Achieving ultrahigh electrochemical performance by surface design and nanoconfined water manipulation

Haisheng Li1,†, Kui Xu2,†, Pohua Chen1, Youyou Yuan1,3, Yi Qiu1, Ligang Wang1, Liu Zhu4, Xiaoge Wang1, Guohong Cai1, Liming Zheng1, Chun Dai1,5, Deng Zhou6, Nian Zhang6, Jixin Zhu2, Jinglin Xie1,7, Fuhui Liao1, Hailin Peng1, Yong Peng8, Jing Ju1, Zifeng Lin1,9 and Junliang Sun1,*

ABSTRACT

The effects of nanoconfined water and the charge storage mechanism are crucial to achieving the ultrahigh electrochemical performance of two-dimensional transition metal carbides (MXenes). We propose a facile method to manipulate nanoconfined water through surface chemistry modification. By introducing oxygen and nitrogen surface groups, more active sites were created for Ti$_3$C$_2$ MXene, and the interlayer spacing was significantly increased by accommodating three-layer nanoconfined water. Exceptionally high capacitance of 550 F g$^{-1}$ (2000 F cm$^{-3}$) was obtained withoutstanding high-rate performance. The atomic scale elucidation of the layer-dependent properties of nanoconfined water and pseudocapacitive charge storage was deeply probed through a combination of ‘computational and experimental microscopy’. We believe that an understanding of, and a manipulation strategy for, nanoconfined water will shed light on ways to improve the electrochemical performance of MXene and other two-dimensional materials.

Keywords: two-dimensional material, MXene, nanoconfined water, energy storage mechanism, supercapacitors

INTRODUCTION

Supercapacitors (SCs) are expected to possess both high energy density and power density, together with the metrics of prolonged cycle life and low cost [1,2]. Electrical double-layer (EDL) capacitors, such as commercial carbon electrode materials, have excellent rate performance but unsatisfactory mass-specific/volumetric capacitance [3,4]. In contrast, most pseudocapacitive materials exhibit high specific capacitance, but are limited by their high cost or poor durability [5–8]. Two-dimensional transition metal carbides (MXenes) have been considered as fascinating materials for pseudocapacitors, having both high volumetric capacitance and power density thanks to their highly redox surface, conductive carbide core and rapid ionic transport through interlayer water [9,10].

Titanium carbide (Ti$_3$C$_2$T$_x$), the first reported and the best-studied MXene, has also exhibited state-of-the-art electrochemical performance. Through electrode morphology design, high volumetric capacitance (∼1300 F cm$^{-3}$ at 100 mV s$^{-1}$) has been achieved in Ti$_3$C$_2$T$_x$ films [10] while high-rate performance (∼200 F g$^{-1}$ at 10 V s$^{-1}$) has been obtained in macroporous or vertically aligned Ti$_3$C$_2$T$_x$ MXene films [10,11]. Through composites with carbons [12], polymers [13] and transition metal oxides/hydroxides [14,15], inspiring progress was also made with regard to better energy storage applications. This exciting work urgently requires an understanding of charge storage and water confinement in MXenes, and thus the pathways for further enhancing performance.

Recent work has revealed that the capacitance of Ti$_3$C$_2$T$_x$ MXene in acidic electrolytes mainly originates from the surface redox reaction with non-negligible EDL-like capacitance [16,17]. Additional work further predicted that increasing the proportion of terminal Ti-O or introducing heteroatoms could enhance the capacitance of Ti$_3$C$_2$T$_x$ MXene.
Mn\textsuperscript{2+} intercalation

Figure 1. Schematic of modified MXenes synthesis. Structure models of (a) pristine Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}, (b) Mn\textsuperscript{2+} intercalated and partially oxidized Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} and (c) further nitrogen-doped Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}.

[18,19]. On the other hand, experimental and computational approaches studied water confinement and revealed its role in capacitive performance in neutral electrolytes [20,21]. In acidic electrolytes, where superior capacitance was achieved, proton redox and transport in MXene-confined water were also theoretically investigated [22]. However, the literature did not often give a complete picture of how the interactions of surface chemistry, protons and nanoconfined water contribute to the high capacitance and high-rate performance. In addition, many researchers ignored the effect of nanoconfined water when modifying MXenes, probably due to the difficulties in characterization [23]. As a result, manipulating nanoconfined water in between MXene layers to maximize the electrochemical performance of MXene electrodes is still a formidable task.

