High-Performance Ultrathin Flexible Solid-State Supercapacitors Based on Solution Processable Mo1.33C MXene and PEDOT:PSS

Qin Leiqiang, Quanzheng Tao, Ahmed El Ghazaly, Julia Fernandez-Rodriguez, Per OA Persson, Johanna Rosén and Fengling Zhang

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Processable Mo$_{1.33}$C MXene and PEDOT:PSS

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Keywords: solid-state supercapacitor, MXene, Mo$_{1.33}$C, PEDOT:PSS, composite film

MXenes, a young family of two-dimensional (2D) transition metal carbides/nitrides, show great potential in electrochemical energy storage applications. Herein, a high performance ultra-thin flexible solid-state supercapacitor is demonstrated based on a Mo$_{1.33}$C MXene with vacancy-ordering in an aligned layer structure MXene/PEDOT:PSS composite film post-treated with concentrated H$_2$SO$_4$. The flexible solid-state supercapacitor delivers a maximum capacitance of 568 F cm$^{-3}$, an ultrahigh energy density of 33.2 mWh cm$^{-3}$ and a power density of 19470 mW cm$^{-3}$. The Mo$_{1.33}$C MXene/PEDOT:PSS composite film shows a reduction in resistance upon H$_2$SO$_4$ treatment, a higher capacitance (1310 F cm$^{-3}$) and improved rate-capabilities than both pristine Mo$_{1.33}$C MXene and the non-treated Mo$_{1.33}$C/PEDOT:PSS composite films. The enhanced capacitance and stability are attributed to the synergistic effect of increased interlayer
spacing between Mo$_{1.33}$C MXene layers due to insertion of conductive PEDOT, and surface redox processes of the PEDOT and the MXene.

1. Introduction
With the fast growing demand for portable and wearable electronic devices, flexible lightweight high-performance energy storage devices are urgently needed.\cite{1} Among various energy storage devices, supercapacitors have attracted tremendous attention due to their outstanding features, such as high power density, ultra-fast charge and discharge rate, low maintenance, long cycle life, excellent stability and safe operation. These features make supercapacitors promising candidates for the next generation of energy storage applications such as portable electronics, power back-up systems and hybrid vehicles.\cite{2} Based on their charge storage mechanisms, supercapacitors can be divided into two groups; electric double-layer capacitors (EDLCs), where the charge is stored via electrosorption of ions, or pseudocapacitors, where surface redox reactions enable charge storage.\cite{3} Current limitation of commercially available supercapacitors is an energy density much lower than those of batteries and fuel cells, which is a hot research topic in search the next generation of energy storage devices. One strategy to increase the energy density of supercapacitors is to use pseudocapacitive materials, such as metal oxides or electrochemically active organic molecules/polymers as electrodes.\cite{4}

In the past decade, 2D materials have been extensively studied for diverse applications due to their excellent electronic, mechanical and optical properties.\cite{5} One example is MXenes, a young family of 2D transition metal carbides, carbonitride and nitrides with metallic conductivity and hydrophilic surfaces. They are produced by selective etching of the A layers in the atomically laminated M$_{n+1}$AX$_n$ phases,\cite{6} where M is a transition metal, A is an A-group element (mostly Al), and X is C and/or N. MXenes have already shown great potential for applications in Li-ion batteries,\cite{7} Li-S batteries,\cite{8} metal ion capacitors,\cite{9} and aqueous supercapacitors.\cite{10} An increased MXene surface area upon delamination of the multilayered
MXenes into single- or few-layer nanosheets, together with a pseudocapacitive charge storage mechanism and a high conductivity, gives capacitance values outperforming carbon electrodes. Most recently, we reported a route for synthesis of vacancy-ordered Mo$_{1.33}$C MXene, with a resulting capacitance of 1153 F cm$^{-3}$, and a higher conductivity of 29674 s m$^{-1}$ for a MXene film.$^{[11]}$ Furthermore, previous research have shown that an improved charge transport in MXene supercapacitor electrodes can be facilitated by an increase in the interlayer spacing of Ti$_3$C$_2$T$_x$ MXene by using intercalation of a polymer (PVA and PDDA).$^{[12]}$ In addition, the structure of the macroporous electrode was constructed by using MXene hydrogel enable exceptional high-rate performance by fast ion transport and reaching the highest volumetric capacitance of ~1500 F cm$^{-3}$. In the light of these promising results, attention is directed towards poly(3,4-ethylenedioxythiophene):poly(styrenesulfonic acid) (PEDOT:PSS) (Scheme S1), one of the most widely used solution processable conducting polymers, shown to be of great significance for both fundamental research and commercial applications. PEDOT prepared by electrochemical polymerization or chemical oxidation is insoluble and difficult for solution processing. To make PEDOT soluble, PSS is added as a surfactant and dopant to promote the suspension of PEDOT in aqueous solutions. In addition, the complex of PEDOT:PSS possesses outstanding advantages of excellent solution processing properties, good film-forming properties, high flexibility, and excellent thermal stability, especially the tunable and improved conductivity achieved by physical or chemical treatment. It is also worth noting that the conductivity of PEDOT:PSS can reach a maximum value of 4380 S cm$^{-1}$ via H$_2$SO$_4$ post-treatment, four orders of magnitude larger than untreated PEDOT:PSS films, which is impossible for pure PEDOT.$^{[14]}$ Therefore, PEDOT:PSS is the most widely studied conducting polymer as a supercapacitor material for its high stability, high conductivity in the doped state, and its fast redox protonation/deprotonation reactions. Combining the outstanding properties of the MXene and the polymer suggests a potential unique electrode architecture of
improved stability and elasticity for solution processed flexible high performance stable solid-state supercapacitors.

