.
- 20 P. Moradihamedani, 2022, preprint, DOI: 10.1007/s00289-021-03603-2.
- S. Sarkar, N. T. Ponce, A. Banerjee, R. Bandopadhyay, S. Rajendran and E. Lichtfouse, 2020, preprint, DOI: 10.1007/s10311-020-01021-w.
- 22 M. Hasanpour and M. Hatami, 2020, preprint, DOI: 10.1016/j.molliq.2020.113094.
- 23 M. Khatami and S. Iravani, *Comments on Inorganic Chemistry*, DOI:10.1080/02603594.2021.1922396.
- 24 K. T. Kubra, M. S. Salman and M. N. Hasan, J Mol Liq, DOI:10.1016/j.molliq.2021.115468.
- 25 M. Naushad, A. A. Alqadami, A. A. Al-Kahtani, T. Ahamad, M. R. Awual and T. Tatarchuk, *J Mol Liq*, DOI:10.1016/j.molliq.2019.112075.
- 26 C. Cai, R. Wang, S. Liu, X. Yan, L. Zhang, M. Wang, Q. Tong and T. Jiao, *Colloids Surf A Physicochem Eng Asp*, DOI:10.1016/j.colsurfa.2020.124468.
- 27 R. Xing, W. Wang, T. Jiao, K. Ma, Q. Zhang, W. Hong, H. Qiu, J. Zhou, L. Zhang and Q. Peng, *ACS Sustain Chem Enq*, DOI:10.1021/acssuschemeng.7b00343.

- G. Rathee, A. Puertas-Segura, J. Blair, J. Rathee and T. Tzanov, *Prog Mater Sci*, 2025, 150 View Article Online 101403.
- 29 G. Rathee, N. Singh and R. Chandra, ACS Omega, DOI:10.1021/acsomega.9b03785.
- 30 G. Rathee, S. Kohli, A. Awasthi, N. Singh and R. Chandra, RSC Adv, DOI:10.1039/d0ra02766a.
- 31 G. Rathee, A. Awasthi, D. Sood, R. Tomar, V. Tomar and R. Chandra, *Sci Rep*, DOI:10.1038/s41598-019-52849-4.
- 32 M. J. K. Ahmed and M. Ahmaruzzaman, 2016, preprint, DOI: 10.1016/j.jwpe.2016.01.014.
- 33 M. Ahmaruzzaman, 2011, preprint, DOI: 10.1016/j.cis.2011.04.005.
- S. A. Khan, D. Hussain, N. Abbasi and T. A. Khan, *Chemosphere*, DOI:10.1016/j.chemosphere.2021.133232.
- S. A. Khan, N. Abbasi, D. Hussain and T. A. Khan, *Langmuir*, DOI:10.1021/acs.langmuir.2c00702.
- 36 K. S. Novoselov, A. K. Geim, S. V Morozov, D. Jiang, Y. Zhang, S. V Dubonos, I. V Grigorieva and A. A. Firsov, *Electric Field Effect in Atomically Thin Carbon Films*, 2000, vol. 404.
- L. Y. Yuan, Z. Q. Bai, R. Zhao, Y. L. Liu, Z. J. Li, S. Q. Chu, L. R. Zheng, J. Zhang, Y. L. Zhao, Z. F. Chai and W. Q. Shi, *ACS Appl Mater Interfaces*, DOI:10.1021/am405584h.
- 38 R. Ma and T. Sasaki, 2010, preprint, DOI: 10.1002/adma.201001722.
- J. N. Coleman, M. Lotya, A. O'Neill, S. D. Bergin, P. J. King, U. Khan, K. Young, A. Gaucher, S. De, R. J. Smith, I. V. Shvets, S. K. Arora, G. Stanton, H. Y. Kim, K. Lee, G. T. Kim, G. S. Duesberg, T. Hallam, J. J. Boland, J. J. Wang, J. F. Donegan, J. C. Grunlan, G. Moriarty, A. Shmeliov, R. J. Nicholls, J. M. Perkins, E. M. Grieveson, K. Theuwissen, D. W. McComb, P. D. Nellist and V. Nicolosi, *Science* (1979), DOI:10.1126/science.1194975.
- 40 K. S. Novoselov, A. K. Geim, S. V. Morozov, D. Jiang, M. I. Katsnelson, I. V. Grigorieva, S. V. Dubonos and A. A. Firsov, *Nature*, DOI:10.1038/nature04233.
- 41 K. V. Wong and B. Bachelier, J Energy Resour Technol, DOI:10.1115/1.4024917.
