**In-situ Investigation of Expansion during Lithiation of Pillared MXene with Ultralarge Interlayer Distance**

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**Abstract**

Pillared Ti3C2Tz MXene with large interlayer spacing (1.75 nm) is shown to be promising for high-power Li-ion batteries. Pillaring dramatically enhances the electrochemical performance, with superior capacities, rate capability and cycling stability compared to the non-pillared material. In particular, at a high rate of 1 A g-1, the SiO2-pillared MXene has a capacity over 4.2 times that of the non-pillared material. For the first time, we apply in-situ electrochemical dilatometry to study the volume changes within the MXenes during (de)lithiation. The pillared MXene has superior performance despite larger volume changes compared to the non-pillared material. These results give key fundamental insights on the behaviour of Ti3C2Tz electrodes in organic Li electrolytes and demonstrate that MXene electrodes should be designed to maximise interlayer spacings, and that MXenes can tolerate significant initial expansions. After 10 cycles, both MXenes show nearly reversible thickness changes after the charge-discharge process, explaining the stable long-term electrochemical performance.

**Introduction**

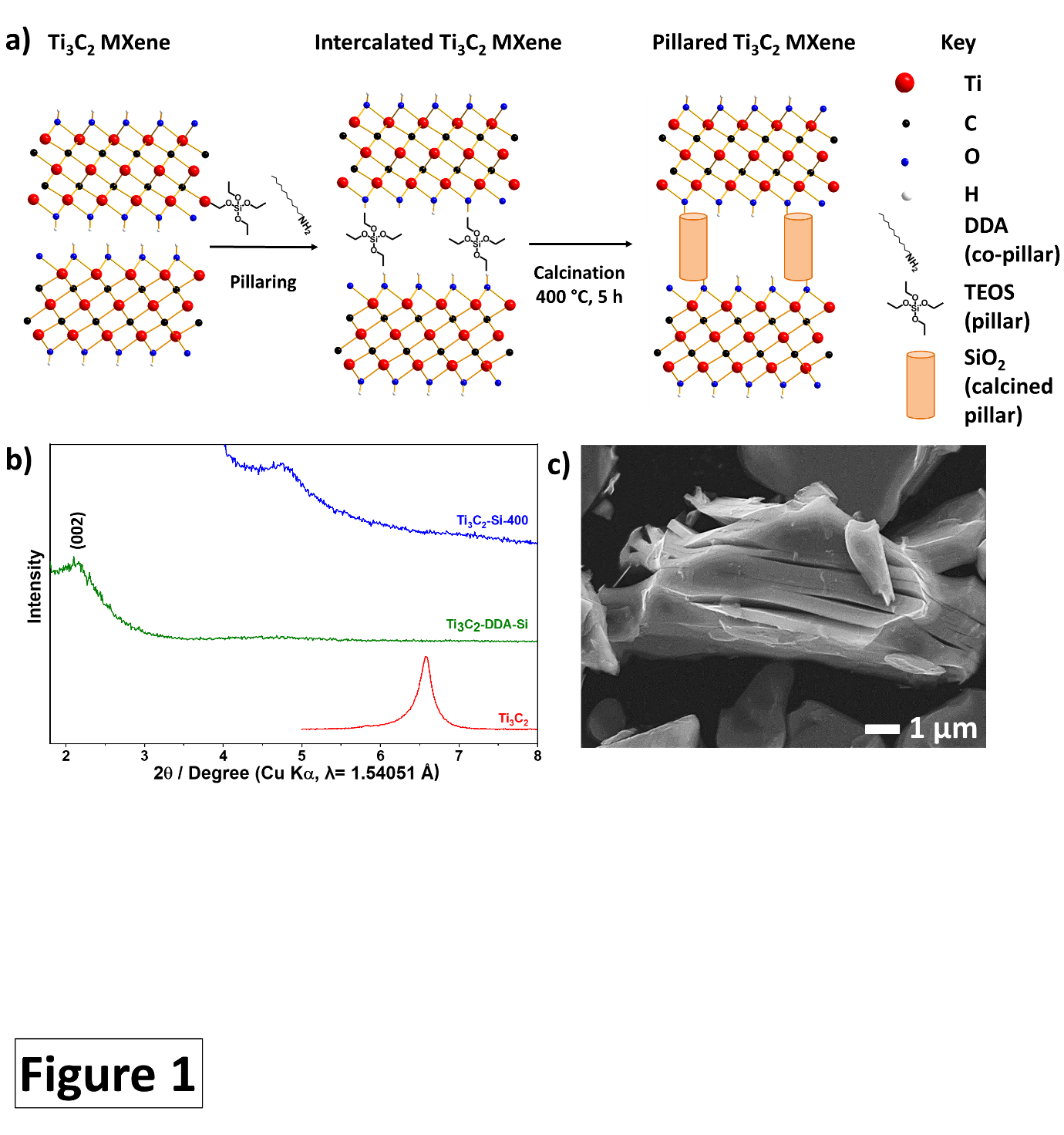
MXenes are a family of two-dimensional materials first reported in 2011, which have shown promising performance in electrochemical energy storage applications such as metal-ion batteries and supercapacitors.1–3 However, their electrochemical performance is highly dependent on the electrode architecture, with multilayered stacked MXene suffering from low capacities and poor rate capabilities and cycling stabilities.4,5 This has increased focus on developing porous MXenes with controlled and open architectures to enhance electrochemical performance, such as flocculation, freeze drying and pillaring.6–8