To date, the few reports about MXene and polymer composites are limited to the evaluation of electrode materials for supercapacitor applications, and the composites have not yet been applied to devices.\[15\] Herein, we demonstrate high performance ultra-thin solid-state supercapacitors based on flexible aligned Mo\(_{1.33}\)C MXene/PEDOT:PSS composite films obtained from solution processable Mo\(_{1.33}\)C MXene and PEDOT:PSS hybrid ink followed by immersion in concentrated H\(_2\)SO\(_4\) solutions for 24 hours. The representative flexible solid-state supercapacitor exhibits a capacitance up to 568 F cm\(^{-3}\), an ultrahigh energy density of 33.2 mWh cm\(^{-3}\), an excellent cycling stability and outstanding deformation tolerance due to intercalating conductive PEDOT:PSS chains in the MXene/PEDOT:PSS composite films.

2. Results and Discussion

The procedures for preparing the composite films and fabricating the solid-state supercapacitors are schematically illustrated in Figure 1 (see details in the Experimental Section). In order to get the optimal Mo\(_{1.33}\)C MXene to PEDOT:PSS ratio, a series of flexible composite films are obtained by vacuum filtration of the Mo\(_{1.33}\)C MXene and PEDOT:PSS dispersion in mass ratios of 50:1, 20:1, 10:1, 5:1, 2:1, 1:1, 1:2, and 1:5. Hereafter the flexible composite films will be referred as M:P (50:1, 20:1, 10:1, 5:1, 2:1, 1:1, 1:2, 1:5). Pure Mo\(_{1.33}\)C MXene and pure PEDOT:PSS "papers" are also prepared for comparison. The electrochemical properties of the different as-fabricated composite electrodes are first inspected using cyclic voltammetry (CV) experiments in 1 M H\(_2\)SO\(_4\) at various sweep rates in the potential window from -0.35 to 0.3 V vs. Ag/AgCl (Figure S1). The volumetric capacitances of ten different composite electrodes are summarized in Table 1 where the pristine Mo\(_{1.33}\)C film exhibits the highest volumetric capacitance of 1187 F cm\(^{-3}\) (at 2 mV/s), followed by 1095 F cm\(^{-3}\) for 20:1, 949 F cm\(^{-3}\) for 10:1, 939 for 50:1 and then a monotonically decrease with increasing the loading of PEDOT:PSS.
To increase the conductivity of PEDOT:PSS in the composite films, all samples are immersed into concentrated H$_2$SO$_4$ for 24 hours as reported in the literature.[14] Encouragingly, significantly improved conductivity and electrochemical properties are observed for the treated composite films (Figure S2). Notably, the H$_2$SO$_4$ treated composite electrode with a mass ratio of 10:1 (referred to as M:P=10:1-24h) exhibits a maximum volumetric capacitance of 1310 F cm$^{-3}$ (2 mV s$^{-1}$, 452 F g$^{-1}$), which is superior to the literature reports on MXene based electrodes.[6,9,10] Consequently, this investigation is from this point forward focused on the M:P=10:1-24h films. The pristine Mo$_{1.33}$C films and M:P=10:1 (without H$_2$SO$_4$ post-treatment) will also be investigated for reference. To study the effect of PEDOT:PSS and H$_2$SO$_4$ treatment on the capacitance of the MXene films, the capacitances of the three electrodes (Mo$_{1.33}$C, M:P=10:1 and M:P=10:1-24h) are recorded by CV at the scan rate of 2 mV s$^{-1}$. It is clear that the M:P=10:1-24h has the highest capacitance, while M:P=10:1 has the lowest capacitance, see Figure 2A. The reduced capacitance of M:P=10:1 may be attributed to the known electrochemically inactivity of PSS in the PEDOT:PSS. This is supported by Table 1, showing that increasing PEDOT:PSS loading results in a decline in capacitances due to electrochemical inactivity and poor conductivity of PSS. As previously reported,[14] when the composite electrodes are treated with concentrated H$_2$SO$_4$, the dense PEDOT networks will be significant changed in both the crystallographic and morphological structures via the formation of crystalline nanofibril structures. This is accompanied by excess uncoupled PSS being removed. [14,16] In the present work a significant removal of the PSS content is confirmed by FT-IR and absorption spectra (Figure S3). Interaction between the presumed negatively charged Mo$_{1.33}$C Mxene[17] and the positively charged PEDOT is expected, which in turn is likely to influence charge transport as well as charge storage capabilities. Furthermore, the PEDOT nanofibril network enlarges the interlayer spacing between the MXene flakes and likely facilitate efficient intra- and inter-chain charge transport, potentially also increasing the accessibility to deep traps, which may enhance the net capacitances of the composite electrodes.
The nearly triangular galvanostatic charge/discharge curves, see Figure 2B, of the M:P=10:1-24h composite film at different current densities confirm the high reversibility of the redox reactions and good coulomb efficiency (~96%) of the tested electrodes. A comparison of the rate performance of the M:P=10:1 composite film with pristine Mo$_{1.33}$C is presented in Figure 2C. The volumetric capacitance of the M:P=10:1-24h is higher than the untreated composite film and the pristine Mo$_{1.33}$C at various sweep rates. In addition, the post-treated composite film shows a good rate performance. Even at a scan rate of 1000 mV s$^{-1}$, it yields a high capacitance of 589 F cm$^{-3}$. This may be due to dual redox contributions from the MXene and the PEDOT, and the aligned PEDOT nanofibril network between the metallic Mo$_{1.33}$C layers providing conductive channels for facile ionic transport. Electrochemical impedance spectroscopy was performed to study the charge transfer resistance ($R_{ct}$) and ion transport in the tested electrodes. Nyquist plots, see Figure 2D, confirms fast ion transport in the M:P=10:1-24h composite film. At low frequency, a large slope close to 90° is present in all films, which may be attributed to the highly accessible surface of the layer-stacked film. Furthermore, in the high-frequency region, the M:P=10:1-24h composite film shows a quite small charge transport resistance with an invisible semicircle, emphasizing the superiority of the composite films.