- 42 O. P. Chen, Y. J. Lin, W. Z. Cao and C. T. Chang, *Mater Lett*, DOI:10.1016/j.matlet.2017.01.030.
- 43 S. Wang, H. Sun, H. M. Ang and M. O. Tadé, 2013, preprint, DOI: 10.1016/j.cej.2013.04.070.
- 44 F. Perreault, A. Fonseca De Faria and M. Elimelech, *Chem Soc Rev*, DOI:10.1039/c5cs00021a.
- 45 Y. Fan, L. Li, Y. Zhang, X. Zhang, D. Geng and W. Hu, 2022, preprint, DOI: 10.1002/adfm.202111357.
- F. A. Janjhi, I. Ihsanullah, M. Bilal, R. Castro-Muñoz, G. Boczkaj and F. Gallucci, 2023, preprint, DOI: 10.1016/j.wri.2023.100202.
- 47 Y. Ibrahim, M. Meslam, K. Eid, B. Salah, A. M. Abdullah, K. I. Ozoemena, A. Elzatahry, M. A. Sharaf and M. Sillanpää, 2022, preprint, DOI: 10.1016/j.seppur.2021.120083.
- 48 I. Ihsanullah and M. Bilal, 2022, preprint, DOI: 10.1016/j.apmt.2022.101674.
- 49 I. Ihsanullah and M. Bilal, Chemosphere, DOI:10.1016/j.chemosphere.2022.135234.

- 50 S. Yu, H. Tang, D. Zhang, S. Wang, M. Qiu, G. Song, D. Fu, B. Hu and X. Wang, 2022, preprint Article Online DOI: 10.1016/j.scitotenv.2021.152280.
- 51 X. Jiang, A. V. Kuklin, A. Baev, Y. Ge, H. Ågren, H. Zhang and P. N. Prasad, 2020, preprint, DOI: 10.1016/j.physrep.2019.12.006.
- 52 M. Naguib, V. N. Mochalin, M. W. Barsoum and Y. Gogotsi, *Advanced Materials*, DOI:10.1002/adma.201304138.
- Q. Tao, M. Dahlqvist, J. Lu, S. Kota, R. Meshkian, J. Halim, J. Palisaitis, L. Hultman, M. W. Barsoum, P. O. Å. Persson and J. Rosen, *Nat Commun*, DOI:10.1038/ncomms14949.
- 54 Y. C. Zhou, X. H. Wang, Z. M. Sun and S. Q. Chen, J Mater Chem, DOI:10.1039/b101520f.
- 55 K. Huang, Z. Li, J. Lin, G. Han and P. Huang, *Chem Soc Rev*, DOI:10.1039/c7cs00838d.
- I. Persson, A. el Ghazaly, Q. Tao, J. Halim, S. Kota, V. Darakchieva, J. Palisaitis, M. W. Barsoum, J. Rosen and P. O. Å. Persson, *Small*, DOI:10.1002/smll.201703676.
- 57 X. Wang, L. Wu, H. Gao and X. Zhang, SCIENTIA SINICA Chimica.
- F. Kong, X. He, Q. Liu, X. Qi, Y. Zheng, R. Wang and Y. Bai, *Electrochim Acta*, DOI:10.1016/j.electacta.2018.01.196.
- J. Halim, J. Palisaitis, J. Lu, J. Thörnberg, E. J. Moon, M. Precner, P. Eklund, P. O. A. Persson, M. W. Barsoum and J. Rosen, *ACS Appl Nano Mater*, DOI:10.1021/acsanm.8b00332.
- B. Anasori, M. R. Lukatskaya and Y. Gogotsi, 2017, preprint, DOI: 10.1038/natrevmats.2016.98.
- A. Lipatov, H. Lu, M. Alhabeb, B. Anasori, A. Gruverman, Y. Gogotsi and A. Sinitskii, *Sci Adv*, DOI:10.1126/sciadv.aat0491.
- J. Zhang, N. Kong, S. Uzun, A. Levitt, S. Seyedin, P. A. Lynch, S. Qin, M. Han, W. Yang, J. Liu, X. Wang, Y. Gogotsi and J. M. Razal, *Advanced Materials*, DOI:10.1002/adma.202070180.
- 63 V. N. Borysiuk, V. N. Mochalin and Y. Gogotsi, *Nanotechnology*, DOI:10.1088/0957-4484/26/26/265705.
- S. Luo, S. Patole, S. Anwer, B. Li, T. Delclos, O. Gogotsi, V. Zahorodna, V. Balitskyi and K. Liao, *Nanotechnology*, DOI:10.1088/1361-6528/ab94dd.
- 65 Y. Gogotsi, *Matter*, DOI:10.1016/j.matt.2021.12.006.