Pillaring introduces foreign species between the layers to act as pillars, allowing pore sizes to be tuned by pillar choice and heat treatment steps.9 Recently, these techniques have been adapted to create porous MXenes for electrochemical applications, including metal-ion batteries, hybrid metal-ion capacitors, and supercapacitors.10–18 For example, pillar species such as hexadecyl trimethylammonium bromide (CTAB), SnS and Sn cations have been utilised to enlarge the interlayer spacing and improve the performance of MXene electrodes in Li-ion batteries.10,16–19 However, while it is well established that pillaring can significantly enhance electrochemical performance, an in-depth understanding of the processes occurring in pillared structures during cycling is lacking.

In this work, we utilise in-situ electrochemical dilatometry (eD) supported by X-ray diffraction (XRD) to rationalize the enhanced performance of silica-pillared Ti3C2Tz MXene as a negative electrode for Li-ion batteries. In-situ eD has previously been applied to MXene research to study the intercalation of various ions and molecules in several electrolyte systems such as ionic liquids20,21 and aqueous Li, Na and Mg systems.22–24 However, this is the first time this technique has been applied to MXenes for organic Li-ion battery systems, and can provide important information about the effect of pillars during (de)lithiation.

**Methods**

The pillared Ti3C2Tz MXene (Ti3C2-Si-400) was synthesized by our previously reported pillaring method using tetraethylortho silicate (TEOS) as the silica source and dodecylamine (DDA) as the co-pillar amine (**Figure 1a**).25 Full details of all synthesis and characterization methods can be found in the experimental section in Supporting Information. The successful intercalation and pillaring of the synthesized Ti3C2Tz MXene were confirmed by PXRD data (**Figure 1b,** *Supporting Information* **Figure S1a**), while scanning electron microscopy (SEM) confirmed the retention of the layered morphology, (**Figure 1c,** *Supporting Information* **Figure S1b**), which matches our previous report.25 We note that our pillared structure is based on multilayered MXenes, and delamination was not attempted since delamination procedures can suffer fromissues, such as low yield (52-60%), multiple steps, and low concentrations of delaminated nanosheets in solution.26 One of the potential attractions of pillaring techniques is to allow full access of the MXene nanosheet layers in a stable electrode architecture using simple methods.