For structural characterization, the cross-sections of the three films (Mo$_{1.33}$C, M:P=10:1 and M:P=10:1-24h) are imaged by scanning electron microscopy (SEM) and shown in Figure 3. The pristine Mo$_{1.33}$C film exhibits the well-aligned stacked MXene sheets (Figure 3A), and maintains a good layered structure after treated by concentrated H$_2$SO$_4$ (Figure S4). The M:P=10:1 composite film display a less clear layer stacking (Figure 3B). This can be attributed to different mechanical properties, resulting in a different appearance upon breaking the paper. The laminated structure is, however, again evident after H$_2$SO$_4$ treatment (Figure 3C). The X-ray diffraction (XRD) patterns (Figure 3D) of the M:P=10:1 composite film shows a major peak at 8°, which originates from the Mo$_{1.33}$C MXene. For the M:P=10:1 sample it can be assumed that stacks of a few layer Mo$_{1.33}$C MXene sheets are interleaved with a polymer layer, where
the latter is indicated from the bumps at ~5° and ~26°. The treated M:P=10:1-24h film exhibit a clear downshift in the (001) peak from ~8.4° to ~6.8°, indicating an increase of ≈2.4 Å in the spacing between the Mo$_{1.33}$C sheets. This shift is attributed to water and/or polymer intercalation in Mo$_{1.33}$C MXenes.[18] However, the previously shown crystallographic structures of the PEDOT after treatment (Figure S5) cannot be distinguished by XRD. Transmission Electron microscopy (TEM) results confirm the laminated microstructure of M:P=10:1-24h as shown in SEM. Figure 3E shows a cross sectional image (top to bottom) of the M:P=10:1-24h. The higher magnification image of the layered structure in Figure 3F reveals that the Mo$_{1.33}$C MXene sheets (dark lines) appear to be separated into individual 2D sheets. Ignoring local disturbances, somewhat regular interplanar spaces of the sheets can be observed, as caused by the conductive polymer (bright lines).

In an evaluation of mechanical properties, the nanohardness (H) and Young’s modulus (E) of the pristine Mo$_{1.33}$C film is found to be 0.47±0.15 GPa and 6±1.8 GPa, respectively. This can be compared to Ti$_3$C$_2$T$_x$ MXene, with an E of ~3 GPa.[11,19] Adding PEDOT:PSS with an E of 3.3±0.1 GPa[20], results in a reduction of H as well as E to 0.11±0.006 and 4.9±0.17 GPa, respectively (Figure S6). A small deviation in the H and E values for the composite film illustrate a homogeneous distribution of PEDOT:PSS and Mo$_{1.33}$C MXene, as confirmed by the TEM analysis.