Nanoconfined water tends to reside in quasi-integer layers in between MXenes and shows layer-dependent properties as theoretically reported [22]. Increasing the layers of nanoconfined water, for example from two to three, is expected to significantly promote the water mobility and proton diffusion coefficients, and hence the electrochemical performance [22]. Though surface modification by adjusting the proportion of terminal -F, -OH and -O of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} MXene has affected the way nanoconfined water resides, most cases did not yield more than two layers of nanoconfined water, and the capacitance enhancement was also unsatisfactory [17,18]. Nitrogen doping has been reported to improve the hydrophilicity of carbon materials [24], hence it is promising in manipulating nanoconfined water in between MXenes. However, current work on nitrogen-doped MXene mainly focuses on electronic structure modulation [19,25,26], omitting an in-depth and systematic study of nanoconfined water.

Herein, a facile yet effective method is proposed to design surface-modified Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} MXene with outstanding electrochemical performance. By introducing oxygen and nitrogen surface groups, nanoconfined water can be manipulated for faster charge storage, and also to create more active sites for ultrahigh capacitance. Ex-situ cryo high-resolution transmission electron microscopy (cryo-HRTEM) and other advanced characterization methods combined with theoretical studies unveil the charge storage mechanism and the role of nanoconfined water. With the manipulated three-layer nanoconfined water, the deep active sites are fully exploited and the high-rate performance is retained, contributing to an exceptional volumetric capacitance in a wide range of scan rates.

RESULTS AND DISCUSSION

Synthesis and structure analysis

The surface-modified Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} MXene film was prepared by a simple and controllable two-step process: first electrostatic assembly with redox metal cations followed by ammonia treatment (Fig. 1 and Supplementary Fig. S1).

To lower the energy barrier for nitrogen doping [19] and facilitate the entrance of ammonia, we firstly used cation intercalation to modify the surface chemistry and enlarge the interlayer spacing of Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x} MXene. Outstanding among a series of intercalated cations, Mn\textsuperscript{2+} was selected as a mild cation to intercalate Ti\textsubscript{3}C\textsubscript{2}T\textsubscript{x}, and a flexible film (Mn-MXene) was obtained through vacuum filtration. However, current work on nitrogen-doped MXene mainly focuses on electrostatic structure modulation [19,25,26], omitting an in-depth and systematic study of nanoconfined water.
Figure 2. Morphological and structural analysis of Mn-MXene-N. (a) Ex-situ XRD patterns of P-MXene, Mn-MXene and Mn-MXene-N films with the interlayer spacing of 14.0 Å, 14.7 Å and 16.2 Å, respectively. (b) XPS results of N 1s of Mn-MXene-N film. (c) The FTIR spectra of the indicated films. The contact angles of (d) P-MXene and (e) Mn-MXene-N. (f) SEM images of the surface and cross-section morphologies of Mn-MXene-N after cycling. (g) Cryo-HRTEM image of Mn-MXene-N.

(Fig. 2a and Supplementary Fig. S3). Moreover, the surface chemistry of Ti₃C₂Tₓ was also modulated by redox Mn ion with the assistance of atmospheric oxygen, resulting in a higher proportion of terminal -O and less -F, as evidenced by X-ray photoelectron spectroscopy (XPS) and X-ray fluorescence analysis (Supplementary Fig. S5, Tables S2 and S3). Nitrogen was then introduced into Mn-MXene, and P-MXene for comparison, through ammonia annealing. The interlayer spacing of the final film (Mn-MXene-N) was further enlarged (from 14.7 Å to 16.2 Å, ex situ XRD, Fig. 2a), with considerable nitrogen species doping. The nitrogen content of Mn-MXene-N was almost twice that in P-MXene-N, indicating the necessity of cation intercalation pretreatment (elemental analysis, Supplementary Table S4).

The chemical formula of Mn-MXene-N was determined to be Mn₀.₁₂-Ti₃C₁.₉₇F₀.₂₅O₁.₀₁(OH)₀.₄-N₀.₃₃, according to the XPS, X-ray fluorescence spectrum, inductive couple plasma (ICP) and thermogravimetric analysis (Supplementary Tables S2–S5, Figs S6 and S7).