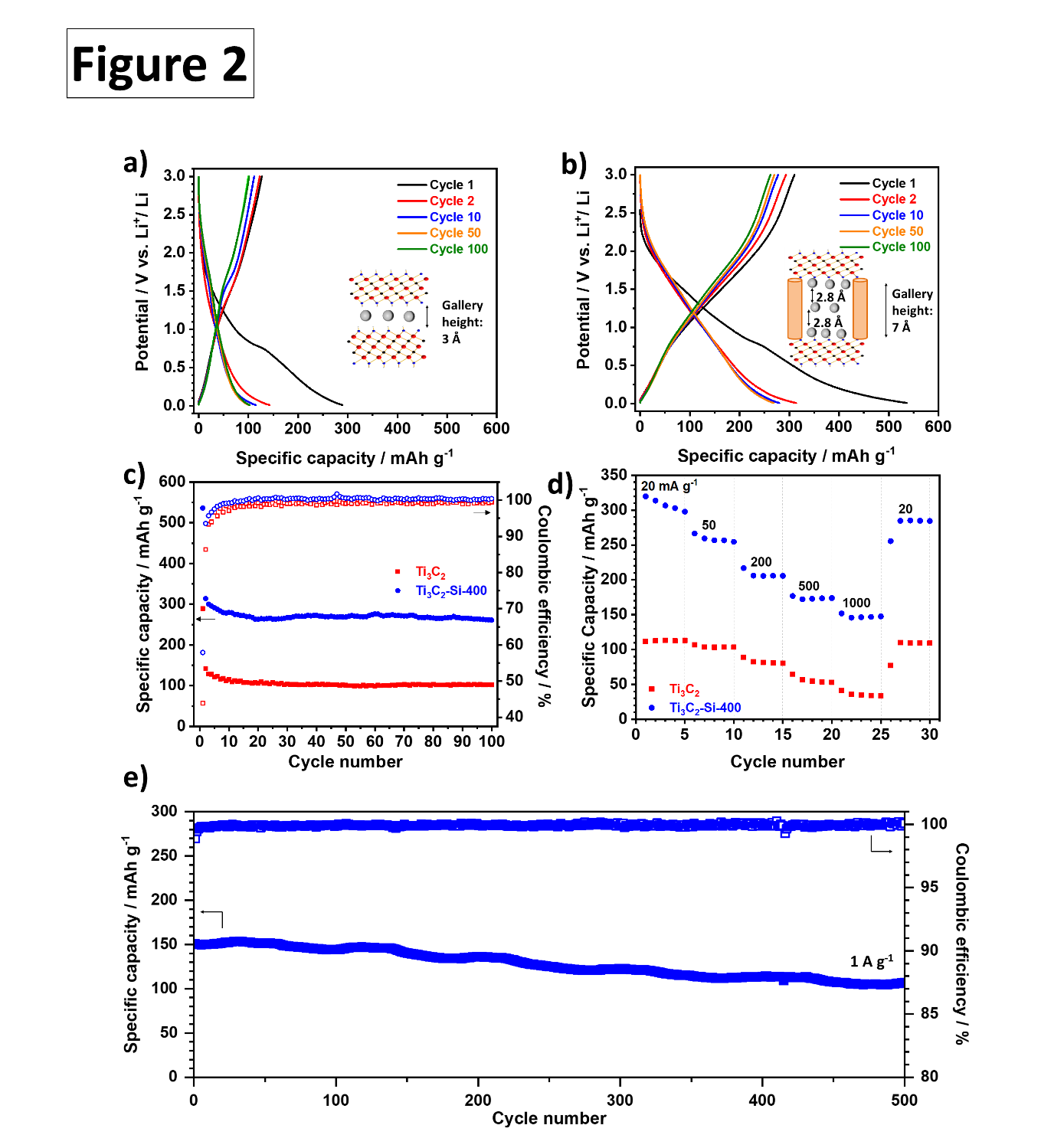
**Figure 1.** **a)** Schematic illustration of the amine-assisted SiO2 pillaring process used to create porous Ti3C2Tz with enlarged interlayer distance. **b)** Low angle (1.8-8° 2θ) PXRD data demonstrating expanded interlayer spacing in the SiO2-pillared MXene (Ti3C2-Si-400), intercalated MXene (Ti3C2-DDA-Si) and non-pillared MXene (Ti3C2). **c)** SEM micrograph of the SiO2-pillared MXene (Ti3C2-Si-400).

