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**TiF₃ catalyzed MgH₂ as a Li/Na ion battery anode**

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**Keywords**: Li ion battery, Na ion battery, MgH₂, TiF₃

**Graphical abstract**

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Abstract

MgH₂ has been considered as a potential anode material for Li ion batteries due to its low cost and high theoretical capacity. However, it suffers from low electronic conductivity and slow kinetics for hydrogen sorption at room temperature that result in poor reversibility, cycling stability and rate capability for Li ion storage. This work presents a MgH₂-TiF₃@CNT based Li ion battery anode manufactured via a conventional slurry based method. Working with a liquid electrolyte at room temperature, it achieves an high capacity retention of 543 mAh g⁻¹ in 70 cycles at 0.2 C and an improved rate capability, thanks to the improved hydrogen sorption kinetics with the presence of catalytic TiF₃. Meanwhile, the first realization of Na ion uptake in MgH₂ has been evidenced in experiments.
1. Introduction

MgH₂ has been intensively researched as an active material for hydrogen storage due to its high capacity (7.6 w.t. %) and low cost [1,2]. It was introduced as a conversion anode material for reversible electrochemical Li ion uptake in 2008 [3], and has attracted growing research interest since then due to its high capacity for Li ion storage and cost-effectiveness [4-8]. Its lithiation undergoes dehydrogenation of MgH₂ with the formation of LiH (Reaction 1) resulting in a Li ion storage capacity of 2038 mAh g⁻¹, which is defined as the theoretical capacity. Li-Mg alloying (Reaction 2) takes place when further lithiation proceeds and the achieved capacity can be even higher.

\[
\begin{align*}
2 \text{Li} + \text{MgH}_2 & \rightarrow \text{Mg} + 2 \text{LiH} \\
x \text{Li} + y \text{Mg} & \rightarrow \text{Li}_x\text{Mg}_y
\end{align*}
\]

However, despite its high capacity, MgH₂ faces tremendous challenges in reaching its potential in Li ion batteries. The most critical issue is the limited H mobility in MgH₂ at room temperature and thus sluggish kinetics for Li ion uptake. Considerable research efforts have been made, among which the utilization of MgH₂ in solid state Li ion batteries has achieved significant progress [8-14]. For instance, a (Nb₂O₅-doped MgH₂)-LiBH₄-acetylene black (4:3:3 in mass) composite achieved a capacity retention of 700 mAh g⁻¹ in 100 cycles at 120 °C [8]. However, the functionality of these solid state batteries entails a high operating temperature (above 100 °C) [8-13], and catalysts (e.g. Al₂O₃ [9] and Nb₂O₅ [8, 10-12]) and a substantial amount of LiBH₄ [8, 11-14] are required to improve H mobility and Li ion diffusivity. The high operational temperature largely limits its potential for practical applications and the addition of large amount of H conducting but storage-inactive LiBH₄ also reduces the overall capacity of the electrode and cost-effectiveness. Very recently, El kharbachi et al. has demonstrated the possibility of a MgH₂ anode based solid state lithium battery operating at room temperature by embedding nanosized MgH₂ in a novel Li₂S-P₂S₅-
Li(BH$_4$)$_{0.75}$I$_{0.25}$ based solid electrolyte, but the reversibility and cycling stability has to be greatly improved [14]. Therefore, the research on MgH$_2$ based solid state batteries is out of the scope of our work.

Intensive research on MgH$_2$ anodes have also been carried out for room-temperature liquid-electrolyte-based Li ion batteries in recent years [3,15-19], and the solutions to improve the kinetic properties include nanosizing MgH$_2$ [3,15,16], producing more cyclable β-MgH$_2$ through gas phase hydrogenation of Mg [17-19], and incorporating a secondary hydride phase (e.g. TiH$_2$) [17,18]. Moreover, both MgH$_2$ and its lithiation product, LiH, are electrically insulating, and the uptake of Li ions in MgH$_2$ is also accompanied with a large volume change (up to 85%), therefore a large content of advanced carbon and binders appear to be necessary to maintain the electrical conduction and structural integrity of the electrode during cycling.