The chemical states of nitrogen were analyzed to be Ti-N-O, Ti-N-OH and Ti-N (Supplementary Discussion 1), with the corresponding N 1s peak position in the XPS (Fig. 2b) located at 401.6, 399.7 and 396.7 eV, respectively. Fourier transform infrared reflection (FTIR) spectra (Fig. 2c, Supplementary Fig. S8) further confirm the existence of Ti-N-O (1400 cm⁻¹). This type of nitrogen-doped surface is proven to be more hydrophilic theoretically and experimentally, and can make aqueous electrolyte and the Mn-MXene-N surface contact closer, which is beneficial for double electrode layer energy storage (Supplementary Fig. S9a–c, Fig. 2d and e). The scanning electron microscopy (SEM) images (Supplementary Fig. S10a, Fig. 2f) of Mn-MXene-N before and after the durability test show that the surface and cross section morphologies are smooth and compact, respectively. Furthermore, ex-situ cryo-HRTEM and rotation electron diffraction (RED) were performed to obtain additional structural information while preserving the interlayer water. The real space image of Mn-MXene-N (Fig. 2g) shows a uniform interlayer spacing of 16.0 Å (cell parameter c = 32.0 Å), consistent with 16.2 Å from the ex-situ XRD. The reconstructed 3D reciprocal lattice (Supplementary Fig. S10b–d) of Mn-MXene-N from the RED data gives the cell parameter a or b with 2.8 Å, close to that of the MAX phase Ti₃AlC₂ [27].

Electrochemical energy storage performance

Electrochemical performances of the as-prepared film electrodes are characterized by a three-electrode configuration in 3M H₂SO₄. Cyclic voltammograms (CV) of Ti₃CₓTₓ MXene films with different
modifications were tested at 5 mV s$^{-1}$ (Fig. 3a). Among them all, the 3-μm-thick Mn-MXene-N film has the largest integral area in the CV profiles, giving an ultrahigh specific capacitance up to 550 F g$^{-1}$ with corresponding volumetric capacitance reaching an unmatched ∼2000 F cm$^{-3}$. Mn cation intercalation (Mn-MXene) or nitrogen doping (P-MXene-N) can also enhance the specific capacitance of Ti$_3$C$_2$T$_x$, independently, but only leads to a similar value of ∼400 F g$^{-1}$ (Fig. 3a, Supplementary Fig. S11a). These results indicated the synergistic effects of Mn$^{2+}$ intercalation and nitrogen doping in terms of improving the performance of Mn-MXene-N. The galvanostatic charge/discharge curves in Supplementary Fig. S11b suggest that Mn-MXene-N has a much longer charge/discharge time at 1 A g$^{-1}$ with a quasi-triangular shape, confirming its excellent specific capacitance.

The CV profiles (Fig. 3b) of 1-μm-thick Mn-MXene-N film were then collected at scan rates from 20 mV s$^{-1}$ to 2000 mV s$^{-1}$. The thinner film was used here to minimize ion transport limitations. The shapes of the CVs were well maintained below 500 mV s$^{-1}$ and a considerable integral area still remained even at 2000 mV s$^{-1}$. To analyze the charge storage kinetics of Mn-MXene-N, the Trassatti analysis method [28] was used here, in which the total amount of stored charge ($q_T$) contained non-diffusion-controlled charges ($q_n$, independent of scan rates ($v$)) and diffusion-controlled charges ($q_d$, proportional to $v^{-1/2}$).

As shown in Fig. 3d and e, 3-μm-thick Mn-MXene-N delivered an exceptional capacitance exceeding 500 F g$^{-1}$ (≈1800 F cm$^{-3}$) at scan rates from 5 mV s$^{-1}$ to 200 mV s$^{-1}$, with a decay <10%. Even at a high scan rate up to 1000 mV s$^{-1}$, a good capacitance of 300 F g$^{-1}$ (≈1100 F cm$^{-3}$) was still obtained. To the best of our knowledge, these results surpass work based on all kinds of materials in volumetric capacitance (Supplementary Discussion 2, Fig. S13a). Mn-MXene-N showed a thickness-dependent electrochemical performance due to the limitation of vacuum filtration. Nonetheless, admirable volumetric capacitance was still obtained with a thickness of 3 μm.
16 μm (5 mg cm⁻²) and 36 μm (10 mg cm⁻²) (Supplementary Fig. S13b–d). It is believed that the intrinsic properties of our MXenes could be fully utilized when combined with state-of-the-art electrode morphology. Moreover, the capacitance of Mn-MXene-N shows ~100% retention and coulombic efficiency when repeating the charge/discharge tests at 10 A g⁻¹ for 10 000 cycles in air (Fig. 3f).

