RAPID COMMUNICATION

Controlling the actuation properties of MXene paper electrodes upon cation intercalation

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Abstract

Atomic force microscopy was used to monitor the macroscopic deformation in a delaminated Ti3C2 paper electrode in situ, during charge/discharge in a variety of aqueous electrolytes to examine the effect of the cation intercalation on the electrochemical behavior and mechanical response. The results show a strong dependence of the electrode deformation on cation size and charge. The electrode undergoes a large contraction during Li+, Na+ or Mg2+ intercalation, differentiating the Ti3C2 paper from conventional electrodes where redox intercalation of ions (e.g. Li+) into the bulk phase (e.g. graphite, silicon) results in volumetric expansion. This feature may explain the excellent rate performance and cyclability reported for MXenes. We

KEYWORDS
MXene; Intercalation; Electromechanical actuator; Atomic force microscopy

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also demonstrated that the variation of the electromechanical contraction can be easily adjusted by electrolyte exchange, and shows interesting characteristics for the design of actuators based on 2D metal carbides.

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Introduction

As the demand for renewable and clean energy sources is ever increasing, interest in electrochemical energy storage devices such as lithium ion batteries and electrochemical capacitors (ECs) has risen in recent years. ECs have received much attention due to their high power densities, but they suffer from low energy densities [1-3]. ECs can be categorized into two types: (i) electric double layer capacitors (EDLCs) that store charge by electrostatic adsorption of ions on the surface of porous conducting materials like carbon, and (ii) pseudocapacitors that store charge by means of fast surface redox reactions [4]. Recently it was shown that fast intercalation of ions between atomic layers of some crystalline transition metal oxides [5] or layered two-dimensional (2D) materials such as the titanium carbide Ti$_3$C$_2$Tx (where Tx stands for surface terminations such as OH, =O and F) [6] can result in high volumetric capacitance. The fast intercalation of ions in these materials happens with no diffusion limitations, which is a key requirement for supercapacitor electrodes. This material is a member of the recently discovered family of 2D materials known as MXenes [7-11]. These materials store energy by rapid ion intercalation between the 2D sheets, as well as Ti$^{4+}$/Ti$^{3+}$ and other surface redox reactions, making them promising ECs electrodes [12,13,41].

In situ X-ray diffraction and Electrochemical Quartz Crystal Admittance (EQCA) measurements during charge/discharge suggested that cation intercalation between MXene sheets in aqueous systems leads to a decrease in the distance between the layers (and the corresponding c-lattice parameter) [6,14]. Depending on the charge/ionic radius ratio of the cation, the screening of electrostatic repulsive forces between charged groups on the Ti$_3$C$_2$Tx sheet surfaces may result in different electrode deformation. Such a feature is appealing for electrochemically-controlled actuators, which convert electrical energy into mechanical energy [15,16]. Moreover, thanks to its metallic electrical conductivity [17], as well as its predicted high elastic constants and bending rigidity [18], Ti$_3$C$_2$Tx MXene appears to be a good candidate for electromechanical applications. Yet, no reports on direct

Figure 1  (a) Scanning electron microscopy image and (b) tapping mode AFM image of the surface of the delaminated Ti$_3$C$_2$Tx paper electrode, (c) picture of the electrochemical cell used for in situ volume change measurements, and (d) cross-section schematic representation of the cell showing how the MXene is electrically connected and how the AFM tip is used to monitor the electrode deformation in the presence of electrolyte.
measurements of the MXene deformation are available, and the effective macroscopic volumetric changes of the electrode during charge and discharge have not been described so far. Furthermore, understanding the intricate interplay of ion insertion, electrode volume changes, and the resulting capacitance can help to optimize and tailor the electrochemical performance and lifetime of electrode materials.

In this work we investigate the macroscopic deformation of the Ti$_3$C$_2$T$_x$ “paper” (additives-free assembly of the few-/single-layer(s) MXene flakes shown in Figure 1a) during cation insertion (charging). Atomic force microscopy (AFM) has the unique advantage of having a Z (height) resolution of <1 nm and high lateral resolution determined by the AFM tip geometry (ca. 20 nm). This technique has been broadly used as an in situ probe to monitor the dimensional changes of battery [19-23], pseudocapacitor [24,25], and EDLC electrodes [26,27]. Another key advantage of these experiments is that they can be performed at various charging and discharging rates. Here, the deformation response of Ti$_3$C$_2$T$_x$ is measured in different aqueous electrolytes to determine the effect of varying the electrolyte cation size and charge on the electrochemical and mechanical response.

