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Letter

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1 Overcoming the High-Voltage Limitations of Li-Ion Batteries Using a 2 Titanium Nitride Current Collector

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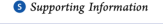
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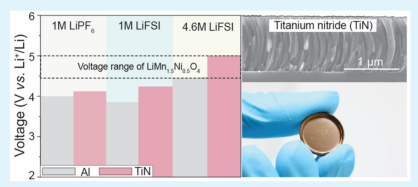
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ABSTRACT: The major obstacle to commercialization of high-voltage Li-ion batteries is the lack of oxidatively stable and inexpensive current collectors that can operate at potentials of up to 5 V vs Li⁺/Li. In this work, we present titanium nitride as a compelling cathode current collector for high-voltage Li-ion batteries exhibiting higher oxidative stability in LiPF₆ and lithium bis(fluorosulfonyl)imide electrolytes than aluminum or stainless steel current collectors. Its high oxidative stability has been assessed with a high-voltage LiMn_{1.5}Ni_{0.5}O₄ cathode. TiN/LiMn_{1.5}Ni_{0.5}O₄ half cells demonstrated a high Coulombic efficiency of 98.5% at a low C-rate of 0.2 C after 100 cycles.

KEYWORDS: titanium nitride, current collector, high-voltage Li-ion battery, LiMn_{1.5}Ni_{0.5}O₄, energy density

echargeable Li-ion batteries (LIBs) have become a key energy storage technology that has served humanity for 21 over the past two decades in a variety of applications, e.g., 22 portable electronic devices and electric vehicles, owing to the 23 high energy density, long cycling lifetime, and high power 24 performance. 1-5 The practical energy density of present-day 25 LIBs composed of a graphite anode and LiCoO2, LiMn2O4, $_{26}$ LiNi $_{0.33}$ Mn $_{0.33}$ Co $_{0.33}$ O $_{2}$, and LiFePO $_{4}$ cathodes, however, is $_{27}$ approaching the practical limit of 250 Wh kg^{-1} . In this 28 context, alternative high-voltage cathode materials have been 29 proposed, such as spinel $LiMn_{1.5}Ni_{0.5}O_4$ (LMNO), $^{9-12}$ $_{30}$ polyanionic $LiVPO_4F$, 13,14 Li-rich layered oxide $Li_{1.2}Ni_{0.2}$ 31 $Mn_{0.6}O_{2}$, 15,16 and Ni-rich layered oxide LiNi_{0.8}Co_{0.1}-32 $Mn_{0.1}O_{0.2}$. These materials offer average discharge voltages 33 of 4.2-4.7 V vs Li⁺/Li, which enable an increase in energy 34 densities by up to 25% or 80%, assuming graphite and metallic 35 lithium, respectively, as anodes.^{2,18} An LMNO cathode is 36 considered to be a particularly attractive candidate for the next 37 generation of LIBs due to its high voltage of 4.7 V vs Li⁺/Li

and relatively large theoretical capacity of 147 mAh g⁻¹ owing 38 to two reversible redox processes, Ni²⁺/Ni³⁺ and Ni³⁺/Ni⁴⁺, 39 occurring at 4.70 and 4.75 V vs Li⁺/Li, respectively.

However, the overcharging problem of high-voltage 41 cathodes, which involves the oxidation and corrosion of the 42 current collectors, is still not fully resolved. Aluminum (Al) 43 and stainless steel (SS), commonly used cathode current 44 collectors, are not suitable for high-voltage applications, as they $_{45}$ often start oxidizing at voltages as low as $3.9^{19,20}$ and $3.1\ V^{21,22}$ $_{46}$ vs Li⁺/Li (depending on the electrolyte used), respectively, 47 and they are fully unusable at voltages above 4.2-4.5 V vs Li⁺/₄₈ Li. Al can still be applied for most research purposes because 49 its corrosion is often slow enough, but it cannot be used in 50 commercial high-voltage batteries. On the other hand, 51 alternative metals/oxides proposed earlier, such as silver 52

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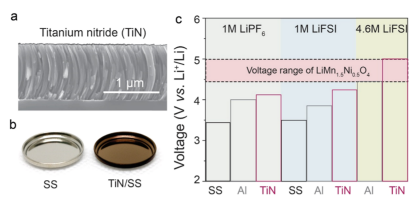


Figure 1. (a) SEM cross-section of a TiN film prepared by magnetron sputtering. (b) Photographs of bare and TiN-coated stainless steel substrates. (c) Illustration of the oxidative stabilities of various current collector materials in 1 M LiPF $_6$, 1 M LiFSI, and 4.6 M LiFSI in EC/DMC electrolytes in terms of the voltage versus Li $^+$ /Li. Oxidative stabilities were derived from cyclic voltammetry curves of the corresponding current collectors obtained at a scan rate of 0.1 mV s $^{-1}$ (Figure S3).