**Results and Discussion**

Galvanostatic charge-discharge (GCD) experiments were used to investigate the performance of the pillared Ti3C2Tz as a Li-ion battery electrode (**Figure 2**), with a specific current of 20 mA g‑1 and a potential window of 0.01-3.0 V vs. Li+/Li. All further potentials in the manuscript are referenced vs. Li+/Li. The voltage profiles of the pillared and non-pillared MXenes display similar features, with a linear region between 3.0-0.3 V followed by a short plateau feature below 0.3 V. This signifies that Li storage in Ti3C2Tz is a two-stage process, as previously reported.27 The first stage corresponds to Li+ intercalation between Ti3C2Tz nanosheets forming Li2Ti3C2Tz (theoretical capacity of 260 mAh g-1), followed by Li+ adsorption above the first intercalated layer, resulting in Li3Ti3C2Tz, in the lower voltage region. An additional short plateau is observed on the initial discharge for both materials, which is commonly assigned to solid electrolyte interphase (SEI) formation, and irreversible reactions between Li and MXene surface groups.27–29 Both materials also show typical linear charge profiles, with a small plateau around 1.5 V, resulting from Li de-intercalation.

The pillared MXene has a significantly larger first discharge capacity (536 mAh g-1) compared with the non-pillared (289 mAh g-1), and a larger initial Coulombic efficiency (58% and 43%, respectively). The enhanced initial Coulombic efficiency can be explained by lower levels of Li trapping due to the enlarged interlayer spacings which would provide open Li diffusion channels. Also, surface groups such as -OH, which have been reported to react irreversibly during lithiation, are likely to have already reacted with the co-pillars during the pillaring process. This would further reduce irreversible capacity losses during (de)lithiation.27 For the second discharge cycle, the pillared Ti3C2Tz shows capacities around 2.2 times greater than the non-pillared material (314 mAh g-1 and 142 mAh g-1, respectively), likely as a result of the larger interlayer spacing facilitating greater ion accessibility to the MXene redox sites.

Significantly, the 2nd discharge capacity (314 mAh g-1) is considerably higher than expected based on monolayer lithium coverage, which is 235 mAh g-1 when accounting for the pillar mass (which is estimated to be 11 wt. % based on SEM-EDS analysis, *Supporting Information,* **Figure S2**), further implying that multilayer adsorption is contributing to the capacity. However, the full formation of a third lithium layer to form Ti­3C2O2Li3 would give a capacity of around 350 mAh g-1, which suggests that the third layer does not fully form. Xie et al. calculated that the extra adsorbed Li layer would be located around 2.8 Å above the Ti-O-Li lithium,27 which requires an interlayer gallery space of at least 6 Å. This is less than the ca. 7 Å interlayer gallery space calculated from XRD data in our pillared MXene, demonstrating the feasibility of this charge storage mechanism, illustrated by the insets in **Figure 2a**-**b**. Note that the interlayer gallery space is defined as the free pore space between layers of a pillared material, and is not equivalent to the *d*-spacing, which includes the sheet thickness. Our pillared Ti3C2Tz has a *d*-spacing of 1.75 nm, as calculated from the (*002*) peak using the Bragg equation. An extra increase of 2.3 Å gallery space would be required for the adsorption of a further layer,27 taking the total to at least 8.3 Å, which is larger than our spacings, revealing why further adsorption does not occur. Although the average interlayer gallery space was calculated to be 0.7 nm from XRD data using a monolayer Ti3C2Tz thickness of just under 1 nm, in line with the method utilized by Luo et al,10 our previous work using transmission electron microscopy (TEM) revealed local variations in the gallery spacings across the pillared material,25 with some areas having spacings less than 0.7 nm, explaining why a full extra adsorption layer was not formed.