As far as is known, this combination of ultrahigh volumetric capacitance and outstanding rate performance in Mn-MXene-N is very rare in electrochemical energy storage (EES) materials. The reason can be attributed to the dilemma presented by the fact that improving ion transport of EES materials has been mainly achieved through increasing surface area, inevitably sacrificing the volumetric capacitance [23]. Nevertheless, another way to significantly accelerate proton transport in a minimum volume was explored by manipulating the nanconfined water and simultaneously introducing extra active sites by modifying the surface chemistry, which together contribute to the ultrahigh performance of Mn-MXene-N as discussed below.

The origin of increased capacitance

To identify the origin of the increased capacitance and probe the charge storage mechanism, a preliminary investigation was initially undertaken based on the electrochemical analysis. The CV profiles (Fig. 3a) showed that Mn-MXene has similar redox peaks to P-MXene near -0.15 V versus Ag/AgCl, with a slightly higher current density. In addition, a new pair of redox peaks appeared at 0 V, which was also reported in pristine Ti₃C₂Tx under a very low scan rate [16] or in vertically aligned electrodes [11]. These redox peaks were independent of Mn ions, since the cations would be lost due to ion exchanges after pre-cycles (ICP, Supplementary Table S5). The role of Mn²⁺ intercalation was to facilitate the surface modification of Ti₃C₂Tx, i.e. increase the proportion of -O (XPS, Table S2), thus enhancing the pseudocapacitance as predicted in previous work [16,18]. In the case of Mn-MXene-N (Fig. 3a), another pair of redox peaks appeared in the potential range from −0.1 V to 0.1 V, which had a larger peak separation when compared to that in P-MXene or Mn-MXene, implying that the introduction of N species might create new active sites. The current density of Mn-MXene-N film above 0.1 V was also obviously increased, where the electrochemical behavior was supposed to be capacitive (Supplementary Fig. S15).

To further investigate the capacitance origin of Mn-MXene-N, in-/ex-situ experiments combined with theoretical computations were conducted. As shown in the ex-situ XPS spectra of N 1s (Fig. 4a, Supplementary Fig. S16), significant enhancement of the peak intensity of Ti-N-OH with a decrease in Ti-N-O was observed when the applied potential was reduced from 0.4 V to −0.5 V, confirming the valence reduction of nitrogen. Ex-situ soft X-ray absorption spectroscopy was used to study the valence change of the element Ti. The peak intensity ratio of Ti L₃ 2p₂ to e₈ progressively decreased, with the pre-edge region shifting to a higher energy (Supplementary Fig. S17) when the applied potential was increased, suggesting an increased Ti oxidation state [29]. These results also imply that terminal N species could provide extra capacitance without blocking the redox reaction of Ti-O.

Furthermore, an in-situ XRD experiment was conducted to trace the interlayer spacing change of Mn-MXene-N during the CV test at 0.5 mV s⁻¹ (Fig. 4b, Supplementary Figs S18 and S19). During charging (cathodic polarization), a sharp d-spacing transition of ~2.8 Å at -0.05 V and a gradual increase of ~0.5 Å in the overall potential range from 0.4 V to -0.5 V were observed. Upon discharging, this process was reversed, indicating that the cycling process was highly reversible. Molecular dynamic (MD) simulation and density functional theory (DFT) calculations were conducted to investigate the interlayer spacing change induced during the charging/discharging process and by the transmission from Ti-N-O to Ti-N-OH. The MD simulations observe a reversible intercalation/deintercalation of H₃O⁺ and H₂O between the MXene layers during the charging/discharging process, which directly leads to an increase/decrease in the interlayer spacing. The change of interlayer H₃O⁺ and H₂O numbers, Δnᵦ, exhibited a high correlation with the change of the inner surface distance Δdᵦ (Fig. 4c and e), with the average normalized correlation coefficient reaching ~96.8%. Also, the static DFT calculation shows a ~0.6 Å interlayer spacing change (0.3 Å × 2, on both sides of MXene) when the Ti-N-O transforms to Ti-N-OH (Fig. 4d).