- B. Liang, X. Liao, Q. Zhu, M. Yu, J. Li, B. Geng, K. Liu, D. Jia, Z. Yang and Y. Zhou, *Ceram Int*, DOI:10.1016/j.ceramint.2021.06.198.
- 67 J. Yin, N. Liu and Y. Gao, 2024, preprint, DOI: 10.15541/jim20230397.
- 68 R. Li, L. Zhang, L. Shi and P. Wang, ACS Nano, DOI:10.1021/acsnano.6b08415.
- 69 K. Wang, Y. Zhou, W. Xu, D. Huang, Z. Wang and M. Hong, *Ceram Int*, DOI:10.1016/j.ceramint.2016.02.059.
- O. Mashtalir, M. Naguib, V. N. Mochalin, Y. Dall'Agnese, M. Heon, M. W. Barsoum and Y. Gogotsi, *Nat Commun*, DOI:10.1038/ncomms2664.

- X. H. Zha, J. Yin, Y. Zhou, Q. Huang, K. Luo, J. Lang, J. S. Francisco, J. He and S. Du, Journal Vigo Article Online Onli
- H. Wang, Y. Wu, J. Zhang, G. Li, H. Huang, X. Zhang and Q. Jiang, *Mater Lett*, DOI:10.1016/j.matlet.2015.08.046.
- C. J. Zhang, B. Anasori, A. Seral-Ascaso, S. H. Park, N. McEvoy, A. Shmeliov, G. S. Duesberg, J.
 N. Coleman, Y. Gogotsi and V. Nicolosi, *Advanced Materials*, DOI:10.1002/adma.201702678.
- 74 X. Tang, X. Guo, W. Wu and G. Wang, 2018, preprint, DOI: 10.1002/aenm.201801897.
- 75 M. Ghidiu, S. Kota, V. Drozd and M. W. Barsoum, Sci Adv, DOI:10.1126/sciadv.aao6850.
- 76 E. S. Muckley, M. Naguib, H. W. Wang, L. Vlcek, N. C. Osti, R. L. Sacci, X. Sang, R. R. Unocic, Y. Xie, M. Tyagi, E. Mamontov, K. L. Page, P. R. C. Kent, J. Nanda and I. N. Ivanov, ACS Nano, DOI:10.1021/acsnano.7b05264.
- 77 E. S. Muckley, M. Naguib and I. N. Ivanov, Nanoscale, DOI:10.1039/c8nr05170d.
- J. Halim, S. Kota, M. R. Lukatskaya, M. Naguib, M. Q. Zhao, E. J. Moon, J. Pitock, J. Nanda, S. J. May, Y. Gogotsi and M. W. Barsoum, *Adv Funct Mater*, DOI:10.1002/adfm.201505328.
- 79 G. Ying, S. Kota, A. D. Dillon, A. T. Fafarman and M. W. Barsoum, *FlatChem*, DOI:10.1016/j.flatc.2018.03.001.
- M. Lu, H. Li, W. Han, J. Chen, W. Shi, J. Wang, X. M. Meng, J. Qi, H. Li, B. Zhang, W. Zhang and W. Zheng, *Journal of Energy Chemistry*, DOI:10.1016/j.jechem.2018.05.017.
- T. Voisin, M. D. Grapes, T. T. Li, M. K. Santala, Y. Zhang, J. P. Ligda, N. J. Lorenzo, B. E. Schuster, G. H. Campbell and T. P. Weihs, *Materials Today*, 2020, **33**, 10–16.
- Y. Ying, Y. Liu, X. Wang, Y. Mao, W. Cao, P. Hu and X. Peng, *ACS Appl Mater Interfaces*, DOI:10.1021/am5074722.
- A. Shahzad, K. Rasool, W. Miran, M. Nawaz, J. Jang, K. A. Mahmoud and D. S. Lee, *ACS Sustain Chem Eng*, DOI:10.1021/acssuschemeng.7b02695.
- M. Bilal, I. Ihsanullah, M. Younas and M. Ul Hassan Shah, 2022, preprint, DOI: 10.1016/j.seppur.2021.119510.
- 85 M. Sajid, 2021, preprint, DOI: 10.1016/j.aca.2020.08.063.
- 86 L. P. Yu, L. Lu, X. H. Zhou and L. Xu, 2023, preprint, DOI: 10.1002/admi.202201818.
- A. Maleki, M. Ghomi, N. Nikfarjam, M. Akbari, E. Sharifi, M. A. Shahbazi, M. Kermanian, M. Seyedhamzeh, E. Nazarzadeh Zare, M. Mehrali, O. Moradi, F. Sefat, V. Mattoli, P. Makvandi and Y. Chen, 2022, preprint, DOI: 10.1002/adfm.202203430.