**Figure 2.** Galvanostatic data from half-cell tests against Li metal at 20 mA g-1 in the voltage range 0.01-3.0 V using 1 M LiPF6 in EC:DEC (1:1 mass ratio) as the electrolyte. Load curves for selected cycles for **a)** non-pillared Ti3C2Tz and **b)** SiO2-pillared Ti3C2Tz (Ti3C2-Si-400). **Insets** illustrate the lithium layers (grey circles) in the respective MXene (blue rectangles) after lithiation (0.01 V). **c)** Coulombic efficiency and discharge capacities of the pillared and non-pillared samples over 100 cycles. **d)** Rate capability tests for the pillared and non-pillared MXene at rates of 20, 50, 200, 500 and 1000 mA g-1. Five cycles are shown for each rate. **e)** Galvanostatic charge-discharge data for SiO2-pillared Ti3C2Tz in a Li-ion half-cell tested for 500 cycles at a rate of 1 A g-1 in a potential window of 0.01-3.0 V.

The pillared MXene also showed improved capacity retention (83%, 262 mAh g-1) compared with the non-pillared MXene (72%, 101 mAh g-1) over 100 cycles (**Figure 2c)**. The majority of the capacity fade occurred over the initial 15 cycles, with capacity retentions of 96% (pillared) and 91% (non-pillared) between cycles 15 to 100. These results demonstrate that the SiO2 pillars significantly improve electrode stability during cycling.

The rate capability of the MXenes was investigated by cycling at rates of 20, 50, 200, 500, 1000 and 20 mA g-1, with five cycles conducted at each rate (**Figure 2d**). The pillared MXene out-performed the non-pillared material at all rates tested, with discharge capacities of 307, 260, 206, 173, 147 and 285 mAh g-1 at the respective rate compared to 113, 104, 82, 55, 35 and 109 mAh g-1, respectively. Interestingly, the capacity enhancement is more considerable at higher rates than lower ones, with an increase of 4.2 times that of the non-pillared Ti3C2Tz observed at 1 A g-1, compared with just 2.7 times at 20 mA g-1. This demonstrates that the pillared architecture benefits the high rate performance in particular, and implies that the enlarged interlayer distance increases the lithium storage capacity and creates channels that allow fast Li+ ion diffusion through the structure. Cycling the pillared MXene at 1 A g-1 for a further 500 cycles after the rate capability tests resulted in a discharge capacity of 151 mAh g-1 (**Figure 2e**) with a capacity retention of 71%, which shows reasonable stability at high rates. The impressive high rate performance can be explained by the significant contribution of a surface-limited capacitive-like process to the charge storage, revealed by *b*-value analysis conducted on cyclic voltammetry (CV) data collected at multiple scan rates (*Supporting Information,* **Figure S4 a-b**). At potentials above 0.5 V, the *b*-values are at least 0.85, consistent with mostly capacitive charge storage behaviour. An ideal capacitor would display a *b*-value of 1, whereas an ideal diffusion-limited battery-like electrode would have *b* = 0.5.30 Furthermore, deconvolution of the capacitive and diffusion-limited charge storage processes, following the method described by Wang et al.,31 revealed that at the scan rate of 5 mV s-1, 78% of the total current was due to capacitive-like processes, demonstrating the importance of capacitive processes at high rates (*Supporting Information,* **Figure S4c**). A scan rate of 5 mV s-1 corresponds to a sweep time of 10 min, similar to the sweep time of 8 min for the GCD testing at 1 A g-1. Even at the low scan rate of 0.2 mV s-1, 43% of the total charge storage was a result of capacitive-like processes (*Supporting Information*, **Figure S4d**). As observed in previous studies of MXenes in organic Li-ion systems, the capacitive contribution to the charge storage increased with increasing scan rate (*Supporting Information*, **Figure S4d**).32,33 Further details on this analysis are given in the experimental section in the Supporting Information. This analysis uses a range of sweep rates, similar to other reported studies on pillared MXenes in organic electrolytes.10,34

