



Title	A widely applicable strategy to convert fabrics into lithiophilic textile current collector for dendrite-free and high-rate capable lithium metal anode
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1 **A widely applicable strategy to convert fabrics into lithiophilic textile current collector for**
2 **dendrite-free and high-rate capable lithium metal anode**

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10

11 **ABSTRACT** Hindered by dendrite growth, unceasing volume change and repeated regeneration
12 of solid-state interface, the practical application of lithium (Li) metal anode is still facing
13 challenges from low Coulombic efficiency (CE), insufficient safety performance and poor cyclic
14 stability. Current collector plays a key role in regulating Li deposition and suppressing dendrite
15 growth. In this report, through a simple bio-template method, a textile-structured nickel (Ni)
16 framework is fabricated as current collector for Li metal anode, whose unique micro-nano
17 hierarchical structure is adequate for accommodating Li. A good performance after more than 200
18 cycles at 3 mA cm⁻² during repeated Li plating/stripping is remained in virtue of this unique
19 structure design. By further introducing Ag₂S nanoparticles uniformly to the current collector, a
20 dendrite-free and high reversible Li metal anode is achieved, showing low over-potential (~24 mV
21 at 1 mA cm⁻²), high CE (~98%) and excellent quick charging/discharging stability (up to 350

22 cycles at 10 mA cm^{-2} in symmetric cell). Furthermore, this new strategy for constructing textile-
23 structured metallic framework opens a foreground for various applications of porous metals.

24 **KEYWORDS** Lithium metal anode, Li metal battery, textile structure, bio-template, metallic
25 current collector.

26

27 **1. Introduction**

28 Lithium (Li) metal has been considered as one of the most promising anode materials for
29 rechargeable Li-based batteries, such as Li-O₂ battery and Li-S battery, because of its ultrahigh
30 theoretical capacity (3860 mA h g^{-1}), lowest negative potential (-3.040 V vs the standard hydrogen
31 electrode) and a low mass density (0.53 g cm^{-3}) [1-3]. However, the practical utilization of Li metal
32 anode is hindered by two pivotal problems: i) dendritic Li growth due to the inhomogeneous Li
33 deposition; ii) unceasing side reactions induced by the thermodynamically unstable interface
34 between Li and electrolyte. Those will lead to the disastrous consequences for Li metal batteries,
35 including the increasing of internal resistance, low Coulombic efficiency (CE) and hazard of short
36 circuit [4,5].

37 In the past several years, considerable efforts have been devoted to enhancing the cycling
38 stability of Li metal anode by employing a variety of strategies. In one approach, due to the fact
39 that the protogenous solid-state electrolyte interface (SEI) on Li metal is always heterogeneous
40 and fragile, stable artificial SEI layers with high conductivity, such as lithium nitride (Li₃N) or
41 lithium sulfide (Li₂S), were pre-coated onto Li metal, and the significantly stabilized
42 electrochemical behavior has been confirmed by several groups [6-13]. However, the irregular
43 deposition of Li still occurs underneath the SEI which inevitably deteriorates the electrode

44 performance as the cycling proceeds. The initial nucleation of dendritic Li is correlated to both the
45 Li ion migration rate and the effective electrode current density, which is depicted as the diffusion
46 model called “Sand’s Time” [14, 15]. In addition, the growth rate of dendritic Li is determined by
47 Li ion migration direction and rate [16]. Therefore, a three-dimensional (3D) porous current
48 collector with large surface area will decrease the local current density and inhibit the formation
49 of Li dendrite [17-19]. In particular, the hierarchically micro-nano structured 3D current collector,
50 which can disperse the electronic field and Li-ion flux simultaneously, is efficient for regularizing
51 the electrodeposition behavior of Li and alleviating the dendritic Li growth [20-23].

52 Considering that metallic nickel (Ni) and copper (Cu) own good electronic conductivity and
53 high mechanical strength, they are suitable current collector which have been widely used in
54 commercial batteries. However, as a 3D container to incorporate Li metal, Ni/Cu framework is not
55 lithiophilic to reduce the severe Li nucleation over potential, resulting in non-uniform plating of
56 Li nuclei [24, 25], thus Li metal cannot be infiltrated into the interior of the porous current collector.
57 In addition, the pathway of Li ion migration can be gradually blocked by previously deposited Li
58 on the surface and the subsequent Li deposition can only proceed on the top of the 3D current
59 collector, especially when a high deposition capacity is involved, making insufficient utilization
60 of the 3D porous structure [20, 26, 27]. For this reason, most of the currently reported metallic 3D
61 current collectors can only work stably for less than 250 cycles at a low current density and a low
62 deposition capacity, *i.e.*, $\leq 1 \text{ mA cm}^{-2}$ and $\leq 1 \text{ mA h cm}^{-2}$ [21, 27-29]. Zhao et. al. developed a 3D
63 Cu current collector through an electrochemical dealloying strategy, and a stable Li
64 plating/stripping performance was observed up to 200 cycles at 1 mA cm^{-2} to a capacity of 1 mA
65 h cm^{-2} [30]. Zhang and his co-workers pre-stored Li into a 3D Ni foam, and stable cycle
66 performance could be maintained more than 100 cycles at 5 mA cm^{-2} to a capacity of 1 mA h cm^{-2}

67 ² [31]. Even with additional decoration strategies applying to the metallic 3D current collectors,
68 such as modifying Ni foam with lithiophilic Ni nitrides layer (illustrating a tested performance up
69 to 160 cycles at 5 mA cm⁻² to 1 mAh cm⁻²) [32] or constructing a lithiophilic Cu-CuO-Ni hybrid
70 3D structure (with a tested performance up to 300 cycles at 1 mA cm⁻² to 1 mA h cm⁻²) [33], the
71 cycle performance is still unsatisfying, especially when the cells were cycling at high current
72 densities.

73 Employing cellulose based materials as templates for developing bio-inspired materials is
74 regarded as an efficient and convenient strategy [23, 34-38]. Herein, we develop a simple cotton
75 fabric template method to produce 3D metallic current collector with textile structure for Li metal
76 anode. By using the cotton canvas template method, the canvas structured Ni current collector
77 (Canvas Ni) is obtained. When the hierarchically micro-nano porous Canvas Ni is used for
78 accommodating Li metal, a performance that is far superior to previously reported 3D metallic
79 current collectors is observed, *i.e.*, > 200 cycles at 3 mA cm⁻² with a capacity of 1 mA h cm⁻². With
80 further decoration with lithiophilic Ag₂S nanoparticles (NPs) to the metallic framework, a stable
81 cycle performance under high current density is observed (up to 350 cycles at 10 mA cm⁻² to 1 mA
82 h cm⁻²), which is rarely reported in literatures for Li metal anode using 3D metallic current
83 collectors.

