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Three-dimensional bimodal pore-rich G/MXene sponge amalgamated with vanadium diselenide nanosheets as a high-performance electrode for electrochemical water-oxidation/reduction reactions†

Khadija Chaudhary, Da Sonia Zulfiqar, Db,c Zeid A. ALOthman, Dd Imran Shakir, Muhammad Faroog Warsi Da and Eric W. Cochran Da c

Exploring new strategies to design non-precious and efficient electrocatalysts can provide a solution for sluggish electrocatalytic kinetics and sustainable hydrogen energy. Transition metal selenides are potential contenders for bifunctional electrocatalysis owing to their unique layered structure, low band gap, and high intrinsic activities. However, insufficient access to active sites, lethargic water dissociation, and structural degradation of active materials during electrochemical reactions limit their activities, especially in alkaline media. In this article, we report a useful strategy to assemble vanadium diselenide (VSe₂) into a 3D MXene/rGO-based sponge-like architecture (VSe₂@G/MXe) using hydrothermal and freeze-drying approaches. The 3D hierarchical meso/macro-pore rich sponge-like morphology not only prevents aggregation of VSe₂ nanosheets but also offers a kinetics-favorable framework and high robustness to the electrocatalyst. Synergistic coupling of VSe₂ and a MXene/rGO matrix yields a heterostructure with a large specific surface area, high conductivity, and multi-dimensional anisotropic pore channels for uninterrupted mass transport and gas diffusion. Consequently, VSe₂@G/MXe presented superior electrochemical activity for both the HER and OER compared to its counterparts (VSe2 and VSe2@G), in alkaline media. The overpotentials required to reach a cathodic and anodic current density of 10 mA cm⁻² were 153 mV (Tafel slope = 84 mV dec⁻¹) and 241 mV (Tafel slope = 87 mV dec⁻¹), respectively. The R_{ct} values at the open circuit voltage were as low as 9.1 Ω and 1.41 Ω for the HER and OER activity, respectively. Importantly, VSe₂@G/MXe withstands a steady current output for a long 24 h operating time. Hence, this work presents a rational design for 3D microstructures with optimum characteristics for efficient bifunctional alkaline water-splitting.

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† Electronic supplementary information (ESI) available: EDX spectra of (a) VSe $_2$, (b) VSe $_2$ @G, and (c) VSe $_2$ @G/MXe (Fig. S1), XRD patterns of MXene and rGO (Fig. S2), I–V profiles of VSe $_2$, VSe $_2$ @G, and VSe $_2$ @G/MXe (Fig. S3), LSV curve of G/MXe for the HER (Fig. S4), LSV curve of G/MXe for the OER (Fig. S5), $C_{\rm cl}$ results for VSe $_2$, VSe $_2$ @G, and VSe $_2$ @G/MXe (Fig. S6), ECAS-normalized LSV curves of VSe $_2$ and VSe $_2$ @G/MXe for the HER (Fig. S7), ECAS-normalized LSV curves of VSe $_2$ and VSe $_2$ @G/MXe for the OER (Fig. S8), comparison of the HER performance of VSe $_2$ @G/MXe with that of similar reported electrocatalytic materials (Table S1), and comparison of the OER performance of VSe $_2$ @G/MXe with that of similar reported electrocatalytic materials (Table S2). See DOI: https://doi.org/10.1039/d4dt00602j

1. Introduction

The global energy crisis, as a consequence of the shortage of fossil fuel reserves and the increase in fuel costs, is one of the most prevailing issues affecting various sectors of the world's economies. In addition, global campaigns against CO₂ emission are putting pressure on different countries to limit their use of fossil fuels to combat climate change. Consequently, the issue of energy security should be at the forefront of each country's policy for a secure and self-sufficient future. For this, we need alternative, low-cost, clean energy resources. In this context, hydrogen fuel is a clean and efficient source of energy because of its high gravimetric energy density and non-harmful by-product (*i.e.*, H₂O). Though hydrogen is considered a green fuel, 95% of the

^aInstitute of Chemistry, Baghdad-ul-Jadeed Campus, The Islamia University of Bahawalpur, Bahawalpur, 63100, Pakistan. E-mail: farooq.warsi@iub.edu.pk ^bDepartment of Chemistry, Faculty of Science, University of Ostrava, 30. Dubna 22, Ostrava 701 03, Czech Republic

^cDepartment of Chemical and Biological Engineering, Iowa State University, Sweeney Hall, 618 Bissell Road, Ames, Iowa 50011, USA. E-mail: ecochran@iastate.edu ^dDepartment of Chemistry, College of Science, King Saud University, P. O. Box 2455, Riyadh 11451, Saudi Arabia

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production of hydrogen fuel itself consumes carbon-based raw materials, mostly coming from fossil fuels.^{6,7} On the other hand, the production of hydrogen from water electrolysis through the hydrogen evolution reaction (HER) includes the CO₂ neutral reactant.⁸ Besides, the second half reaction during water electrolysis, i.e., oxygen evolution reaction (OER), results in the formation of the by-product oxygen which is an important industrial gas and finds applications in combustion, semiconductor production, and wastewater treatment. 9,10 The state-of-the-art electrocatalysts applied in water electrolysers are RuO2, IrO2, and Pt.11 Due to the scarcity and high prices of these noble metal-based electrocatalysts, hydrogen production from water electrolysis is accompanied by high production costs that hinder its market penetration. In this regard, several attempts have been made in recent years to design inexpensive Earth-abundant transition metal based electrocatalysts for the HER, OER, and overall water-splitting.