 $_{53}$ (Ag), $_{23}^{23}$ gold (Au), $_{24}^{24}$ platinum (Pt), $_{25}^{25}$ molybdenum (Mo), $_{26}^{26}$ tungsten oxide (WO₃), $_{27}^{27}$ and platinum—rhodium alloy (Pt– $_{55}$ Rh), $_{28}^{28}$ are prohibitively costly.

Herein, we report that titanium nitride (TiN), a highly selectrically conductive material, is highly suited for commercial use as a high-voltage current collector due to its high oxidative stability in LiPF₆- and LiFSI-based electrolytes at potentials of up to at least 5.0 V vs Li⁺/Li. Its practical advantages are the high natural abundance of the constituting elements and low cost of fabrication. Its superior stability toward electrochemical oxidation is presented herein with cells utilizing an LMNO cathode. TiN/LMNO half cells exhibited a high Coulombic efficiency of ~98.5% at a C-rate of 0.2 C after 100 cycles. A side-by-side comparison of the electrochemical oxidation stability of TiN with those of Al and SS current collectors is also presented.

Titanium nitride current collectors were prepared on a stainless steel bottom coin-type cell substrate by magnetron sputtering using a Ti target under a nitrogen—argon atmosphere (Figure 1a,b). Alternatively, for the pouch-type cell configuration, sputtering of TiN on a flexible polyimide film with a relatively large surface area of 0.1 m² was also demonstrated (Figure S1). X-ray diffraction measurements (Figure 2) confirmed the formation of a highly crystalline cubic TiN film on stainless steel (space group Fm3m, a = 4.241 %Å, JCPDS 038-1420) oriented in the [111] direction. The cross-sectional scanning electron microscopy (SEM) image

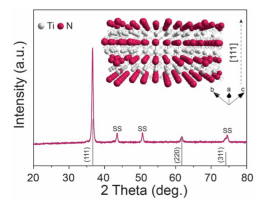


Figure 2. X-ray pattern of TiN film deposited on a stainless steel substrate (inset: crystal structure of TiN).

shown in Figure 1a reveals a columnar morphology with a 80 thickness of approximately 1 μ m. Top-view SEM images of the 81 samples show that the stainless steel substrate is fully covered 82 by the pinhole-free continuous TiN film (Figure S2). 83

The electrochemical stability of the TiN current collector 84 was evaluated using 1 M LiPF₆ and 1 M lithium bis-85 (fluorosulfonyl)imide (LiFSI) in ethylene carbonate/dimethyl 86 methyl carbonate (EC/DMC) electrolytes. Figure S3a,b shows 87 the cyclic voltammetry (CV) curves for various current 88 collectors in both electrolytes, and the results are summarized 89 in Figure 1c. The onset of electrochemical oxidation in LiPF₆/ 90 LiFSI electrolytes occurs at 3.44 V/3.49 V, 4.0 V/3.85 V, and 91 4.12 V/4.24 V vs Li⁺/Li for Al, SS, and TiN current collectors, 92 respectively. As shown in Figure 1c, the oxidative stability of 93 TiN in both electrolytes exceeds those of Al and SS. The 94 increase in the current above 4.12 V/4.24 V vs Li⁺/Li in 95 LiPF₆/LiFSI electrolytes for the TiN current collector could be 96 associated with the oxidation of the LiPF₆ and LiFSI 97 electrolytes. The high oxidative stability of the TiN current 98 collectors might be attributed to the preferred (111) 99 orientation (Figure 2) of the TiN film, as discussed later. 100 We note that, in our previous studies on a TiN current 101 collector for aluminum batteries, the high oxidation stability of 102 TiN was demonstrated even in highly corrosive aluminum- 103 based ionic liquids, outperforming SS, molybdenum, and 104 tungsten current collectors.²⁹