The high correlation between the Δnᵦ and Δdᵦ qualitatively revealed that the sharp change of interlayer spacing up to 2.8 Å originated from sudden H₃O⁺/H₂O intercalation at the corresponding potential. When considering that the intercalated number of H₂O is a tenth that of H₃O⁺, it is reasonable to mainly attribute water intercalation to the interlayer spacing change (Supplementary Figs S20 and S21). This larger interlayer spacing change of Mn-MXene-N allows the in-layer water structure to directly transform from one layer to three layers, indicating that more H₃O⁺/H₂O...
Natl Sci Rev, 2022, Vol. 9, nwac079

Figure 4. Surface evolution and interlayer spacing change of Mn-MXene-N during CV test. (a) Ex-situ XPS spectra of N 1s for Mn-MXene-N after holding at different potentials. The proportion of Ti-N-O is reduced with the applied potential decreased. (b) In-situ XRD patterns during CV cycles at 0.5 mVs⁻¹. (d) Change of lattice parameter calculated from static DFT. (c and e) Change of the interlayer H₃O⁺ and H₂O numbers Δnw and change of the inner surface distance Δdi observed during charge-discharge MD simulations.

The investigation and manipulation of nanoconfined water

Nanoconfined water in between MXene layers provides a major way for protons to move to and from the redox surface, and hence the properties of nanoconfined water will strongly affect proton transport. Experiments were designed and theoretical calculations conducted to manipulate the nanoconfined water and probe its role in electrochemical performance.

Firstly, how the nanoconfined water resides in between Mn-MXene-N was investigated. A common phenomenon was noticed whereby MXenes (and also some 2D and layered materials) usually need dozens of CV pre-cycles, or to rest in the electrolytes, to achieve their best electrochemical performance in aqueous electrolytes, and this is referred to as the activation process. This process is supposed to be a result of water intercalation, though this lacks solid evidence. Thus, in-situ XRD was performed during the CV pre-cycles (Fig. 5a) of Mn-MXene-N to investigate this process. As shown in Fig. 5b and Supplementary Fig. S24, the initial phase...
Figure 5. Layer-dependent properties of nanoconfined water. (a) CV curves of Mn-MXene-N during CV pre-cycles at 20 mV s\(^{-1}\). (b) In-situ XRD patterns and (c) the ex-situ cryo-HRTEM image of Mn-MXene-N captured during the CV pre-cycles. (d) The atomic resolution HAADF images of three distinct phases with interlayer spacing of 1.29, 1.45 and 1.57 nm, respectively. All scale bars are 2 nm. The experimental results are highly consistent with MD simulations. (e) Local snapshot of the Mn-MXene-N model with three-layer (15.8 Å), two-layer (14.4 Å) and one-layer (13.0 Å) water in between the layer of MD simulation, with the probability density profiles of water molecules along the \( \text{c} \) lattice direction displayed on the left side. (f) The trends of the initial phase residue and capacitance enhancement of Mn-MXene-N during CV pre-cycles. (g) Low field \(^1\)H time-domain nuclear magnetic resonance spectra of the indicated samples. The shaded region represents the time-domain region of interlayer water.

(partially dehydrated, at 7.0°) was evolved into the activated phase (fully hydrated, at 5.6°) during the activation process, with the corresponding interlayer spacing increased from 12.7 Å to 15.7 Å. The two peaks are distinct, implying that the change of interlayer spacing due to water intercalation might not be continuous.

To provide complementary evidence at the atomic level, ex-situ cryo-HRTEM was used to observe the interphase between the initial and activated phase during pre-cycles. As shown in Fig. 5c and Supplementary Fig. S25, a clear interlayer spacing change from 13.0 Å to 16.0 Å was observed, corresponding to the transformation of the initial phase to the activated phase. Surprisingly, in such a small domain, the change of interlayer spacing was still discrete. Moreover, the atomic resolution high-angle annular dark-field (HAADF) images of these distinct phases were also captured individually (Fig. 5d). Atomically imaging water molecules in between MXenes using the current TEM technique was extremely difficult (Supplementary Discussion 3). Thus, we conducted MD simulations to shed light on this unusual phenomenon, which indicated that water molecules tend to be intercalated in quasi-integer layers with particular interlayer spacing (Supplementary Fig. S26). Furthermore, the three phases with interlayer spacing of 12.9 Å, 14.5 Å and 15.7 Å were assigned to be Mn-MXene-N with one, two and three layers of confined water respectively, according to the simulation results (Fig. 5c). The cryo-HRTEM image also indicated a fast transition of the initial phase to the activated phase, which was completed within dozens of nanometers, accounting for the distinct peaks observed in the in-situ XRD.