To further explore the potential of MXene-polymer composite films for energy storage, flexible solid-state supercapacitors are fabricated based on M:P=10:1-24h. Similar to the composite films, the corresponding devices show superior mechanical flexibility, and can be bent or twisted without damaging the structure and morphology. The typical thickness of the entire device is ~67 μm, which is thinner than a commercial standard printer paper. The performance of the all-solid-state supercapacitor is investigated based on CV curves (Figure 4A and Figure S7,8). For M:P=10:1-24h, the shapes of the CV curves remained pseudorectangular, even at very high scan rates (1000 mV s$^{-1}$), which indicates a low resistance
and good reversibility. Figure 4B shows the galvanostatic charging/discharging curves of the M:P=10:1-24h device. The near triangular shapes of these curves indicate good reversibility of the charging/discharging process, which is in good agreement with the CV curves shown in Figure 4A. The current density dependence of the volumetric capacitance can be calculated from the discharge curves of the flexible supercapacitor. The capacitance of the device reaches 568 F cm\(^{-3}\) at a current density of 0.5 A cm\(^{-3}\). Similar to the results from a three-electrode system (Figure 3C), the M:P=10:1-24h device shows superior rate performance compared with supercapacitor devices made from pristine Mo\(_{1.33}\)C and M:P=10:1 (Figure 4C). With a current density increase, the volumetric capacitance of the M:P=10:1-24h device remains high. Even at the highest current density of 30 A cm\(^{-3}\), the capacitance is 421.2 F cm\(^{-3}\). This outstanding charging/discharging rate stability of the M:P=10:1-24h device may be explained by the aligned layer structure formed by the intercalated PEDOT:PSS nanofibrils between the Mo\(_{1.33}\)C sheets, which increases the layer spacing and facilitates ion transport between the MXene sheets. To show further the merits of the composite film as the electrode material, the electrochemical impedance spectra (EIS) of devices from M:P=10:1-24h, M:P=10:1, and pristine Mo\(_{1.33}\)C are compared in Figure 4D. It can be seen that the complex plane plots of M:P=10:1-24h and pristine Mo\(_{1.33}\)C show a larger slope of close to 90° compared to M:P=10:1 at a low frequency, which indicates fast ion diffusion. At the high frequency, see inset, the M:P=10:1-24h shows a smaller charge-transport semicircle than M:P=10:1 and pristine Mo\(_{1.33}\)C, which is consistent with the result that the crystalline PEDOT:PSS nanofibril network enhance the conductivity of the composite film upon H\(_2\)SO\(_4\) treatment.

To demonstrate the application prospects of the composite films of M:P=10:1-24h, the devices are directly integrated in series and/or parallel connections to increase the output voltage or total capacitance (Figure 5A and B). Figure 5C shows that an applied voltage of \(\approx 2.6\) V obtained from four integrated all-solid-state supercapacitors in series, can essentially power a LED for tens of seconds. Furthermore, the cycling stability of the fabricated all-solid-
state supercapacitors are investigated by galvanostatic charge/discharge tests at the current of 5 A cm$^{-3}$ (Figure 5D). After 10000 cycles, the M:P=10:1-24h device exhibit a capacitive retention of 90%, which compares favorably to the M:P=10:1 (80%), and pristine Mo$_{1.33}$C (49%) device. The latter value can be compared to the previously reported 84% retained capacitance for a Mo$_{1.33}$C film.[11] The morphology of the M:P=10:1-24h device after 10 000 cycles is almost maintaining the aligned structure while the pristine Mo$_{1.33}$C device changed significantly after cycling (Figure S9). Two main reasons for the much improved cycling performance of the post-treated composite electrodes compared with the untreated composite electrodes or pristine Mo$_{1.33}$C electrode are suggested: First, after treatment with concentrated H$_2$SO$_4$, the PEDOT:PSS nanofibril network will strengthen the composite films and suppress the volume change and counter-ion draining during the charging/discharging process. Second, the M:P=10:1-24h device exhibit lower internal resistance and charge transport resistance, which has been discussed in detail in the EIS discussion part. This means that the M:P=10:1-24h films will display smaller charge loss during long-term cycling, which further increases the cycling stability. Furthermore, the devices based on M:P=10:1-24h shows exceptional electrochemical stability under different bending angles. It is shown that CV curves from different bending angles show nearly the same capacitive behavior (Figure 5E). Also, the devices of M:P=10:1-24h shows lower self-discharge, it is decaying to above 0.35 V after 100 h (Figure S11).

Energy density is a critical parameter used to compare different types of energy storage systems, with a high volumetric energy density being particularly important for supercapacitors. The volumetric energy and power densities of the M:P=10:1-24h devices, as well as a comparison with previously reported devices and commercial supercapacitors, are plotted in the Ragone plot in Figure 5F. A maximum energy density of 33.2 mWh cm$^{-3}$ obtained in M:P=10:1-24h devices is higher than that of most recently reported electric double-layer capacitors or pseudocapacitors.[21] This high energy density is likely due to the dual fast pseudocapacitive behavior of the Mo$_{1.33}$C MXene and the PEDOT:PSS. Also, the M:P=10:1-24h
24h device can maintain a surprisingly high energy density of 24.72 mWh cm$^{-3}$ at the high power density of 19470 mW cm$^{-3}$. The outstanding performance of the M:P=10:1-24h likely originates from highly conducting Mo$_{1.33}$C MXene sheets with pseudocapacitive charge storage mechanisms being separated by a nanofibril structure from H$_2$SO$_4$ treated PEDOT:PSS. This, in turn facilitate MXene-PEDOT interaction, which seems to improve charge storage (active surface area) and effectively promote electron and ion transport, both inter- and intra-layer, during the charge/discharge cycle. Finally, the PEDOT network is likely to minimize the volume change and counter-ion draining during the charging/discharging process, which greatly improve the long-term performance of the composite electrodes.