Next, the oxidative stability of the TiN current collector was 106 probed in highly concentrated LiFSI-based solutions. These 107 represent a novel class of Li electrolytes with advantageous 108 properties, such as high oxidative stability, thermal stability, 109 and low volatility, over those of conventional 1 M LiPF₆ 110 electrolytes.³⁰ In addition, it was recently demonstrated that 111 LiFSI electrolytes sustain reversible plating/stripping of 112 metallic Li, as required for Li metal/high-voltage cathode 113 batteries.³⁰ According to the cyclic voltammetry curves 114 measured at 0.1 mV s⁻¹ (Figures S3c) in highly concentrated 115 4.6 M LiFSI electrolyte, the onset of electrochemical oxidation 116 occurs at >5.0 and 4.45 V vs Li⁺/Li for the TiN and Al current 117 collectors, respectively, highlighting the superior stability of the 118 TiN current collector over that of Al. The electrochemical 119 stabilities of both the TiN and Al current collectors in 120 combination with a high-voltage LMNO cathode were probed. 121 Specifically, a series of galvanostatic cycling measurements of 122 TiN/LMNO and Al/LMNO electrodes was performed to 123 examine the effect of LMNO active material loading on the 124

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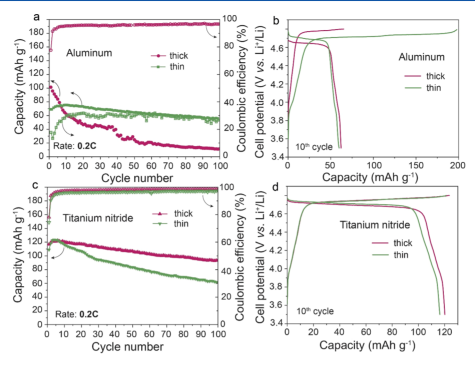


Figure 3. Cyclability/Coulombic efficiency and typical voltage profiles of thick (\sim 55–60 μ m) and thin (\sim 15–20 μ m) LMNO cathodes deposited on Al (a, b) and TiN (c, d) current collectors. The measurements were performed in 4.6 M LiFSI in EC/DMC electrolyte at a C-rate of 0.2 C (1 C = 147 mA g⁻¹) using a voltage range of 3.5–4.8 V vs Li⁺/Li.

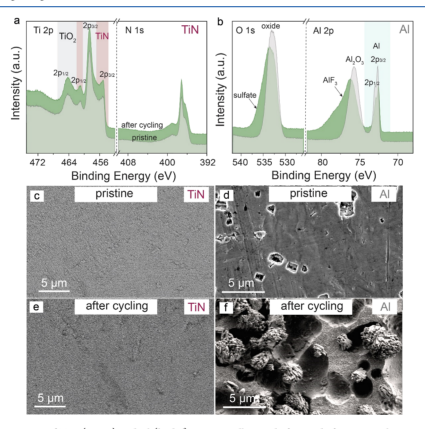


Figure 4. XPS data and SEM images of TiN (a, c, e) and Al (b, d, f) current collectors before and after 1000 galvanostatic charge—discharge cycles with the LMNO cathode in the 4.6 M LiFSI in EC/DMC electrolyte (see Figure S6 for XPS survey spectra). XPS peaks of Ti^{4+} (TiO_2) and Ti^{3+} (TiN) are indicated by gray and rose bands, respectively. Prior to these measurements, the LMNO cathode was carefully removed from the electrodes and rinsed off with pure EC/DMC solvent.

125 Coulombic efficiency and cyclic stability of half cells. In fact, a 126 high electrode mass loading (high thickness) can somewhat mask the oxidative instability of current collectors. Thus, two 127 different mass loadings of the LMNO cathode over the TiN 128

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129 and Al current collectors were investigated: a lower mass 130 loading of 1 mg cm⁻² (thickness of 15–20 μ m) and a higher 131 mass loading of \sim 4 mg cm⁻² (thickness of 55–60 μ m). 132 Further details of electrode preparation and battery assembly 133 can be found in the Supporting Information.

Figure 3a-d and Figure S4a-d summarize the results of the 135 electrochemical tests of the LMNO cathode with TiN and Al 136 current collectors at a C-rate of 0.2 C in 4.6 M LiFSI and 1 M 137 LiPF₆ electrolytes, respectively. As follows from Figure 3a-d 138 (4.6 M LiFSI electrolyte) for thick LMNO cathodes with TiN 139 and Al current collectors, relatively high initial Coulombic 140 efficiencies of 78% and 77% were observed, which gradually increased after 100 cycles up to 98.5% and 96.7%, respectively. 142 For thin electrodes, however, a severe decrease of Coulombic 143 efficiency down to 42% was revealed for the Al current 144 collector, in contrast to TiN (Figure 3c). The higher 145 Coulombic efficiency of thick LMNO electrodes can be 146 explained by the decrease in the proportion of parasitic 147 irreversible oxidation reactions of the electrolyte on the current 148 collector to the reversible oxidation of LMNO cathode with 149 higher LMNO loading.