- Z. Othman, H. R. Mackey and K. A. Mahmoud, *Chemosphere*, DOI:10.1016/j.chemosphere.2022.133849.
- 89 C. J. Zhang, S. Pinilla, N. McEvoy, C. P. Cullen, B. Anasori, E. Long, S. H. Park, A. Seral-Ascaso, A. Shmeliov, D. Krishnan, C. Morant, X. Liu, G. S. Duesberg, Y. Gogotsi and V. Nicolosi, *Chemistry of Materials*, DOI:10.1021/acs.chemmater.7b00745.

- 90 K. Rasool, R. P. Pandey, P. A. Rasheed, S. Buczek, Y. Gogotsi and K. A. Mahmoud, 2019 View Article Online preprint, DOI: 10.1016/j.mattod.2019.05.017.
- 91 S. P. Sreenilayam, I. Ul Ahad, V. Nicolosi and D. Brabazon, 2021, preprint, DOI: 10.1016/j.mattod.2020.10.025.
- 92 M. Soleymaniha, M. A. Shahbazi, A. R. Rafieerad, A. Maleki and A. Amiri, 2019, preprint, DOI: 10.1002/adhm.201801137.
- 93 A. M. Jastrzębska, A. Szuplewska, T. Wojciechowski, M. Chudy, W. Ziemkowska, L. Chlubny, A. Rozmysłowska and A. Olszyna, *J Hazard Mater*, DOI:10.1016/j.jhazmat.2017.06.004.
- 94 A. M. Jastrzębska, A. Szuplewska, A. Rozmysłowska-Wojciechowska, M. Chudy, A. Olszyna, M. Birowska, M. Popielski, J. A. Majewski, B. Scheibe, V. Natu and M. W. Barsoum, *2d Mater*, DOI:10.1088/2053-1583/ab6a60.
- 95 W. Wu, H. Ge, L. Zhang, X. Lei, Y. Yang, Y. Fu and H. Feng, *Chem Res Toxicol*, 2020, **33**, 2953–2962.
- 96 J. Wu, Y. Yu and G. Su, *Nanomaterials*, DOI:10.3390/nano12050828.
- 97 J. Yoon, S. Kim, K. H. Park, S. Lee, S. J. Kim, H. Lee, T. Oh and C. M. Koo, *Small Methods*, DOI:10.1002/smtd.202201579.
- 98 X. Wu, J. Gong, H. Zhang, Y. Wang and F. Tan, *Science of the Total Environment*, DOI:10.1016/j.scitotenv.2023.169227.
- 99 S. F. Hansen, M. B. Nielsen, L. M. Skjolding, J. Kaur, N. Desivyana, F. Hermansson, J. Bird, S. Barg, A. Khort, I. Odnevall, B. Fadeel and R. Arvidsson, *Sci Rep*, 2024, **14**, 31030.
- 100 G. K. Nasrallah, M. Al-Asmakh, K. Rasool and K. A. Mahmoud, *Environ Sci Nano*, DOI:10.1039/c7en01239j.
- 101 Q. Xiang, Z. Wang, J. Yan, M. Niu, W. Long, Z. Ju and X. Chang, *Aquatic Toxicology*, DOI:10.1016/j.aquatox.2024.106904.
- M. Dadashi Firouzjaei, S. K. Nemani, M. Sadrzadeh, E. K. Wujcik, M. Elliott and B. Anasori, *Advanced Materials*, DOI:10.1002/adma.202300422.
- 103 K. S. Novoselov, D. Jiang, F. Schedin, T. J. Booth, V. V. Khotkevich, S. V. Morozov and A. K. Geim, *Proc Natl Acad Sci U S A*, DOI:10.1073/pnas.0502848102.
- 104 M. S. Cao, Y. Z. Cai, P. He, J. C. Shu, W. Q. Cao and J. Yuan, 2019, preprint, DOI: 10.1016/j.cej.2018.11.051.
- 105 M. Naguib, M. Kurtoglu, V. Presser, J. Lu, J. Niu, M. Heon, L. Hultman, Y. Gogotsi and M. W. Barsoum, in *MXenes: From Discovery to Applications of Two-Dimensional Metal Carbides and Nitrides*, 2023.
- J. Liang, C. Ding, J. Liu, T. Chen, W. Peng, Y. Li, F. Zhang and X. Fan, *Nanoscale*, DOI:10.1039/c9nr02085c.
- 107 M. Naguib, J. Halim, J. Lu, K. M. Cook, L. Hultman, Y. Gogotsi and M. W. Barsoum, *J Am Chem Soc*, DOI:10.1021/ja405735d.