### 3. Conclusion

In summary, we demonstrate an effective strategy to realize a high-performance ultra-thin flexible solid-state supercapacitor by using aligned layer structured composite films as active electrodes, obtained from vacuum filtrated Mo$_{1.33}$C MXene and PEDOT:PSS based solution processable aqueous suspensions. The flexible solid-state supercapacitors exhibit a maximum capacitance of 568 F cm$^{-3}$, an ultrahigh energy density of 33.2 mWh cm$^{-3}$ and a power density of 19470 mW cm$^{-3}$. We attribute the excellent electrochemical performance and cycle life to the self-assembled layered architecture with aligned PEDOT nanofiber network confined between the highly conducting Mo$_{1.33}$C layers, which leads to fast reversible redox reactions and improved ion transport through short diffusion paths. The results of XRD and TEM indicates that the higher volumetric capacitances (1310 F cm$^{-3}$) and excellent rate performances compared to pure Mo$_{1.33}$C and untreated Mo$_{1.33}$C/PEDOT:PSS composite films is facilitated by the formation of PEDOT nanofibers in the sandwich-like Mo$_{1.33}$C/PEDOT:PSS composite film after treatment with concentrated H$_2$SO$_4$. Furthermore, the enhanced capacitance is likely due to the synergistic effect of increased interlayer spacing between the Mo$_{1.33}$C layers and the surface redox processes of both the PEDOT and MXene. Our demonstrated results provide a
4. Experimental Section
Synthesis of the Mo$_{1.33}$C MXene was described previously.\cite{10} Briefly, 1 g of (Mo$_{2/3}$Sc$_{1/3}$)$_2$AlC is added to 20 ml 48% HF, stirring for 24 h at RT. After removing the acid, multilayer MXene is delaminated into single/few layer MXene by intercalating with TBAOH. The Mo$_{1.33}$C/PEDOT:PSS composites is prepared by dropwise addition of a PEDOT:PSS (Clevios PH 1000) into the colloidal solution of Mo$_{1.33}$C. The mixture is stirred for 5 min and then vacuum-filtered onto nanoporous polypropylene membranes (Celgard 3501, 0.064 µm pore size, Celgard LLC) in air. The H$_2$SO$_4$ treatment is performed by soaking the Mo$_{1.33}$C/PEDOT:PSS composite film into the concentrated H$_2$SO$_4$ solution for 24 h at room temperature. Then the treated film is rinsed with deionized water three times. Finally, the composites film is dried at 110 °C for about 3 min.

In the three-electrode configuration, the Mo$_{1.33}$C or the composite film is used as working electrodes, platinum foils as counter electrodes and Ag/AgCl in 1 M KCl as the reference electrode using the aqueous solution containing H$_2$SO$_4$ (1 M) as the electrolyte. The all solid-state supercapacitors is fabricated using Au-coated polyethylene terephthalate (PET) substrate as the flexible conductive substrate and PVA/H$_2$SO$_4$ gel electrolyte as the solid electrolyte (~45 µm). First, 1 g of H$_2$SO$_4$ is mixed with 10 ml deionized water, and then 1 g of polyvinyl alcohol power is added. The mixture is stirred under 85 °C until the mixture becomes clear. When the electrolyte is cooled down, it is cast onto the electrodes (~2 µm), then left in the fume hood at room temperature for 1 h to vaporize the excess water. Then the two pieces of electrodes are pressed together. The electrolyte was solidified and functioned as a glue holding all the device components together, and improving the mechanical stability.
The electrochemical tests are performed using a VSP potentiostat (BioLogic, France). The impedance measurements are performed with a 5 mV amplitude in a frequency range from 10 mHz to 100 kHz. The characterizations of the film morphology are carried out using SEM (LEO 1550 Gemini) and TEM (LEO 912 OMEGA). XRD is carried out on a PANalytical X’Pert diffractometer using Cu Kα radiation (45 KV and 40 mA). A Hysitron TI950 triboindenter device equipped with a Berkovich indenter is used to measure mechanical properties of the specimens. An array of 4×4 indentations with a trapezoid loading profile is performed at 50μN peak load for each specimen. A standard fused silica is used for calibration before each measurement.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