84

85 **2. Experimental Section**

86 *2.1. Preparation of Canvas Ni framework*

87 A piece of cotton canvas fabric (size 7.0 cm×2.5 cm) was firstly cleaned in acetone by
88 consecutive ultrasonication for 30 min and then dried in vacuum oven. Nickel acetate tetrahydrate
89 ($\text{Ni}(\text{CH}_3\text{COO})_2 \cdot 4\text{H}_2\text{O}$, Labchem-Wako) was dissolved in ammonia solution with a concentration
90 of 28% ~ 30% (Kanto Chemical) to prepare a 2.5 mol L^{-1} Ni source solution. The dried cotton
91 fabric was firstly soaked in the Ni source solution for 12 h, followed by calcining at 600 °C for 2
92 h in air to remove the cotton fabric template. The as-prepared Canvas NiO precursor was
93 subsequently reduced by hydrogen gas in a tubular reactor at 650 °C for 1 h to obtain the Canvas
94 Ni framework. The optimized production condition of using 2.5 mol L^{-1} Ni solution is shown in
95 **Fig. S16**.

96

97 *2.2. Preparation of Ag_2S nanoparticles coated Canvas Ni electrode*

98 A silver mirror reaction was used to deposit Ag nanoparticles on the Canvas Ni electrode. An
99 ammonia solution which contained $6.375 \times 10^{-3} \text{ mol L}^{-1}$ silver nitrate (AgNO_3 , Labchem-Wako,
100 99.8%), $3.405 \times 10^{-2} \text{ mol L}^{-1}$ glucose ($\text{C}_6\text{H}_{12}\text{O}_6$, Kanto Chemical, 98%) and $4.005 \times 10^{-3} \text{ mol L}^{-1}$
101 tartaric acid ($\text{C}_4\text{H}_6\text{O}_6$, Kanto Chemical, 99.5%) was used as the silver mirror reagent. The Canvas
102 Ni electrode was soaked in the freshly prepared silver mirror reagent for 30 min. In this way, Ag
103 nanoparticles were formed on the Canvas Ni, which was subsequently washed by
104 ethanol/deionized water for several times and dried in a vacuum oven. The determination of a
105 reaction time of 30 min is shown in **Fig. S17**.

106 Dimethyl sulfoxide (DMSO, Kanto Chemical, 99%) which contained $2 \times 10^{-2} \text{ mol L}^{-1}$ sulfur
107 was used as the sulfurization reagent. The Ag nanoparticles coated Canvas Ni was put into the
108 sulfurization reagent and kept at 100 °C for 1 h. The Ag_2S coated Canvas Ni electrode was washed

109 by toluene and deionized water carefully afterward, which was finally dried in a vacuum oven at
110 100 °C for 24 h.

111

112 *2.3. Electrochemical Measurements*

113 To investigate the electrochemical deposition behavior and the Coulombic Efficiency of Li
114 stripping/plating on different electrodes, a half-cell measurement in a Swagelok cell was conducted
115 with a Li planar foil (Alfa Aesar, 99.9%) as the counter/reference electrode and the Canvas Ni-
116 Ag₂S electrode with a 10 mm disk shape or Canvas Ni electrode or planar Cu foil (as control group)
117 as the working electrode.

118 To investigate the voltage hysteresis during long-term plating/stripping process, we firstly
119 deposited 8 mA h cm⁻² of Li onto the Canvas Ni-Ag₂S electrode or Canvas Ni electrode at a current
120 density of 1 mA cm⁻² in a Swagelok cell to prepare Li@Canvas Ni-Ag₂S electrode or Li@Canvas
121 Ni electrode. The pre-plated electrodes were taking out from the cells and washed with 1,2-
122 dimethoxyethane (DME). Symmetrical cells were assembled with two identical Li@Canvas Ni-
123 Ag₂S electrodes or Li@Canvas Ni electrodes or Li planar foils (as control group).

124 All of the mentioned cells in this paper were assembled/disassembled in an argon filled
125 glovebox. The galvanostatic discharge-charge cycle performance was carried out on a multi-
126 channel battery tester (Hokuto Denko). The electrolyte was 1 mol L⁻¹ lithium
127 bis(trifluoromethanesulfonyl)imide (LiTFSI, Kishida Chemical, 99.90%) and 1 wt% lithium
128 nitrate (LiNO₃, Kanto Chemical) in a mixed solution of 1,3-dioxolane (Sigma-Aldrich, 99.8%)
129 and 1,2-dimethoxyethane (Sigma-Aldrich 99.5%) (DOL/DME, 1:1 v/v). Dosage of electrolyte

130 used in each cell was 125 μL . A glass fiber separator was placed between two electrodes. In order
131 to remove the surface contaminations and make the electrodes contain the same areal capacity of
132 Li metal, the half-cell measurement was first cycled at 0-1 V versus Li at 50 μA for six cycles. The
133 pre-plating process started when the voltage dropped to 0 V at the sixth discharge step. The cut off
134 voltage was set to 1 V for Li stripping process and -2 V for Li plating process.

135 To investigate the impedance change during long-term plating/stripping process, Li foil | Li
136 foil, Canvas Ni | Li@Canvas Ni, and Canvas Ni-Ag₂S | Li@Canvas Ni-Ag₂S cells were assembled.
137 Here, the Li foil, Canvas Ni and Canvas Ni-Ag₂S were used as the working electrodes. Li was
138 firstly plated onto the working electrodes to a controlled capacity of 4 mA h cm⁻² at 1 mA cm⁻². In
139 this way, the cells were changed to symmetric cells, which were subsequently cycled for repeated
140 plating/stripping at 1 mA cm⁻² with a capacity of 1 mA h cm⁻². The electrochemical impedance
141 spectra (EIS) at different cycles were obtained by a Princeton apparatus (Princeton VersaSTAT 3)
142 with a frequency range of 100 kHz to 0.1 Hz.

143 To evaluate the performance of full cells, lithium cobalt oxide (LiCoO₂) was employed as
144 cathode, while Li@Canvas Ni-Ag₂S electrode or Li@Canvas Ni electrode or Li planar foil was
145 used as anode. LiCoO₂ slurry was prepared by mixing the active material, *poly-vinylidene*
146 *difluoride* (PVDF) binder and carbon black with a weight ratio of 8:1:1 in *N*-methyl-2-pyrrolidone
147 (NMP) (Chameleon Reagent, LBG-50481). After pasting the slurry to an Al foil, the LiCoO₂
148 cathode was dried at room temperature for 2 h and 110 °C for 12 h in a vacuum oven. The areal
149 mass load on the Al foil current collector for the cathode with a disk size of 10 mm was 2.5 mg
150 cm⁻². The electrolyte used was the binary carbonate solution which contained 1 mol L⁻¹ LiPF₆ in
151 ethylene carbonate/dimethyl carbonate (EC/DMC 1:1 v: v) (Chameleon Reagent, LBG-00022).
152 Dosage of electrolyte used in each cell was 100 μL and the separator used in all cells was glass

153 fiber. The galvanostatic charge/discharge measurement was conducted in the voltage range of 3.0
154 V~ 4.2 V for the full cells. The specific capacity of discharge was calculated based on the mass of
155 active material.