Material and methods

MXene synthesis

Delaminated Ti$_3$C$_2$T$_x$ electrode synthesis and preparation are described elsewhere [6]. In short, Ti$_3$C$_2$T$_x$ was synthesized by etching Al from Ti$_3$AlC$_2$ using 48% concentrated hydrofluoric acid at room temperature (RT) for 18 h. After washing to remove the etching products followed by drying, Ti$_3$C$_2$T$_x$ powder was intercalated by dimethylsulfoxide (DMSO). Then, the intercalated MXene powder was sonicated in deionized water and centrifuged. The resulted colloidal solution of delaminated MXene was vacuum-filtered to produce free-standing MXene “paper”. The latter, consisting of parallel oriented flakes (Figure 1a and b), was used directly as the working electrode (mass density of 4 mg cm$^{-2}$ and thickness of 25 ± 2 μm).

Electrochemical characterization

Electrochemical experiments were performed using a BioLogic SP-200 potentiostat (Bio-Logic, USA). A three-electrode commercial in situ electrochemical AFM cell (Asylum Research, Oxford Instrument Company, USA) was used in this study and shown in Figure 1c. A 14 mm$^2$ Ti$_3$C$_2$T$_x$ paper used as the working electrode was attached to a glassy carbon current collector using colloidal graphite and clamped with a PolyEtherEtherKetone (PEEK) cap to ensure a good electrical contact. A 12 cm$^2$ activated carbon ring was used as the counter electrode (1300 m$^2$ g$^{-1}$, Y-Carbon, USA). The electrolytes used in this study were 1 M Li$_2$SO$_4$, 1 M Na$_2$SO$_4$, 0.5 M K$_2$SO$_4$, and 1 M MgSO$_4$ ($\geq$ 99.0%, all purchased from Sigma Aldrich, USA). Cyclic voltammetry (CV) curves were recorded at sweep rates between 1 and 20 mV s$^{-1}$. Prior to AFM measurements, each working electrode was precycled at 10 mV s$^{-1}$ for 24 h to reach a steady state. All potentials are vs Ag/AgCl reference electrode. The potential range for electrochemical characterization was chosen to avoid unwanted reactions such as MXene oxidation at high potentials or hydrogen evolution at low potentials.

Atomic force microscopy

The relative electrode deformation was monitored using an MFP-3D AFM (Asylum Research, Oxford Instrument Company, USA). Measurements were conducted by keeping the working electrode with a constant contact force while performing CV at various sweep rates (see Figure 1d). Pt coated Nanosensors™ (PPP-EFM-50, k = 0.5-9.5 N m$^{-1}$) AFM tips were used.

X-ray diffraction

X-ray diffraction (XRD) experiments on the Ti$_3$C$_2$T$_x$ paper were performed on a Panalytical XPert diffractometer under copper K-alpha radiation. The electrochemical cell was charged and discharged by linear sweep voltammetry at 1 mV s$^{-1}$. The potential was held during 20 min before collecting the XRD patterns.

Results and discussion

Figure 2a and b shows the CV curves and the corresponding deformation profile demonstrated by the Ti$_3$C$_2$T$_x$ paper electrode during intercalation of various cations. Interestingly, insertion of Li$^+$, Na$^+$ and Mg$^{2+}$ ions results in contraction of the electrode, whereas for K$^+$ electrolyte a small expansion is observed. The values of the relative height change scale appropriately with the charge/ionic radius ratio values (1.67 Å$^{-1}$, 1.05 Å$^{-1}$, 0.75 Å$^{-1}$ and 2.78 Å$^{-1}$ for Li$^+$, Na$^+$, K$^+$ and Mg$^{2+}$, respectively) [28]. For monovalent cations, the contraction amplitude decreases for larger ionic radius, since the steric hindrance contribution due to larger size becomes more important. The attractive forces between negatively charged MXene sheets and cations in the interlayer spaces are likely to dominate the deformation, as represented in Figure 2c. In K$^+$ case however, the steric occupancy between MXene sheets dominates the charge storage due to the large ionic radius. It is interesting to note that the effective radii of hydrated ions (ca. 3.82 Å, 3.58 Å, 3.31 Å and 4.28 Å for Li$^+$, Na$^+$, K$^+$ and Mg$^{2+}$, respectively) [29] do not correlate with the deformation profiles: the larger size of the hydrated Li$^+$ ion should result in a smaller contraction than hydrated Na$^+$ or K$^+$ ions. Most likely, the hydration shell is partially removed or rearranged when ions are intercalated between the layers [14,29,30]. Hence adsorption of Li$^+$ ions, 4-coordinated in water, results in stronger interactions with the MXene layers than Mg$^{2+}$ (6-coordinated), leading to a larger contraction.