Acknowledgements
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References


**Figure 1.** Schematic illustration of the preparation of composite films and the fabrication of a solid-state supercapacitor.
Figure 2. (A) Volumetric capacitance of pristine Mo$_{1.33}$C and composite films (M:P=10:1) at scan rate 2 mV/s. (B) Galvanostatic charge/discharge curves of M:P=10:1-24h at different current densities. (C) Gravimetric rate performances of pristine Mo$_{1.33}$C and composite films. (D) Electrochemical impedance spectroscopy of pristine Mo$_{1.33}$C and composite films.
Figure 3. Cross-sectional SEM image of (A) pristine Mo$_{1.33}$C, (B) M:P=10:1 and (C) M:P=10:1-24h. Scale bars: 2 µm. (D) XRD pattern of the PEDOT:PSS intercalated MXene compared with delaminated MXene film. (E) Low-magnification (scale bar: 200 nm) and (F) high-magnification (scale bar: 40 nm) cross-sectional TEM images of M:P=10:1-24h with aligned PEDOT:PSS chains between MXene sheets. Inset of (C): Digital photographs showing a flexible and free-standing sandwich-like MXene/PEDOT:PSS paper wrapping of around a 5-mm-diameter glass rod.
Figure 4. (A) Typical cyclic voltammograms (inset: thickness of the as-prepared flexible solid-state supercapacitor) and (B) galvanostatic charge/discharge curves of the flexible solid-state device. (C) The comparison of volumetric capacitances of Mo$_{1.33}$C, M:P=10:1 and M:P=10:1-24h at different current density. (D) Nyquist plot of the device tested at the open-circuit potential within the frequency range from $10^{-2}$ to $10^5$ Hz. The inset shows the enlarged plot in the high frequency region.
Figure 5. (A) CV curves of devices (M:P=10:1-24h) connected in series and in parallel at a scan rate of 100 mV s⁻¹. A single device is shown for comparison. (B) GCD curves of devices (M:P=10:1-24h) connected in series and in parallel at a current density of 1 A cm⁻². (C) Optical image of a light-emitting diode (LED) powered by four tandem devices. (D) Cyclic stability of Mo₁.₃₃C, M:P=10:1 and M:P=10:1-24h devices. (E) CV curves of the M:P=10:1-24h device bended with different angles at 100 mV s⁻¹. Inset: Optical image of the flexible ultra-thin supercapacitor. (F) Energy and power density of the M:P=10:1-24h device compared with previously reported devices.

Table 1. The capacitance performance of the composite film for different MXene:polymer mass ratio

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<th>5 mV/s [F cm⁻³]</th>
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a) M=Mo₁.₃₃C, P=PEDOT:PSS; b) The film did not undergo any treatment; c) The film was treated with concentrated H₂SO₄ for 24 hours.
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A MXene-based solution processable flexible solid-state supercapacitor with high performance is developed from a MXene/PEDOT:PSS composite film. After post-treatment with concentrated H₂SO₄, the PEDOT nanofiber network is aligned between the MXene sheets, leading to highly improved flexibility and, most importantly, improved capacitances (1310 F cm⁻³), rate-capabilities and stability.

Keyword
solid-state supercapacitor, MXene, Mo₁.₃₃C, PEDOT:PSS, composite film

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High Performance Ultrathin Flexible Solid-State Supercapacitors based on Solution Processable Mo₁.₃₃C MXene and PEDOT:PSS
Supporting Information

High Performance Ultrathin Flexible Solid-State Supercapacitors based on Solution Processable Mo$_{1.33}$C MXene and PEDOT:PSS

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The volumetric capacitance from the cyclic voltammetry data is determined from the following equation (1):

\[ C = \frac{1}{\Delta V} \int \frac{j \, dV}{s} \]  

(1)

C is the normalized capacitance (in units of F cm\(^{-3}\)), \( j \) is the current density (in A cm\(^{-3}\)), \( s \) is the rate (in V s\(^{-1}\)), \( V \) is the voltage (in V), \( \Delta V \) is the voltage window (in V). Calculations of the gravimetric capacitance (in F g\(^{-1}\)) is performed using current density per electrode weight (in A g\(^{-1}\)).

The volumetric specific capacitance from galvanostatic charge/discharge data is calculated from the following equation (2):

\[ C = \frac{2It}{\Delta V \, v} \]  

(2)

where \( I \) and \( t \) represent the discharge current (A) and time (s), respectively; \( \Delta V \) is the voltage during the discharge process after iR drop (V), \( v \) is the volume of one electrode or device.

The volumetric energy density of the device is obtained from the formula given in Equation (3):

\[ E = \frac{1}{2} \times C_{volumetric}^{device} \times \frac{(\Delta V)^2}{3600} \]  

(3)

where \( E \) is the energy density (in Wh/cm\(^{3}\)), \( C_{volumetric}^{device} \) is the volumetric capacitance obtained from Equation (1) and \( \Delta V \) is the discharge voltage range (in volts, V).

The power density of the device is calculated from the formula given in Equation (4):

\[ P = \frac{E}{\Delta t} \times 3600 \]  

(4)

where \( P \) is the power density (in W/cm\(^{3}\)), \( E \) is the volumetric energy density obtained from Equation (3) and \( \Delta t \) is the discharge time (in seconds, s).
**Scheme S1.** The chemical structure of PEDOT:PSS.