- 108 R. B. Rakhi, B. Ahmed, M. N. Hedhili, D. H. Anjum and H. N. Alshareef, *Chemistry of Material* Article Online DOI:10.1021/acs.chemmater.5b01623.
- Z. Sun, S. Li, R. Ahuja and J. M. Schneider, *Solid State Commun*, DOI:10.1016/j.ssc.2003.12.008.
- 110 X. Sang, Y. Xie, M. W. Lin, M. Alhabeb, K. L. Van Aken, Y. Gogotsi, P. R. C. Kent, K. Xiao and R. R. Unocic, *ACS Nano*, DOI:10.1021/acsnano.6b05240.
- 111 M. Ghidiu, M. R. Lukatskaya, M. Q. Zhao, Y. Gogotsi and M. W. Barsoum, *Nature*, DOI:10.1038/nature13970.
- 112 A. Lipatov and A. Sinitskii, in *2D Metal Carbides and Nitrides (MXenes): Structure, Properties and Applications*, 2019.
- J. Zhao, L. Zhang, X. Y. Xie, X. Li, Y. Ma, Q. Liu, W. H. Fang, X. Shi, G. Cui and X. Sun, *J Mater Chem A Mater*, DOI:10.1039/c8ta09840a.
- 114 M. Naguib, R. R. Unocic, B. L. Armstrong and J. Nanda, *Dalton Transactions*, DOI:10.1039/c5dt01247c.
- 115 I. R. Shein and A. L. Ivanovskii, *Comput Mater Sci*, DOI:10.1016/j.commatsci.2012.07.011.
- P. Urbankowski, B. Anasori, T. Makaryan, D. Er, S. Kota, P. L. Walsh, M. Zhao, V. B. Shenoy, M. W. Barsoum and Y. Gogotsi, *Nanoscale*, DOI:10.1039/c6nr02253g.
- T. Li, L. Yao, Q. Liu, J. Gu, R. Luo, J. Li, X. Yan, W. Wang, P. Liu, B. Chen, W. Zhang, W. Abbas, R. Naz and D. Zhang, *Angewandte Chemie International Edition*, DOI:10.1002/anie.201800887.
- 118 T. Li, X. Yan, L. Huang, J. Li, L. Yao, Q. Zhu, W. Wang, W. Abbas, R. Naz, J. Gu, Q. Liu, W. Zhang and D. Zhang, *J Mater Chem A Mater*, 2019, **7**, 14462–14465.
- S. Yang, P. Zhang, F. Wang, A. G. Ricciardulli, M. R. Lohe, P. W. M. Blom and X. Feng, Angewandte Chemie - International Edition, DOI:10.1002/anie.201809662.
- M. R. Lukatskaya, J. Halim, B. Dyatkin, M. Naguib, Y. S. Buranova, M. W. Barsoum and Y. Gogotsi, *Angewandte Chemie International Edition*, DOI:10.1002/anie.201403559.
- W. Sun, S. A. Shah, Y. Chen, Z. Tan, H. Gao, T. Habib, M. Radovic and M. J. Green, *J Mater Chem A Mater*, DOI:10.1039/c7ta05574a.
- 122 Z. Zhang, F. Zhang, H. Wang, C. Ho Chan, W. Lu and J. Y. Dai, J Mater Chem C Mater, DOI:10.1039/c7tc03652c.
- 123 X. Xiao, H. Yu, H. Jin, M. Wu, Y. Fang, J. Sun, Z. Hu, T. Li, J. Wu, L. Huang, Y. Gogotsi and J. Zhou, *ACS Nano*, DOI:10.1021/acsnano.6b08534.
- J. Jia, T. Xiong, L. Zhao, F. Wang, H. Liu, R. Hu, J. Zhou, W. Zhou and S. Chen, *ACS Nano*, DOI:10.1021/acsnano.7b06607.
- 125 C. Xu, L. Wang, Z. Liu, L. Chen, J. Guo, N. Kang, X. L. Ma, H. M. Cheng and W. Ren, *Nat Mater*, DOI:10.1038/nmat4374.
- 126 Y. Gogotsi, *Nat Mater*, DOI:10.1038/nmat4386.

- Y. Yan, H. Han, Y. Dai, H. Zhu, W. Liu, X. Tang, W. Gan and H. Li, *ACS Appl Nano Mater* View Article Online DOI:10.1021/acsanm.1c02339.
- 128 C. Hao, G. Li, G. Wang, W. Chen and S. Wang, *Colloids Surf A Physicochem Eng Asp*, DOI:10.1016/j.colsurfa.2021.127730.