For instance, a composite of ultra-small (~ 5 nm) MgH$_2$ embedded within a high surface area graphite matrix (MgH$_2$ : C = 1:1 in mass), synthesized with a bottom-up approach, showed a retained capacity of around 500 mAh g$^{-1}$ for Li ion storage after 20 cycles [16]. A composite of nanostructured MgH$_2$, carboxymethyl cellulose formate (CMC-f) binder and preground C$_{t,x}$ carbon (MgH$_2$ : CMC-f : C$_{t,x}$ = 1:1:1 in mass) reached a capacity retention of 542 mAh g$^{-1}$ in 40 cycles [19].

However, in the above-mentioned studies, besides the limited commercial viability due to the complex synthesis and/or high material cost, the kinetic properties of Li ion transport were still limited, and the cycling rates were extremely low (usually 0.1 Li per hour, i.e. 0.05 C with C = 2038 mA g$^{-1}$) limited by the sluggish ionic transport. As a result, the reversibility of Li ion storage was rather low, leading to a poor cycling stability and rate capability. Moreover, in experimental studies, mechanically pressed disc-like MgH$_2$ electrodes were commonly prepared for the electrochemistry measurement. In such electrodes the Li ion diffusion kinetics throughout the electrode is limited because the pellet should remain intact, which is a
further handicap for sustained cycling stability and high rate capability. Recently, a
magnetron-sputtered MgH2-C thin-film anode (~270 nm) with Pd coating has also been
reported, however, its active mass loading is extremely low (< 0.1 mg cm⁻²), and the electrode
is non-porous resulting in limited Li ion diffusion in the solid and poor cycle life (10 cycles)
[20]. Recently El kharbachi et al. has introduced the conventional slurry-based method to
produce porous, thin-film MgH₂ electrodes using N-Methyl-2-pyrrolidone (NMP) as a
solvent, but the cycling performance is still to be improved due to the aggregation issue of
MgH₂ that limits both the electronic and ionic transport [21]. Furthermore, the formation of
solid electrolyte interphase (SEI) is unavoidable in organic liquid electrolyte based Li ion
batteries. The SEI layer is mechanically unstable, and therefore continuous SEI growth occurs
during cycling resulting in the loss of active materials and capacity decay. This issue is even
aggravated taking into account the pulverization of active materials caused by the large
volume expansion/contraction during de-/lithiation.

In this work, we present a MgH₂-TiF₃@carbon nanotubes (CNT) composite based Li ion
battery anode, fabricated through a slurry-based approach using anhydrous tetrahydrofuran
(THF), which has a lower boiling point (66 °C, compared to 202 °C for NMP) and is thus
easier to be removed, as a solvent. 5 mol. % TiF₃ is present in the composite as a catalyst to
enhance the kinetics for hydrogen sorption and Li ion uptake in MgH₂. Transitional metal
halides (e.g. TiF₄ [22], ZrCl₄ [23] and NbF₅ [24]) have been proven to be able to improve the
hydrogen sorption kinetics in MgH₂; and it has been shown by some of us that, in gas phase
experiments, the hydrogen absorption/release is possible at reduced temperatures with the
catalysis of TiF₃ [25]. The role of CNT in the composite is to address the poor electronic
conductivity of MgH₂ and LiH, providing sufficient electronic charge transport throughout the
electrode and enabling its electrochemical function. Furthermore, to suppress the SEI growth,
10 w.t. % fluoroethylene carbonate (FEC) was utilized as an electrolyte additive. The MgH₂-
TiF$_3$@CNT anode exhibits a capacity retention of 543 mAh g$^{-1}$ in 70 cycles at 0.2 C and outstanding rate capability in liquid electrolyte based Li ion batteries. Meanwhile, this work has, for the first time, experimentally achieved Na ion uptake in a MgH$_2$ based electrode.