FTIR and Absorption spectra were used to estimate the relative PEDOT/PSS ratio in the composite. For the sample preparation, the PEDOT:PSS was spin coated on quartz. The composite film of M:P=10:1 was drip casted on the quartz. The samples of PEDOT:PSS and M:P=10:1 were soaked in concentrated H₂SO₄ for about 5 h, denoted as PEDOT:PSS-SA and M:P=10:1-SA.

FT-IR spectra of PEDOT:PSS and the composite films of M:P=10:1 are shown in Figure S4A. The PEDOT:PSS spectrum shows a strong peak at 1367 cm⁻¹ which is attributed to the aromatic ring vibration in the thiophene ring. Bands at 1261 cm⁻¹ are attributed to the C–O–C stretching modes and the peak at 858 cm⁻¹ is related to C–S stretching vibration of the EDOT ring. The peaks at 1124 and 1161 cm⁻¹ are attributed to the stretching vibrations of the merged aromatic ring of PSS. Here we use the peak of 1261 cm⁻¹ belonging to PEDOT as a standard to compare the contention of PSS before and after treatment with concentrated H₂SO₄. It is shown that the H₂SO₄ treatment induces a change in the PEDOT:PSS composition ratio, which is revealed by the different transmittance observed for the different samples. After the H₂SO₄ treatment, the peak intensity at 1161 cm⁻¹ is significantly reduced, whereas the peak at 1261 cm⁻¹ exhibits almost no change. These observations clearly demonstrate that the H₂SO₄ treatment induces the selective removal of PSS without influencing the PEDOT part in PEDOT:PSS. When the M:P=10:1 is treated, the peak intensity at 1161 cm⁻¹ decreases further than the treated PEDOT:PSS, which means that more PSS is being removed. This conclusion is in consistent with the weight loss after treatment.

The PEDOT/PSS ratio change after H₂SO₄ treatment is also revealed by the absorption spectra. As shown in Figure S4B, the strong absorption features in the UV range are attributed to the phenyl moieties in PSS, and the broad absorption band in the visible and IR regions is associated with the free charge carriers in PEDOT. When the H₂SO₄ treatment, two strong absorption peaks in the UV range are significantly reduced, whereas the absorption features with a lower
energy (>500 nm) exhibit almost no change. These observations clearly demonstrate that the H$_2$SO$_4$ treatment induces the selective removal of PSS without influencing the PEDOT part in PEDOT:PSS. For M:P=10:1, when treated with H$_2$SO$_4$, a more prominent reduction of the peak in the UV region can be seen comparing to treated PEDOT:PSS. Combining the FT-IR and absorption spectra, we conclude that the PEDOT to PSS ratio in the treated composite film is larger than the treated PEDOT:PSS.

From previous report, a certain amount of negatively charged PSS is necessary for the charge balance of the positively charged PEDOT, and for the formation of the highly crystalline form of PEDOT:PSS (PEDOT/PSS ratio ~2) upon H$_2$SO$_4$ treatment.[2] In the current study, we don’t observe the formation of the highly crystalline form of PEDOT:PSS based on XRD, likely due to the large PEDOT:PSS ratio (PEDOT/PSS ratio >2) in the composite. We speculate that the presumed negatively charged MXene may, at least in part, take the role of PSS, so most of the PSS is removed from the composite film after H$_2$SO$_4$ treatment.

**Figure S4.** Cross-sectional SEM image of (A) pristine Mo$_{1.33}$C, (B) Mo$_{1.33}$C-24h, (C) PH1000 and (D) PH1000-24h (scale bar: 1 µm).
At the same time we also study the change of pristine Mo\textsubscript{1.33}C and PEDOT:PSS after treatment by concentrated H\textsubscript{2}SO\textsubscript{4}. For Mo\textsubscript{1.33}C, as shown in the scanning electron microscopy (SEM) images (Figure S4A and B), the Mo\textsubscript{1.33}C maintains the layered structure after treatment. In addition, the X-ray diffraction (XRD) patterns (Figure S5A) of the pristine Mo\textsubscript{1.33}C film show a major peak at $8.4^\circ$. The Mo\textsubscript{1.33}C films treated with concentrated H\textsubscript{2}SO\textsubscript{4} for 24 hours (Mo\textsubscript{1.33}C-24h) exhibit a clear shift in the (001) peak from $\sim 8.4^\circ$ to $\sim 7.1^\circ$, indicating an increase in the spacing between the Mo\textsubscript{1.33}C sheets. This shift may be attributed to water or proton intercalation.

Table 1 (in the manuscript) shows that the capacitance of Mo\textsubscript{1.33}C after treatment with concentrated H\textsubscript{2}SO\textsubscript{4} only display a small drop compared to untreated Mo\textsubscript{1.33}C.