To better understand the influence of interlayer gallery spacing on enhancing electrochemical lithium-ion storage, the volume change of pillared/non-pillared Ti3C2Tz MXene during ion uptake and release was investigated by in-situ eD, which can observe volume change and CV simultaneously. A scan rate of 0.1 mV s-1 was applied with a narrower potential window of 0.01-2.5 V to focus on expansion/contraction processes related directly to (de)lithiation of the MXene. The CV shape for the pillared material closely matched with what was recorded in typical coin cells, with similar broad redox features, demonstrating that the cell used for in-situ eD35 does not change the electrochemical processes. **Figure 3** shows the CV in the dilatometry cell and *Supporting Information,* **Figure S3** shows the comparison in a typical coin cell. The non-pillared material shows more pronounced redox-peaks compared with the pillared MXene, which are very broad, as was observed in our previous work in a Na-ion system and supports the expanded nature of the MXene.25,36 As expected from the GCD data, multiple irreversible reduction peaks are observed in the pillared material during the first discharge, with peaks at around 1.3 V and 0.6 V attributed to the SEI formation and trapping of lithium-ions,37 while lithiation of the conductive additive contributes to the peak at 0.01 V.38,39 In the following delithiation sweep, two oxidation peaks were observed at 1.0 V and 1.8 V, representing the extraction of lithium ions, whose intensities continuously decrease slightly in the following cycles. For the non-pillared Ti3C2Tz similar redox features occur, with pronounced reversible peaks at 1.8 V and 1.0 V (reduction) and at 0.9 V and 2.0 V (oxidation) throughout cycling. The fact that the CV and load curves for the non-pillared and pillared MXenes display similar features with no extra redox peaks/plateaus present in the data for the pillared MXene strongly implies that the enhanced performance results from the enlarged interlayer spacing, rather than electrochemical activity of the silica pillars. This is consistent with the results of our previous work on pillared Mo2TiC2, where ex-situ x-ray photoelectron spectroscopy (XPS) and nuclear magnetic resonance spectroscopy (NMR) studies showed that the silica pillars in that system did not undergo lithiation during cycling.15

The unprocessed dilatometry data for the non-pillared MXene electrode reveals slight volume changes (~0.5 µm) during the initial 72 h of stabilization time, which is related to the electrode wetting process (**Figure 3c**). A relatively large expansion (1.84 µm, 5.7% strain) occurs during the first discharging step, consistent with Li+ intercalation **(Figure 3a)**. At the end of the first lithiation sweep (0.01 V), a large jump is observed in the dilatometry data (**Figure 3a**). Comparison with the corresponding CV curve (**Figure 3a**) shows that while there is a large peak here, which likely results from a combination of Li intercalation, lithiation of the carbon additive,39 and changes in MXene termination groups, the magnitude of this jump is significantly larger than corresponding CV peak. This implies that there is also significant contribution from device artifacts, which cannot be removed from the data in **Figure 3a** without losing physical meaning. However, it is clear that the contribution from artifacts is low overall and does not affect the conclusions that can be drawn from the data.