156 *2.4. Materials characterization*

157 Morphologies of the samples were characterized by a scanning electron microscope (SEM, ZEISS
158 Sigma 500). The Li composite electrodes after electrochemical test at different status were firstly
159 taken out from the cells and rinsed by DOL/DME to remove the residue of electrolyte. After drying
160 in the glovebox, the electrodes were kept in Ar-filled containers for further ex-SEM or ex-XPS
161 observation. Crystal structures of the materials were measured by X-ray diffractometer (XRD,
162 Rigaku MiniFlex600) using Cu K α radiation. Energy dispersive spectroscopy (EDS) characterized
163 by a scanning electron microscope (SEM, JSM-7500F, JEOL) was also performed on the current
164 collector to confirm the chemical composition. X-ray photoelectron spectroscopy (XPS)
165 measurement were conducted by using an Al-K α X-ray source (JEOL, JPS-9200). The Brunauer-
166 Emmett-Teller (BET) specific surface area and the Barrette-Joynere-Halenda (BJH) pore size
167 distribution of the porous frameworks were characterized on a nitrogen adsorption surface area
168 analyzer (BELSORP-mini).

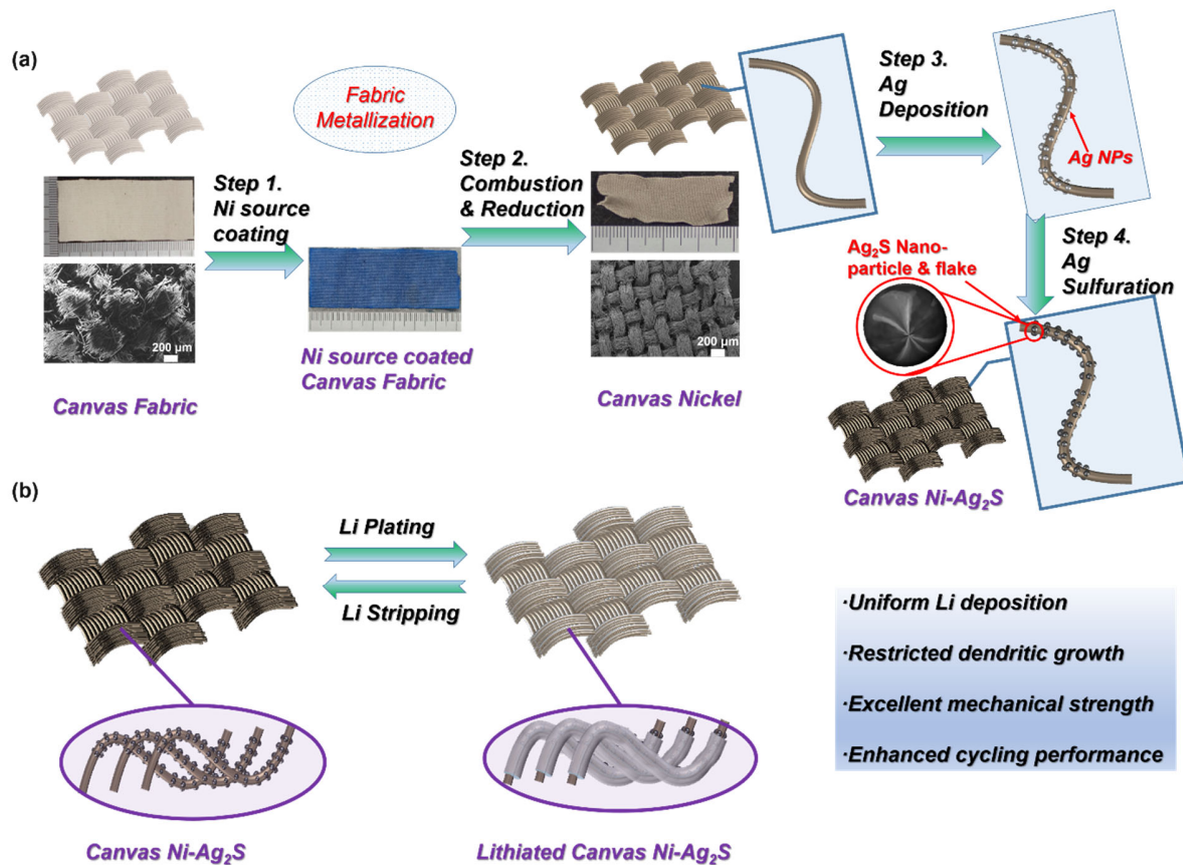
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171 **3. Results and discussion**

172 As shown in **Fig. 1a**, through a series of treatments, including immersion of Ni source to
173 cotton canvas fabric, calcination under air to remove the canvas template and reduction under

174 hydrogen, metallic Canvas Ni was obtained which can preserve the original canvas textile structure.
175 Afterward, uniform deposition of Ag NPs on the Ni substrate was achieved by a silver mirror
176 reaction, and subsequently Ag NPs were sulfurized in a DMSO solution containing sulfur to form
177 Ag₂S NPs decorated Canvas Ni (named as Canvas Ni-Ag₂S). The Canvas Ni framework is served
178 as a 3D current collector with high mechanical strength and high electronic conductivity for
179 accommodating Li. The porous structure allows the 3D current collector to have a weight even
180 lower than Cu foil (see **Table S1**), indicating that this kind of material has a large potential in
181 developing high energy density batteries. The presence of Ag₂S NPs not only act as lithophilic
182 sites for inducing Li deposition into the interior pores of 3D current collector, but also act as a
183 precursor for generating artificial Li₂S SEI. As depicted in **Fig. 1b**, Canvas Ni-Ag₂S electrode
184 shows mainly three advantages: i) the enlarged surface area and the hierarchical micro-nano porous
185 structure of Ni skeleton can effectively decrease the local current density (as shown **Fig. S1**); ii)
186 the Ag₂S NPs can guide homogeneous Li nucleation and prevent dendrite growth of Li; iii) the 3D
187 and hierarchically porous Canvas Ni framework can provide adequate space to alleviate the huge
188 volume change during Li plating/stripping. These characteristics finally ensure the Li anode with
189 a stabilized and long cycle performance at a high current density.



190

191 **Fig. 1.** (a) Scheme of the preparation of Canvas-Ni and Canvas Ni-Ag₂S frameworks. Optical
 192 photos of the fabric template and the Canvas Ni are included. (b) Graphical representation of the
 193 uniform electrodeposition behavior of Li on the Canvas Ni-Ag₂S electrode during plating and
 194 stripping.