Interestingly, two different regions are clearly observed in the CV curves and height change profile when the MXene paper is cycled in Li$_2$SO$_4$. At potentials higher than ~0.25 V, Li$^+$ intercalation capacitance is higher than 300 F cm$^{-3}$ and it drops to around 200 F cm$^{-3}$ when E < ~0.25 V. At the same time, the electrode contraction is small at first and increases strongly when the capacitance changes during the charging process. During discharge, the contraction rate is small at first and increases strongly again when the capacitance
changes, resulting in a hysteresis between charge and discharge (Figure 2b).

The decrease in the Li$^+$ intercalation capacitance along with the change of contraction rate at potentials lower than $-0.25$ V remains unclear at this point, and contrasts with the increase in the capacitance in the same potential region for Mg$^2+$ intercalation. Nevertheless, this strongly points to a 2 steps storage process, either due to adsorption in deep and shallow sites, or to reorganization/ordering of the layered structure. In turn, the hysteresis could be associated with the large positive overpotential required to extract strongly bonded cations that may be exchanging with protons on MXene surface, or to a partial intercalation of anions at potentials higher than 0 V. In order to shed light on this observation, in situ XRD measurements were performed. The XRD pattern of the dry Ti$_3$C$_2$Tx paper (Figure 3a) shows the flakes ordering in the c direction. The (0002) peak is located at 8.83°, corresponding to a d-spacing of 10.04 Å. Figure 3b shows that the (0002) peak shifted to higher angles during Li$^+$ intercalation, and shifted back to lower angles as Li$^+$ ions are extracted. As expected, after cycling the corresponding d-spacing values in Figure 3d show the shrinkage of the MXene layers upon Li$^+$ intercalation, from 12.84 Å to 12.56 Å, which is equivalent to ~2.1%. Since the decrease in the d-spacing is lower than the macroscopic contraction recorded previously, additional electrostatic forces must play a role in the interactions between MXene sheets. The volume change profile is consistent with the dilatometry experiments, with the same hysteresis shape between charge and discharge. Upon cation intercalation, the diffraction peak becomes sharper and more intense, highlighting that the carbide layers become more ordered as the Li$^+$ content increases. As a comparison similar experiments were performed in K$_2$SO$_4$ electrolyte (Figure 3c). Here again the d-spacing evolution matches the macroscopic deformation profile, since only 0.3% volume change was measured. In this case as well the peak shows higher intensity at larger K$^+$ content. Surprisingly, intercalation of large K$^+$ ion led to smaller d-spacing than Li$^+$. Although the reason for this is unclear at this point, the hydration shell structure, the possibility of anion co-intercalation and/or changes in the MXene oxidation state can contribute to variations in the interlayer distances.

The unusual electrode contraction differs from the well-established swelling of other 2D materials such as carbon [31-33] or reduced graphene oxide [34] when they are intercalated with
molecules and/or ions. Moreover, the amplitude of the shrinkage can be higher than 10% in the case of intercalation of Li\(^+\). To get more insight in the two-step process observed for Li\(^+\) intercalation, Figure 4a and b shows the CV curves and corresponding height change profiles plotted vs normalized time for scan rates from 1 to 20 mV s\(^{-1}\). As the capacitance decreases with increasing sweep rate, the electrode undergoes important deformation changes. During intercalation, the electrode first expands (Figure 4b), and the amplitude of the expansion increases with the sweep rate. The change in the electrode deformation may be the result of complex interactions between Ti\(_3\)C\(_2\)Tx sheets and adsorbed metal ions. At high rates, adsorption of the positively charged cations at the Ti\(_3\)C\(_2\)Tx surface may not be fast enough to shield the electrostatic repulsions between negatively charged MXene sheets. The subsequent contraction is measured once a sufficient number of Li\(^+\) ions are intercalated. This is consistent with the slight expansion observed at high rate during the Li\(^+\) extraction process, caused by repulsive interactions between the residual cations and the positively charged layers. Such mechanism can contribute to the important capacitance loss at high sweep rates. Overcrowding by the presence of water molecules and possibly anions between the layers may also contribute to the expansion at high rates. Such species might not have enough time to make room for cation adsorption, hence leading to steric hindrance between MXene layers.