2. Experimental details

**Sample Preparation:** MgH$_2$ (hydrogen-storage grade, Aldrich) was mixed with 5 mol. % TiF$_3$ (Alfa Aesar) and CNT (industrial standard, 150 – 210 m$^2$ g$^{-1}$) (MgH$_2$-TiF$_3$ : CNT = 3:1 in mass), and subsequently was ball milled with a Fritsch Planetary Mono Mill PULVERISETTE 6 for 50 hours at 400 rpm under Ar atmosphere using a Tungsten Carbide grinding bowl and a ball-to-powder mass ratio of 50:1.

**Characterization:** The micro-morphology of MgH$_2$-TiF$_3$@CNT and the as-prepared electrode were imaged with a JEOL JSM 6010F scanning electron microscope working at 5 kV. XRD measurements were performed with a PANalytical X’Pert Pro PW3040/60 diffractometer (Cu K$_\alpha$ radiation) operating at 45 kV and 40 mA. The lithiated electrode for XRD measurement was washed 3 times with diethyl carbonate (DEC) to remove soluble electrolyte residuals in the electrode, and were dried in a glovebox before the measurements.

**Electrode preparation and battery assembly:** The MgH$_2$ based Li ion battery anodes were prepared inside an Ar-filled glovebox using a conventional slurry based method using anhydrous THF (≥ 99.9%, Sigma-Aldrich) as the solvent. and polyvinylidene difluoride (PVDF, Kynar flex) worked as the binder and super P carbon black (TIMCAL) was used to integrate the electrical conduction throughout the electrode. The slurry (MgH$_2$-TiF$_3$@CNT: super P carbon black : PVDF = 70:15:15 in mass) was casted on a planar Cu foil (12.5 μm, Goodfellow) with a doctor blade and dried inside the glovebox. The thickness of active materials was ~ 20 μm and the mass loading was ~ 0.5 mg cm$^{-2}$. The composition/preparation of the MgH$_2$ based Na ion battery anodes was similar to that of the Li ion batteries except that
super P carbon black instead of CNT was utilized as the conducting additive in the ball milled composite. Half-cell Li/Na ion batteries were assembled inside an Ar-atmosphere glovebox with the O₂ and H₂O levels < 0.1 ppm. A Li/Na foil (Aldrich) was applied as the counter electrode. The working electrolyte for Li ion batteries was 1 M LiPF₆ dissolved in ethylene carbonate (EC), DEC and FEC (EC : DEC = 1:1 in volume with 10 % FEC); and 1 M NaPF₆ dissolved in EC and dimethyl carbonate (DMC) and (EC : DMC = 1:1) was applied as the electrolyte in Na ion batteries. A borosilicate glass micro fibre (Whatman) was used as the separator.

Electrochemistry measurements: The galvanostatic electrochemical properties were tested with a MACCOR 4600 battery cycler. The cut-off voltages for Li ion batteries were 0.005 V and 3.0 V versus Li/Li⁺ for discharge and charge, respectively. The cycling voltage range for Na ion batteries was 0.005 – 2.8 V versus Na/Na⁺, and the cyclic voltammograms were obtained with an Autolab potentiostat scanning within the same voltage range.

Subtraction of the carbon capacity: The specific capacities reported in this paper were calculated based on the mass of MgH₂ (excl. carbon additives). The capacity of CNT was determined by measuring its electrochemical performance (Fig. S1) in Li ion batteries; and the capacity of super P carbon black has been determined by us in previous work [26]. The capacity contributions from these carbon additives were then subtracted to obtain the specific capacity of MgH₂.