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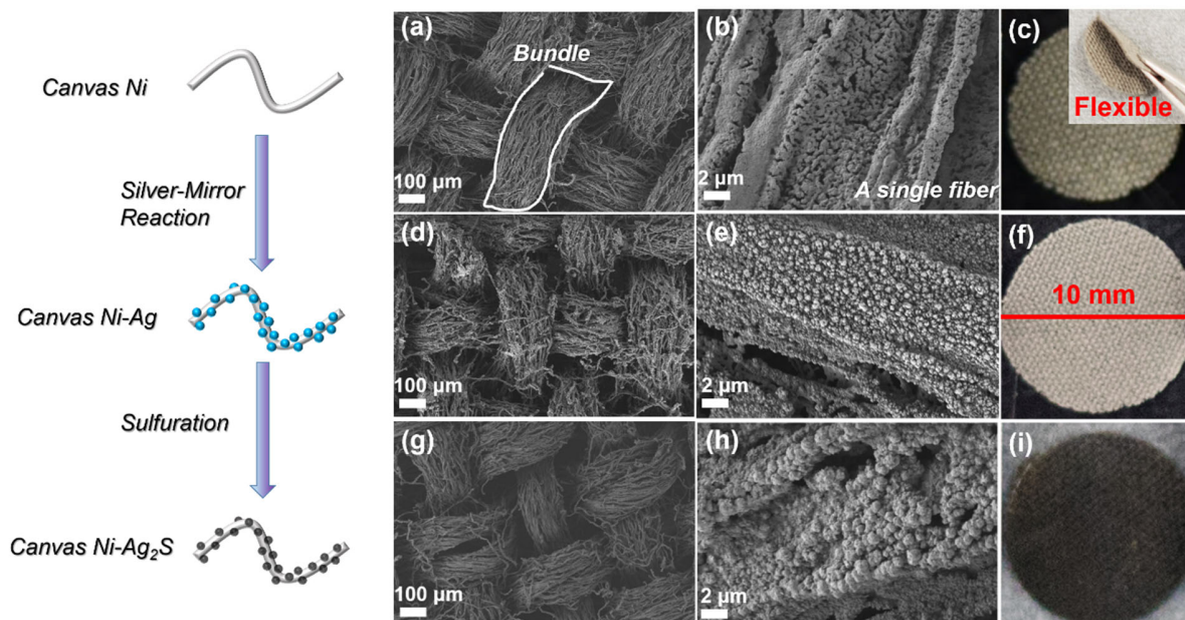
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201 Scanning electron microscope (SEM) images and optical photos of the Canvas Ni disk
202 electrode are presented in **Fig. 2a-c**, and it is worth mentioning that the Canvas Ni is flexible.
203 Similar to the morphology of canvas fabric, metallic Ni fibers are converged to form a “bundle”,
204 which leaves a plenty of voids inside. Careful observation of a single metallic fiber indicates that the
205 fiber is porous containing pores of around 1 μm . More SEM images at different scales are also
206 shown in **Fig. S3**. As previously reported, this pore size is suitable for Li deposition [28]. By a
207 modified silver mirror reaction [39], Ag NPs are evenly deposited onto the surface of Ni fibers
208 through the entire Canvas Ni to form the Ag NPs coated Canvas Ni. After deposition of Ag NPs,
209 color of the Canvas Ni disk changes from gray to silver white (**Fig. 2c** and **Fig. 2f**). The SEM
210 images are shown in **Fig. 2d,e** and **Fig. S4**. There are no structure changes of the textile skeleton
211 except the coating of Ag NPs on Ni fibers, indicating that the destruction of the active Ni substrate
212 by the replacement reaction between Ni and Ag^+ can be avoided through our modified silver mirror
213 reaction. Finally, the Ag NPs are transformed to Ag_2S through a mild sulfurization strategy. The
214 color of the disk electrode changes to black after sulfurization treatment (**Fig. 2i**). The textile
215 structure is well retained and the Ag_2S NPs are constructed with nanoflakes, as shown by the SEM
216 images in **Fig. 2g,h** and **Fig. S5**. The composition analysis by X-ray diffraction (XRD) of the
217 Canvas electrodes treated at different stages is shown in **Fig. S6**, confirming the formation of
218 targeted substances at each step. Energy dispersion spectroscopy (EDS) analysis results of the
219 Canvas Ni (**Fig. S7**) treated by the same sulfurization solution in comparison with EDS results of
220 Canvas Ni-Ag (**Fig. S8**) and Canvas Ni- Ag_2S (**Fig. S9**) confirm that the mild sulfurization reaction
221 is only performed on Ag NPs.

222



223

224 **Fig. 2.** SEM images and optical images for the textile electrode treated at different stages. a,b,c)
 225 as-prepared Canvas Ni electrode, d,e,f) Canvas Ni-Ag and g,hi) Canvas Ni-Ag₂S. The inset in Fig.
 226 2c is the optical photo for showing the flexibility of Canvas Ni electrode.

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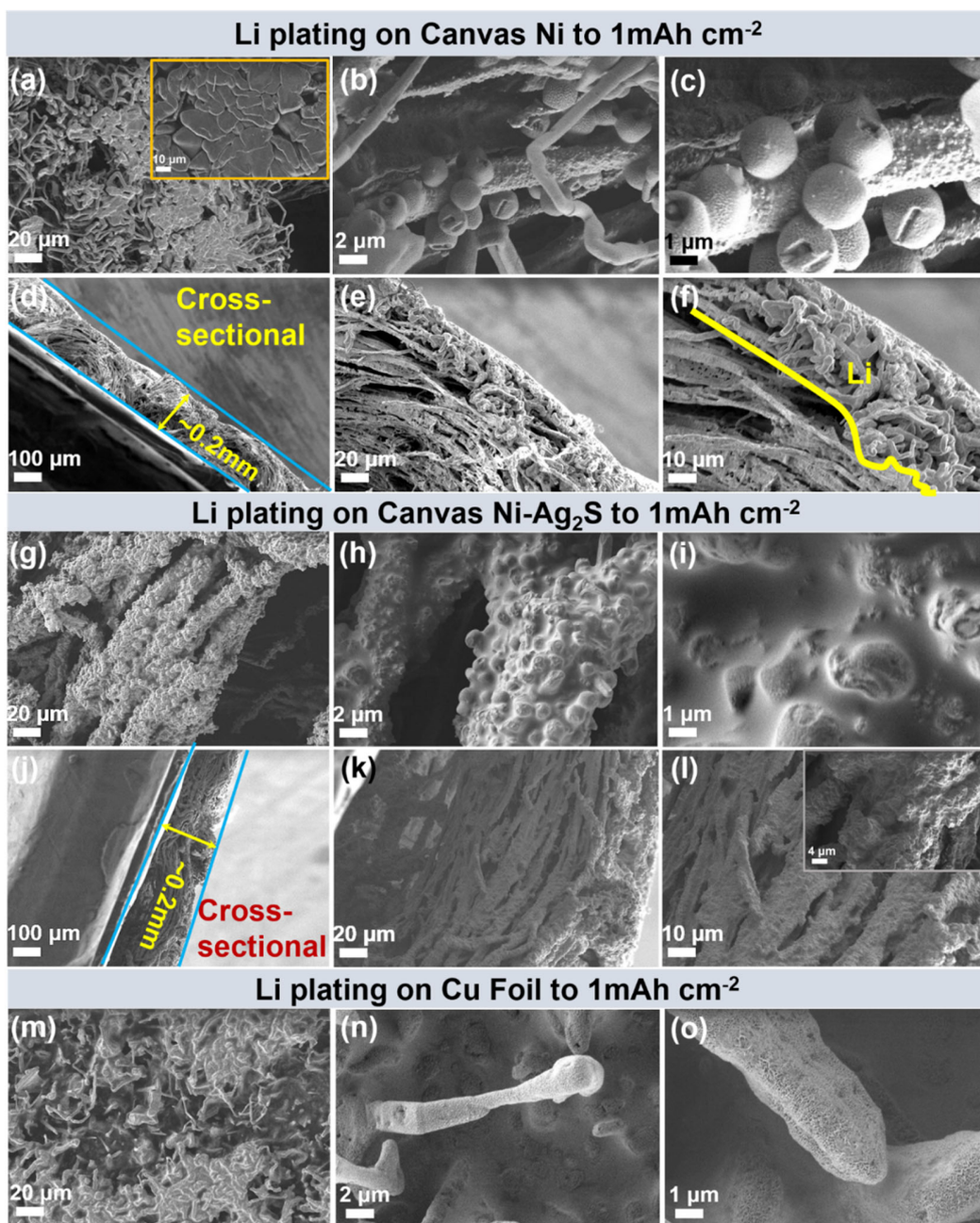
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235 To preliminarily estimate whether the obtained materials are suitable for Li metal anode, the
236 electrodeposition behavior of Li on Canvas Ni, Canvas Ni-Ag₂S and planar Cu foil were
237 investigated, respectively. Li was firstly electrodeposited to an areal capacity of 1 mA h cm⁻² at a
238 current density of 1 mA cm⁻² onto those electrodes after SEI formation pre-treatment, and the
239 morphologies of the Li pre-plated electrodes were captured by SEM. As can be seen in **Fig. 3a**,
240 most of the Li metal is deposited on the current collector to form a flat surface, and spherical Li
241 crystals are deposited around the Ni fibers of the Canvas Ni substrate (**Fig. 3b and 3c**), indicating
242 that the 3D porous Canvas Ni current collector is useful to store Li. However, due to the lack of
243 lithiophilicity for the pure Ni substrate, dendritic Li with a “whisker growth mode” is also observed
244 (**Fig. 3a**) which suggests a high over-potential of Li nucleation on the bare metallic current
245 collector [40]. The as-grown Li dendrites can be merged together and are confined within the
246 spaces and voids among Ni bundles and fibers (**Fig. 3a**), leading to the flat surface morphology on
247 the Canvas Ni electrode (**Fig. 3d**). The nucleation of Li crystals is preferred on the surface side of
248 the bare current collector. As the growth of Li crystals, the diffusion pathway for Li⁺ is obstructed
249 by the previously deposited Li crystals near the surface, therefore, the continuing Li deposition
250 can only proceed at the top surface of Canvas Ni (**Fig. 3e and Fig. 3f**). Therefore, a lithiophilic
251 surface treatment of the Ni substrate is needed to lead the deposition of Li inside the porous
252 electrode. It is also noted that the dendritic Li crystals mainly form at the edges of Ni fiber bundles.
253 This is because of that the top of the electrode is compressed by the separator, hence the dendrites
254 are spread to the spaces between bundles [41]. As a result, the growth of Li crystals can be
255 restricted inside the porous structure, preventing the quick piercing of the separator, which is one