During de-lithiation, the expansion is only partially reversed, with a decrease in volume of ca. 1%. This demonstrates that Ti3C2Tz remains in an expanded state even after de-lithiation, which is likely related to partial Li trapping, changes in MXene surface chemistry, SEI formation and solvent co-intercalation within the electrode layers. A similar pattern can be seen in the second cycle, but with reduced amplitude of volume changes on charge/discharge, while having a relative strain during the third cycle of -0.36%. Between cycles 3-10, the volume changes in the electrode appear to stabilize, with the magnitude of volume expansion on lithiation matching the magnitude of contraction on de-lithiation, so that the relative strain during the cycle tends towards zero. The pattern of expansion during lithiation and contraction during delithiation matches well with previous in-situ XRD studies carried out on multilayered Ti-based MXenes, which supports our dilatometry data.40

Since the volume changes in the stabilization time are artifacts of the device, a background/baseline subtraction was used in **Figure 3 c-d**. For the calculation of the relative and absolute expansion in **Table 1**, only the lithiation process was considered in each case. During the cycles, the non-pillared MXene behaves as a low strain material (<0.53% strain within the entire lithiation cycle). The expansion levels for this sample increase slightly continuously up to the 10th cycle. Since the galvanostatic charge-discharge tests showed that the discharge capacity fades over these first 10 cycles, this may indicate the presence of irreversible side reactions, such as Li trapping. This would lead to a consistent expansion of the non-pillared material during cycling and explain the lower cycling stability for this material compared to the pillared MXene. The overall volume changes are very small for all cycles, which agrees well with the stable cycling performance observed during the GCD tests after the initial few cycles.

Post-mortem XRD recorded after 2 cycles of CV in custom-built polyether ether ketone (PEEK) cells of the non-pillared samples shows a slight shift in the (002) diffraction peak to lower angles (from 8.4° to 7.8° 2θ), supporting the small expansion observed in the dilatometry data (*Supporting Information,* **Figure S5 a-b)**).

For the pillared MXene (**Figure 3b and 3d**), a large initial expansion occurs over the first few hours under open circuit potential, with a gradual relaxation towards a minimum level of ca. 1.6 µm lower compared to the initial volume over the remainder of the rest period.

As observed for the non-pillared MXene, a large expansion (~1.02 µm) occurs in the first lithiation cycle (**Table 1**).The initial contraction has a smaller magnitude compared to the initial expansion. This suggests that the pillars aid in keeping the MXene electrode expanded, allowing wide Li-diffusion pathways rather than fixing a constant interlayer spacing.The pillared electrode continues to expand and contract over the next few cycles with approximately constant absolute and relative expansion. After the first cycle, the strain decreases slightly in the short term, but when looking at the continuous change, consistent values of around 3-4% relative strain can be obtained for all cycles. This correlates with the electrode’s stable performance during galvanostatic charge/discharge cycling(**Figure 2d**). Simultaneously, slight capacity fading was observed over the first few cycles where the amplitude of contraction (de-lithiation) was consistently more minor than the previous expansion during lithiation, suggesting small quantities of irreversible side processes, such as Li trapping, were occurring, albeit significantly less than observed for the non-pillared material.Post-mortem XRD on the pillared MXene after two cycles shows a loss of the (002) diffraction peak (*Supporting Information,* **Figure S5d**), suggesting the expansions observed in the dilatometry arise from inhomogeneous changes in the interlayer spacing during the initial (de)lithiation cycles. This early loss of observable diffraction peaks demonstrates the advantages of techniques such as in-situ dilatometry to study the (de)lithiation processes, which does not rely on long-range order.

Table 1. Comparison of the displacement and strain measured using in-situ electrochemical dilatometry for the non-pillared (Ti3C2) and pillared (Ti3C2-Si-400) MXenes. Background subtraction has been carried out to remove artifacts that are unrelated to the electrode shrinking and expansion and the data normalized by electrode thickness, according to the method reported by Budak et al.41 These values were calculated by referring to the height change between the start of the cycle (2.5 V) and the opposite end of the lithiation process (0.01 V).