3. Results and discussions

X-ray Diffraction (XRD) patterns of the MgH₂-TiF₃@CNT composite (Fig. 1a) show the characteristic diffraction peaks of MgH₂ (110), (101), (200) and (211) lattice planes at 27.7°, 35.6°, 39.6° and 54.4°, respectively. Weak but distinguishable peaks of TiF₃ are present at 23.1°, 31.4°, 41.0° and 48.3°. The peaks at 25.6°, 42.7° and 78.2° can be allocated to the
Bragg reflections on CNT [27-28] The average crystalline domain size of MgH₂ is ~ 10 nm calculated based on the line shape of MgH₂ in the Rietveld refinement (Fig. S2). The scanning electron microscopy (SEM) images of the sample (Fig. 1b – 1d) show aggregates of sub-micro grains induced by the high energy ball milling. Energy-dispersive X-ray spectroscopy (EDX) (Fig. S3) demonstrates that the distribution of TiF₃ next to MgH₂ is homogeneous on the basis of the superposition of Mg and F element distribution at the submicro-/nanoscale. The nature of the small grains is thus the agglomeration of nanosized MgH₂ and TiF₃ which warrants an optimal catalytic activity of TiF₃.

Morphological images of the as-prepared electrode (Fig. 1e – 1f & S4) show that the active material loading is uniform with micro-/nanoscale porosities all over the electrode, and the active material grains are integrated by the binder and carbon matrix. The porous nature of the electrode allows the access of liquid electrolyte throughout the electrode and reduces the Li ion diffusion pathway through the solid phase fractions.
**Fig. 1.** (a), XRD patterns of the ball-milled MgH$_2$-TiF$_3$@CNT sample. (b) – (d), SEM images of the sample at different magnifications. (e), a photo and (f), a SEM image of the as-prepared electrode (12.7 mm in diameter).

The electrochemical performance of the MgH$_2$ based anodes was measured within half-cell Li ion batteries. **Fig. 2a** shows that, cycling at 0.2 C, a capacity of 3254 mAh g$^{-1}$ was achieved for the first lithiation, and the following delithiation reaches a capacity of 1959 mAh g$^{-1}$. This relatively low Coulombic efficiency (61.7 %) mainly originates from the irreversible SEI formation. The reversible capacity declines during the first few cycles and 1139 mAh g$^{-1}$ is retained after 5 cycles. The Coulombic efficiency increases to 86.3 % and 97.4 % at the 2$^{nd}$ and 5$^{th}$ cycle, respectively, indicating that though further SEI growth still takes place but the amount is increasingly limited, and most of the Li ions absorbed during lithiation can be released during the following delithiation. It also indicates that the increasingly reduced capacity during Li ion uptake is the main reason of the rapidly decaying reversible capacity retention. The capacity retention slips to 707 mAh g$^{-1}$ after 20 cycles, and reaches 543 mAh g$^{-1}$ in 70 cycles. **Fig. 2b** shows that the initial reversible capacity for Li ion storage amounts to 2293, 2291 and 824 mAh g$^{-1}$ at 0.05, 0.1 and 0.5 C, respectively. Compared with the published results on MgH$_2$ based anode cycling in liquid electrolytes [5,6,8,17], the MgH$_2$-TiF$_3$@CNT electrode in this work exhibits improved rate capability. Such improved kinetics supports that TiF$_3$ is an effective catalyst for hydrogen absorption and release in Mg/MgH$_2$.