256 advantage of the 3D current collector, particularly, that our textile electrodes have multi-scaled
257 pores.



258
259 **Fig. 3.** SEM images of the Li electrodeposition behavior on different electrodes. a-c) Top view
260 and d-f) cross-sectional observation for the Canvas Ni after Li plating to a restricted capacity of 1

261 mA h cm⁻². g-i) Top view and j-l) cross-sectional observation for the Canvas Ni-Ag₂S after Li
262 plating to a restricted capacity of 1 mA h cm⁻². m-o) Li dendrites on Cu foil after Li plating to 1
263 mA h cm⁻².

264

265 The Li deposition behavior in the porous electrode can be greatly improved by further
266 introducing lithiophilic substances (Ag₂S) on Ni surface. The low magnification SEM image (**Fig.**
267 **3g**) of lithiated Canvas Ni-Ag₂S electrode indicates a dendrite-free morphology. For the Canvas
268 Ni-Ag₂S electrode, Ag₂S NPs are homogeneously covered on the surface of the Ni fibers. Two
269 reactions occur during the initial pre-SEI process in the potential range of 0-2 V vs. Li (**Fig. S10b**):



272 The conversion reaction between Ag₂S and Li at around 2 V induces the formation of Li₂S, which
273 can be observed in XPS spectra of the S 2p species and the Li 1s species shown in **Fig. S11a**. It is
274 regarded that Li₂S is a good SEI component for Li metal anode. [11, 42] The following potential
275 plateau near 0 V is the alloying reaction between Ag and Li. After the stabilization of SEI layer,
276 Li is plated into the porous substrate. Since that Ag-Li alloy is adequately lithiophilic with a high
277 bonding energy to Li, the pre-formed Ag-Li seeds the Li nucleation homogeneously on Ni
278 substrate and guides the Li deposition into 3D porous structure [43, 44]. In this manner, a dendrite-
279 free Li coating layer underneath the dense SEI is formed on the Canvas Ni-Ag₂S electrode, as
280 evidenced by SEM images in **Fig. 3g-i**. This is further confirmed by the cross-sectional SEM
281 observation in **Fig. 3j-l**. The plated Li can be guided into the inner porous structure and
282 homogeneously coated on the Ni fibers, since that the Ag-Li is evenly covered on the Ni fibers and

283 plays a key role of drainage for Li deposition. As a comparison, SEM images (**Fig. 3m-o**) of the
284 Li deposition on a planar Cu foil illustrate needle-like Li dendrites. The dendrites grow vertically
285 on the top of planar Cu foil, posing a huge safety hazard, which may pierce the separator and make
286 a short circuit of the battery at any time. Benefiting from the dendrite-free nature of Canvas Ni-
287 Ag₂S, even after that Li is completely stripped from the electrode, the SEI layer is perfectly
288 remained on the Ni fibers, as shown in **Fig. S12a**. As a comparison, the SEI layer on Canvas Ni is
289 partially destroyed, and it is observed that a small amount of dead Li is remained in the structure
290 (**Fig. S12b**). For Cu foil, a large amount of porous Li clusters are found on the surface after the
291 stripping of Li (**Fig. S13**), indicating that Li@Cu foil anode must have a low Coulombic efficiency
292 (CE).

293 As-encouraged by the above excellent Li deposition behavior on the Canvas Ni-Ag₂S electrode,
294 the repeated Li plating/stripping performance was further investigated. The CE of different
295 electrodes, as shown in **Fig. S14a**, were measured by assembling Li | Cu Foil, Li | Canvas Ni and
296 Li | Canvas Ni-Ag₂S half-cells, respectively. The CE of Li | Canvas Ni-Ag₂S cell reaches a high
297 value of 98% after several cycles, which can be maintained throughout the entire test upon 150
298 cycles, exhibiting an outstanding cyclic reversibility. In contrast, Li | Cu foil cell shows a low CE
299 of less than 95%, and because of the uncontrolled dendrite growth, an internal short-circuit can be
300 observed after 60 cycles. In addition, owing to its micro-nano 3D porous structure which can
301 somewhat confine the Li crystals inside the porous structure, Canvas Ni electrode also shows better
302 cycle performance than planar Cu foil. However, since that large dendritic crystals can be still
303 observed in the top surface and the large pores of the Canvas Ni frameworks (**Fig. 3a,f**), the risk
304 of piercing the separator is obvious for the un-modified 3D current collector. The potential profiles
305 of the plating and stripping processes at different cycles for these cells are shown in **Fig. S14b,c,d**.