- 129 Y. Lei, Y. Cui, Q. Huang, J. Dou, D. Gan, F. Deng, M. Liu, X. Li, X. Zhang and Y. Wei, *Ceram Int*, DOI:10.1016/j.ceramint.2019.05.331.
- P. Najibikhah and A. Rahbar-Kelishami, *Chemosphere*, DOI:10.1016/j.chemosphere.2023.141058.
- O. Mashtalir, K. M. Cook, V. N. Mochalin, M. Crowe, M. W. Barsoum and Y. Gogotsi, *J Mater Chem A Mater*, DOI:10.1039/c4ta02638a.
- Z. Wei, Z. Peigen, T. Wubian, Q. Xia, Z. Yamei and S. ZhengMing, *Mater Chem Phys*, DOI:10.1016/j.matchemphys.2017.12.034.
- 133 C. Peng, P. Wei, X. Chen, Y. Zhang, F. Zhu, Y. Cao, H. Wang, H. Yu and F. Peng, *Ceram Int*, DOI:10.1016/j.ceramint.2018.07.124.
- 134 M. Vakili, G. Cagnetta, J. Huang, G. Yu and J. Yuan, *Molecules*, DOI:10.3390/molecules24132478.
- L. Zhang, D. Huang, P. Zhao, G. Yue, L. Yang and W. Dan, *Sep Purif Technol*, DOI:10.1016/j.seppur.2022.120718.
- 136 W. Li, J. Tong and G. Li, *Chemosphere*, 2024, **360**, 142376.
- 137 K. Li, G. Zou, T. Jiao, R. Xing, L. Zhang, J. Zhou, Q. Zhang and Q. Peng, *Colloids Surf A Physicochem Eng Asp*, DOI:10.1016/j.colsurfa.2018.05.044.
- 138 Y. Li, G. Qu, H. Zhang, L. Xie and Y.-F. Zhang, *Chem Eng Sci*, 2024, **300**, 120648.
- Y. Xu, Z. Zhang, Z. Cui, L. Luo, P. Lin, M. Xie, Q. Zhang, B. Sa and C. Wen, *Chemical Engineering Journal*, 2024, **488**, 151078.
- L. Li, X. Y. Shi, T. Huang, N. Zhang and Y. Wang, *J Mater Chem A Mater*, DOI:10.1039/d3ta04869a.
- 141 R. Wang, H. Cao, C. Yao, C. Peng, J. Qiu, K. Dou, N. Tsidaeva and W. Wang, *Appl Surf Sci*, DOI:10.1016/j.apsusc.2023.157091.
- 142 Z. Zhu, M. Xiang, L. Shan, T. He and P. Zhang, *J Solid State Chem*, DOI:10.1016/j.jssc.2019.120989.
- 143 G. S. Elgarhy, G. M. El-Subruiti, A. M. Omer and A. S. Eltaweil, *J Mol Liq*, DOI:10.1016/j.molliq.2023.123889.
- 144 Y. Zheng, H. Zhang, S. Yu, H. Zhou, W. Chen and J. Yang, *Sep Purif Technol*, 2024, **349**, 127908.
- 145 R. Wang, C. Yao, C. Peng, J. Qiu, Q. Wang, X. Liu, J. Meng and W. Wang, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2024.149916.
- 146 X. Wang, A. Zhang, M. Chen, M. K. Seliem, M. Mobarak, Z. Diao and Z. Li, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2023.145385.

- H. Yang, J. Lin, Z. Wen, Z. Li, J. Zeng, L. Wang, Y. Tao, D. Gao and D. Wang, *Chemosphere* View Article Online DOI: 10.1016/j.chemosphere.2023.139797.
- 148 A. Bukhari, I. Ijaz, A. Nazir, S. Hussain, H. Zain, E. Gilani, A. A. Ifseisi and H. Ahmad, *RSC Adv*, 2024, **14**, 3732–3747.
- 149 I. Ijaz, A. Bukhari, E. Gilani, A. Nazir, H. Zain, A. Bukhari, A. Shaheen, S. Hussain and A. Imtiaz, *Process Biochemistry*, DOI:10.1016/j.procbio.2023.03.029.
- 150 R. Imsong and D. Dhar Purkayastha, Sep Purif Technol, DOI:10.1016/j.seppur.2022.122636.
- 151 M. Li, P. Zhang, Q. Wang, N. Yu, X. Zhang and S. Su, *Polymers (Basel)*, DOI:10.3390/polym15092110.