The initial Li ion uptake (**Fig. 2c**) shows a voltage plateau at ~ 1.2 V corresponding to the SEI formation, which is not observed in the following cycles, and a lithiation capacity of around 200 mAh g$^{-1}$ is reached during this process. The main lithiation of MgH$_2$ takes place at a lower voltage range including two stages: (i), a voltage plateau at ~ 0.2 V related to the hydrogen extraction from MgH$_2$ with the formation of LiH (**Reaction 1**); and (ii), a low voltage plateau ~ 0.05 V associated with the alloying process between Mg and Li (**Reaction
At the beginning of the first stage, the voltage undergoes a dip to 0.13 V, which is consistent with the other studies [3,15-17], possibly originating from the large energy barrier of initial LiH nucleation on the surface of Mg, before it rises to the main voltage plateau at ~0.2 V due to the co-existence of Mg and MgH₂. Upon the complete depletion of MgH₂, the presence of a single MgH₂ phase results in a declining voltage slope. When the voltage goes down to 0.08 V the cumulative lithiation capacity reaches above 2150 mAh g⁻¹ indicating that the Li ion uptake in MgH₂ achieves a capacity of ~ 1950 mAh g⁻¹, which is close to the theoretical capacity. In the second stage, the voltage also exhibits a plateau originating from the co-existence of Li and Mg solid solutions and a following sloped voltage attributed to one single phase (Li solid solution) [3,15]. The cumulative lithiation capacity reaches ~ 3250 mAh g⁻¹ when the cut-off voltage is reached revealing a Li ion storage capacity of ~ 1100 mAh g⁻¹ (i.e. Li : Mg ≈ 1 : 1) in the alloying reaction. In the following delithiation, the voltage exhibits a reverse course of lithiation. It begins with a de-alloying process from LiₓMgy that occurs at a voltage range < 0.25 V achieving a capacity of about 600 mAh g⁻¹. Subsequently, it undergoes a plateau at ~ 0.6 V associated with the delithiation from LiH and the hydrogenation of Mg, which obtains a capacity of ~ 1400 mAh g⁻¹. The Coulombic efficiency (72 %) of the delithiation from LiH is higher than that of the de-alloying process (55 %), indicating that the Li ion extraction from the Li-Mg alloys is less reversible and may cause the active material loss and capacity deterioration over cycling.

The overpotential between the Li-Mg de-alloying and alloying is less than 0.2 V; while it rises up to 0.5 V between the lithiation of MgH₂ (forming Mg) and the reverse reaction. This overpotential increase stems from the significant conductivity reduction from conductive metallic Mg to electrically insulating MgH₂. Over cycling, the voltage plateaus become increasingly sloped with growing overpotentials, which results from the build-up of internal resistance due to the gradual structural deformation over repetitive volume expansion and
contraction. As a result, the lithiation cut-off voltage has been reached earlier and before the Li-Mg alloying occurs, therefore, the achieved lithiation capacity drops. Given that the low lithiation capacity is the main limiting factor for the reversible capacity retention as explained before, we can conclude that the drastic capacity drop within the first few cycles mainly originates from the increasing overpotential resulting in the decreasing amount of Li ion uptake during lithiation.

The voltage profiles at different current rates (Fig. 2d) show growing overpotentials when the current rate increases. It leads to a more rapid voltage collapse to the cut-off voltage and thus a lower capacity when it is lithiated at a higher current rate; and hence in the following Li ion extraction step, the achieved capacity is lower. It is also observed that the Li-Mg alloying reaction (near 0 V) reduces significantly as the overpotential grows.

**Fig. 2.** (a), The capacity retention and Coulombic efficiency at 0.2 C (C = 2038 mA g\(^{-1}\)). (b), The initial reversible capacity of the MgH\(_2\)-TiF\(_3\) anode at different C-rates. (c), Voltage
profiles of the 1st, 2nd, 20th and 70th cycles of the MgH2-TiF3 anode at 0.2 C. (d), Voltage profiles of the MgH2-TiF3 anode for the 1st cycle at different C-rates.