306 The polarization of Li plating on Canvas Ni-Ag₂S electrode in the 10th, 50th and 100th cycles
307 manifest potentials of -28 mV, -25 mV and -24 mV vs. Li, respectively. The low value of over-
308 potential for Canvas Ni-Ag₂S in the initial cycles confirms that the pre-generated Li₂S SEI owns
309 high ion conductivity. Moreover, benefiting from the dendrite-free characteristic, a stabilized
310 polarization potential of 25 mV suggests that the SEI layer is maintained during repeated
311 plating/stripping. The sudden increase of potential occurs at the end of each stripping step,
312 indicating that the reversible part of Li metal has been completely stripped from the electrode. It
313 can be seen from these profiles that reversible areal capacity of Li is increasing and stabilizing
314 upon cycling for the Canvas Ni-Ag₂S. This phenomenon also demonstrates that the high
315 conductive SEI layer remains stable during cycling.

316 It is worth pointing out that the half-cell test is generally unable to withstand a large number of
317 cycling due to unsatisfactory cycle performance of Li metal foil used as the counter/reference
318 electrode [45]. In order to further verify the superior cyclic performance of Li on Canvas Ni-Ag₂S
319 electrode, symmetric Li@Canvas Ni-Ag₂S | Li@Canvas Ni-Ag₂S cells were assembled to avoid
320 the instability caused by using Li metal foil. Symmetric cells of bare Li foil and Li@Canvas Ni
321 were also assembled as control groups. Due to that the working electrode and the reference
322 electrode are identical, the voltage hysteresis, namely the offset value between Li stripping and
323 plating, can reasonably reflect the interfacial stability and charge transfer resistance of an
324 electrochemical system at constant current densities. **Fig. 4a** shows the voltage profiles when the
325 cells are running under mutative current densities and capacities. This measurement is useful for
326 simulating the alterable charging/discharging circumstance in actual use. The Li foil | Li foil
327 symmetric cell exhibits a high voltage hysteresis of 40 mV at a current density of 0.1 mA cm⁻².
328 After then, a sudden internal short-circuit in the bare Li foil cell occurs when the current density

329 is increased to 1 mA cm⁻². The size and number density of Li nuclei are directly proportional to
330 current density and capacity respectively [18], thereby, both the diameter and number of Li nuclei
331 would show an extremely uneven distribution under variable current densities and capacities,
332 leading to more severe dendrite growth and easy piercing of the separator. Benefiting from the
333 hierarchical micro-nano structure which can effectively decentralize the local current density,
334 Li@Canvas Ni symmetric cell shows lower voltage hysteresis than the Li foil cell, and the voltage
335 hysteresis is even stable and lower for the Li@Canvas Ni-Ag₂S cell.

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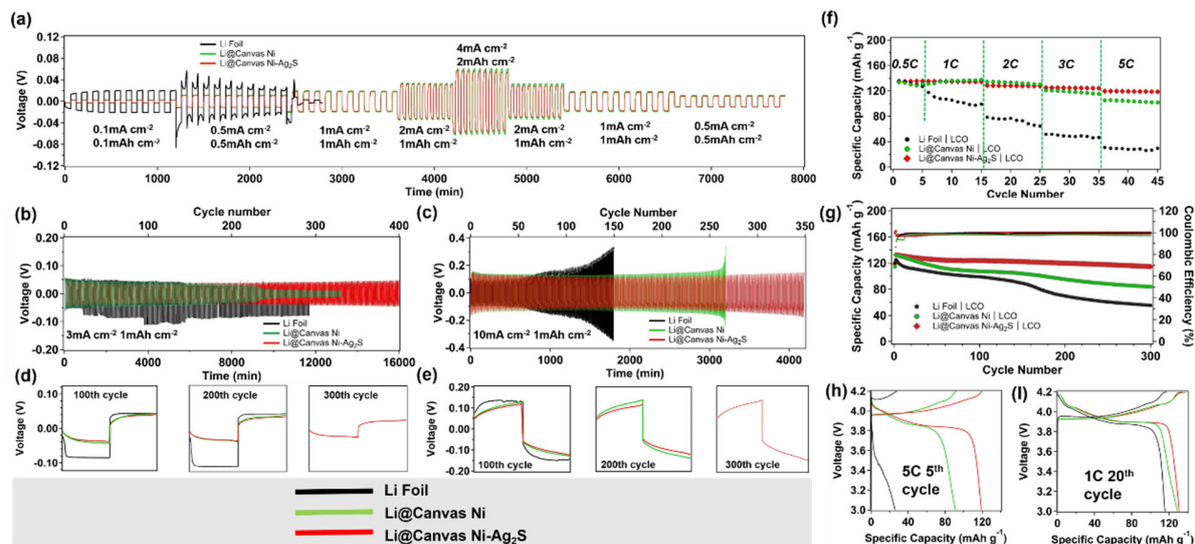
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 345 **Fig. 4.** Electrochemical cycle performance of different electrodes containing Li metal. a, b, c)
 346 Voltage profiles of bare Li foil symmetric cells (black), Li@Canvas Ni symmetric cells (green)
 347 and Li@Canvas Ni-Ag₂S symmetric cells (red) a) at different current densities with different
 348 capacities; b) at 3 mA cm⁻² with a capacity of 1 mA h cm⁻² and c) at 10 mA cm⁻² with a capacity
 349 of 1 mA h cm⁻², d) shows the zoomed-in voltage profiles at different cycles correspond to (b); e)
 350 shows the zoomed-in voltage profiles at different cycles correspond to (c). f) Rate performance
 351 and g) long-term cycle performance of full cells using LCO as cathode. Voltage profiles
 352 corresponding to Li Foil | LCO full cell (black), Li@Canvas Ni | LCO full cell (green) and
 353 Li@Canvas Ni | Ag₂S-LCO full cell (red) at h) 5th cycle with the rate of 5 C and i) 20th cycle with
 354 the rate of 1 C.

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359 The voltage profiles of different symmetric cells measured at 3 mA cm^{-2} with a restricted areal
360 capacity of 1 mA h cm^{-2} are shown in **Fig. 4b**. The details of the voltage-time curves at the 100th,
361 200th and 300th cycles are also captured and exhibited in **Fig. 4d**. In comparison to the high over-
362 potential ($>100 \text{ mV}$) presented by bare Li foil symmetric cell, both Li@Canvas Ni and Li@Canvas
363 Ni-Ag₂S exhibit over-potentials lower than 60 mV at the beginning of cycling. Upon cycling, the
364 over-potentials gradually decrease and finally stabilize to around 40 mV . However, after 200
365 cycles, voltage hysteresis of Li@Canvas Ni cell starts to decrease to an abnormally low value with
366 a flat voltage profile, indicating that a short circuit happens due to the inevitable Li dendrite growth
367 on the un-modified metallic 3D current collector. Significantly, Li@Canvas Ni-Ag₂S cell
368 illustrates stable voltage plateaus even after 300 cycles, demonstrating its superior cycle
369 performance enabled by the uniform deposition of Li as-accelerated by the Ag₂S-induced nuclei.
370 Additionally, the Canvas Ni-Ag₂S electrode can maintain its morphology after 400 cycles (**Fig.**
371 **S18**), indicating that this kind of electrode owns superior mechanical strength to withstand the
372 volume expansion during cycling. Moreover, to meet the demands of fast charging/discharging,
373 cycle performance of the cells under a high current density of 10 mA cm^{-2} were evaluated, as
374 shown in **Fig. 4c** and **Fig. 4e**. A continuously aggravated polarization of Li plating/stripping on
375 bare Li foil is observed, which is represented by the gradual rising trend of voltage hysteresis that
376 starts from around 50th cycle. As for Li@Canvas Ni cell, this trend is delayed until 150th cycle,
377 confirming that a better performance can be achieved by using 3D current collector. The sharp
378 increase of voltage hysteresis in Li@Canvas Ni cell after 260 cycles is observed, implying that the
379 “electrical disconnection” is the failure mechanism of the cell rather than short circuit. As it is well
380 known that upon cycling some Li fragments may detach from the Li substrate and are wrapped in
381 an electronically insulating SEI coating, thus losing their electrochemical activity, and forming the