The investigation of electrode deformation reveals interesting insights into the intercalation process for cations of different sizes and charges and their interaction with the MXene sheets. Due to the high reproducibility of the observed behavior, it can also be used to tailor the electrode deformation for specific applications. For example, the insertion of K\(^+\) ions reveals an almost zero electrode deformation which will guarantee low mechanical stresses during charge and discharge, important for a long cyclic lifetime. Cyclability is an important issue for energy storage materials, since the volume changes associated with the electrochemical reaction are often responsible for the capacity fading, as for example Li\(^+\) intercalation in metal oxides [35,36] or Li alloying reaction with metal anodes [19,37]. One of the principal benefits of low volume changes or even contraction materials is that large number of charge/discharge cycles can be achieved along with minimal capacitance loss. Interestingly, electrodes with different interlayer distances can easily be obtained, with the advantage of keeping the electronic conductivity metallic, as well as increasing both the tensile strength and the capacitance values. To date, such MXene-based composites have been fabricated with carbon nanotubes (CNTs), reduced graphene oxide (rGO), onion-like carbon (OLCs) [38], polyvinyl alcohol (PVA) or polydiallyldimethylammonium chloride (PDDA) [39].

Figure 3 (a) X-ray diffraction patterns of the dry Ti\(_3\)C\(_2\)Tx paper, evolution of the (0002) diffraction peak at different potentials during electrochemical cycling in (b) 1 M Li\(_2\)SO\(_4\) and (c) 0.5 M K\(_2\)SO\(_4\) electrolytes, and (d) corresponding d-spacing values as a function of the potential.

Figure 4 (a) CV curves from 1 to 20 mV s\(^{-1}\) and (b) corresponding height change profiles plotted as a function of normalized time in Li\(_2\)SO\(_4\) electrolyte.
In each case the interlayer spacing strongly depends on the nature and quantity of the hosted intercalant, and different deformation behaviors are expected for each material. With such a variety of composites in hands, designing a proper electrode/electrolyte couple becomes more accessible to further increase the cycling stability.

On the other side, certain electrochemical capacitors are considered attractive for mechanical actuation applications [40]. Depending on the actuation requirements, the need to change the actuation properties during utilization may arise. To investigate the tunability of the electrochemical capacitor deformation for actuator applications, the same paper electrode was cycled after the electrolyte was switched in the electrochemical cell in the following order: 1 M MgSO₄, 1 M Li₂SO₄, 0.5 M K₂SO₄ and 1 M MgSO₄. Figure 5a and b shows the CV curves and corresponding electrode deformation profiles recorded after each electrolyte switch, respectively. The typical electrochemical signature for Li⁺ intercalation along with its height change profile confirms that Li⁺ ions replace Mg²⁺ ions in the 2D spaces between the MXene sheets. When K⁺ ions are subsequently intercalated, the typical CV curve is observed, with very small electrode deformation (compare with Figure 2b). Finally, when Mg²⁺ ions are intercalated again, the CV curves show a less resistive behavior and the relative electrode height change, although of lower amplitude, shows the same profile. The faster Mg²⁺ transport between MXene sheets may come from several effects including (i) different adsorption energies due to residual Li⁺ and/or K⁺ cations, or (ii) larger interlayer spaces created by large intercalated K⁺ ions, making more room for Mg²⁺ transport in the 2D channels. This demonstrates that the actuation properties can be recovered when switching the electrolyte, enabling to tune the mechanical behavior of the MXene. It also shows a pathway to optimize the intercalation of multivalent ions of bigger size for increased charge storage. Even though mechanical deformation tests were not performed in a traditional setup, the observed behavior of MXene paper upon adsorption of cations shows its unique actuation properties and promises for applications such as controllable artificial muscles. Strong free-standing MXene electrodes with gel electrolyte were recently reported and can be used as actuators, especially when combined with easily

Figure 5  (a) CV curves at 2 mV s⁻¹, (b) corresponding relative deformation profiles in different electrolytes for the same Ti₃C₂Tx paper electrode.

Figure 6  Potential examples of actuators based on MXene paper electrode, (a) electrical switch, (b) micro-elevator, or (c) circuit breaker that shrinks or expands depending on the chosen electrolyte.
expanding rGO paper electrodes (Figure 6a and b) [39]. Moreover, controlling the actuation behavior of a device by simple switching of the electrolyte can be a great asset for electromechanical applications (Figure 6c).

Conclusions

In situ monitoring of the Ti3C2Tx paper electrode deformation during electrochemical intercalation of various cations using AFM showed that charging of MXene electrode in Li+, Na+ and Mg2+ electrolytes results in contraction, or light expansion, in the case of K+. This was confirmed by in situ XRD measurements. Although further experiments must be performed to fully understand the charge storage mechanism of the different cations, we can conclude that Ti3C2Tx MXene electrode deformation strongly depends on the charge/ionic radius ratio. The unusual reversible contraction during Li+ and Mg2+ intercalation may enable the MXene electrode to sustain a very large number of charging cycles. Finally, delaminated Ti3C2Tx paper shows promising properties for asymmetric electromechanical actuators, which can be tuned to cater to specific actuation requirements.

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References

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