Here, the layer-dependent properties of nanoconfined water (with additional investigations presented in Supplementary Discussion 4) was probed. Firstly, three-layer nanoconfined water...
in between Mn-MXene-N showed much improved water mobility and proton diffusion coefficient than one-layer water, close to bulk water as evidenced by the MD simulations results (Supplementary Fig. S29). This enhancement of ionic mobility could boost the charging and discharging process and effectively shorten its charging time, which is consistent with the calculated time constant (\( \tau \)) of resistance capacitance transmission line model equivalent circuit (Supplementary Fig. S21). In addition to the kinetic perspective, the increased layer spacing could markedly reduce the free energy barrier (\( \Delta G \)) for \( H_3O^+ / H_2O \) migrating from the bulk electrolyte into the MXene galleries via the edges of the MXene layers, while the \( \Delta G \) for three layer case is 2.5 kJ mol\(^{-1}\), smaller than the \( \Delta G \) for one layer case. Meanwhile, the calculated hydrogen bonding number density \( \rho \) in the three-layer water structure is \(~29.2 \text{#/nm}^3\), while \( \rho \) in one-layer water is only \(~10.2 \text{#/nm}^3\). This enhanced denser hydrogen bonding network could effectively promote the proton-hopping transport, likely via the Grotthuss mechanism. The reduction of the energy barrier and the enhanced denser hydrogen bonding network jointly provide a solution to the main bottleneck for the charging process [22].

Thus, superior electrochemical performance in activated MXenes is expected and confirmed by in-situ experiments. In the in-situ XRD during the activation process, it was noticed that the peak intensity of the initial phase gradually decreased and further disappeared in the end, with the obtained activated phase appearing in the reverse trend. An enhancement of the integral area of CVs was simultaneously observed (Fig. 5a). When combining the two trends, an interesting result was produced: the increased specific capacitance during activation was highly correlated with the proportion of the initial phase converted (Fig. 5f, Supplementary Fig. S30). This semi-quantitative relationship suggested that the specific capacitance of Mn-MXene-N can be fully exploited with three-layer water, thereby nearly doubling the capacitance of the initial phase.

In-situ electrochemical impedance spectroscopy (EIS) was also performed to investigate the effect of water intercalation. As illustrated in Supplementary Fig. S31, an obvious decrease of the 45-degree linear part (related to ion transport resistance) and a steeper slope of the Nyquist plot in the low-frequency range (related to specific capacitance) were observed when the CV cycle numbers were increased, indicating the fruitful role of intercalated water on both ion accessibility and the exploitation of active sites.

In addition, in the potential range (below 0.05 V versus Ag/AgCl) that pseudocapacitance dominates, Mn-MXene-N can accommodate three-layer confined water while P-MXene can only accommodate two layers, as shown in this work and a previous work [16]. Hence, protons and water in between Mn-MXene-N layers can be transported more quickly, leading to a superior rate performance (Supplementary Fig. S32). It was also noticed that Mn-MXene has a worse rate retention than P-MXene despite both having two-layer water. This may be attributed to other factors affecting proton mobility, including surface terminals, intercalated cations and defects [30–32]. Nevertheless, layers of nanoconfined water are still the dominant factor, which was experimentally confirmed by low field \(^1H\) nuclear magnetic resonance.

Figure 5g depicts the continuous distribution of the spin-spin relaxation time, \( T_2 \), for the three samples. Generally, there are three recognizable states of protons in these films, that is, protons in bonded terminals, interlayer water and free water, based on the feature that protons with higher mobility have a longer relaxation time. The time-domain region between 0.5 ms and 20 ms is ascribed to interlayer water (including hydrated proton). In this region, Mn-MXene film has the lowest relaxation time of all, and is responsible for the worst rate performance. Mn-MXene-N film has the longest relaxation time of all, and further with higher mobility of protons in mesosplit water (time-domain near 100 ms) and freer surface proton (time-domain below 0.5 ms), which together account for the much better rate performance of Mn-MXene-N. On the other hand, MD simulations indicate that Mn-MXene-N with three-layer confined water can store more charge and simultaneously accelerate the charging/discharging process (Supplementary Fig. S21), further confirming the superior electrochemical performance. Unlike conventional methods that increase porosity, another way was demonstrated here to maximize the capacitive performance of Ti_xC_yT_z. MXene through manipulating the behavior of confined water, in which high volumetric capacitance of bulk layered materials was perfectly combined with the fast ion diffusion of exfoliated 2D materials.