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Cycle number** | **Ti3C2** | | **Ti3C2-Si-400** | |
| **Displacement (µm)** | **Relative strain (%)** | **Displacement (µm)** | **Relative strain (%)** |
| **Cycle 1** | 0.08 | 0.10 | 1.02 | 3.91 |
| **Cycle 3** | 0.15 | 0.35 | 0.69 | 2.69 |
| **Cycle 5** | 0.18 | 0.41 | 0.82 | 3.14 |
| **Cycle 10** | 0.23 | 0.53 | 0.80 | 3.07 |

The non-pillared MXene undergoes more minor volume changes than the pillared counterpart, which may be explained by the significantly larger quantities of Li intercalated into the structure (ca. 2.5 times) in the pillared electrode compared to the non-pillared. This implies that the key to unlocking high and persistent performance for MXenes is to engineer large interlayer spacings, which would avoid narrow interlayer spacings limiting Li storage capacity and diffusion. Large but reversible expansions do not seem to be detrimental to MXene performance. Pillared MXene undergoes large but consistent volume changes whilst displaying high cycling stability compared to the non-pillared MXene, where the expansion gradually increased in magnitude during cycling. This finding suggests that MXene architectures should be designed to minimise irreversible volume changes during cycling.

Diagram

Description automatically generated

**Figure 3.** Cyclic voltammograms and corresponding in-situ dilatometric measurements (electrode height change/initial height = strain) from half-cell setup versus Li metal at 0.1 mV s-1 in the voltage range 0.01-2.5 V for **a)** non-pillared Ti3C2Tz **b)** pillared MXene (Ti3C2-Si-400). The electrochemical response of the working electrode (continuous line) can be simultaneously tracked with the height change (dashed line). **c-d)** Potential and relative height change vs. time of MXene electrodes measured by in situ electrochemical dilatometry **c)** for non-pillared Ti3C2Tz (electrode thickness of 43 μm) **d)** for pillared MXene (Ti3C2-Si-400, electrode thickness of 26 μm). The artifacts which are unrelated to the electrode response had been subtracted as background signals.

**Ti3C2**

**Ti3C2-Si400**

**b)**

**Conclusions**

In conclusion, silica-pillared Ti3C2Tz MXene with an ultralarge interlayer spacing displayed significantly enhanced performance as the negative electrode of a Li-ion battery. In particular, at the high rate of 1 A g-1, the SiO2-pillared MXene had a capacity over four times that of the non-pillared material (over 150 mAh g-1), and retained over 71% of its initial capacity after 500 cycles. In-situ dilatometry combined with XRD data revealed that the superior cycling performance occurred despite larger volume changes and a decrease in crystallinity, suggesting inhomogeneous interlayer expansion. During lithiation, a constant relative expansion of 3-4% in each cycle is obtained for the pillared MXene sample. In contrast, for the non-pillared MXene, a slightly increasing expansion is obtained with increasing cycle numbers. These results provide key insights for the design of MXene electrodes demonstrating that large interlayer spacings and consistent volume changes, as provided by pillaring techniques, are key for high performance. This is the case even for large (3-4%) expansions, which are well tolerated by pillared MXene.

**Supporting Information**

Experimental methods for the synthesis and pillaring of MXenes. Experimental details for the XRD characterisation, SEM characterisation, electrochemical characterisation, dilatometry experiments and ex-situ X-ray diffraction. X-ray diffractograms and scanning electron micrographs of the non-pillared Ti3C2Tz. SEM-EDS analysis for the pillared MXene. Cyclic voltammetry and capacitive contribution from CV data for the pillared MXene. Ex-situ XRD and corresponding CV data for the pillared and non-pillared material.

**Acknowledgments**

P.A.M. gratefully acknowledges support from the EPSRC Graphene NOWNANO Centre for Doctoral Training. The INM authors thank Eduard Arzt (INM) for his continued support. V.P. acknowledges funding of the MXene-CDI project (PR-1173/11) by the German Research Foundation (DFG, Deutsche Forschungsgemeinschaft). NTR acknowledges the Royal Society (RG170150), Energy Lancaster and Lancaster University for financial support.

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**TOC Graphic**

**Diagram, schematic

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