- 152 Y. Feng, H. Wang, J. Xu, X. Du, X. Cheng, Z. Du and H. Wang, *J Hazard Mater*, DOI:10.1016/j.jhazmat.2021.125777.
- 153 Q. Wang, J. Qiao, Y. Xiong, F. Dong and Y. Xiong, *Environ Res*, DOI:10.1016/j.envres.2023.117568.
- 154 G. Song, W. Fan, J. Zhang, T. Xue, Y. Shi, Y. Sun and G. Ding, *Appl Surf Sci*, 2024, **661**, 160036.
- S. yu Chen, Y. fan Deng, T. Huang, N. Zhang and Y. Wang, *Sep Purif Technol*, DOI:10.1016/j.seppur.2023.125040.
- T. Lv, F. Wu, Z. Zhang, Z. Liu, Y. Zhao, L. Yu, J. Zhang, C. Yu, C. Zhao and G. Xing, *ACS Appl Nano Mater*, DOI:10.1021/acsanm.4c00038.
- 157 X. Wang, Q. Xu, L. Zhang, L. Pei, H. Xue and Z. Li, *J Environ Chem Eng*, DOI:10.1016/j.jece.2022.109206.
- H. Xue, X. Gao, M. K. Seliem, M. Mobarak, R. Dong, X. Wang, K. Fu, Q. Li and Z. Li, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2022.138735.
- 159 W. Zhao, H. Chi, S. Zhang, X. Zhang and T. Li, Molecules, DOI:10.3390/molecules27134243.
- 160 M. Khazaei, M. Arai, T. Sasaki, C. Y. Chung, N. S. Venkataramanan, M. Estili, Y. Sakka and Y. Kawazoe, *Adv Funct Mater*, DOI:10.1002/adfm.201202502.
- 161 M. A. Hope, A. C. Forse, K. J. Griffith, M. R. Lukatskaya, M. Ghidiu, Y. Gogotsi and C. P. Grey, *Physical Chemistry Chemical Physics*, 2016, **18**, 5099–5102.
- M. R. Lukatskaya, O. Mashtalir, C. E. Ren, Y. Dall'Agnese, P. Rozier, P. L. Taberna, M. Naguib, P. Simon, M. W. Barsoum and Y. Gogotsi, *Science* (1979), DOI:10.1126/science.1241488.
- I. Persson, L. Å. Näslund, J. Halim, M. W. Barsoum, V. Darakchieva, J. Palisaitis, J. Rosen and P. O. Å. Persson, 2d Mater, DOI:10.1088/2053-1583/aa89cd.
- A. Abdul Ghani, B. Kim, M. Nawaz, K. C. Devarayapalli, Y. Lim, G. Kim and D. S. Lee, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2023.143473.
- A. A. Ghani, A. Shahzad, M. Moztahida, K. Tahir, H. Jeon, B. Kim and D. S. Lee, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2020.127780.
- 166 M. Xu, C. Huang, J. Lu, Z. Wu, X. Zhu, H. Li, L. Xiao and Z. Luo, *Molecules*, DOI:10.3390/molecules26113150.

- 167 M. Szlachta and P. Wójtowicz, Water Science and Technology, DOI:10.2166/wst.2013.487/iew Article Online
- T. Lv, J. Zhang, L. Yu, Y. Zhao, T. Zhao, Y. Yang, C. Yu, C. Zhao and G. Xing, *Journal of Industrial and Engineering Chemistry*, 2025, **144**, 454–462.
- 169 R. Li, X. Tang, J. Wu, K. Zhang, Q. Zhang, J. Wang, J. Zheng, S. Zheng, J. Fan, W. Zhang, X. Li and S. Cai, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2023.142706.
- 170 Y. Hu, H. Wang, X. Ren, F. Wu, G. Liu, S. Zhang, H. Luo and L. Fang, *Nanomaterials*, 2024, **14**, 1925.
- J. J. Gao, Y. B. Qin, T. Zhou, D. D. Cao, P. Xu, D. Hochstetter and Y. F. Wang, *J Zhejiang Univ Sci B*, DOI:10.1631/jzus.B12a0225.
- 172 H. M. El-Bery, M. Saleh, R. A. El-Gendy, M. R. Saleh and S. M. Thabet, *Sci Rep*, DOI:10.1038/s41598-022-09475-4.
- 173 K. A. Mohamad Said, M. A. M. Amin, I. Yakub, M. R. Rahman, A. B. H. Kueh, S. Hamdan and M. M. Rahman, *Bioresources*, DOI:10.15376/biores.18.3.5120-5132.