As follows from Figure S4a-d, almost no difference was 151 observed in Coulombic efficiency between LMNO electrodes 152 deposited on TiN and Al current collectors for 1 M LiPF₆ 153 electolyte. The Coulombic efficeency was about 80% and 90% 154 for thin and thick LMNO cathodes deposited on Al and TiN 155 current collectors, respectively. The latter can be explained by 156 lower oxidation stability of 1 M LiPF₆ electrolyte with an upper 157 oxidation limit of 4.3 V vs Li⁺/Li in comparison with 4.6 M 158 LiFSI electrolyte (see Figure S3a,c for tungsten electrode). It 159 should be noted that degradation of the LMNO cathode 160 deposited on both Al and TiN current collectors is caused 161 primarily by the Mn/Ni dissolution. Although substantial 162 cyclic stability improvements of the LMNO cathode have been 163 made in the past several years using various surface 164 functionalization approaches such as surface modification/ 165 coating/doping, degradation of LMNO cathode during its 166 cycling is still not fully solved. We suggest that future work 167 should be possibly focused on the combination of LMNO 168 cathodes with Li-ion solid-state electrolytes, which can solve 169 the Mn/Ni dissolution issues and have a high oxidative 170 stability window of >5.0 V vs Li⁺/Li.

To obtain a closer look at the chemical processes at the TiN 172 and Al surface under the LMNO thick electrodes, their 173 surfaces were examined before and after 1000 galvanostatic 174 charge—discharge cycles (Figure S5) with the LMNO cathode 175 in the 4.6 M LiFSI in EC/DMC electrolyte by X-ray 176 photoelectron spectroscopy (XPS) and SEM. XPS of the 177 TiN current collector suggests that the outermost layers of Ti 178 atoms are in the Ti⁴⁺ state, presumably with oxygen anions as 179 the plane terminating the surface. Nearly identical Ti 2p and N 180 1s XPS bands before and after cycling of TiN revealed the 181 absence of reactivity (Figure 4a). Furthermore, no F 1s or S 2p 182 signals were observed for the cycled TiN current collector 183 (Figure S6b). The N/Ti atomic ratios derived from XPS 184 spectra were practically identical before and after cycling (1.0 ± 0.1 and 1.0 ± 0.1 , respectively). Apparently, no obvious 186 changes in the surface morphology of TiN current collectors 187 were observed even after 1000 cycles (Figures 4c,e), thus 188 indicating the excellent corrosion resistance of the TiN current 189 collector. In contrast, the surface morphology of Al was 190 completely changed after cycling (Figures 4d,f) and was 191 composed of relatively large amounts of S and F according to

XPS measurements (Figure 4b and Figure S6a). The products 192 of the Al oxidation in the highly concentrated LiFSI electrolyte 193 could be AlF₃, Al₂(SO₄)₃, and Al(FSI)₃. As follows from XRD 194 and XPS measurements, we assume that the high oxidative 195 stability of TiN current collectors is caused by the preferred 196 (111) orientation (Figure 2) of the TiN film. Obviously, due 197 to the alternation of Ti and N planes, this crystallographic 198 direction is highly suited for passivation of Ti³⁺ ions 199 (protecting them from deep oxidation to Ti⁴⁺).

In summary, we report on titanium nitride (TiN) as a 201 compelling current collector material for high-voltage cathode 202 materials that can be easily fabricated by magnetron sputtering 203 on stainless steels or polyimide surfaces on a large scale. 204 Comprehensive characterization of the TiN current collector 205 by XRD, XPS, and SEM revealed its high oxidative stability in 206 LiPF₆ and LiFSI electrolytes, exceeding those for SS and Al 207 current collectors. The high oxidative stability of TiN in 208 combination with a high-voltage LMNO cathode has been 209 assessed. TiN/LMNO electrodes demonstrated a high 210 Coulombic efficiency of 98.5% at a low C-rate of 0.2 C after 211 100 cycles. This work indicates further avenues for the 212 commercialization of high-voltage cathode materials such as 213 LMNO for next-generation high-voltage LIBs.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the 217 ACS Publications website at DOI: 10.1021/acsaem.8b01771. 218

Additional experimantal details and characterization 219 information presented as Figures S1-S6 (PDF) 220

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Author Contributions

The manuscript was written through the contribution of all 231 authors. All authors have given approval to the final version of 232 the manuscript. 233

The authors declare no competing financial interest.

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