The high performance of the MgH2-TiF3@CNT based Li ion battery anode may benefit from the synergistic effects of several factors. Firstly, the slurry-casting manufactured electrode, compared with a pellet electrode, is more porous and allows a facile access for the liquid electrolyte to the active materials and may enable faster Li ion transport throughout the electrode. Therefore, a lower internal ionic resistance is expected resulting in a lower overpotential, and the rate capability could be promoted as well. Moreover, the voids in the electrode can accommodate the volume change upon Li ion insertion. Secondly, the active material grains are embedded within the CNT matrix warranting intimate electronic conduction between the active materials and the carbon network induced by high energy ball milling, and enabling sustained electrical conduction between the grains over cycling. Thirdly, The FEC additive in the electrolyte induces a thin, stable and compact LiF-dominant SEI layer that prevents further SEI growth, and therefore an enhanced Coulombic efficiency and cycling stability can be achieved [29-31]. This is also evidenced by comparing with the results with a FEC-free electrolyte (Fig. S5c). Last but not least, the catalytic TiF3 accelerates the hydrogen sorption kinetics in MgH2 and leads to facilitated lithium ion uptake. Specifically, during the initial lithiation, apart from the dehydrogenation of MgH2 (Reaction 1), the presence of TiF3 next to MgH2 leads to the (irreversible) formation of MgF2 and TiH2 (Reaction 3). Subsequently, the small MgF2 crystallites act as a grain refiner (i.e. seed crystals for the growth of MgH2) for MgH2/Mg, keeping their crystallites small [25,32]. This grain refinement mechanism results in a destabilized system facilitating the hydrogen sorption of Mg/MgH2, and in this way the Li ion uptake is also accelerated. Meanwhile, TiH2 may also be lithiated (Reaction 4) and vanishes, but it is much more difficult since the enthalpy of
formation of TiH$_2$ ($\sim 136$ kJ mol$^{-1}$) is more negative than MgH$_2$ ($\sim 74$ kJ mol$^{-1}$) and there is no alloying between Li and Ti [33].

2 Li + 3 MgH$_2$ + 2 TiF$_3$ → 3 MgF$_2$ + 2 TiH$_2$ + 2 LiH \hspace{1cm} (3)

2 Li + TiH$_2$ → Ti + 2 LiH \hspace{1cm} (4)

The advantageous catalytic effect is also evidenced by measuring the performance of a TiF$_3$-free electrode at the same conditions as the MgH$_2$-TiF$_3$ electrode (Fig. S5c). Recently such grain refinement has been employed as well in Li-O$_2$ electrode systems [34].

To further investigate the behaviours of TiF$_3$ and MgH$_2$ during cycling, XRD patterns of a MgH$_2$-TiF$_3$@CNT electrode (Fig. 3) were acquired at different stages: pristine, lithiated to 0.05 V and 0.005 V, respectively, at 0.1 C. In the pristine electrode, apart from the peaks from MgH$_2$, the main diffraction peaks of TiF$_3$ are weak but distinctively recognizable. Bragg diffractions on the CNT are visible as well. When it is lithiated to 0.05 V, MgH$_2$ disappears while Mg and LiH show up, and the Li-Mg alloy emerges as well. It indicates that the dehydrogenation of MgH$_2$ (Reaction 1) is accomplished and the lithiation has proceeded to the Mg-Li alloying (Reaction 2). It is also observed that TiF$_3$ vanishes due to Reaction 3, but the peaks of MgF$_2$ and TiH$_2$ are hardly visible due to their much lower presence together with much smaller grain sizes ($\sim 3$ nm) compared with MgH$_2$/Mg ($\sim 10$ nm) based on the Rietveld refinement (Fig. S6 & Table S1 – S2), and the peak broadening is significant in the XRD signals. These findings are in good agreement with the previous studies [25,32]. During the further lithiation till 0.005 V, LiH remains unaffected while Mg alloys with Li forming Li$_x$Mg$_y$; the stoichiometry is $x : y \approx 1 : 1$, and the average alloy grain size amounts to 18.3 nm (Table S3) The grain size growth from Mg (10.1 nm) to Li$_x$Mg$_y$ (18.3 nm) originates from the absorption of Li ions in the Mg structure. The Rietveld refinement also indicates the crystal structure transformation from a hexagonal lattice (Mg, space group: $p63/mmc$) to cubic (Li$_x$Mg$_y$, space group: $Im-3m$) upon Li ion uptake. Moreover, the larger size of Li$_x$Mg$_y$
compared to LiH (< 8 nm) results in a slower Li ion desorption and thus worse reversibility. This is also consistent with the higher Coulombic efficiency of Li ion extraction from LiH compared to the dealloying of intermetallic Li$_x$Mg$_y$ (as discussed in Fig. 2c).