Driven by their high electrical conductivity and redox activity, transition metal selenides (TMSes) such as NiSe₂, ¹² Ni_{0.85}Se, ¹³ Co₃Se₄, ¹⁴ FeSe₂, ¹⁵ MoSe, ¹⁶ and Cu₂Se¹⁷ have been widely explored for water-splitting applications. The selenium atom in TMSes may bind with other metal atoms due to the close energy level of the 3d orbital with those of 3s and 3p. 18 This unique electronic structure induces an enhanced metallic characteristic in TMSes that is beneficial for electron transport and the occurrence of redox reactions. 19 Apart from the abovementioned TMSe electrocatalysts, 2D vanadium diselenide (VSe₂) with a group-five transition metal (with an enhanced metallic characteristic) shows a very high electrical conductivity (~1000 S m⁻¹ at 300 K).²⁰ Considering the inherent electrochemistry intrinsic to transition metal (V) and chalcogen (Se), VSe₂ has been predicted to have strong potential for electrochemical applications.21 In addition, the unique 2D layered chalcogenide-like (MX2, M = metal and X = S, Se, and Te) structure of VSe₂ can offer it high concentration of active sites desirable for efficient catalytic activity.²² Lately, VSe₂ nanosheets prepared by the colloidal route showed impressive HER activity with a very low overpotential of 206 mV at a current density of 10 mA cm⁻².²³ In a similar report, Yan and co-workers reported sufficient potential of VSe2 nanosheets for HER activity.²⁴ Most recently, Liu et al. studied the efficiency of hydrothermally synthesized layered VSe2 for overall water splitting.25 The as-synthesized electrocatalyst revealed an optimal electrocatalytic potential for the HER and OER activity with overpotentials of 1.011 V and 2.467 V at 10 mA cm⁻², respectively. However, the problem with 2D layered structures is the interlayer interactions that stimulate restacking among layers and conceal the active sites.²⁶ Furthermore, a high electron density of unoccupied d-orbitals of vanadium in VSe2 suggests a high Gibbs free energy ($\Delta G_{\rm H}$) for hydrogen adsorption (large Heyrovsky and Tafel steps), which leads to sluggish reaction kinetics.20 Constructing heterostructures of TMSes with carbon materials can be a useful approach to overcome the stacking issues. The synergistic interactions among the pristine TMSe and second phase optimize ΔG_H for hydrogen adsorption, thus facilitating the catalytic process.²⁷

MXenes $(M_{n+1}X_nT_r, M = \text{transition metal}, X_n = C \text{ and } N, \text{ and}$ T_x = surface terminal groups) are a new family of 2D transition metal carbides, nitrides, or carbonitrides that have promising potential to be used as hybridizing matrices.²⁸ MXenes possess high metallic conductivity and hydrophilicity that are seldom realized by other hybridizing materials.²⁹ Reports on MXene composites such as CoS-MXene, FeS2-MXene, 1 CuS-Cl terminated MXene, 32 CoSe₂@MXene, 33 and NiCo₂Se₄-MXene³⁴ have shown significant improvement in the electrocatalytic activity. Recently, the focus has been shifted to designing 3D architectures of MXenes (hydrogel, foams, and sponges). 35,36 Despite many recent reports on 3D MXenes, their composite systems still need to be explored. To the best of our knowledge, the combination of 2D layered MX₂ with a 3D MXene system has not yet been reported.

In view of the above discussion, herein, we report a novel electrocatalyst (VSe₂@G/MXe) where 2D VSe₂ nanosheets are assembled into a Ti₃C₂T_x MXene (hereafter represented as MXene) based 3D sponge. Due to the stiff nature of MXene sheets, graphene oxide (GO) is used as a gelation agent to construct a 3D MXene sponge (henceforth represented as G/MXe). The 3D VSe₂@G/MXe sponge-like hybrid exhibits outstanding performance towards both the HER and OER. rGO not only served as the gelation agent to assemble the VSe₂@G/MXe monolith but also provided a high specific surface area to the electrocatalyst. The high conductivity of the MXene combined with rGO provided fast charge transport. The active sites of VSe₂ amalgamated into the highly porous G/MXe sponge were easily accessible to the electrolyte for electrochemical activity. Intriguingly, VSe₂@G/MXe exhibited a significant improvement in the overpotential for the HER (153 mV@10 mA cm⁻²) and the OER (241 mV@10 mA cm $^{-2}$) compared with VSe₂ and VSe₂@G. Consequently, this study provides insight into the development of MXene-based 3D composite materials as a useful strategy to improve the energy conversion performance of electrocatalysts.

Experimental section 2.

Chemicals

Ammonium metavanadate (NH₄VO₃; 99.99%), selenium dioxide (SeO₂; 999.99%), formic acid (HCOOH; \geq 99%), MAX powder (Ti₃AlC₂; ≥90%, ≤200 μm particle size), and hydrofluoric acid (HF; 40%) were purchased from Sigma Aldrich. All chemicals were used as procured. Nickel foam (NF; 150 × 150 mm and thickness 1.6 mm) from Sigma Aldrich was used as a substrate to prepare electrodes. Distilled water from a selfregulating plant was used throughout the experimental work.

2.2. Synthesis of vanadium diselenide (VSe₂)

For the synthesis of VSe₂, NH₄VO₃ (118.4 mg) and SeO₂ (220.9 mg) were added to 40 mL of distilled water and stirred for 5 min. 5 mL of HCOOH was added to the above stirring solution in a dropwise manner, and the mixture was allowed to stir for further 5 min. Next, the solution was transferred to a

50 mL autoclave and incubated at 220 °C for 24 h. After completion of the reaction, the autoclave was naturally cooled down and precipitates were separated through ultracentrifugation. The precipitates were washed with ethanol and water to remove unreacted salts and dried at 60 °C overnight to obtain

VSe₂ powder. The formation of VSe₂ from the precursors can be represented by the following eqn (1)–(3):²⁵

$$VO^{-3} + HCOOH + 2H^{+} \rightarrow VO^{+2} + CO_{2} + 2H_{2}O$$
 (1)

$$SeO_2 + HCOOH + 2H^+ \rightarrow H_2Se + CO_2 + 2H_2O$$
 (2)

$$VO^{+2} + 2H_2Se \rightarrow VSe_2 + 2H^+ + H_2O$$
 (3)

2.3. Synthesis of VSe₂@G/MXe

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For the synthesis of VSe₂@G/MXe, first, GO and MXene were prepared. Synthesis of GO was carried out by the Hummers method reported elsewhere.37 To synthesize MXene, 1 g of MAX powder was slowly added to 20 mL of HF solution with continuous stirring. The reaction mixture was left to stir at room temperature for 48 h. Next, the HF-etched MXene was washed with distilled water to neutralize MXene particles. Exfoliation of the MXene was carried out in distilled water under ultrasonication conditions. Finally, the MXene was dried at 60 °C to obtain MXene powder. Next, a suspension of VSe₂ was ultrasonically prepared by dispersing VSe₂ (20 mg) in 10 mL of distilled water. A second suspension of GO: MXene (20:10 mg) was also prepared ultrasonically in 10 mL of distilled water. VSe₂ suspension prepared during the first step was added to the GO: MXene suspension and the mixture was further treated under ultrasonication conditions for 30 min to obtain a homogeneous solution. Next, 20 mg of ascorbic acid was added to the obtained suspension, and the mixture was processed hydrothermally at 70 °C for 4 h. The as-obtained monolith was freeze-dried at -90 °C to obtain VSe₂@G/MXe. For comparison, VSe₂@G was also prepared by following the same procedure without the addition of the MXene.