- 174 Z. Li, H. Hanafy, L. Zhang, L. Sellaoui, M. Schadeck Netto, M. L. S. Oliveira, M. K. Seliem, G. Luiz Dotto, A. Bonilla-Petriciolet and Q. Li, *Chemical Engineering Journal*, DOI:10.1016/j.cej.2020.124263.
- 175 N. N. Marnani, F. H. Tezel and O. D. Basu, *Applied Sciences*, 2024, **14**, 9811.
- N. A. Travlou, G. Z. Kyzas, N. K. Lazaridis and E. A. Deliyanni, *Langmuir*, DOI:10.1021/la304696y.
- T. Madrakian, A. Afkhami, M. Ahmadi and H. Bagheri, *J Hazard Mater*, DOI:10.1016/j.jhazmat.2011.08.078.
- T. Aysu and M. M. Küçük, *International Journal of Environmental Science and Technology*, DOI:10.1007/s13762-014-0623-y.
- B. M. Jun, J. Heo, N. Taheri-Qazvini, C. M. Park and Y. Yoon, *Ceram Int*, DOI:10.1016/j.ceramint.2019.09.293.
- 180 E. Ahmed, A. Deep, E. E. Kwon, R. J. C. Brown and K. H. Kim, *Sci Rep*, DOI:10.1038/srep31283.
- 181 X. Liu, H. Pang, X. Liu, Q. Li, N. Zhang, L. Mao, M. Qiu, B. Hu, H. Yang and X. Wang, 2021, preprint, DOI: 10.1016/j.xinn.2021.100076.
- 182 I. Ihsanullah, 2020, preprint, DOI: 10.1016/j.cej.2020.124340.
- 183 M. Alhabeb, K. Maleski, B. Anasori, P. Lelyukh, L. Clark, S. Sin and Y. Gogotsi, *Chemistry of Materials*, DOI:10.1021/acs.chemmater.7b02847.
- N. Bhalla, *Advanced Physics Research*, DOI:10.1002/apxr.202400205.
- 185 K. T. Butler, D. W. Davies, H. Cartwright, O. Isayev and A. Walsh, 2018, preprint, DOI: 10.1038/s41586-018-0337-2.
- 186 G. R. Schleder, A. C. M. Padilha, C. M. Acosta, M. Costa and A. Fazzio, *JPhys Materials*, DOI:10.1088/2515-7639/ab084b.

- 187 M. Zhong, K. Tran, Y. Min, C. Wang, Z. Wang, C. T. Dinh, P. De Luna, Z. Yu, A. S. Rasouli, P. View Article Online Brodersen, S. Sun, O. Voznyy, C. S. Tan, M. Askerka, F. Che, M. Liu, A. Seifitokaldani, Y. Pang, S. C. Lo, A. Ip, Z. Ulissi and E. H. Sargent, *Nature*, DOI:10.1038/s41586-020-2242-8.
- 188 B. Sanchez-Lengeling and A. Aspuru-Guzik, 2018, preprint, DOI: 10.1126/science.aat2663.
- 189 G. Pilania, A. Mannodi-Kanakkithodi, B. P. Uberuaga, R. Ramprasad, J. E. Gubernatis and T. Lookman, *Sci Rep*, DOI:10.1038/srep19375.
- 190 C. T. Chen and G. X. Gu, MRS Commun, DOI:10.1557/mrc.2019.32.
- 191 M. Hamadache, J. H. Jung, J. Park and B. D. Youn, *JMST Advances*, DOI:10.1007/s42791-019-0016-y.
- 192 M. Grieves, White Paper.
- 193 B. L. Decost and E. A. Holm, Comput Mater Sci, DOI:10.1016/j.commatsci.2015.08.011.
- 194 S. Hochreiter and J. Schmidhuber, *Neural Comput*, DOI:10.1162/neco.1997.9.8.1735.
- 195 S. Li, L. Da Xu and S. Zhao, 2018, preprint, DOI: 10.1016/j.jii.2018.01.005.
- 196 L. Da Xu, W. He and S. Li, 2014, preprint, DOI: 10.1109/TII.2014.2300753.
- 197 K. Deb, A. Pratap, S. Agarwal and T. Meyarivan, *IEEE Transactions on Evolutionary Computation*, DOI:10.1109/4235.996017.
- 198 J. M. Granda, L. Donina, V. Dragone, D. L. Long and L. Cronin, *Nature*, DOI:10.1038/s41586-018-0307-8.
- 199 A. Dey, M. Billinghurst, R. W. Lindeman and J. E. Swan, *Frontiers Robotics AI*, DOI:10.3389/frobt.2018.00037.

Nanoscale Accepted Manuscript

View Article Online DOI: 10.1039/D5NR02336J

Data Availability Statement

No new data were created in this study