382 so called “Dead Li” [46]. Meanwhile, a porous and high impedance interphase will be formed on
383 Li metal surface due to the formation of “Dead Li”. This is especially serious when the cell is
384 tested at high current density, appearing as the aggravated polarization in the voltage profiles [45].
385 As for the Li@Canvas Ni-Ag₂S symmetric cell, an alleviated polarization is observed. The
386 hysteresis voltage increases from 240 mV at the 100th cycle to 270 mV at the 300th cycle. Another
387 noteworthy point is the shape of voltage-time curve in each cycle which exhibits a well-defined
388 plateau with smooth upward tilting, indicating the uniform Li plating/stripping behavior for
389 Li@Canvas Ni-Ag₂S electrode [47]. In comparison, the “arch” voltage curve is observed for Li
390 foil cell, indicating that a porous structure is formed on Li foil caused by the formation of “Dead
391 Li”, leading to the non-uniform plating/stripping kinetics on the porous surface.

392 Combining LiCoO₂ (LCO) cathode with Li metal anodes for evaluating the actual
393 performance in full cells, Li@Canvas Ni | LCO, Li@Canvas Ni-Ag₂S | LCO and Li foil | LCO
394 batteries were assembled with traditional binary carbonate electrolyte and the galvanostatic
395 charge/discharge test was conducted. As shown in **Fig. 4f**, Li@Canvas Ni-Ag₂S | LCO exhibits
396 prominent advantages in rate performance, which displays a high discharge capacity of 139 mA h
397 g⁻¹ at 1 C and a satisfying value of 120 mA h g⁻¹ at 5 C, however the discharge capacities of the
398 Li@Canvas Ni | LCO and Li foil | LCO batteries show lower values of 92 mA h g⁻¹ and 28 mA h
399 g⁻¹, respectively. Severe polarization is observed in the charging/discharging voltage curve of Li
400 foil | LCO battery as shown in **Fig. 4h**, which can be ascribed to the sluggish Li⁺ transmission at
401 high current density. **Fig. 4g,l** shows the long-term cycle performance at 1 C. The Li@Canvas Ni-
402 Ag₂S | LCO battery displays an initial capacity of 136 mA h g⁻¹, and the value is slightly decreased
403 to 126 mA h g⁻¹ after 20 cycles due to the process of SEI formation. 96.8% of the initial capacity
404 can be reserved after 300 cycles with a value of 122 mA h g⁻¹. Moreover, a high CE over 99.9% is

405 observed throughout the entire testing stage. As a comparison, the capacities for the Li@Canvas
406 Ni | LCO and Li foil | LCO batteries decrease severely, especially for the Li foil battery. The
407 superior cyclic stability for the Li@Canvas Ni-Ag₂S | LCO battery can be mainly explained by
408 two reasons: i) The pre-generated Li₂S serves as the SEI layer which can mitigate the consumption
409 of electrolyte for generating SEI in the binary carbonate electrolyte. As a contrast, the SEI
410 formation process in Li@Canvas Ni | LCO lasts for a long period, which can be evidenced by the
411 continuous loss of discharge capacity during the initial 100 cycles. ii) The dendrite-free surface
412 can protect the as-generated SEI layer from repeated destruction and regeneration. In contrast, Li
413 foil | LCO battery shows continuous fading of capacity from the initial 125 mA h g⁻¹ to the final
414 48 mA h g⁻¹ due to the unceasing renovation of SEI.

415 The effective principle of Canvas Ni-Ag₂S electrode is depicted in detail in **Fig. 5a**. It is well
416 known that electrolyte is consumed in electrochemical cycle as SEI formation [48]. As confirmed
417 hereinbefore, when dendritic Li is repeatedly formed on the top of a un-modified 3D current
418 collector (Canvas Ni), SEI layer with a low shear modulus experiences incessant destruction and
419 reparation [49], finally resulting in depletion of electrolyte, formation of an ultra-thick SEI layer
420 and creation of large “dead Li”. However, by introducing Ag₂S NPs onto the 3D current collector,
421 Li⁺ can be transported towards inner surface of 3D framework and deposited inside the framework,
422 forming a dendrite-free morphology. Hence, the thickening of SEI and “Dead Li” layer can be
423 greatly alleviated, guaranteeing a high cyclic stability of battery. [15, 50]

424 Electrochemical impedance spectroscopy (EIS) was conducted with different symmetric cells
425 to support these viewpoints, with the spectra shown in **Fig. 5b-d**. The equivalent circuit for fitting
426 the EIS curves is shown in **Fig. 5e**. Here, R_s refers to the electrolyte ohmic resistance, and Z_w
427 denotes Warburg resistance sourcing from the diffusion process of Li⁺ in the bulk of electrode;

428 R_{CT} and R_{SEI} represent resistances of charge transfer process and SEI, respectively; C_{CT} and C_{SEI}
429 are two constant phase elements of electrode and SEI film respectively. The enlarged polarization
430 on the surface of electrodes during cycling significantly influence the value of R_{SEI} , which can be
431 used for evaluating the impedance caused by thickening of SEI and accumulation of “Dead Li”.
432 As shown in **Fig. 5f**, R_{SEI} value of the bare Canvas Ni-Ag₂S electrode is 127.9 Ω for the fresh cell
433 and values at the 50th and 150th cycles are 2.1 and 14.99 Ω , while these values for Canvas Ni are
434 97.6, 5.8, and 56.3 Ω , respectively. The rapid decrease in R_{SEI} after cycling demonstrates the
435 formation of the SEI layer that provides an efficient Li⁺ transfer pathway. Benefiting from the
436 micro-nano structure of Canvas Ni framework, electrolyte is able to contact the active sites
437 sufficiently, leading to a further reduction of R_{SEI} in Canvas Ni and Canvas Ni-Ag₂S as compared
438 with the Li foil. Moreover, the conversion of Ag₂S to metallic Ag can somewhat increase electronic
439 conductivity of Canvas Ni-Ag₂S electrode. Therefore, both the Canvas Ni-Ag₂S and the Canvas
440 Ni showed low R_{SEI} value after 50 cycles. But when the number of cycles is increased to 150, R_{SEI}
441 value of Canvas Ni electrode gradually increased to 56.3 Ω , revealing a severe deterioration of
442 cyclic stability. Meanwhile, R_{SEI} value of Canvas Ni-Ag₂S remains stable for the following cycles,
443 confirming the effectiveness of Canvas Ni-Ag₂S electrode for enhancing cyclic stability of Li
444 metal anode as we depicted hereinbefore. The SEM images after 150 cycles that are shown in **Fig.**
445 **5a**, providing the evidence for the above description. As for the Canvas Ni electrode after EIS
446 measurement, large “Dead Li” agglomerate and fractured SEI can be observed on the surface. The
447 uneven and porous surface leads to non-uniform electrochemical kinetics and high impedance.
448 However, even after a long-term cycling, on the Canvas Ni-Ag₂S electrode, metallic Li is still
449 evenly located underneath the SEI, and the well-preserved electrode without damage can be clearly
450 observed. All of these strongly demonstrate that the Canvas Ni-Ag₂S electrode can significantly

451 enhance the cyclic stability of Li metal anode. Table S2 shows the comparison with the recent
452 frontier reports.

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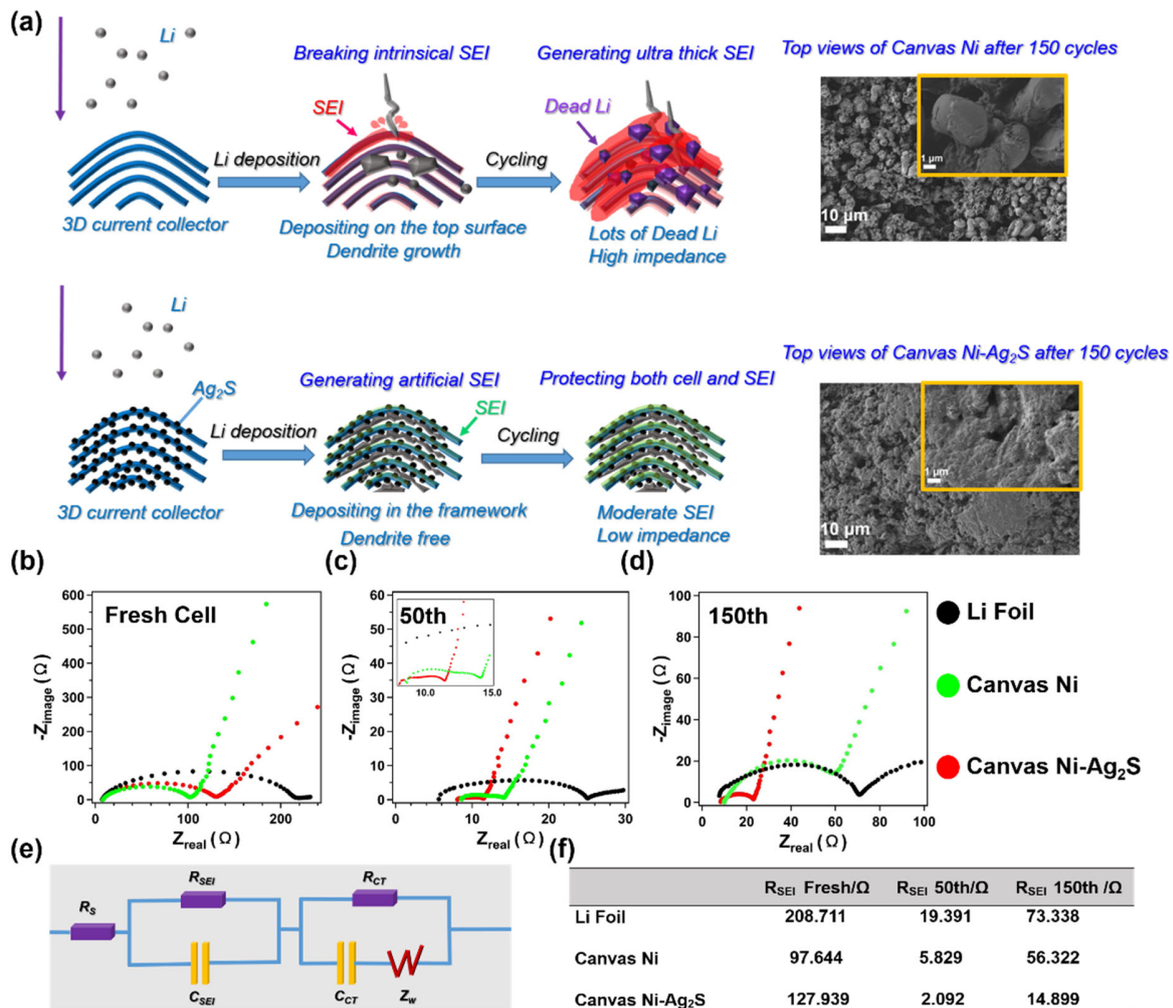
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Fig. 5. (a) Schematic diagram of the multifaceted cycle performance degradation when Li repeatedly plating/ stripping on a common 3D current collector (top), and the enhanced stable cycle performance of the Canvas Ni-Ag₂S (below); morphologies of the electrodes after 150 cycles are included as illustrations. The comparison of EIS of different electrodes (b) before cycling, (c) after 50 cycles and (d) after 150 cycles. (e) The corresponding equivalent circuit drawn for simulating EIS curves. (f) The SEI resistance of different electrodes after different cycles. Results were obtained from symmetric cells measurement at 1 mA cm⁻² with cycling capacity of 1 mA h cm⁻².

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474 **4. Conclusion**

475 In summary, we have fabricated a canvas textile structured Ni current collector for Li metal
476 anode and a new method for promoting uniform electrodeposition of Li metal into 3D current
477 collector by decorating with Ag₂S was introduced. The Canvas Ni 3D current collector whose
478 structure was inspired by cotton textile fabric showed a unique hierarchical micro-nano structure,
479 by which the local current density could be reduced and Li metal could be accommodated
480 becomingly. Moreover, through a modified silver-mirror reaction, Ag₂S NPs were uniformly
481 decorated on Canvas Ni, which offered nucleation sites for Li metal and surface-covered Li₂S pre-
482 SEI. Our Ag₂S-decorated and textile-structured Canvas Ni 3D current collector presented obvious
483 advantages compared to conventional 3D current collector, such as smooth and homogenous Li
484 deposition, stabilized SEI, reduced over-potential and alleviated polarization. Beneficial from
485 these characteristics, the Canvas Ni-Ag₂S electrode showed high CE of 98%, low voltage
486 hysteresis during cycling, superior cyclic stability at high current densities, *i.e.*, up to 400 cycles
487 at 3 mA cm⁻², 1 mA h cm⁻² and 350 cycles at 10 mA cm⁻², 1 mA h cm⁻². When Li@Canvas Ni-
488 Ag₂S was used as the anode in Li | LCO battery, high CE (>99.9%), long lifespan (90.4% capacity
489 was remained after 300 cycles) and high rate capability (120 mA h g⁻¹ at 5 C) were achieved. All
490 of those reinforced that our new 3D current collector could be applied to Li metal anode.
491 Furthermore, this paper also presents a new method for preparing textile metallic framework with
492 a unique porous structure which has never been reported, and we anticipate that many studies
493 requiring the use of bio-structured metals would benefit from this work.

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496 **Appendix A.**

497 Supplementary data

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