CONCLUSION

This work shows how surface chemistry modification and nanoconfined water manipulation can enhance electrochemical performance (improving its specific capacitance and rate performance, simultaneously). The designed Mn-MXene-N delivers an ultrahigh volumetric capacitance at wide scan rates (above 1800 F cm\(^{-3}\) at 200 mV s\(^{-1}\) and 1100 F cm\(^{-3}\) at 1000 mV s\(^{-1}\)), far superior to most state-of-the-art electrode materials. The origin of the increased capacitance was also investigated and the role of
interlayer water in ultrahigh electrochemical performance was understood. In particular, through the first use of cryo-HRTEM in MXene fields associated with other advanced characterization methods and computational simulations, a clear picture of how nanoconfined water resides and acts in MXene layers was achieved during the electrochemical process. Precisely manipulating and understanding the nanoconfined water of MXenes is of wide interest but remains a cutting-edge challenge. Nevertheless, our work can open up opportunities in this field. Moreover, on account of the facile synthesis strategy and fundamental mechanism study, this work might also provide new insights into other 2D and layered materials with nanoconfined fluids beyond MXenes, and be applied beyond energy storage, to areas such as water desalination and ion-selective membranes.

SUPPLEMENTARY DATA

Supplementary data are available at NSR online.

ACKNOWLEDGEMENTS

The authors thank Prof. Ying Jiang, Prof. Peng Gao and Dr. Jihan Zhou at Peking University for their discussions on the nature of confined water and suggestions on confined water characterization.

FUNDING

This work was supported by the Ministry of Science and Technology of China (2020YFA0210700 to J.S.), the National Natural Science Foundation of China (22125102, 21871009 and 2020YFA0210700 to J.S.), the National Natural Science Foundation of Jiangsu Province (BK20200702 to K.X.).

AUTHOR CONTRIBUTIONS

H.L. and J.S. conceived the idea and designed the experiments. H.L. conducted the material preparations and most of the characterization. K.X. and J.Z. conducted the computational calculations. P.C. designed an in-situ XRD cell, conducted XRD experiments and analyzed the data. Y.Y., L.W., C.D. and Z.L. helped with the electrochemical tests and analyzed the data. Y.Q., X.W. and J.J. helped with cryo-HRTEM and RED. I.Z. and Y.P. helped with atomic-resolution cryo-STEM. Liming Z. and H.P. provided the Graphene Cu grid. G.C., D.Z. and N.Z. helped with the ex-situ XAS experiment. J.X. helped with the ex-situ XPS experiment. F.L. conducted thermogravimetric analysis and analyzed the data. H.L., K.X., P.C., Z.L. and J.S. co-wrote the manuscript. All authors contributed to the discussion of the data and writing the paper.

Conflict of interest statement. None declared.