![Fig. 3. XRD patterns of the MgH$_2$ electrode at different stages: pristine, lithiated to 0.05 V and 0.005 V, respectively. (a), experimental data; (b), Refined XRD patterns after subtraction of background; (The 2 theta ranges of carbon and Li-C alloy has been subtracted since it is out of the scope of this work. Details on the refinement are displayed in Fig. S6.) Here the electrode is a pressed pellet (~ 70 μm in thickness), instead of a slurry based electrode, in order to avoid the strong Bragg reflections from the Cu current collector and to produce XRD patterns with a higher signal-to-noise ratio. The promising results of the TiF$_3$ catalyzed MgH$_2$ anode in Li ion batteries encourage us to investigate its potential for Na ion batteries. Though a sodiation voltage has been predicted theoretically [35,36], MgH$_2$, or any other metal hydride, has not been reported to have a capability of Na ion storage probably due to the poor kinetics of Na ion transport. The improved kinetics of Li ion uptake in the TiF$_3$ catalyzed MgH$_2$ suggests that the Na ion diffusion may be facilitated with the catalytic TiF$_3$ and thus enable the realization of Na ion uptake in MgH$_2$.](image-url)
In this work, the possibility of Na ion insertion in MgH$_2$ was studied by testing MgH$_2$ based anodes within Na ion half-cells. The experimental result illustrates that, judging from the comparison of voltage profiles between the MgH$_2$ based electrode (Fig. 4a) and the carbon additive based electrode (Fig. S7a), the voltage plateau at 0.15 V and part of the sloping voltage between 0.15 V and the sodiation cut-off voltage originate from the sodiation of MgH$_2$, and the achieved capacity in this process is only ~ 350 mAh g$^{-1}$; during desodiation, the sloping voltage range of 0.4 – 0.9 V may be partially related to the Na ion extraction from the product of sodiated MgH$_2$ achieving a capacity of merely ~ 80 mAh g$^{-1}$ (excl. the capacity contribution from the carbon additive [26]). Although the achieved capacity is low, the realization of Na ion insertion in MgH$_2$ is evident. It is also evidenced by the cyclic voltammograms (Fig. 4b & S7b) and XRD patterns of the sodiated MgH$_2$ electrode (Fig. S8).

![Voltage profiles and cyclic voltammograms](image)

**Fig. 4.** (a), Voltage profiles at 0.05 C and (b), Cyclic voltammograms (scan rate: 0.02 mV s$^{-1}$) of the MgH$_2$-TiF$_3$ electrode tested in a Na ion half-cell.

### 4. Conclusions

This work presents a novel MgH$_2$-TiF$_3$@CNT composite based anode which achieves promising electrochemical performance in liquid electrolyte based Li ion batteries. MgF$_2$ derived from the TiF$_3$ during the initial lithiation process work as the catalytic species to
improve the electrochemical hydrogen sorption of Mg/MgH₂ and therefore to facilitate the Li ion uptake in MgH₂. We also find out that the dealloying of intermetallic Li-Mg is less reversible than the delithiation of LiH due to its larger grain size. Moreover, Na ion uptake in the MgH₂-TiF₃ electrode has been, for the first time, witnessed in experiments even though the cycling performance is poor and the reaction mechanism is still to be elucidated. It may stimulate considerable follow-up research on metal hydrides for reversible Li/Na ion uptake.

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Declarations of interest

None.

Appendix A. Supplementary data

Supplementary information related to this article can be found at [XXXXXXXXX].

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