2.4. Characterization

A LabX XRD-6100 diffractometer (CuK α , $\lambda = 0.154$ nm) was used to elucidate the crystalline structure and composition of the prepared materials. FT-IR spectra were recorded on a Shimadzu FT-IR affinity/1S. The morphology was recorded using a ZEISS LEO SUPRA 55 field emission scanning electron microscope. A JEOL JCM-6000 Plus SEM instrument was used to collect the elemental data. A Micromeritics ASAP 2020 physisorption analyzer was used to determine the surface properties of electrocatalysts. A Keithley voltage source 6487/E was used to investigate the electrical behavior. The electrochemical water-splitting application study was carried out on a GAMRY Potentiostat interface 5000/E.

2.5. Water-splitting measurements

For electrochemical water-splitting, a slurry was prepared from the synthesized electrocatalysts. 10 mg of each electrocatalyst (VSe₂, VSe₂@G, and VSe₂@G/MXe) was mixed with Nafion

binder and ethanol and ultrasonicated for 5 min. The slurry was deposited on Ni foam (fixed area = $1 \times 1 \text{ cm}^2$) and dried at room temperature. The prepared electrodes were used as working electrodes in a three-electrode workstation. Platinum wire and silver/silver chloride were applied as the auxiliary and reference electrodes, respectively. 1 M KOH was used as the electrolyte medium to collect linear sweep voltammetry (LSV), cyclic voltammetry (CV), impedance, and chronoamperometry data. LSV and CV curves were recorded at 3 mV s⁻¹ and 20-120 mV s⁻¹, respectively. Impedance or EIS measurements were conducted at an AC voltage of 5 mV with initial and final frequencies of 0.1 Hz and 100 kHz, respectively. Chronoamperometry stability tests for the HER and the OER were performed for 24 h at a fixed potential. In addition, all the potential (E) values were referenced to the reversible hydrogen electrode (RHE) using the following eqn (4):

$$E(RHE) = E(Ag/AgCl) + 0.197 + (0.0591 \times pH)$$
 (4)

3. Results and discussion

3.1. Microstructure assembly and morphological and compositional analyses

Fig. 1 schematically illustrates the synthesis of the VSe₂@G/ MXe microstructure. For hydrothermal assembly, the $Ti_3C_2T_x$ MXene was obtained through acidic etching of the Al layer from Ti₃AlC₂ MAX powder. First, the MXene suspension was treated with GO ultrasonically in the presence of the hydrothermally synthesized VSe2. Due to the surface-terminated polar groups, MXene sheets get attracted and stick to the surface of GO sheets. Next, when the ultrasonically treated homogeneous suspension was treated under the hydrothermal conditions in the presence of ascorbic acid, it resulted in the generation of π - π interactions leading to the hydrophobic effect among the macroscopic rGO sheets. As the hydrophobic interactions dominate the hydrophilic interactions at the rGO interface, the partially reduced rGO with attached MXene sheets undertook phase separation into a 3D monolith that is then freeze-dried into a cross-linked porous microstructure.

FESEM analysis was used for a comprehensive structural and morphological study of the prepared electrocatalysts. The HF-etched MXene in Fig. 2(a) shows a few-layer accordion-like structure with an obvious interlayer spacing, whereas the FESEM image of bare VSe₂, shown in Fig. 2(b), shows the existence of thin VSe2 nanosheets that stack on each other to form quasi-lamellar structures with a relatively smooth surface. FESEM images in Fig. 2(d-f) and (g-i) display different sectional views of VSe₂@G and VSe₂@G/MXe microstructures, respectively. A pore-rich 3D sponge-like structure can be observed for VSe₂@G and VSe₂@G/MXe. Fig. 2(d and g) shows the preferred direction of growth of ice crystals for the prepared microstructures. Certainly, during the freeze-drying process, cross-linked walls of VSe₂@G and VSe₂@G/MXe monoliths under the action of the ice template are repelled by the growing ice crystals leading to the accumulation of cell

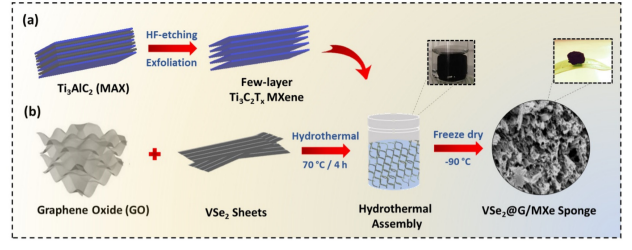


Fig. 1 Schematic illustrations of (a) HF-etching of the MAX phase and (b) hydrothermal assembly of VSe₂, MXene, and rGO into VSe₂@G/MXe.

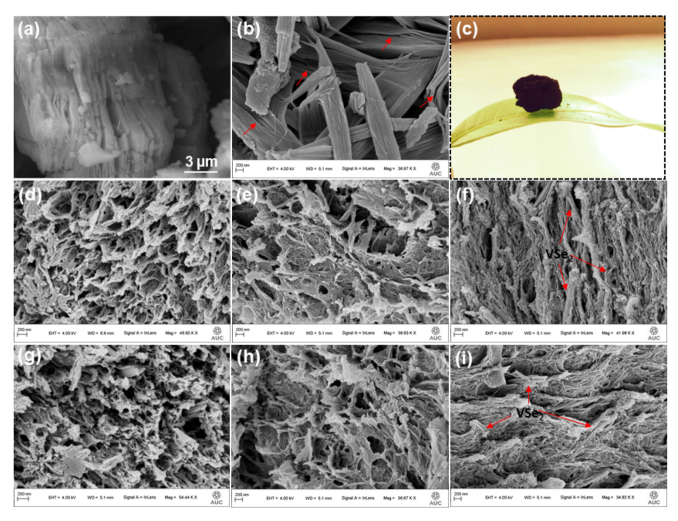


Fig. 2 FESEM images of (a) MXene and (b) VSe₂ nanosheets, and (c) photograph of VSe₂@G/MXe on a leaf; different sectional view images of (d-f) VSe₂@G and (g-i) VSe₂@G/MXe.

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walls parallel to the ice template.³⁸ The cross-section views in Fig. 2(e and h) confirm that the pores generated as a result of the cross-linking and the ice-templating effect cumulate in anisotropic porous channels within VSe₂@G and VSe₂@G/MXe. Meanwhile, VSe₂ nanosheets simultaneously intervene in rGO and rGO/MXene sponge, in their respective microstructures (side view in Fig. 2(f and i)). This integration of VSe₂ nanosheets into the cross-linked 3D microstructure restricts the aggregation of VSe₂ nanosheets exposing the maximum active sites for the electrochemical process. Fig. 2(c) shows a photograph of the 3D VSe₂@G/MXe microstructure on a leaf which confirms its ultralight density due to its pore-rich interior.

In addition, to gain insight into the elemental compositions, the EDX spectra of VSe₂, VSe₂@G, and VSe₂@G/MXe were recorded and are shown in Fig. S1(a-c).† The EDX profiles of all electrocatalysts exhibit signals from V and Se along with other elements, i.e., C, O, and Ti, in the corresponding electrocatalyst. The appearance of peaks from all conforming elements (V, Se, Ti, C, and O) in VSe₂@G/MXe confirms the successful assembly of VSe2, MXene, and rGO into a 3D microstructure.

3.2. Phase analysis

XRD analysis was carried out to evaluate the phase purity and crystal structure of the synthesized samples. Multiple diffraction peaks of the MXene (Fig. S2†) at 8.4°, 17.8°, 26.7°, 36.1°, 39.7°, and 60.3° can be indexed to the (002), (004), (006), (103), (104), and (110) planes of the Ti₃C₂T_x MXene phase (JCPDS card: 52-0875), respectively.³⁹ Fig. 3(a) shows the XRD patterns of VSe2, VSe2@G and VSe2@G/MXe. For VSe2, the peaks observed at 14.6°, 29°, 30.2°, 33.9°, 43.1°, 53.9°, and 54.8° are attributed to the (001), (002), (100), (011), (102), (110), and (103) planes of the hexagonal crystalline phase of VSe₂ (JCPDS:

01-089-1641), respectively. 40 The XRD patterns of VSe₂@G show an additional distinctive broad bump at around 23.9° that corresponds to the (002) plane of rGO (Fig. S2†) in addition to the peaks of VSe₂. This broad bump at 23.9° corresponds well to hydrothermally reduced rGO previously reported in the literature. 41 Likewise, for VSe₂@G/MXe, in addition to VSe₂ and rGO, two new peaks appeared from the (002) and (004) planes of the MXene. Noticeably, the (002) peak of the MXene gets shifted from 8.4° to 7.7° in VSe₂@G/MXe. This shift is presumably related to the expansion in the interlayer spacing of the MXene from 1.05 nm to 1.14 nm, after its incorporation into the rGO based sponge. 42 Furthermore, the absence of any extra peak from impurity confirms the phase purity of the prepared samples.

3.3. Functional group analysis

Different vibrational modes associated with functional groups present in VSe2, VSe2@G, and VSe2@G/MXe were examined by FT-IR analysis, as shown in Fig. 3(b). The FT-IR spectrum of VSe₂ exhibits two IR bands in the fingerprint region at 585 cm⁻¹ and 742 cm⁻¹ corresponding to different vibrational modes of the V-Se bond. 43 A broad band at around 3461 cm⁻¹ is contributed by the symmetric and asymmetric stretching vibrations of the O-H group associated with physically adsorbed water on the surface of VSe2.44 In contrast to the spectrum of VSe₂, the FT-IR spectra of VSe₂@G and VSe₂@G/ MXe exhibit several new bands in the ~1000-2000 region, corresponding to the functional groups of rGO and the MXene. Bands that appeared at around 1392 cm⁻¹ and 1569 cm⁻¹ belong to C-H bond bending and C=C bond stretching vibrations from graphitic domains of rGO, respectively. 45 The characteristic bands at 1005 cm⁻¹, 1222 cm⁻¹, and 1685 cm⁻¹ are adhered by the oxygen containing functional groups in VSe₂@G and VSe₂@G/MXe microstructures.

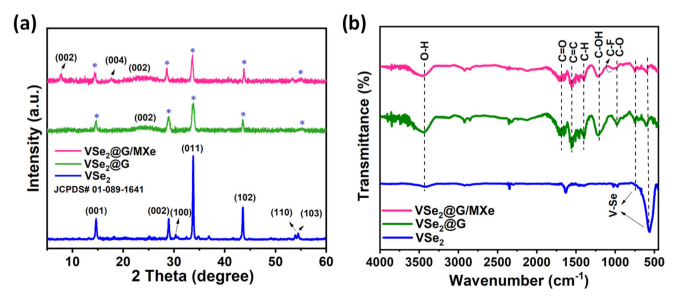


Fig. 3 (a) XRD patterns and (b) FT-IR spectra of VSe₂, VSe₂@G, and VSe₂@G/MXe.

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Specifically, bands at 1005 cm⁻¹ and 1685 cm⁻¹ can be assigned to the stretching modes of C-O and C=O bonds.46 An intense band identified at 1204 cm⁻¹ represents a combined contribution from C-O stretch and O-H deformation vibrations.47 Additionally, a small band at 1033 cm⁻¹ in VSe₂@G/MXe corresponds to the C-F stretch of the HF-etched MXene.48 Moreover, the considerably deeper O-H bond for VSe₂@G and VSe₂@G/MXe infers the hydrophilic surface characteristic of the prepared electrocatalysts which is also evident from the existence of vibrational modes from different hydrophilic groups (C-O, C=O, O-H, and C-F). These hydrophilic groups are particularly desirable for electrocatalysts as they tend to improve the interaction with electrolyte ions and in turn boost up the electrochemical performance of the electrocatalytic material.49 In addition, these functional groups on the surface of 3D sponges are involved in interfacial interactions with VSe2 for enhanced charge transport and stability during the electrochemical operation.⁵⁰

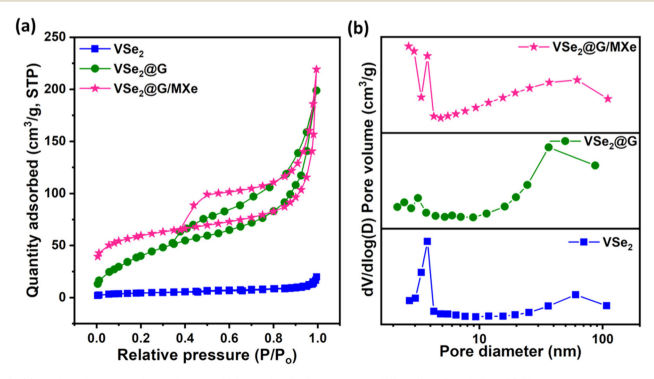
3.4. Physisorption analysis

 N_2 adsorption-desorption measurements were performed at 77 K to analyze the surface area and porous characteristics of VSe₂@G and VSe₂@G/MXe microstructures and bare VSe₂. As displayed in Fig. 4(a), all samples exhibited a type-IV adsorption isotherm. Well defined hysteresis loops can be observed in the adsorption-desorption isotherms for VSe₂@G and VSe₂@G/MXe microstructures with no limiting adsorption at a high relative pressure, indicating the characteristic of the H_3 -type hysteresis loop. ⁵¹ Capillary condensation in porous VSe₂@G and VSe₂@G/MXe microstructures occurred at a rela-

tive pressure $P/P_0 = \sim 0.4$ which directs to the presence of a large number of mesopores.⁵² The BET specific surface area (SSA) for VSe2 was found to be 15 m2 g-1 which increased to 40.4 and 43.9 m² g⁻¹ for VSe₂@G and VSe₂@G/MXe, respectively. Apparently, rGO contributes significantly to the enhancement of the SSA. Even though the SSAs of VSe₂@G and VSe₂@G/MXe are comparable, VSe₂@G/MXe showed a higher SSA than that of the previously reported 2D MXene.⁵³ This implies better surface accessibility of the MXene after its assembly into a 3D microstructure. Fig. 4(b) shows the pore size distribution curves of VSe2, VSe2@G, and VSe2@G/MXe. In contrast to VSe₂, VSe₂@G and VSe₂@G/MXe porous microstructures exhibit wide pore size distribution with mean pore diameters in the mesoporous and macroporous range. The BJH desorption average peak pore diameter for VSe₂@G was found to be 16.33 nm which shifted to 11.18 nm for VSe₂@G/ MXe. The result suggests that the addition of the MXene most likely affects the crosslinking ability and density of ice crystals formed during the freeze-drying process, which ultimately affects the textural properties and average pore size of the resulting microstructure.

3.5. Conductivity measurements

The electrical behavior of VSe₂, VSe₂@G, and VSe₂@G/MXe was analyzed by the two-probe current-voltage (I–V) measurements by converting the samples into a pellet with the help of a hydraulic press. Fig. S3 \dagger shows the I–V profiles of all samples under the applied voltage range of -10 to 10 V. Current responses for all the samples followed Ohm's law with a typical linear relationship between the current and voltage



 $\textbf{Fig. 4} \quad \text{(a) N_2 adsorption-desorption isotherms and (b) pore size distribution curves of VSe_2, $VSe_2@G$, and $VSe_2@G/MXe$. }$

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axes. Evidently, the current response for the electrocatalysts under the same applied voltage was in the order $VSe_2@G/MXe > VSe_2@G > VSe_2$. This indicates that $VSe_2@G/MXe$ exhibited the lowest electrical resistance and high conductivity among all electrocatalysts. The electrical conductivity (σ) for each electrocatalyst was calculated using eqn (5).

$$\sigma = \frac{w}{R \times A} \tag{5}$$

In the above equation, w and A denote the width and area of the pellet, respectively, and R is the resistance obtained from the slope of the I-V profile. The electrical conductivities of VSe₂, VSe₂@G, and VSe₂@G/MXe were found to be 8.84 × 10^{-8} , 2.15×10^{-7} , and 5×10^{-7} S cm⁻¹, respectively. The low electrical resistance and significant improvement in the electrical conductivity of VSe₂@G/MXe as compared to VSe₂ and VSe₂@G are reasonably ascribed to the integration of highly conductive MXene sheets into the 3D matrix and generation of ohmic contacts among the individual components of the VSe₂@G/MXe microstructure.⁵⁴ The higher electrical conductivity of VSe₂@G/MXe is expected to promote the transport of electrons during the electrochemical activity.

3.6. Water-splitting experiments

To evaluate the overall water-splitting performance, HER and OER catalytic activities of bare VSe2 and sponge-like VSe2@G and VSe₂@G/MXe microstructures were analyzed in 1 M KOH as the alkaline medium. Fig. 5(a) exhibits IR-corrected polarization curves for the HER activity measured at 3 mV s⁻¹. For comparison, the HER performance of commercial Pt/C and the NF substrate is also shown. Notably, Pt/C showed the highest HER activity and NF was the least active. The onset potential for the HER in the presence of VSe2, VSe2/G, and VSe₂@G/MXe electrocatalysts was observed at -0.19, -0.15, and -0.11 V, respectively. To achieve a cathodic current density of 10 mA cm⁻², VSe₂ required an overpotential of 261 mV that decreased significantly for VSe₂@G and VSe₂@G/MXe to 202 and 153 mV, respectively to reach the same current density. This finding suggests that integration of VSe₂ into the 3D rGO and rGO/MXene based sponge is beneficial for the electrochemical activity. Nonetheless, the overpotential for VSe₂@G/ MXe is the lowest which signifies its greater HER activity among all electrocatalysts. The LSV curve depicted in Fig. S4† illustrates a discernable HER activity for pure G/MXe (overpotential = 307 mV at 10 mA cm⁻²) which is anticipated to be contributed by the redox-active Ti⁺³/Ti⁺⁴ in the MXene. However, this activity when solely relying on G/MXe is notably lower than that of VSe2, indicating that the active sites for the HER activity are predominantly contributed by VSe₂. In this way, the enhanced electrocatalytic performance of VSe₂@G/ MXe can be explained by its 3D sponge-like structure, rich in mesopores and macropores, coupled with the high electrical conductivity of G/MXe. This structure exposes more active sites of VSe₂ and facilitates rapid ion and electron transport, thereby increasing the active site efficiency of VSe₂@G/MXe during the HER. To comprehend the kinetics of HER activity,

Tafel slopes (b) for all electrocatalysts were obtained from the corresponding polarization curves according to the following equation for the half-cell reaction (6):

$$\eta = b \log (j) \tag{6}$$

In the above equation, η denotes the overpotential and iis the current density. As depicted in Fig. 5(b), Tafel slopes for VSe2, VSe2@G, and VSe2@G/MXe are 140, 107, and 84 mV dec⁻¹, respectively. Generally, the HER activity in alkaline media proceeds through the water dissociation Volmer step (i.e., $H_2O + e^- \leftrightarrow H_{ads} + OH^-$) and thereupon by the Heyrovsky step (i.e., $H_2O + H_{ads} \leftrightarrow H_2 + OH^-$) or the Tafel reaction (i.e., $2H_{ads} \leftrightarrow H_2$). The theoretical values of the Tafel slopes for the Volmer, Heyrovsky, and Tafel reactions are 120, 40, and 30 mV dec⁻¹, respectively. 20 For VSe₂@G/MXe, a value of 84 mV dec⁻¹ for the Tafel slope indicates a typical Volmer-Heyrovsky mechanism, where electrochemical desorption or the Heyrovsky reaction is the rate-limiting step.55 For Tafel slopes close to 120 mV dec⁻¹, primary water dissociation (the Volmer reaction) is the rate limiting step, suggesting different H2 evolution kinetics for VSe2 and VSe₂@G from that of VSe₂@G/MXe. Clearly, a small value of Tafel slope for VSe₂@G/MXe demonstrates better kinetics for H₂ evolution in the presence of the VSe₂@G/MXe electrocatalyst.

Furthermore, considering the current density and concentration of V ions, the turnover frequency (TOF) was determined for all electrocatalysts at different overpotentials, as shown in Fig. 5(c). For bare VSe₂, the TOF can reach up to 0.027 s⁻¹ at 0.3 V, which elevated to 0.093 s⁻¹ for VSe₂@G and was the maximum, *i.e.*, 0.15 s⁻¹, in the presence of VSe₂@G/MXe. However, the calculated value of the TOF can be the upper hold as not all the V ions are necessarily accessible for catalytic activity. Nonetheless, the least overpotential and Tafel slope for VSe₂@G/MXe are convincing for its highest TOF value and it must have provided the maximum number of active sites and accessibility of electrolyte to the active sites for the significant HER activity.⁵⁶

The low overpotential (153 mV), Tafel slope (84 mV dec⁻¹), and high TOF (0.15 s⁻¹) for VSe₂@G/MXe can be ascribed to its enhanced intrinsic characteristics. In addition to the 3D spongy framework, the hydrophilicity of VSe₂@G/MXe combined with its high electrical conductivity improves the electrical contact of reactants to the active sites of VSe₂@G/MXe and hence in succession tailors the free energy for H-adsorption through the synergistic effects of VSe₂ and the compositing material (G/MXe).⁵⁷

Moreover, electrochemical impedance measurements were performed to analyze the charge transfer kinetics during the electrochemical reaction. The corresponding Nyquist plots (Re (Z) vs. -Im(Z)) derived from the impedance data show a characteristic Nyquist arc along the real impedance axis for all the prepared electrocatalysts (Fig. 5(d)). The equivalent fitted circuit (Randle's circuit) for Nyquist was used to determine the charge transfer resistance (R_{ct}). Obviously, VSe₂@G/MXe shows a smaller Nyquist arc with a low R_{ct} of 9.1 Ω that is far less

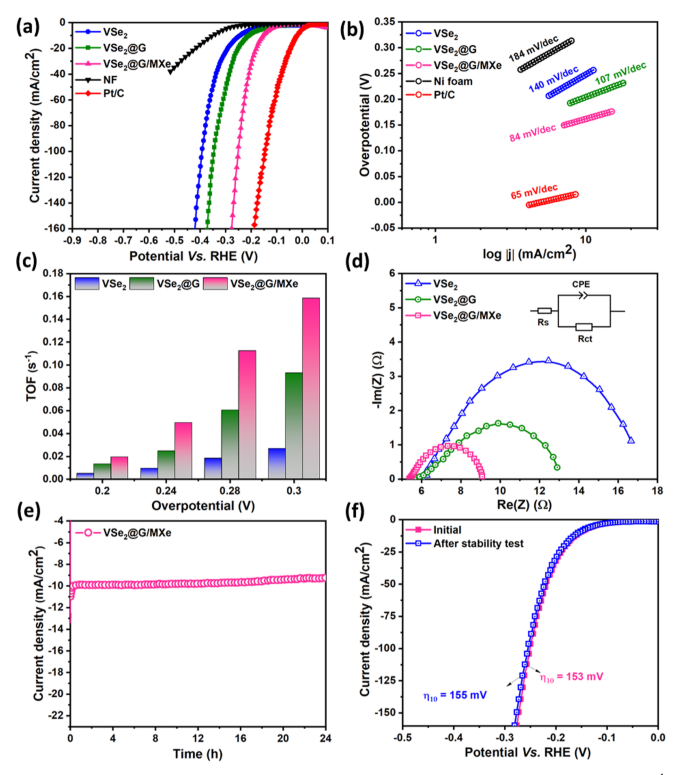


Fig. 5 Electrochemical measurements for the HER activity in 1 M KOH of VSe₂, VSe₂@G, and VSe₂@G/MXe: (a) LSV curves measured at 3 mV s⁻¹, (b) Tafel plots, (c) TOF at 0.2, 0.24, 0.28, and 0.3 V, (d) Nyquist plots at 0.1 Hz to 100 kHz, (e) chronoamperometry test of VSe₂@G/MXe for 24 h, and (f) LSV tests of VSe₂@G/MXe for 1000 cycles at 100 mV s⁻¹.

than those of VSe₂@G (12.8 Ω) and VSe₂ (16.5 Ω), indicating a low resistance to charge transfer on the surface of VSe₂@G/MXe than those of the counterpart electrocatalysts.

Besides the catalytic performance, stability is also an important parameter to assess the potential practical application of an electrocatalyst. The stability of VSe₂@G/MXe was evaluated

through the chronoamperometric and LSV measurements. The chronoamperometry results, displayed in Fig. 5(e), show that at a constant overpotential, the current density fluctuates with a very small loss after a long period of 24 h. This small fluctuation in the current density can possibly be caused by the hydronium consumption or lingering H₂ bubbles that impede

the HER activity, as also previously reported by Zhu *et al.*²⁰ Alternatively, LSV tests were conducted at a high scan rate of 100 mV s⁻¹ for 1000 cycles to further confirm the stability of VSe₂@G/MXe. As shown in Fig. 5(f), the VSe₂@G/MXe electrocatalyst maintains the initial overpotential with a slight shift from 153 mV to 155 mV over continuous 1000 LSV scans.

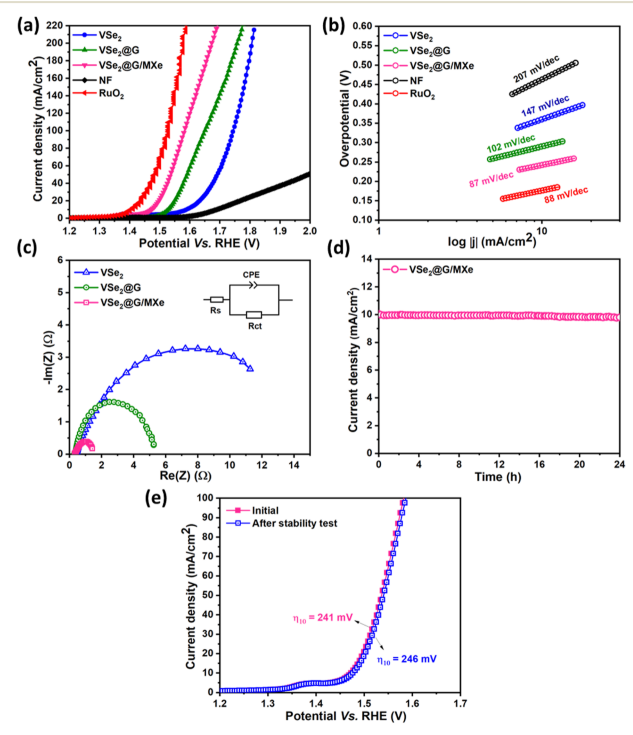


Fig. 6 Electrochemical measurements for the OER activity in 1 M KOH of VSe₂, VSe₂@G, and VSe₂@G/MXe: (a) LSV curves measured at 3 mV s⁻¹, (b) Tafel plots, (c) Nyquist plots at 0.1 Hz to 100 kHz, (d) chronoamperometry test of VSe₂@G/MXe for 24 h, and (e) LSV tests of VSe₂@G/MXe for 1000 cycles at 100 mV s⁻¹.

These results confirm the mechanical robustness and chemical durability of the VSe₂@G/MXe electrocatalyst. This enhanced stability of VSe₂@G/MXe can be anticipated to be contributed by the integration of VSe₂ into G/MXe which (i) prevents the aggregation of VSe₂ sheets and (ii) provides better dispersion of active sites, thus minimizing the mechanical stress concentrations. Additionally, as mentioned beforehand, the interfacial interactions between G/MXe and VSe₂ impede the percolation of VSe₂ sheets during the electrocatalytic activity.

Driven by the promising HER performance, the catalytic activities of the prepared electrocatalysts for the OER were also evaluated in 1 M KOH within the same electrochemical setup. Fig. 6(a) and Fig. S5† show IR-corrected polarization curves for the OER in the presence of the NF substrate, VSe₂, VSe₂@G, VSe₂@G/MXe, benchmark RuO₂ and G/MXe, respectively.

Similar to the HER, commercial RuO2 showed the highest activity while the NF substrate and G/MXe exhibited the lowest activity for the OER. The onset potentials for VSe₂, VSe₂@G, and VSe₂@G/MXe were recorded at 0.34, 0.23, and 0.19 V, respectively. Obviously, VSe2@G/MXe offered an earlier onset of the anodic current as compared to its analogs after the benchmark RuO2. Following this, VSe2@G/MXe also outperformed other electrocatalysts in terms of the overpotential for the OER with an optimum potential of 241 mV to drive an anodic current density of 10 mA cm⁻². The overpotentials for G/MXe, VSe₂, and VSe₂@G to drive the same current density were found to be 419, 347, and 293 mV, respectively. To compare the OER kinetics, the Tafel slope of each electrocatalyst was obtained from the corresponding polarization curve, as shown in Fig. 6(b). Tafel slopes for VSe2, VSe2@G and VSe₂@G/MXe were 147, 102, and 87 mV dec⁻¹. Notably, the

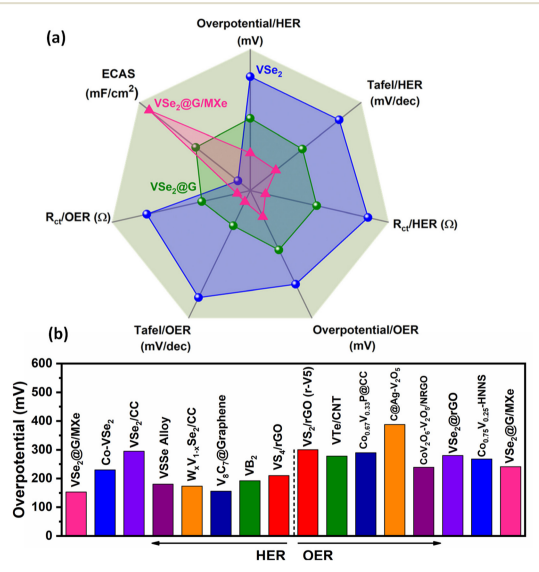


Fig. 7 (a) Radar plot showing a comparison of different electrochemical parameters obtained for VSe_2 , $VSe_2@G$, and $VSe_2@G/MXe$ from the HER and OER activity and (b) bar graph showing a comparison of the performance of $VSe_2@G/MXe$ for the HER and OER activities in terms of overpotential with that of similar reported electrocatalytic materials.

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Tafel slope of VSe₂@G/MXe approaches the Tafel slope of benchmark RuO₂. It is well reported that the OER mechanism embraces multiple steps of electron/proton transfer reactions. The value of the Tafel slope for the OER (87 mV) in the presence of VSe₂@G/MXe suggests that the rate-limiting step should be toward the end of the O2 evolution process as MOOH + HO $^ \rightarrow$ MOO $^-$ + H₂O, where M represents a site on the surface of the electrocatalyst.⁵⁸ Meanwhile, impedance measurement in Fig. 6(c) shows R_{ct} values in descending order as we proceed from VSe_2 (11.2 Ω), $VSe_2@G$ (5.25 Ω), and $VSe_2(\Omega G/MXe (1.41 \Omega))$. As previously observed for the HER, a considerably small diameter of a Nyquist arc and a small value of the R_{ct} indicate an enhanced electron transfer across the electrode-electrolyte interface VSe₂@G/MXe. Chronoamperometric measurements at a constant overpotential (Fig. 6(d)) show that VSe₂@G/MXe maintains an almost steady current output for 24 h with a small loss in current density. Likewise, LSV measurements in Fig. 6(e) at 100 mV s⁻¹ for continuous 1000 LSV cycles resulted in a small overpotential shift from 241 mV to 246 mV, which confirms the durability of VSe₂@G/MXe for the OER activity.

To get a deeper understanding of the intrinsic electrochemical activity of VSe₂@G/MXe, the electrochemical double layer capacitance (Cdl) was determined for all the electrocatalysts from the CV measurements at different scan rates (20–120 mV s⁻¹). Fig. S6† displays CV curves for VSe₂, VSe₂@G, and VSe₂@G/MXe recorded within a voltage range of 0.4-0.5 V where the output current is believed to be contributed by the electrical double layer charging. Linear plots of scan rate vs.

anodic and cathodic current densities were fitted to obtain the Cdl value. The average Cdl values for VSe2, VSe2@G, and $VSe_2@G/MXe$ were found to be 5.1, 8.9, and 13.1 mF cm⁻². C_{dl} is usually considered to be proportional to the electrochemical active surface area (ECAS) of the electrode-electrolyte interface for charge transfer during the electrochemical reaction.⁵⁹ A high C_{dl} or ECAS of VSe₂@G/MXe is due to the synergistic combination of VSe₂ and G/MXe that engenders an architecture that optimally exposes VSe₂ active sites while concurrently promoting efficient charge transfer, subsequently enhancing the available electroactive surface area for electrocatalytic activity.

Moreover, the ECAS-normalized current density displayed in Fig. S7 and S8† shows a higher current density of VSe₂@G/ MXe than that of VSe₂ for both the HER and OER, as is also observed in the case of geometric surface area-normalized current density. This approves the superior intrinsic electrocatalytic activity of VSe₂@G/MXe for the HER and the OER.

Fig. 7(a) shows a radar plot comparing different electrochemical parameters (overpotential, Tafel slope, ECAS, and R_{ct}) obtained from water-splitting measurements. For both the HER and OER, the electrocatalytic activity of VSe₂@G/MXe was significant as compared to those of VSe₂@G and VSe₂. Moreover, the HER and OER activities of VSe₂@G/MXe were also comparable or even better than previously reported similar electrocatalysts (Fig. 7(b) and Tables S1 and S2†).

The superior electrochemical performance of VSe₂@G/MXe can be accredited to its large SSA, large number of exposed electroactive sites, hydrophilicity, and high conductivity of G/ MXe. These characteristics of VSe₂@G/MXe are the outcomes

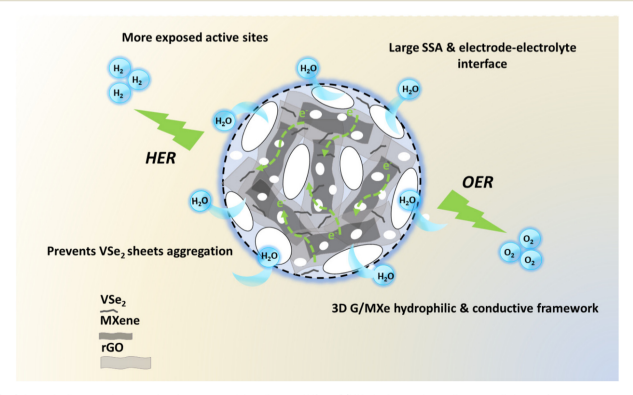


Fig. 8 Schematic diagram of structural and componential attributes of VSe₂@G/MXe for boosted overall water-splitting performance.

that resulted from the individual components (VSe₂, rGO, and MXene) and their synergistic effects, and most importantly from the morphological engineering of VSe₂@G/MXe into the 3D sponge like framework (Fig. 8).

4. Conclusions

Paper

In summary, an effective strategy was used to assemble VSe₂ into a MXene/rGO matrix-based 3D microstructure as VSe₂@G/ MXe. Physiochemical characterization of VSe₂@G/MXe exhibited a sponge-like pore-rich skeleton for VSe2@G/MXe with a high SSA of 43.9 m 2 g $^{-1}$ and an electrical conductivity of 5 \times 10⁻⁷ S cm⁻¹. Assembly of VSe₂ in a 3D sponge prevented the aggregation of VSe2 nanosheets, thereby exposing maximum active sites for the electrochemical reaction. Intrinsic characteristics of individual components (VSe₂, MXene, and rGO) and the unique morphology of VSe2@G/MXe synergistically worked together to improve the inherent electrocatalytic activity of the VSe₂@G/MXe microstructure. As a bifunctional electrocatalyst, VSe2@G/MXe exhibited significant electrochemical activity with low overpotentials of 153 mV and 241 mV to attain 10 mA cm⁻², for the HER and OER activities, respectively. Chronoamperometry and LSV stability tests confirmed the long-term stability of VSe₂@G/MXe. Hence, this work can pave the way to develop high performance 3D electrocatalysts for efficient energy conversion and water-splitting applications.

Conflicts of interest

There are no conflicts to declare.

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