REFERENCES

1. Simon P and Gogotsi Y. Perspectives for electrochemical capacitors and related devices. Nat Mater 2020; 19. 1151–63.
2. Simon P, Gogotsi Y and Dunn B. Where do batteries end and supercapacitors begin? Science 2014; 343. 1210–1.
3. Lukatskaya M-R, Dunn B and Gogotsi Y. Multidimensional materials and device architectures for future hybrid energy storage. Nat Commun 2016; 7. 12647.
4. Zhang C, Lv W and Tao Y et al. Towards superior volumetric performance: design and preparation of novel carbon materials for energy storage. Energy Environ Sci 2015; 8. 1390–403.
5. Zheng J-P, Cygan P J and Jow T-R. Hydrous ruthenium oxide as an electrode material for electrochemical capacitors. J Electrochem Soc 1995; 142. 2699–703.
6. Oh S-H and Nazar L-F. Direct synthesis of electroactive mesoporous hydrous crystalline RuO2 templated by a cationic surfactant. J Mater Chem 2010; 20. 3834–9.
7. Wang X, Xu M and Fu Y et al. A highly conductive and hierarchical PANi micro/nanostructure and its supercapacitor application. Electrochim Acta 2016; 222. 701–8.
8. Simon P and Gogotsi Y. Materials for electrochemical capacitors. Nat Mater 2008; 7. 845–54.
9. Anasori B, Lukatskaya M-R and Gogotsi Y. 2D metal carbides and nitrides (MXenes) for energy storage. Nat Rev Mater 2017; 2. 16098.
10. Lukatskaya M-R, Kota S and Lin Z et al. Ultra-high-rate pseudocapacitive energy storage in two-dimensional transition metal carbides. Nat Energy 2017; 2. 17105.
11. Xia Y, Mathis T-S and Zhao M-Q et al. Thickness-independent capacitance of vertically aligned liquid-crystalline MXenes. Nature 2018; 557. 409–12.
12. Zhao M-Q, Ren C-E and Ling Z et al. Flexible MXene/carbon nanotube composite paper with high volumetric capacitance. Adv Mater 2015; 27. 339–45.
13. Boota M, Anasori B and Voigt C et al. Pseudocapacitive electrodes produced by oxidant-free polymerization of pyrrole between the layers of 2D titanium carbide (MXene). Adv Mater 2016; 28. 1517–22.
14. Zhou J, Yu J and Shi L et al. A conductive and highly deformable all-pseudocapacitive composite paper as supercapacitor electrode with improved areal and volumetric capacitance. Small 2018; 14. 1803786.
15. Zhao R, Wang M and Zhao D et al. Molecular-level heterostructures assembled from titanium carbide MXene and Ni-Co-Al layered double-hydroxide nanosheets for all-solid-state flexible asymmetric high-energy supercapacitors. ACS Energy Lett 2018; 3. 132–40.
16. Mu X, Wang D and Du F et al. Revealing the pseudointercalation charge storage mechanism of MXenes in acidic electrolyte. Adv Funct Mater 2019; 29. 1902953.
17. Shao H, Xu K and Wu Y-C et al. Unraveling the charge storage mechanism of Ti3C2Tx MXene electrode in acidic electrolyte. ACS Energy Lett 2020; 5. 2873–80.
18. Chen X, Zhu Y and Zhang M et al. N-butyl lithium-treated Ti3C2Tx MXene with excellent pseudocapacitor performance. ACS Nano 2019; 13. 9449–56.
19. Lu C, Yang L and Yan B et al. Nitrogen-doped Ti3C2 MXene: mechanism investigation and electrochemical analysis. Adv Funct Mater 2020; 30. 2000982.
20. Sugahara A, Ando Y and Kajiyama S et al. Negative dielectric constant of water confined in nanosheets. *Nat Commun* 2019; 10: 850.
21. Gao Q, Sun W and Ilani-Kashkouli P et al. Tracking ion intercalation into layered Ti$_3$C$_2$ MXene films across length scales. *Energy Environ Sci* 2020; 13: 2549–58.
22. Sun Y, Zhan C and Kent P-R-C et al. Proton redox and transport in MXene-confined water. *ACS Appl Mater Interfaces* 2020; 12: 763–70.
23. Augustyn V and Gogotsi Y. 2D materials with nanoconfined fluids for electrochemical energy storage. *Joule* 2017; 1: 443–52.
24. Li S, Yu C and Yang J et al. A superhydrophilic ‘nanoglu’ for stabilizing metal hydroxides onto carbon materials for high-energy and ultralong-life asymmetric supercapacitors. *Energy Environ Sci* 2017; 10: 1958–65.
25. Le T-A, Bui Q-V and Tran N-Q et al. Synergistic effects of nitrogen doping on MXene for enhancement of hydrogen evolution reaction. *ACS Sustainable Chem Eng* 2019; 7: 16879–88.
26. Wen Y, Rufford T-E and Chen X et al. Nitrogen-doped Ti$_3$C$_2$Tx MXene electrodes for high-performance supercapacitors. *Nano Energy* 2017; 38: 368–76.
27. Wu E, Wang J and Zhang H et al. Neutron diffraction studies of Ti$_2$Si$_{1.9}$Al$_{0.1}$C$_2$ compound. *Mater Lett* 2009; 59: 2715–9.
28. Ardizzone S, Fregonara G and Trasatti S. ‘Inner’ and ‘outer’ active surface of RuO$_2$ electrodes. *Electrochim Acta* 1980; 35: 263–7.
29. Al-Temimy A, Prenger K and Golnak R et al. Impact of cation intercalation on the electronic structure of Ti$_3$C$_2$Tx MXenes in sulfuric acid. *ACS Appl Mater Interfaces* 2020; 12: 15087–94.
30. Osti N-C, Naguib M and Ostadhossein A et al. Effect of metal ion intercalation on the structure of MXene and water dynamics on its internal surfaces. *ACS Appl Mater Interfaces* 2016; 8: 8859–63.
31. Raviv U, Laurat P and Klein J. Fluidity of water confined to subnanometre films. *Nature* 2001; 413: 51–4.
32. Wittekindt C and Marx D. Water confined between sheets of mackinawite FeS minerals. *J Chem Phys* 2012; 137: 054710.