The cross-sections of the PH1000 and PH1000-24h are imaged by SEM and shown in Figure S4C and D. The PH1000 film displays a uniform compact section, with no clear microstructure. However, after treatment with concentrated H\textsubscript{2}SO\textsubscript{4} for 24 hours, the PH1000-24h film displays a laminated structure. This can be attributed to an excessive amount of uncoupled PSS being removed by H\textsubscript{2}SO\textsubscript{4}. In this process, the strong $\pi-\pi$ stacking of PEDOT and the rigidity of its backbone induce dense PEDOT networks with significant changes in both the crystallographic and morphological structures via the formation of a crystalline nanofibril laminated structure.

The XRD patterns (Figure S5B) of PH1000-24h film shows a major sharp peak at $6^\circ$, however the untreated PH1000 did not show clear peak, which indicates that concentrated H\textsubscript{2}SO\textsubscript{4}
treatment significantly improved the crystallinity of the PH1000 films. This is consistent with
the results from SEM. In addition, after treatment with concentrated H₂SO₄, the capacitance of
PH1000-24h was significantly improved compared to PH1000 (Table 1 in the manuscript). The
above results show that the treatment with concentrated H₂SO₄ has little effect on Mo₁.₃₃C, but
significantly improves the crystallinity and capacitance of PH1000.

Figure S6. The mechanical properties of the Mo₁.₃₃C and composite film.

Table S1. The electrical conductivity of pristine Mo₁.₃₃C and composite film was measured
with a 4-point-probe.

We are aware of the limitations of this technique for analysis of composite materials, however,
for the present study we focus on the qualitative trend in conductivity before/after H₂SO₄
treatment.

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<td>σ [S m⁻¹]</td>
<td>129</td>
<td>121</td>
<td>1843</td>
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a) The high conductivity (29674 S m⁻¹) of Mo₁.₃₃C reported by Tao et al. [3] was measured with a Physical
Property Measurement System (Quantum Design, San Diego). Here, four-point probe (Jandel RM3000
Resistivity Meter) was used to measure the RT resistivity of the composite films and pristine Mo₁.₃₃C.
Figure S7. (A) Cyclic voltammetry (CV) profiles of the device (pristine MXene) obtained at different scanning rates. (B) Galvanostatic charge/discharge (GCD) curves of the device (pristine MXene) at different current densities.

Figure S8. (A) Cyclic voltammetry (CV) profiles of the device (M:P=10:1) obtained at different scanning rates. (B) Galvanostatic charge/discharge (GCD) curves of the device (M:P=10:1) at different current densities.
Figure S9. Cross-sectional SEM image of the device after cycling test. (A) M:P=10:1-24h, (C) M:P=10:1 and (E) pristine Mo$_{1.33}$C. The left side SEM image (B, D and F) is an enlarged view of the active materials part of the device.
After cycling, the Mo$_{1.33}$C MXene/PEDOT:PSS devices were studied with SEM (Figure S9) and XRD (Figure S10, which also includes the three films before cycling for comparison). Figure S9 shows cross-sectional SEM image of the device ((A) M:P=10:1-24h, (C) M:P=10:1 and (E) pristine Mo$_{1.33}$C), respectively. After 10000 cycles, the morphology of the M:P=10:1-24h and M:P=10:1 films (Figure S9B and D) almost maintain the aligned structures to those before the cycling, indicating that the Mo$_{1.33}$C were effectively stabilized by the PEDOT nanofiber network. However, the layered structure of the pristine Mo$_{1.33}$C disappeared as shown in Figure S9F, indicated that the morphology of the pristine Mo$_{1.33}$C film significantly changed. The XRD patterns (Figure S10) of the pristine Mo$_{1.33}$C after cycling (Pristine Mo$_{1.33}$C-c) shows a reduced peak at 8 degrees, which indicates that the order of Mo$_{1.33}$C has decreased. Also, the main peak for M:P=10:1-24h did not shift, suggesting a stable interlayer distance. Furthermore, a new peak appeared at 10.8 degrees for all devices after cycling, which may originate from oxidation of some of the Mo$_{1.33}$C MXene, to form MoO$_x$, during the cycling. The above data suggest that the stabilization of the composite film may be attributed to the constraint of intercalated PEDOT nanofiber network during the cycling process and the strong bond with the Mo$_{1.33}$C sheets.
Figure S11. Self-discharge behaviour of M:P=10:1-24h composite device.

Figure S12. The cycling stability of the fabricated all-solid-state supercapacitors by CV at the scan rate of 100 mV s⁻¹. (A) Pristine Mo₁.₃₃C. (B) M:P=10:1. (C) M:P=10:1-24h.
Figure S13. (A) Cyclic voltammetry (CV) profiles of PH1000 device obtained at different scanning rates. (B) Galvanostatic charge/discharge (GCD) curves of PH1000 device at different current densities.

Figure S14. (A) Cyclic voltammetry (CV) profiles of PH1000-24h device obtained at different scanning rates. (B) Galvanostatic charge/discharge (GCD) curves of PH1000-24h device at different current densities.
**Figure S15.** Nyquist plot of the device tested at the open-circuit potential within the frequency range from $10^{-2}$ to $10^5$ Hz. The inset shows the enlarged plot in the high frequency region.

Reference: